Laminin-5 and type I collagen promote adhesion and osteogenic differentiation of animal serum-free expanded human mesenchymal stromal cells

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Abstract

Mesenchymal stromal cells (MSC) are differentiation competent cells and may generate, among others, mature osteoblasts or chondrocytes in vitro and in vivo. Laminin-5 and type I collagen are important components of the extracellular matrix. They are involved in a variety of cellular and extracellular activities including cell attachment and osteogenic differentiation of MSC. MSC were isolated and expanded using media conforming good medical practice (GMP)-regulations for medical products. Cells were characterized according to the defined minimal criteria for multipotent MSC. MTT- and BrdU-assays were performed to evaluate protein-dependent (laminin-5, laminin-1, type I collagen) metabolic activity and proliferation of MSC. MSC-attachment assays were performed using protein-coated culture plates. Osteogenic differentiation of MSC was measured by protein-dependant (laminin-5, and vitronectin, have been identified as regulators of MSC-adhesion and inducers of osteogenic differentiation in vitro.1,10 Fetal calf serum (FCS) is widely used for cell culture,4,7 but in the context of possible further clinical applications animal serum does not meet the European Good Manufacturing Practice (GMP) regulations for medical products. The risk of immunological reactions, prion transmission and potential metabolic changes in cultured cells must be considered.1,13 Therefore, we use a GMP-compatible protocol employing MSC-expansion and differentiation media in which FCS is replaced by human plasma and platelet extract as described previously.11-13 In this study, we examined the effects of laminin-5, laminin-1 and type I collagen on MSC, their in vitro proliferation, adhesion to these matrix proteins and their osteogenic differentiation.

Introduction

Mesenchymal stem cells (MSC) were originally described by Friedenstein et al. in 1966 as clonal, plastic adherent cells with a high proliferation rate and the potential of differentiation into cells of mesodermal origins including osteocytes, chondrocytes and adipocytes in vitro.1 According to the consensus in nomenclature, these cells were termed multipotent mesenchymal stromal cells.2,3 For both cell populations, the acronym MSC may be used.1 The ability of MSC to differentiate into numerous tissues reveals promising potential for tissue engineering, cell therapies and future clinical applications.

Cell differentiation is regulated by soluble factors (e.g. TGF-β, BMPs, insulin-like growth factor, dexamethasone) and the extracellular matrix (ECM) surrounding the cells in which both mechanisms are closely connected. Several ECM-proteins, such as type I collagen, laminin-5, and vitronectin, have been identified as regulators of MSC-adhesion and inducers of osteogenic differentiation in vitro.4 Cell differentiation rate and the potential of differentiation changed at least twice a week. Cells were changed at least twice a week. Cells were

Materials and Methods

Isolation, expansion and cell culture of human bone marrow mesenchymal stromal cells

Bone marrow samples were obtained from 14 patients (age 45 to 72 years) undergoing total hip arthroplasty (femur shafts) or pelvic osteotomy at the Department of Orthopaedic Surgery at UKT (Tübingen, Germany) or the BGU Trauma Center (Tübingen, Germany), after written consent was given. This study has been approved by the local ethics committee of the University of Tübingen (Germany).

Bone marrow was washed with phosphate-buffered saline (PBS, BioWhittaker) and a lipid layer was removed after centrifugation.

MSC were isolated using Ficoll density-gradi- ent centrifugation and mononucleated cells were collected in the interphase (Ficoll-Paque Premium, GE Healthcare, Munich, Germany). The cells were washed with PBS, centrifuged and seeded at a density of 1x10^6 to 1x10^7 cells/cm² in 75 cm² cell culture flacons (BD Biosciences, Heidelberg, Germany) in expansion medium containing low-glucose Dulbecco’s modified Eagle medium (DMEM, Lonza, Basel, Switzerland) supplemented with 5% human fresh frozen plasma (FFP), 5% platelet concentrate (10⁹ platelets/mL medium, Blood Donor Center, University of Tübingen), 2 mM L-glutamine (Lonza, Basel, Switzerland), 1000 IU heparin sodium (Roth, Karlsruhe, Germany), 100 U/mL penicillin G, and 100 μg/mL streptomycin (Gibco, Invitrogen, Karlsruhe, Germany) at 37°C in a humidified atmosphere with 5% air and 5% CO₂. After 24 h of incubation, non-adherent cells were removed and adherent cells were detached by trypsin-EDTA, passedaged and expanded in fresh cell culture medium. The medium was changed at least twice a week. Cells were
detached with Accutase® (PAA, Pasching, Austria) and passaged when reaching 80% confluence. They were seeded at a density of 1.5x10^5 cell/flask. Cells were used for experiments after three passages.

