Pressureless Mechanical Induction of Stem Cell Differentiation Is Dose and Frequency Dependent

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Abstract

Movement is a key characteristic of higher organisms. During mammalian embryogenesis fetal movements have been found critical to normal tissue development. On the single cell level, however, our current understanding of stem cell differentiation concentrates on inducing factors through cytokine mediated biochemical signaling. In this study, human mesenchymal stem cells and chondrogenesis were investigated as representative examples. We show that pressureless, soft mechanical stimulation precipitated by the cyclic deformation of soft, magnetic hydrogel scaffolds with an external magnetic field, can induce chondrogenesis in mesenchymal stem cells without any additional chondrogenesis transcription factors (TGF-β1 and dexamethasone). A systematic study on the role of movement frequency revealed a classical dose-response relationship for human mesenchymal stem cells differentiation towards cartilage using mere mechanical stimulation. This effect could even be synergistically amplified when exogenous chondrogenic factors and movement were combined.

Introduction

Chemical signaling organizes the structure of biological tissues[1] and stem cell differentiation during growth or repair. Diffusion, local release of growth factors and concentration gradients shape an organism’s 3D structure. Unfortunately, liquid filled pockets and particularly movement destroy such local information pattern (e.g. in a joint, see Figure 1) and challenge our present understanding of static cell-cell assemblies’ signaling[2,3]. We therefore hypothesize that soft movement is a key local tissue structuring factor. The role of movement has probably been best studied in tissue engineering for the growth of articular cartilage in vitro[4-8] using mechanical top load (i.e. pressure)[9-11]. Chondrocyte growth has been associated with demanding physical input such as cell deformation[12], hydrostatic pressure gradients[13], fluid flow[14], streaming currents[15] and physicochemical changes[16,17]. Similarly, substrate strain has also been used to improve hMSC differentiation into vascular smooth muscle cells for vascular tissue repair[18]. Clinical evidence provides a clear argument for the necessity of movement in function-guided, local cell differentiation[19,20]: Mammalian embryogenesis is characterized by an early start of intense movement (for humans, typically in the first third of gestation), a prerequisite to normal organ development[21] and takes place in a pressure-free environment. A second piece of evidence comes from the required absence of movement for healing bone fractures. Without adequate traumatic surgery or fixation, pseudoarthrosis may occur at non-fixed bone fracture sites[22]. Continued movement of such non-fixed primary callus then promotes local stem cell differentiation into cartilage and ligament tissue instead of bone tissue. A third argument arises when normal movement is impeded. If a joint is not moved anymore (e.g. due to arthritis), even an otherwise fully functional joint stiffens (ankylosis) and the tissue locally transforms into a bone-type material[23]. A fourth argument is counter-intuitive: Mechanical stress on ligaments regeneration has recently been shown by Altmann et al[24] to improve the healing process.

These clinical facts underline the importance of movement (Figure 1) and suggest that movement may follow a dose/response type relationship, similar to classical chemicals. Static mechanical properties (porosity, matrix elasticity/stiffness) have recently emerged as key factors in cell lineage specification[25,26]. In order to test our hypothesis, we took...
normal embryonic movement frequency as a starting dose[21] to run an in vitro study on the role of movement as a differentiation factor. We used pressure-free, soft and smooth deformations of a magnetic hydrogel scaffolds by time-programmed application of external magnetic fields. Human mesenchymal stem cells (hMSC) and chondrogenic differentiation were chosen as representative examples[27-29].

We demonstrate that purely mechanical stimulation provokes correct differentiation even in vitro and in the complete absence of any exogenous biochemical differentiation factors. This shows that mechanical stimulation is a key local organization factor at the cellular level. In a second series of experiments, we demonstrated that the frequency of stimulation and overall number of applied stimulations follow a dose-response relationship for these stem cells. This is similar to the present (biochemical) understanding where specific concentrations of the differentiation factor TGF-β indeed induce chondrogenesis[30] to a specific level of differentiation[31].

Materials and Methods

Detailed methods on nanomagnet functionalization, characterization of functionalized nanomagnets and magnetic hydrogel, cell isolation, expansion and characterization, biochemical analysis and immunohistochemistry are provided in the Information in File S1.

