Neurogenesis-on-Chip: Electric field modulated transdifferentiation of human mesenchymal stem cell and mouse muscle precursor cell coculture

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A number of bioengineering strategies, using biophysical stimulation, are being explored to guide the human mesenchymal stem cells (hMScs) into different lineages. In this context, we have limited understanding on the transdifferentiation of matured cells to another functional-cell type, when grown with stem cells, in a constrained cellular microenvironment under biophysical stimulation. While addressing such aspects, the present work reports the influence of the electric field (EF) stimulation on the phenotypic and functionality modulation of the coculture of murine myoblasts (C2C12) with hMScs [hMSc:C2C12=1:10] in a custom designed poly-methylmethacrylate (PMMA) based microfluidic device with in-built metal electrodes. The quantitative and qualitative analysis of the immunofluorescence study confirms that the cocultured cells in the conditioned medium with astrocytic feed, exhibit differentiation towards neural-committed cells under biophysical stimulation in the range of the endogenous physiological electric field strength (8 ± 0.06 mV/mm). The control experiments using similar culture protocols revealed that while C2C12 monoculture exhibited myotube-like fused structures, the hMScs exhibited the neurosphere-like clusters with SOX2, nestin, βIII-tubulin expression. The electrophysiological study indicates the significant role of intercellular calcium signalling among the differentiated cells towards transdifferentiation. Furthermore, the depolarization induced calcium influx strongly supports neural-like behaviour for the electric field stimulated cells in coculture. The intriguing results are explained in terms of the paracrine signalling among the transdifferentiated cells in the electric field stimulated cellular microenvironment. In summary, the present study establishes the potential for neurogenesis on-chip for the coculture of hMSc and C2C12 cells under tailored electric field stimulation, invitro.

1. Introduction

In the expanse of regenerative medicine, the adult mesenchymal stem cell is of great importance due to its potential to transdifferentiate into both mesenchymal and non-mesenchymal cell types under the influence of biophysical cues [1]. While reprogramming the stemness, a number of studies have reported the osteogenesis and chondrogenesis, in vitro [2]. Many researchers have further investigated mesenchymal stromal cell capacity to differentiate into multiple cell types of mesodermal and non-mesodermal origin, including endothelial cells [3], cardiomyocytes [4], hepatocytes [5] and neural cells [6,7].

The targeted manipulation of stem cells is considered a promising approach in the field of therapeutic tissue regeneration. The elucidation of the governing molecular pathways has made the process of transdifferentiation possible by exploiting the plasticity of adult stromal cells to generate various cellular lineages. The conversion of hMScs into neural cells is an excellent example of transdifferentiation and reprogramming of the cellular genetics. Several studies have also reported the genetic engineering approaches as well as microenvironmental influences on transdifferentiation [8-10].

The endogenous electric field is one of the biophysical cues to regulate the cellular behaviour. Yamada et al. reported that mild electrical stimulation strongly influences embryonic stem cells to assume neuronal fate [11]. Han et al. suggested that the surface charge
influences the cells more than the wettability and roughness of the substrate [12]. As an extension of their observation, the researchers have acknowledged that the differentiation of stem cells under the influence of external cues can open up possibilities for studying novel mechanisms underlying cellular differentiation and more importantly, suggested the possibilities for adaption in clinical contexts.

The proliferative potential and the propensity to differentiate towards multiple lineages makes the myoblast cell line suitable for generative engineering [13]. The myoblasts could be differentiated into osteocytes or adipocytes when cultured with bone morphogenetic proteins (BMPs) or adipogenic inducers, respectively. Such prior studies suggest myoblasts are a multipotent cell type with the capability to modulate its genetic decision to a non-muscular lineage [14–16]. Further, the induction of BMP into muscular tissues induces ectopic bone formation at the site of implantation, in vivo [16].

The present study is based on the hypothesis that both the stromal origin cells of C2C12 and hMSC exhibit some degree of plasticity to support the genetic reprogramming [17,18]. The hypothetical pathway, leading to neural-like cells from the committed adult cells (C2C12), can be considered as dedifferentiation to achieve stemness and transdifferentiation to generate non-muscular cell type. To validate this hypothesis, we have fabricated lab-on-a-chip (LOC) with embedded metal electrodes, which facilitates the application of manipulative biophysical cues. The most important aspect of deploying microfluidic technology is to create a controlled cellular microenvironment, which can closely mimic the physiological conditions. Lindstrom et al. demonstrated that stem cell based bioengineering approaches could be achieved by the combination of the robust TiO2 coating of the microwell chip, as this stem cell based bioengineering approaches could be studied using a to create a controlled cellular microenvironment, which can closely

