Development of a model system to analyze chondrogenic differentiation of mesenchymal stem cells

Anke Ruedel*, Simone Hofmeister*, Anja-Katrin Bosserhoff

Institute of Pathology, University of Regensburg Medical School, Franz-Josef-Strauss-Allee 11, D-93053 Regensburg, Germany. *Equal contributors.

Received September 17, 2013; Accepted October 2, 2013; Epub November 15, 2013; Published December 1, 2013

Abstract: High-density cell culture is widely used for the analysis of cartilage development of human mesenchymal stem cells (HMSCs) *in vitro*. Several cell culture systems, as micromass, pellet culture and alginate culture, are applied by groups in the field to induce chondrogenic differentiation of HMSCs. A drawback of all model systems is the high amount of cells necessary for the experiments. Further, handling of large experimental approaches is difficult due to culturing e.g. in 15 ml tubes. Therefore, we aimed to develop a new model system based on “hanging drop” cultures using 10 to 100 fold less cells. Here, we demonstrate that differentiation of chondrogenic cells was induced as previously shown in other model systems. Real time RT-PCR analysis demonstrated that Collagen type II and MIA/CD-RAP were upregulated during culturing whereas for induction of hypertrophic markers like Collagen type X and AP-2 epsilon treatment with TGF beta was needed. To further test the system, siRNA against Sox9 was used and effects on chondrogenic gene expression were evaluated. In summary, the hanging drop culture system was determined to be a promising tool for *in vitro* chondrogenic studies.

Keywords: Chondrogenic differentiation, hanging drop, human mesenchymal stem cells, collagen

Introduction

Model systems to analyze cartilage development *in vitro* are widely used. With these, detailed analysis of molecular processes during differentiation of mesenchymal stem cells (MSC) is feasible. Here, cells can be manipulated e.g. using expression constructs or siRNA. These aspects made these models indispensable tools for studying chondrocyte differentiation in great detail in addition to *in vivo* studies. Several model systems are used to investigate chondrogenesis *in vitro* following the literature, e.g. micromass culture [1, 2], alginate culture [3, 4] or pellet culture [5, 6]. All models are based on high-density cell culture preventing the MSCs from adhesion and therefore creating a suitable environment for the cells for differentiation and for maintaining the chondrocyte phenotype. In all these model systems the MSCs produce certain differentiation markers like for example Collagen type II, which is a marker for early stage chondrogenesis, or Collagen type X which is produced by hypertrophic chondrocytes. In alginate culture the alginate is used to provide a suitable environment for chondrocytes to e.g. migrate into defects and repair cartilage (Hunziker 1996). Alginate is an unbranched linear copolymer of 1-4 linked β-D-mannuronic acid and α-L-guluronic acid. It forms gels in the presence of Ca$^{2+}$ and is used to entrap cells for *in vitro* cultivation. In this model the cells are able to completely differentiate into hypertrophic chondrocytes and produce extracellular matrix like *in vivo*.

Another high density culture method is the pellet culture in which cells form a tightly aggregated cell mass which mimics embryonic cartilage development [6, 7]. A third system which is widely used is the micromass culture, which was first described by Ahrens et al in 1977 [1]. Zhang et al, were able to show that in this culture method the expression of Col I and Col X
In vitro analysis of chondrogenic differentiation

3043
Int J Clin Exp Pathol 2013;6(12):3042-3048

was down-regulated compared to pellet culture [7]. This indicates that micromass culture is a good tool to mimic hyaline cartilage.

All these model systems nicely exemplify the cellular differentiation in the developing cartilage; however, the biggest disadvantage is the number of cells necessary and the culturing systems. In most cases, cell numbers of around 200,000 to 600,000 cells per treatment are needed. Further, several systems, like the pellet culture, are performed in 15 ml tubes making the handling and the space needed rather problematic and expensive. Therefore, we aimed to transfer a model known from analysis of tumor cells, the hanging drop model, to analysis of cartilage tissue.