**Characterization of mesenchymal stromal cells**

The MSC were characterized by the defined minimal criteria for multipotent mesenchymal stromal cells, such as adherence to plastic, expression or lack of expression of defined cell-surface antigens analyzed by flow cytometry (CD11b, CD14, CD34, CD45, CD73+, CD90+, and CD105+) and the ability for in vitro differentiation into osteoblasts, adipocytes and chondroblasts.14

**Attachment assays**

For cell attachment assays, laminin-1 (Sigma), laminin-5 (Abcam, Cambridge, MA, USA) and type I collagen (BD Bioscience) were added to 1 ml aliquots of the protein solutions (about 1 µg µL or 5000 µg µL). Tissue culture plates were coated with 1 µL aliquots of the protein solutions (about 1 µg µL of plate area). BSA (bovine serum albumin) served as control. These spots were air-dried and the plates were blocked with 200 µL 1% BSA/PBS (Sigma) for 1 h and washed with PBS prior to the assay. Then 200 µL of MSC suspension in PBS were loaded on the spot for attachment for 15 min, washed with PBS, and adherent cells were microscopically photographed (Zeiss Axiovert 13 S, Jen, Nikon camera).

**MTT cell proliferation assay**

For measurement of protein-dependant differences in metabolic activity of MSC, we used the MTT assay to quantify mitochondrial enzymatic activity; 3000 MSC per well were seeded in a 96-well plate (coated with laminin-5, laminin-1, type I collagen or plastic as control) for 24 h and the activity of mitochondrial reductase was quantified as described by the supplier by measuring the absorbance of the produced agent at λ=490 nm and λ=620 nm after 2.5 h of incubation using the ELISA reader (EL800, Bio Tek, Bad Friedrichshall, Germany).

**BrdU cell proliferation assay**

For further measurement of a possible time-dependent proliferation of MSC we used the BrdU assay quantifying DNA synthesis after three, seven, and 24 h of incubation (BrdU-Kit I (Roche)). Three thousand MSC per well were seeded in a 96-well plate (coated with laminin-5, laminin-1, type I collagen or plastic as control), 10 µmol 5-bromo-2'-deoxy-uridine (BrdU) were added. Analysis of incorporated BrdU was performed with anti-BrdU antibodies which were then linked with anti-mouse IgG fluorescein. Counterstaining was performed with DAPI and visualized using a fluorescence microscope (DM IRBE, Leica).

**Osteogenic differentiation and von Kossa stain**

To explore osteogenic differentiation of the cells, MSC of 10 different patients were seeded at 5x10^4 cells/2 mL/well (coated with 5 µL of 1 µg/µL laminin-5, laminin-1, type I collagen or plastic as control) in 6-well plates. MSC in starvation medium and in non-coated plates/flasks served as controls. After four days of MSC expansion, growth medium was replaced by osteogenic induction medium consisting of high glucose DMEM, FCS 10%, 0.1 µM dexamethasone, 10 nM β-glycerolphosphate, 0.17 mM L-ascorbic acid, 100 U/mL penicillin G, 100 µg/mL streptomycin and fungicides. Progress of osteogenic differentiation was analyzed at various time points after three, seven and 28 days of differentiation. The cultures were stained according to the von Kossa staining protocol to quantify mineralization. The plates were scanned and images were analyzed for staining density (Image J, NIH, USA). The level of confluent blackness was given as a number between 0 and 200, with 0 being black and 200 being white. For visualization, reciprocal value was calculated and normalized to the control.