2. Materials and methods

2.1. Fabrication of PMMA microfluidic devices with in-built electrodes

The development of microfluidic culture system involves the design of lab-on-a-chip (LOC) layout, selection of materials, fabrication process, and sterilization technique. In the current work, the microfluidic channel layout was designed using the CorelDRAW X6 software. Poly(methylmethacrylate) (PMMA) was selected as the material for the device fabrication, as it is non-cytotoxic, biocompatible and autoclavable. The LOC devices with implanted electrodes were fabricated using three polymethyl methacrylate (PMMA, Plexiglas® USA) sheets of thickness 1 mm (Fig. 1a). The bottom-most layer of the device was used as a substrate for the cells. The middle layer of the PMMA sheet contained parallel through-cut channels of 2 mm-width, 35 mm-length and 1 mm-height, made using a CO2 laser-cutting machine (Universal LASER, USA). The channel height is kept as 1 mm, in commensurate with the thickness of the PMMA sheet. The top layer of PMMA sheet is used for sealing the channels from external environment, and it includes the inlet and outlet channels (both 1 mm in diameter), which are flush mounted with micropits to avoid leakage. The PMMA sheets are bonded together using two 3 M® pressure sensitive adhesive (PSA) films of 50 μm. It is important to mention that conducting PSA tape was used for bonding the top layer with middle layers and to enhance the electric field stimulation of the cells. To avoid misalignment of the patterns, these layers were stacked using the alignment screws.

Two parallel inbuilt electrodes (at a distance of 1 mm from the edge of the central channel) were placed on both sides of a central cell-loading channel (which is width of 2 mm and length of 35 mm). These electrodes were built with Field’s metal, (Tm ~−62 °C), which is a eutectic alloy with nominal composition 51 wt% In - 32.5 wt% Bi - 16.5 wt % Sn. Molten metal (T ~70 °C) was filled into the electrode-channels of the LOC. The metal was then cooled, resulting in channels with conducting metal, serving as electrodes on the both sides of the central channel. These electrodes were connected to a direct current source (Digilog DC Source) (Fig. 1b). The advantages of using the in-built electrodes are four fold. First, the in-built electrodes provide good reproducibility as they were permanently fixed in the LOC. Secondly, the designed biomicrofluidic device can facilitate a better quantification of the electric fields, because of identical electrode dimension. Thirdly, the entire cell population could simultaneously be stimulated with uniform electric field, unlike the conventional petridish culture. The fourth advantage is the microfluidic design, which can be implemented to have a close resemblance to physiological endogenous electric field [25].

2.2. Electric field application

In this subsection, the methodology of electric field stimulation of cells within microfluidic device is briefly discussed. While mimicking the endogenous low continuous electric field, we have designed the experimental protocol to apply an electric field of strength ~8 ± 0.06 mV/mm, continuously for 9 days at an exposure of 20 h/day, with a short intermittent resting phase of 4 h for each 24 h of the culture period. In all our experiments, the cell were initially grown for 3 day, with a short intermittent resting phase of 4 h for each 24 h of the culture period. In all our experiments, the cell were initially grown for 3 days, before they experienced electrical stimulation (Fig. 1b). The electric field strength has been measured by recreating exact simulation of the experimental conditions. In such scenario, a microfluidic device has been filled with cell culture media, while the two parallel electrodes were connected to the DC source at 10 V. In addition, two silver electrodes were placed just at the edges of the central channel, at different locations along the length (Fig. 1c and d). Two electrodes were connected to a high precision multimeter (resolution up to the 4th decimal place). The electric field strength (E) at these locations across the channel was then determined by the ratio of potential difference (V) to the distance (D) between the electrodes. In Fig. 1d, the plot of E as a
function of distance along the channel is provided. The electric field varied within the range of $8 \pm 0.06$ mV/mm.

2.3. Cell culture experiments in LOC device

The non-hematopoietic stem/progenitor cell source of human bone marrow is commonly called as human mesenchymal stem cells or hMSc. As the name suggests, hMScs are produced from the mesenchymal tissue of the developing fetus. It is isolated from Wharton's jelly and umbilical cord blood as an embryonic source as well as available from molar tooth bud and bone marrow, as an adult source. The hMScs used in this study were characterised by expression of positive and negative markers (Fig. S1). These multipotent stromal cells, can be differentiated into both mesenchymal and non-mesenchymal origin cells, including neural lineage [26] with stem cell behaviour being regulated by bio-physical cues [1,27].

The C2C12 cell line was the mouse pluripotent mesenchymal precursor, obtained from the American Type Culture Collection (ATCC). The rationale behind choosing the C2C12 cell line includes the fact, that (a) they are the multipotent cells with the ability to undergo differentiation to adipocytes, osteocytes, neural lineage in the presence of suitable cues and, that (b) they have good adherence onto the surfaces without compromising the reproducibility of the experimental outcomes.