Material and methods

Primary human mesenchymal stem cells

Human mesenchymal stem cells (HMSC; Provitro, Berlin, Germany) were cultivated in HMSC proliferation medium (Provitro) under a humidified atmosphere of 8% CO₂ at 37°C. Cells of four different donors were used in passages 4 to 10. Transfection of the HSMCs with siRNA against Sox9 (CTCAC-CTTCACCTCATGAA, Qiagen, Hilden, Germany) or the respective control was performed using Lipofectamine 2000 (Invitrogen, Darmstadt, Germany). The cells were cultured in T75 culture flasks to 30-50% confluency. For transfection 750 pmol siRNA were used in the transfection solutions. The transfection was performed following the manufacturer’s instructions. The cells were harvested 48 h later, and used for the hanging drops.

Hanging drop assay

To generate spheroids the HMSC were detached from the culture flask by adding 1 ml Trypsin-EDTA (Provitro). After incubation for 5 minutes 1 ml neutralizing solution (Provitro) and 8 ml HMSC proliferation medium (DMEM with glucose (4.5 g/L), penicillin (400 units/ml) streptomycin (50 µg/ml), L-glutamine (300 µg/ml), sodium pyruvate (1 mM), L-ascorbic acid 2-phosphate (0.17 mM), L-proline (0.35 nM), dexamethasone (1 µM) and ITS premix (BD Biosciences, Heidelberg, Germany)), the cells were counted and

Table 1. Primer sequences

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer</th>
</tr>
</thead>
</table>
| 3043
| ß-actin forward | 5'-CTA CGT CGC CCT GGA CTT CGA GC |
| ß-actin reverse | 5'-GAT GGA GCC GCC GAT CCA CAC GG |
| Coll X forward | 5'-AGCCAGGGTTGCCAGAGCAGCA   |
| Coll X reverse | 5'-TTTTCCCACTCCAGGGAGGC   |
| Mia forward | 5'-CATGCTATCCGTCTATGCAGCACCAGCTG |
| Mia reverse | 5'-GATAAGCTTTCACTGGCAAGTAGAATC |
| Sox9 forward | 5'-CGAAGCAAGCAGACACAGCA   |
| Sox9 reverse | 5'-AGTGAAGTGGAGTAGAGGC |
| Aggrecan forward | 5'-CTCAACAGGAGTAGCAGGC |
| Aggrecan reverse | 5'-TGGTAGGAGCTGGTAC |
| Coll II forward | 5'-AGGGCAATAGCAGGTTCACG |
| Coll II reverse | 5'-GGTCAAGGGTCAGCATTCA   |
| PS4th forward | 5'-TTCTCCAGATCCATCGAG   |
| PS4th reverse | 5'-ATCCAAATCAGACTACGGC |
| AP2c forward | 5'-GGAGTAAGGAGGAGGGTCTCTC |
| AP2c reverse | 5'-GCACCAACTCCACGAGCCTC |

was down-regulated compared to pellet culture [7]. This indicates that micromass culture is a good tool to mimic hyaline cartilage.
adjusted to 50,000 cells/ml. 20% methocel (6 g methyl cellulose (Sigma-Aldrich, Munich, Germany), 250 ml basal medium) was added and 25 µl of the cells suspension were dropped onto the cover of a 9 mm petri dish. For each experimental condition 10 drops were used. The petri dish was filled with PBS and the cover dish was inverted (Figure 1) and incubated for 72 h under a humidified atmosphere of 8% CO₂ at 37°C for 1, 2, 4 and 7 d followed by RNA isolation or alcian blue staining. For treatment, TGF-beta3 (10 ng/ml, Biomol, Hamburg, Germany) was added. The experiment was repeated 4 times.

**Pellet culture assays**

HMSC differentiation was also performed as 3D pellets. For this, 2.5 × 10⁵ HMSCs were seeded into 15 ml polypropylene tubes and pelleted. HMSCs were cultured for 0, 3, 7, 9, 11, 13, 17, 19, 23, 27, or 33 days as 3D pellets in serum-free induction medium (DMEM-high glucose, 0.1 µM dexamethasone, 1 mM sodium pyruvate, 0.17 mM ascorbic acid-2-phosphate, 0.35 mM proline and 1:500 insulin–transferrin–selenite (Becton Dickinson, Heidelberg, Germany)). The experiment was repeated three times.