Magnetic hydrogel synthesis

Appropriate amounts of 2-hydroxy-ethyl-methacrylate (HEMA, 5.1 mL, puriss. ≥99%, Fluka), ethylene glycol dimethacrylate (EGDMA, 4.9 µL, purum ≥97%, Fluka) and styrene maleic anhydride copolymer (SMA® 1000H, 0.75 g, 36%, Sartomer) solution were mixed in water (25 mL, Millipore). Subsequently, 4-vinylbiphenyl functionalized nanomagnets (5 g) were suspended in the mixture using an ultrasonic probe (UP400S, 24 kHz, Hielscher GmbH) during 5 min. After addition of a rheology additive (BYK® 420, 0.782 g, Brenntag Schweizerhall AG), tetramethylethylenediamine (TMEDA, 31.9 µL, 99%, Sigma Aldrich) and ammonium persulfate (APS, 205 mg, 99%, Sigma Aldrich) the mixture was processed again with the ultrasonic probe for 5 min. The reaction mixture was then poured into an electrophoresis casting mold (gel casting mold, kuroGEL Midi 13) and reacted for 1 h at ambient temperature. The obtained gel was treated with deionized water for 24 h while the water was changed 3 times. The procedure was repeated with phosphate-buffered saline (PBS, pH 7.4, GibCO) to remove unreacted monomer and nanomagnets and to obtain a stable swelling behavior of the hydrogel. The obtained deep-black magnetic hydrogel (2 mm thickness) was punched out to the desired shape and sterilized in 70% ethanol. Sterile scaffolds were rinsed 3 times with fresh PBS to remove any residual ethanol. A dog-bone like shape of the scaffold (Figure 2c) was chosen to increase the flexibility and enable a hammock like deformation. This structure allowed for minimization of the required magnetic force.

Scaffold seeding

A homogeneous distribution of seeded cells onto the hydrogel surface was obtained by using small volumes of relatively high concentrated cell suspensions (2.4•10⁶ P3 hMSC mL⁻¹) either in control (DMEM, 10% FBS, 1% antibiotic/antimycotic) or chondrogenic medium (DMEM, 1% antibiotic/antimycotic, 100 nM dexamethasone (water-soluble, cell culture tested, Sigma-Aldrich), 6.25 µg mL⁻¹ insulin (bovine pancreas, Sigma-Aldrich), 50 µg mL⁻¹ ascorbic acid (≥95%, Sigma-Fine Chemicals), 40 µg mL⁻¹ L-proline (≥98.5%, Sigma-Fine Chemicals), 6.25 µg mL⁻¹ ITS (BD ITS+ Premix, BD Biosciences) and 10 ng mL⁻¹ transforming growth factor beta 1 (rTGF-β1, R&D Systems, [30]). Precut and sterilized hydrogel scaffolds were fixed with the aid of sterile stainless steel rings in wells of a 12 well plate (non-tissue-culture-treated, Falcon). The lower ring acted as a spacer with 2 mm thickness to allow hydrogel deformation. 250 µL of the respective cell suspension was fixed with the aid of sterile stainless steel rings in wells of a 12 well plate (non-tissue-culture-treated, Falcon). The lower ring acted as a spacer with 2 mm thickness to allow hydrogel deformation. 250 µL of the respective cell suspension was fixed with the aid of sterile stainless steel rings in wells of a 12 well plate (non-tissue-culture-treated, Falcon). The lower ring acted as a spacer with 2 mm thickness to allow hydrogel deformation.
Cell attachment and viability

Successful cell seeding and adherence was verified by using 4 mM calcein acetoxymethyl ester (calcein AM, Invitrogen) in the corresponding medium for 20 min in a humidified incubator (37°C, 5% CO₂). Fluorescence images from cells cultivated on scaffolds were immediately obtained using an inverted research microscope equipped with reflected fluorescence system (IX51, Olympus). Cell cytotoxicity was directly measured from the supernatant using a cytotoxicity detection kit measuring the lactate dehydrogenase (LDH) activity according to the manufacturer’s instructions (Roche Applied Science).

Cyclical magnetically assisted mechanical stimulation

The vertical motion of the magnetic, soft hydrogel scaffold was controlled by a magnetic field (0.8 T) induced by an external electromagnet (G MH X 025, Magnet-Schultz GmbH). 12 electromagnets were arranged like a 12 well plate (one per well) and anchored on an aluminum plate (18 cm x 18 cm x 0.8 cm) with an incorporated cooling circuit. Solenoids were connected with a switching power supply (PSP 1803, Volcraft) and controlled by a computer. Cyclic deformation was obtained using LabView (Version 8.2) for regulating power on/off and interval time, respectively. Constant temperature (37°C) of the actuating device was attained using a bath and circulation thermostat (polystat cc3, Huber Kältemaschinenbau GmbH).