Both of these cells (hMSc and C2C12) were cocultured in the ratio of 1:10 under the electric field stimulation in the custom designed LOC. The cellular morphology of the coculture could be observed in Fig. S2, as the tagged C2C12 cells have successfully expressed the RFP-tubulin along with untagged hMSc. C2C12 cell tagging was performed with CellLight™ Tubulin-RFP, BacMam 2.0, following manufacturer's protocol. The transdifferentiating potential of myoblasts, together with stem cells, were performed with the prior approval of the Institutional Committee for Stem Cell Research and Therapy (IC-SCRT), Indian Institute of Science, Bangalore.

Three sets of experiments were carried out to understand the cellular behaviour of the cocultured cells under the influence of electric field. They were a) coculture of hMSc and C2C12 under electric field, b) monoculture of hMSc under electric field, c) monoculture of C2C12 under electric field. For every set, the control was considered as the cells growing without the electric field. The days were counted from the day of the applying the electric field. The protocol for electric field application was optimized through several trial attempts and finalized as 20 h a day for 9 days in culture, as mentioned in Subsection 2.2. As control experiments, monoculture of C2C12 and hMSc were also exposed to electric field stimulation in parallel, keeping the culture-protocol identical with experimental group. Before the cells were exposed to electric field, they were allowed to grow for 3 days to allow them to physically adapt inside the device.

The cells were cultured using two different media; (i) proliferation media (before applying electric field for 3 days) and (ii) differentiation media (during application of electric field from day 4 for next 9 days). The control groups were not exposed to electric field, but they were also grown in similar media in parallel to the experimental groups with identical culture timeline. It is worthwhile to mention that the cell culture media plays a supportive role along with the external stimulus to facilitate the differentiation process. The researchers usually prefer to use the conditioned media on top of the external stimuli to differentiate the cells [28,29].

Proliferation media: Complete culture medium containing DMEM (Dulbecco's Modified Eagle's Medium; Invitrogen), supplemented with 20% Mesenchymal Stem Cell - Fetal bovine serum (MSC-FBS; Invitrogen), 1% antibiotic-antimycotic solution (Sigma) was used for the first three days in culture, to allow the cells to adapt in biophysical environment in LOC before applying the electric field. The cells were maintained at 37 °C in a saturated humidified atmosphere, containing 5% CO$_2$. Prior to the seeding, the cells were revived in a tissue culture graded T25 flask (Eppendorf, Germany). Upon reaching 70%-80% confluency, the cells were used for the experiment. They were
harvested using 0.05% Trypsin-EDTA (Invitrogen) and sub-cultured for further use. The cells were centrifuged at 900 rpm for 4 min and then re-suspended in the complete growth medium in order to achieve single cell suspension of cell concentration of $\sim 10^4$ cells per ml of media for further use.

**Differentiation media:** A specialized media was prepared for the differentiation experiments with the neurobasal media (Invitrogen), 200 μM ascorbate and 2 mM glutamine (Invitrogen) along with 2% B-27 supplement. 90% of this media was mixed with 10% of primary astrocytic feed (media collected from astrocytes cultured of 5 days). The astrocytic feed was included to enhance the neural microenvironment.

The single-cell suspension of the hMSCs, was used first for seeding into the microfluidic LOC devices for the coculture experiment. After a gap of 24 h, C2C12 cells were seeded inside the culturing channel. It is worth mentioning that before the cells were seeded, the channel was coated with rat tail type-I collagen for better adherence of the cells.

2.4. Cellular viability and functionality analysis

2.4.1. Cell viability using WST-1 assay

The WST-1 assay was chosen for evaluating cell viability. This specific biochemical assay was performed using water-soluble tetrazolium salts (WST-1, 4-[3-(4-iodophenyl)-2-(4-nitrophenyl)-2H-5-tetrazolio]-1-benzene-disulfonate, Roche) [30]. The WST-1 assay was performed after the prescribed culture durations of 3 days, 5 days and 9 days. To perform the colorimetric WST-1 (Roche) assay, 10% v/v solution of WST-1 salt and complete culture medium was prepared and was injected into each of the devices. The samples were incubated for 4 h at 37 °C in 5% CO₂ with 95% humidity.