**Figure 2.** Expression of marker genes for early chondrogenic differentiation. A: The HMSCs were cultivated in the hanging drop setting for 7 d and mRNA expression of Sox9, Collagen type II, and MIA/CD-RAP was measured on day 1, 2, 4 and 7 by quantitative RT-PCR (n=4). MIA/CD-RAP (p=0.0006) and Col II (p=0.0002) were significantly up-regulated over time, indicating that the HMSC undergo differentiation, although Sox9 expression did not change. B: Western blot analysis of Sox9. Protein levels of Sox9 were similar at all time-points. C: MIA ELISA showed that expression of MIA protein increased over time, therefore indicating higher differentiation states of the chondrocytes at day 7 compared to day 1 (n=3, p=0.02). D: RNAs from classical pellet culture [13, 14] were analyzed for Col II and MIA/CD-RAP mRNA expression at day 0, 3, 7, 9, 11, 13, 17, 19, 23, 27, and 33 by quantitative RT-PCR. Both genes were up-regulated over time. The levels at days 23-30 were comparable to the mRNA levels of day 7 in the hanging drop setting. (*: p<0.05; ***: p<0.001; ns: p>0.05).
In vitro analysis of chondrogenic differentiation

Isolation of RNA and reverse transcription

For RNA isolation 10 spheroids were pooled. Total cellular RNA was isolated from cultured cells using the Micro Elute Total RNA Kit (VWR, Darmstadt, Germany) and cDNAs were generated by reverse transcriptase reaction performed in 20 μl reaction volume containing 4 μl of total cellular RNA, 4 μl of 5x first strand buffer (Invitrogen, Darmstadt, Germany), 2 μl of 0.1 M DTT, 1 μl of dN6-Primer (10 mM), 1 μl of dNTPs (10 mM) and DEPC-water as described previously [8]. The reaction mixture was incubated for 10 min at 70°C, 200 units of Superscript II reverse transcriptase (Invitrogen) were added and RNA were transcribed for 1 h 37°C. Reverse transcriptase was inactivated at 70°C for 10 min and RNA was degraded by digestion with 1 μl RNase A (10 mg/ml) at 37°C for 30 min.

Analysis of mRNA expression by quantitative real time PCR

Quantitative real time PCR was performed on a Lightcycler (Roche, Mannheim, Germany), cDNA template (2 μl), 0.5 μl (20 mM) of forward and reverse primers (Table 1) and 10 μl SybrPremix ExTaq (Takara (Lonza), Cologne, Germany) in a total of 20 μl were applied to the following PCR program: 30 s 95°C (initial denaturation); 20°C/s temperature transition rate up to 95°C for 15 s, 3 s – 60°C, 5 s – 72°C, 80°C acquisition mode single, repeated for 45 times (amplification). The PCR reaction was evaluated by melting curve analysis and checking the PCR products on 1.8% agarose gels. The values were calculated relative to β-actin and relative to the expression of the gene of interest in HMSC monolayer.

Western blot and histological staining

Western blot analysis was performed as described before (Schubert et al., 2009). Sox9 antibody (Merck Millipore, Billerica, MA, USA) was used in a 1:1000 dilution in 5% dry milk/TBS-Tween (0.1%).

Enzyme-linked immunosorbent assay (ELISA)

MIA protein was quantified in cell culture supernatants by an ELISA system as described in 2003 by Bosserhoff et al. [9].

Statistical analysis

In the bar graphs, results are expressed as mean±S.D. (range) or percent. All calculations were made using the GraphPad Prism Software (GraphPad Software, Inc., San Diego, USA).

Results

The hanging drop model, established in the early 1900s, has been adapted to the application in the tumor field [10, 11] in the last years, and was now adapted to HMSCs to approach the problems occurring using other 3D-model systems mentioned above.

As described in detail in the Material and Method section the assay was performed in petri dishes using hanging drops of medium each containing about 1000 HMSCs (see Figure 1 for schematic illustration). Usually for each experimental approach 10 parallels are used. The cells quickly formed spheroids and started to differentiate (data not shown). After incubation for 1, 2, 4 and 7 days RNA was isolated and used for further experiments. Of course, also analysis for protein content or GAGs in the supernatant is feasible.