Figure 2. Magnetic and soft scaffold preparation and human mesenchymal stem cells. a) Carbon protected metal nanomagnets (transmission electron microscopy image) were covalently linked into a hydrogel (b, cryo-section) with high porosity (pore size ~ 10 µm) to facilitate cell attachment. c) Magnetic, soft, cell culture scaffold. The dog-bone shape minimizes the required magnetic force for soft deformation. d) Good adherence of human mesenchymal stem cells seeded on hydrogel surfaces (Calcein-AM staining).

doi: 10.1371/journal.pone.0081362.g002
Movement Based Stem Cell Differentiation

connected to water cooling system. Equilibrated temperature of the electromagnets was obtained with the aid of a programmed ramp function of the bath and circulation thermostat. When electromagnets were activated the cooling medium was tempered to 32°C (experimentally determined) to compensate the generated heat from the electromagnets. That way, the cell culture medium was always kept at a constant temperature of 37°C within a humidified incubator (data not shown, controlled with an infrared thermometer (Scantemp pro 440)). The specific time interval (2 seconds on, 25 seconds off) was adjusted to stimulate the seeded hMSC on hydrogel scaffolds for 2 x 30 min within 3 h per day (5 weeks, total cycles: 4666) when the influence of scaffold type or chondrogenic medium composition was investigated.

For the impact of mechanical stimulation at different intensities regarding to differentiation, the stimulus was performed in 30 min cycles every 1.5 h for 8 h per day during 3 weeks (daytime activity, total number of cycles = 12600, 2016 and 672 respectively) for each group (n = 3). Cell culture samples were taken after 1, 2 and 3 weeks, respectively.

**Statistical analysis**

All quantitative data are presented as average ± standard deviation. Medium composition, biochemical and mechanical stimulation parameters of chondrogenic differentiation after 5 weeks of culture compared to the control were evaluated by One-way ANOVA (SPSS, 19.0.0). Bonferroni corrections were used to account for multiple comparisons. Differences between groups of p < 0.05 were considered as statistically significant.

**Results**

**Magnetic hydrogels of controlled flexibility**

The surface of biochemically inert carbon-coated metal nanomagnets (C/Co, Figure 2a) was chemically functionalized with covalently bound vinyl groups (Supporting Information, Figure S1 in File S1) and crosslinked into a hydrogel polymer backbone[32]. (See Supporting Information, Table S1 and Figure S2 in File S1). The produced hydrogels show a water content close to the one of most mammalian tissues[32]. The here used metal nanomagnets with high saturation magnetization (162 emu g-1)[33] allows rapid and controlled deformation. Analysis (SEM, Figure 2b) of hydrogel cryosections confirmed a homogeneous distribution and porous structure (~ 10 μm), essential for good cell adherence[26]. Elasticity measurements resulted in 21 ± 6 kPa (Supporting Information, Figure S3 in File S1) which is favorable for chondrogenesis[25].

**Magnetic-force controlled stimulation and chondrogenesis**

Movement in cell cultures has a long tradition based on mechanically connected systems to distort a specific culture surface[12,34]. For very porous, soft and elastic materials, however, such design is unfavorable[25] and leads to scaffold distortions. Binding nano-sized metal magnets to the polymer strands of soft and flexible hydrogels, however, allows in principle to apply a force at each polymer strand, since an external magnetic field will pull each metal nanomagnet, and therefore all polymer strands will move in a similar way. This smooth scaffold deformation is schematically depicted in Figure 3 and Figure S4 in File S1 (Supporting Information). Cells were seeded on magnetic scaffolds (Figure 2d) in control medium or serum-free chondrogenic medium, respectively, and optionally subjected to mechanical stimulation (Supporting Information, Figure S4 in File S1). Cells were cultured for 5 weeks. We further introduced various control experiments: hMSC were cultivated on tissue culture plates (no scaffold), pure hydrogels without nanomagnets and non-stimulated magnetic hydrogels (control experiments to rule out any other influences). Chondrogenesis (i.e. successful differentiation) was quantitatively assessed by measuring sulfated glycosaminoglycan (GAG) deposition, a major component of cartilage extracellular matrix. Importantly, GAG deposition was detected only when cells were cultivated with chondrogenic medium or when cultivated in control medium under mechanical stimulation (Figure 3a). This shows that mechanical stimulation is sufficient to induce and promote chondrogenesis in hMSC. In agreement with the reported literature[35,36], control medium (no movement, no differentiation factors) allows bone marrow derived hMSC to proliferate, but does not induce any differentiation. The absence of GAG formation in control experiments further confirmed that all materials, protocols and treatments used in this work indeed did not significantly influence the hMSC differentiation behavior. The only difference between differentiating and non-differentiating hMSC seeded on the scaffolds used here, was the application of the external magnetic field and the resulting reversible, soft deformation of the scaffold.