**Analysis of osteogenic marker genes**

RNA was isolated from cells after osteogenic differentiation for three, seven or 14 days (RNesay-Mini Kit, Qiagen, Hilden, Germany). Cells were detached and the pellets disrupted in RLT buffer (1% β-mercaptoethanol) and RNA was isolated following the manufacturer’s protocol. Reverse transcription of the RNA into cDNA was achieved with 1 µg RNA using a cDNA synthesis kit (Clontech, Mountain View CA, USA). Gene-specific cDNA was amplified by quantitative PCR (LightCycler 1.5, Roche), each according to the manufacturer’s protocol. Primer pairs for alkaline phosphatase (MWG, Ebersberg, Germany), osteopontin, osteocalcin (LCSearch, Heidelberg, Germany, sequences not published) and runt-related transcription factor 2 (Runx-2, MWG) were used. The used primer sequences were:

| Primer | Sequence |
|--------|----------|
| AP up  | 5'- ATT ACC TGG ACA TCG GCA AC -3' |
| AP down| 5'- TTG GGC ACC ACA TCA TAG AA -3' |
| Runx-2  | 5'- TCT GGC CTT CCA CTC GTA GT -3' |
| Runx-2 down | 5'- GAC TGG CGG GGT GTA AGT AA -3' |

In each PCR, GAPDH served as housekeeping reference gene as described.15 To quantify the RNA synthesis the number of RNA copies was divided by the number of GAPDH copies. All gene analyzes were performed on three cultures of MSC and repeated twice.

**Statistical analysis**

Representative data were presented as mean values. Statistical analyses were performed using ANOVA. P<0.05 was considered significant.

**Results**

**Expansion of mesenchymal stromal cells**

MSC were expanded to the second or third passage. The cells presented as fibroblast-like adherent and proliferating cells (Figure 1). All populations (n=14) were characterized by flow cytometry revealing expression of CD73, CD90 and CD105 and the absence or minimal expression of CD14, CD34 and CD45 (data not shown). The proliferation capacity of MSC in GMP-conforming media was high with an average time of 10.6 days till expansion into the next passage (passage doubling time).

**Attachment assay**

Adhesion was measured by incubation of MSC in protein-coated tissue culture plates using cultures of MSC of 9 patients. Coated bovine serum albumin served as blocking agent and as negative control. The assay was performed with protein amounts of 1, 5, 100
and 500 fmol/mL per spot. A protein amount of 5 fmol/mL was the lower limit for attachment to type I collagen and laminin-5, although some MSC attached to spots of 1 fmol/mL type I collagen. Visualization of adherent MSC revealed type I collagen to be most effective, followed by laminin-5 and laminin-1 (Figure 2).

**MTT and BrdU assays**

MTT assay was performed using cultures of MSC of 8 patients. Cells of all cultures were vital without showing significant differences between the proteins used for coating. Growth on cell culture plastic (control) revealed the same cell viability index and metabolic activity (average extinction rate 0.39-0.43; data not shown) compared to the protein-coated samples.

BrdU assay was performed using cultures of 9 patients. MSC were expanded on protein-coated 96-well plates and BrdU-incorporation was measured after three, seven, 16 and 24 h. On average, 73% of MSC of laminin-5 coated well proliferated after 24 h, compared to type I collagen with 70% followed by plastic (67%) and laminin-1 (63%). Differences did not reach statistical significance. After three, seven and 16 h, differences between proteins and plastic were negligible: for example, average proliferation of MSC on laminin-5 coated plates was 25% (3 h), 42% (7 h) and 51% (16 h).

**Osteogenic differentiation, von Kossa stain, and analysis of osteogenic marker genes**

MSC were expanded on protein-coated plates (for von Kossa stain) or in protein-coated cell culture flasks (for transcript analysis of marker genes) with osteogenic differentiation medium for three, seven, 14 and 28 days. Osteogenic differentiation of the MSC was analyzed using von Kossa stain. Plates were evaluated macroscopically and also scanned to explore the degree of mineralization indicated by black silver precipitates. Compared to plastic after 28 days of culture, laminin-5 coated plates contained significant more calcium-phosphate granula (P<0.04), followed by laminin-1 (P<0.09) and type I collagen (Figure 3). No spontaneous osteogenic differentiation of MSC in starvation medium was observed.