To investigate the differentiation of the MSCs several marker genes, like Collagen II, MIA/CD-RAP, Collagen X and AP-2 epsilon were ana...
In vitro analysis of chondrogenic differentiation

In contrast to early induction of cartilage differentiation marker like Collagen II or MIA/CD-RAP the mRNA levels of Collagen X and AP-2 epsilon, both expressed in later phases of cartilage differentiation, stayed only weakly induced until day 7. Here, TGF-beta treatment of the MSCs strongly enhances expression of late differentiation markers starting at day 2 (Figure 3).

In an experimental approach using siRNA against Sox9 the validity of the new assay sys-
In vitro analysis of chondrogenic differentiation

3047

Int J Clin Exp Pathol 2013;6(12):3042-3048

tem was evaluated. The siRNA used for the transfection was already published to decrease Sox9 expression efficiently by Wenke et al. in 2009 [8]. HMSCs were transfected with siRNA and used in the assay 2 days after transfection. The knock-down was stable for 7 d in the spheroids and in monolayer on mRNA and protein levels (Figure 4A, 4B). MessengerRNA expression of several chondrocyte genes was analyzed by quantitative RT-PCR (Figure 4C). At all time points strong reduction of Sox9 mRNA expression was found compared to siRNA control (set as 1). Significant reduction of gene expression was observed for Aggrecan (p=0.0057) in the siSox9 treated cells. MIA/CD-RAP, Col II showed a tendency towards reduced gene expression (Figure 4C). As we assumed this might be due to redundancy mechanisms of p54

Discussion

In this study we aimed to establish a model system for high-density cell culture models of chondrogenesis due to several disadvantages of the established systems. The main disadvantage of these is the high number of cells necessary. According to the literature all model systems require 5 × 10⁵ to 4 × 10⁶ cells per batch [18, 19]. Either for experimental settings with many conditions, with murine or rat primary cells or using transfection these numbers are hard to reach. Further, handling of the established assay systems is time- and space-consuming and in some instances prone to contamination due to long periods of culturing and cell culture devices used.

The new format is based on “hanging drop” culturing. The hanging drop method has first been used to cultivate nerve fibers and fibroblasts isolated from chick embryos [20, 21]. Since then, the method has been modified and applied to several kinds of cell types, e.g. human embryonic stem cells [22] or tumor cells [10]. Here, we applied the hanging drop method to human chondrocytes. The amount of cells is 10 to 100 fold less than in other 3D cell models although the methods can be equivalently used for studies of gene regulation of early chondrogenesis. Comparison of two assays, the new hanging drop and the classical pellet culture, showed that the results were comparable with regards to induction of differentiation and marker gene expression. These findings let us state that the hanging drop system provides a 3D environment similar to cartilage formation during embryonic development and allows HMSCs to undergo chondrogenesis. Interestingly, the cells in the classical pellet culture experiment differentiated much slower whereas differentiation was strongly induced in the hanging drop model already after 4 days. Potentially, this depicts another advantage for experiments where cells are transfected prior to differentiation or expensive, sensitive treatments are performed. Regarding this aspect it is also of importance that only small amounts of substance for treatment are needed because of the culturing volume of 25 µl. As bone marrow-derived MSCs are heterogeneous due to donor differences the quality of the new assay is supported by the low standard deviation (see Figure 2A) although cells from 4 different donors were used in independent experiments. The experimental setting applied in this study using siRNA against Sox9 demonstrated that application of siRNA is feasible and effective in this model. It, further, proves that effects on gene expression can be determined in detail. In summary, we suggest that the hanging drop model is a good alternative for analyzing chondrogenic differentiation which requires less space, time and costs but still provides results comparable to other, established 3D models for chondrocyte differentiation.

Acknowledgements

We thank Sibylla Lodermeyer for excellent technical assistance. This study was supported by a DFG grant to AKB.

Disclosure of conflict of interest

None.

Address correspondence to: Dr. Anja-Katrin Bosserhoff, Institute of Pathology, University of Regensburg Medical School, Franz-Josef-Strauss-Allee 11, D-93053 Regensburg, Germany. Tel: +49 941 944 6705; Fax: +49 941 944 6602; E-mail: anja.bosserhoff@klinik.uni-regensburg.de
In vitro analysis of chondrogenic differentiation

References