The combination of exogenous chondrogenic factors and mechanical stimulation even amplified the chondrogenesis and resulted in a synergistic effect as evidenced by a significantly higher GAG deposition than non-stimulated scaffolds using chondrogenic medium (p < 0.001; Figure 3a). In contrast, no GAG deposition could be observed for cells cultured in 2D on standard tissue culture plates. Terminal DNA quantification assays confirmed normal hMSC proliferation on tissue culture plates or on control samples in the absence of mechanical stimulation (Figure 3b). A high DNA value obtained for all cell cultivation methods indicated good cell expansion and decreasing LDH activity demonstrated good cytocompatibility (Supporting Information, Figures S5 and S6 in File S1) of the scaffolds and no negative impact by soft, magnetic hydrogel deformation. Nanomagnet-free hydrogels showed comparable cell growth and differentiation behavior if compared to magnetic hydrogels without external magnetic fields (i.e. no mechanical stimulation). This confirms the absence of cytotoxicity, in agreement with earlier studies[37,38].

**Chondrogenesis – The influence of mechanical stimulation combined with different chondrogenic factor concentrations**

Chondrogenesis inducing factors (TGF-β1 and dexamethasone[35,39]) and mechanical stimulation together
were investigated at various dilutions and showed a strongly synergistic effect (Figure 3; p < 0.001). The GAG deposition levels indicated a strong influence of mechanical stimulation directing hMSC into chondrogenic lineage even in the absence of any chondrogenesis inducing factors. hMSC cultured at a 50% decreased chondrogenic factors concentration in combination with soft mechanical stimulation displayed a significantly higher degree of chondrogenic differentiation (p < 0.001, Figure 3) compared to standard chondrogenic differentiation medium (classical, non-stimulated cell culture media).

Figure 3. Mechanical stimulation induced chondrogenesis. a) Cell numbers (DNA amount per scaffold) confirmed good cell expansion and growth. Below is the glycosaminoglycan (GAG) deposition per scaffold over a period of 5 weeks. Control medium (white bars) and chondrogenic medium (grey bars) were applied on cells seeded into either tissue culture plate (no scaffold), hydrogel scaffold (no nanomagnets, i.e. no movement is possible) or magnetic hydrogel. Mechanical stimulation (arrow) triggered higher GAG deposition. b) Comparable DNA amount indicated good cell growth for cells seeded into magnetic hydrogels with both medium types and no negative effects from mechanical stimulation. GAG deposition using diluted chondrogenic (grey) versus control medium (white bars). Mechanically stimulated hMSC in control medium showed comparable GAG deposition as in standard chondrogenic medium under magnetic actuation (indicated by []). * p < 0.01 cells cultured with control medium under mechanical stimulation versus non stimulated and mechanically stimulated hydrogel using both cell culture media.

doi: 10.1371/journal.pone.0081362.g003
conditions). In addition, chondrogenesis was quantitatively characterized beyond GAG deposition by immunostaining of the cartilage specific proteoglycan core protein aggrecan[40], collagen type II and transcription factor SOX9[41] (Figure 4) providing an independent confirmation of our results. The effects of exposure to both low concentrations of differentiation factor and mechanical stimulation (Figure 3) are complex and require more detailed studies.

Chondrogenesis – The impact of stimulation frequencies

The impact of different mechanical stimulation intensities were investigated with cells cultured in medium with reduced chondrogenic factors (50%) and with control medium. The external magnetic field pulse was kept constant in all groups for 2 seconds. The break intervals (electromagnets = off) were set to 10, 75 and 225 seconds (Figure 5). Non-stimulated magnetic hydrogel scaffolds (no movements) served as control. After only 3 weeks of cultivation and stimulation, the amount of GAG deposition had reached at least the same level as described above with less actuation and longer cultivation time (5 weeks). hMSC cultivated with control medium and a break interval of 10 seconds showed almost the same amount of GAG deposition compared to cells cultured in chondrogenic medium (Figure 5). Mechanical stimulation increased the chondrogenic lineage considerably. Fluorescence microscopy analysis again confirmed the biochemical assay results and large, correctly differentiated cell assemblies were observed for cultures in control medium stimulated with 10 seconds break intervals (Supporting Information, Figure S7 in File S1). DNA assays (Supporting Information, Figure S8 in File S1) revealed no negative effect on proliferation when the stimulation frequency was increased and the obtained results were again in line with the controls and previous studies.