To analyze osteogenic differentiation capacity, osteogenic marker genes alkaline phosphatase, osteopontin, osteocalcin and Runx-2 were investigated on 3 cultures. All four genes were expressed whereas analysis of gene expression patterns did not reveal significant differences between MSC cultured in coated flasks and the uncoated control after three, seven and 14 days (data not shown).
Discussion

Cells expanded for clinical applications have to meet the quality standards set by the health authorities (e.g. EMA, FDA). We, therefore, isolated and expanded MSC with animal serum-free medium (FFPP: fresh frozen plasma platelet) meeting current GMP-regulations. Due to different donors and preparation techniques, the amounts and the composition of growth factors varies within the plasma employed here. For that reason, serum samples from 15-25 individuals tested for growth factor content and sera from suitable donors were then pooled and stored at -70°C. Comparably, MSC vary considerably depending on health and age of MSC donors, and depend on culture conditions, number of passages, and other variables during in vitro expansion.

MSC expanded in FFPP medium showed a high proliferation capacity and fulfilled the minimal criteria for MSC with expression of CD 90, CD 73 and CD 105 and lack of CD 14, CD 34 and CD 4, as described in previous studies. By MTT and BrdU assays, MSC high proliferation rates were confirmed, but without detectable influences by the coating proteins employed on the metabolic activity. This could be due to the short time of exposure to the proteins (only 24 h). Furthermore, FFPP media activated higher proliferation of MSC compared to standard media. Therefore, minor changes in MSC proliferation facilitated by laminins or collagen possibly remained below detection levels. Alternatively, the cell cycle and mitochondrial metabolism was not directly influenced by ECM-proteins investigated in our study. In contrast, Lindner et al. found a higher proliferation rate of MSC in the presence of ECM proteins when they studied a longer proliferation period compared to our study.

Kees et al. revealed, that laminin-5 induces adhesion, osteogenic gene expression and differentiation of human MSC through an ERK-dependent pathway. In their experiments, MSC were expanded in media containing FCS. Similar results are described for type I collagen. By using FFPP medium for culturing MSC, our study supports the previous findings of increased binding of MSC to the ECM-proteins type I collagen and laminin-5. In that context, we have to consider that we used a specific amount of proteins without noting their exact amount of binding sites, especially when comparing different proteins. Concerning laminin-5, the 3 LG3-domain was described to induce adhesion and osteogenic differentiation of MSC. Isolated use of such domains (peptide fragments) would possibly increase binding strength to MSC and could be part of further studies.

Evaluation of von Kossa stain was performed by scanning the plates and measuring the blackness of the silver precipitates. Evaluating the amount of produced calcium granula is described as an alternative.

Regarding osteogenic differentiation, our study revealed a significant advantage of laminin-5 compared to type I collagen-coated or uncoated plastic growth after 28 days of culture in osteogenic differentiation medium, which is consistent with previous studies. Laminin-1 also promoted osteogenesis. But this did not reach statistical significance (P<0.09).

We found only slight differences in RNA expressions of osteogenic marker genes in MSC after a few days of osteogenic induction on protein-coated versus non-coated dishes, which correlate with a previous study. One explanation would be the small number of observed probes. In addition, extraction of mRNA from mineralizing probes is technically challenging. Therefore, progress in osteogenic differentiation makes it increasingly difficult to extract suitable mRNA for cDNA synthesis and quantitative PCR analyses. Furthermore, MSC from bone marrow are already primed for bone repair. In contrast to, for example, MSC from placenta, they express some genes associated with osteogenesis spontaneously in vitro. RNA analyses are very sensitive and we revealed osteogenic differentiation stain even in non-coated plates after 3 days.

In summary, we report that human bone marrow derived MSC proliferate well in animal serum-free GMP-conforming media, attach to type I collagen, and with lower affinity to laminin-5 and laminin-1. Osteogenic differentiation of MSC is not promoted by type I collagen, but by attachment to laminin-1 and more so by laminin-5. We conclude that coating orthopedic implants with type I collagen may not facilitate regeneration of bony defects, despite its high affinity to MSC. Our data confirm that laminin-5 or laminin-5-derived peptides serve better in bone repair, as this material promotes both firm attachment and osteogenic differentiation of MSC.

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