Discussion

Mesenchymal stem cells in healthy and diseased cartilage retain the potential to regenerate tissue[40,42]. While these multi-potent cells have been investigated in tissue engineering of cartilage or bone[43-45], our results suggest the possibility
Figure 5. Frequency dependent hMSC differentiation. Mechanical stimulation frequency influences the differentiation and formation of tissue-typical extracellular matrix (amount of GAG formed) in both control and chondrogenic medium. Cells were pressure-free stretched on soft scaffolds for 2 seconds (stimulation period). Non-moved scaffolds (left) served as additional controls. The amount of GAG deposition indicated differentiation on all mechanical stimulated scaffolds particularly at high frequency. This behavior shows that mechanical soft movement follows a dose-effect type response similar to a classical response of specific cells to a given biochemical factor.

doi: 10.1371/journal.pone.0081362.g005
identified as a conserved regulator for hematopoietic stem cell (HSC) formation[51].

Another potential discussion point, namely the use of fetal bovine serum (FBS) in standard cell culture medium, can be answered based on the observation that all non-moved control experiments also contained the same amount and batch of bovine serum. FBS was indeed suggested to have an influence on hMSC differentiation[52] but this was not observed during this study. The effect of low frequency electromagnetic fields was suggested to have a positive effect on the chondrogenic differentiation[53]. The controls used here, i.e. hydrogels without nanomagnets in the presence of the same electromagnetic fields did not show any differentiation. This again proves that actual soft movement is indeed the critical stimulus to differentiation even in the absence of any chemical signals or factors.

For all parts of the study, normal proliferation was observed and the absence of cytotoxicity confirmed excellent cytocompatibility of the used chemically stable and reliably bound nanomagnets. Immunohistochemistry underlined the biochemical assay’s results. Moreover, cells cultured in standard control medium combined with stimulation showed more of chondrogenic characteristics compared to cells cultivated with chondrogenic medium only. Chondrogenesis was clearly accelerated by mechanical stimulation of hMSC cultured in chondrogenic medium. The here observed effects motivate for further studies: The purely mechanically induced stimulation might be the result of local production of growth and differentiation factors and subsequent self-influencing stimulation. Such mechanism would be in line with many growth factors’ capability to stably bind to the basolateral matrix of the cells. Alternatively, the soft, repetitive mechanical deformation might directly have promoted expression of pro-chondrogenic factors.

The here presented elastic, soft, mechanical stimulation cell culturing device uses the highly magnetic, active part of hydrogel scaffolds to convey force through external application of electromagnetic fields. This simple setup allows the same easy handling (seeding, cultivation, medium exchange) as in standard cell culture plates and opens the way for more in depth studies on the relationship between physical deformation, force and local development of tissue function, particularly for the musculoskeletal apparatus, but also in understanding organ barriers (basal membranes), the role of peristaltic movement in maintaining homeostasis in the intestine, and cardiac regeneration. Simple bioreactors may be constructed for larger, 3D magnetic scaffolds and driven by external magnetic forces to mechanically stimulate functionally correct development of larger tissues. With respect to cartilage regeneration, soft and highly magnetic scaffolds are suggested to combine the role of cell culture supports and actuators in tissue engineering of 3D artificial cartilage to treat osteoarthritis patients with their own bone marrow derived stem cells.

Supporting Information

File S1. Additional Materials and Methods, Table S1, Figures S1, S2, S3, S4, S5, S6, S7 and S8.

(PDF)

Acknowledgements

We are grateful to Urs Krebs and Max Wohlwend for magnetic stimulation device construction and Dr. Kirill Feldman for performing elasticity measurements. We would like to thank Dr. Tobias Schwarz for the fluorescence microscopy introduction, Joachim Hehl for his support in histology and Dr. Roger Wepf for SEM analysis at the LMC / EMEZ ETH Zurich.

Author Contributions

Conceived and designed the experiments: RF SH RNG WJS. Performed the experiments: RF NH JRV. Analyzed the data: RF IKH SH RNG WJS. Contributed reagents/materials/analysis tools: RF JRV IKH. Wrote the manuscript: RF SH NH WJS.

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