As part of the microfluidic culture, the pooling of the cell samples (maintained under identical conditions) was performed as the fluid volume from a single channel was less for spectroscopic detection. We collected 200 μL of the fluid and the optical density (OD) of the developed color was measured by a microplate reader (iMark, Biorad laboratories) at a wavelength of 450 nm. The absorbance value provides a color was measured by a microplate reader (iMark, Biorad laboratories) at a wavelength of 450 nm. The absorbance value provides a direct correlation to the number of proliferating cells for each sample. The measurements are reported after averaging the data from three wells repeated thrice. Cell viability (or proliferation) was calculated using the following equation;

\[
\text{% cell viability} = \frac{\text{Mean absorbance of the sample} - \text{Absorbance of the blank}}{\text{Mean absorbance of the control} - \text{Absorbance of the blank}} \times 100
\]  

(1)

2.4.2. FDA/PI staining for live/dead cells

For assessing live and dead cells, the cells grown in LOC device were stained with fluorescein diacetate (FDA) and propidium iodide (PI), respectively. It is a double staining method for the live samples. The staining was performed with 1 mL of 25 μg/mL of FDA for 15 min, maintained at 37 °C. Subsequently, the cell samples were stained with the counter red stain with 1 mL of 10 μg/mL of PI for 5 min at room temperature. After staining, the samples were washed thrice with 1 × PBS to remove excess stain and imaged under Nikon fluorescence microscope with FITC and Texas Red filter (Nikon LV 100D, Japan).

2.4.3. Immunostaining of actin filaments and microtubules

The cell samples were washed twice with 1 × PBS and then added to 4% paraformaldehyde solution (PFA, SD Fine-Chem Ltd), before it was held for 30 min. Once the cells are fixed, the covering PMMA sheets were carefully removed and the bottom most sheet, was retained for staining. This minimizes the difficulty of pumping reagents into the channel. The samples were washed three times with 1 × PBS and 0.1% Triton X was used for permeation of the samples. The cell samples were blocked with 1% bovine serum albumin (BSA) for 1 h to prevent unwanted binding of the dye, leading to false positive results as well as to reduce background signal due to non-specific binding. Prior to microscopic observations, actin and microtubules were stained with an anti-α-tubulin antibody, labelled with phalloidin-rodamine (Invitrogen) and Alexa Fluor 488 (Invitrogen), respectively for 1 h. DAPI (Invitrogen) was used to stain the nuclear chromatin. Any excess stain was removed by washing with 1 × PBS and the cells were observed under a fluorescence microscope (Nikon LV 100D, Japan).

2.4.4. Primary staining for selected neural markers

Multi-labelling of combinatorial marker expression establishes the reliability of the results of a canonical stem cell research. In the present study, a combination of specific markers was used to avoid erroneous inferences. For example, nestin is a neural marker, which is perhaps best known to characterize the neuronal precursor cells. The immature or incompletely differentiated cells express the nestin, which is decreased later as the neural cell matures in the lineage [31]. Any study on neurogenesis widely focuses on the class III beta-tubulin protein (βIII-tubulin), as it is being selectively expressed in neuronal cells during the progression of the neurogenesis [32]. The MAP2 proteins serve to stabilize the architecture of the neurites by microtubular interaction and hence, growth by establishing crosslinking with the intermediate filaments. These neuron-specific cytoskeletal associated proteins are found to be more intensely distributed in the dendrites along with the axon, implicating its role in deciding neural protrusion and giving the functional shape during neuron development [33].

The presence of the above-mentioned marker proteins was evaluated by immunocytochemistry at the end of day 9. For this purpose, the cells were fixed with 4% paraformaldehyde (PFA) for 30 min. For immunostaining, the cells were permeabilized with 0.1% Triton-X in PBS for 5 min and 1% BSA was used for 30 min to block non-specific staining of antibodies. The primary antibody labelling with non-fluorescent antibody was carried out by incubating in a 1:200 dilution of antibody solution at room temperature for 1 h. The working solution of antibody was prepared using dilution buffer (1% BSA + 0.01% Triton-X in PBS). After subsequent washing, secondary antibody, labelled with FITC, was added at a dilution of 1:500 (anti-mouse FITC-conjugate). Before adding the secondary antibody, the washing of the cell samples with 1 × PBS is crucial in order to avoid false positive results. Complementary staining of the actin filaments and nucleus were performed using rhodamine-phalloidin conjugate (Invitrogen) and DAPI (Invitrogen). The imaging was carried out using fluorescence microscope (Nikon LV 100D) with appropriate filters of specified wavelengths with oil immersion lenses.

2.4.5. Quantitative gene expression study by qRT-PCR

Quantitative-reverse-transcription PCR (qRT-PCR) was carried out using a one-step cell-to-Ct, qRT-PCR kit procured from Invitrogen (A24571, Ambion). This specific PCR kit was chosen considering the challenge of extracting mRNA from a very low number of cells, even after sample pooling. The protocol was set according to the manufacturer’s instructions with thoughtful, but minor modification to minimize the procedural loss of mRNA. As the channels could accommodate a low number of cells, the cell lysis solution was added directly to the adhered cells, instead of enzymatic lifting. Primer solution at strength of 200 nM was added to 20 μL of reaction volume, which consists of 10 μL of the qRT-PCR mix and 0.16 μL of the RT-PCR mix. The volume was made up to the above-mentioned desired amount by adding nuclease-free water. The specifications for running the PCR reaction, include 40 cycles at 95 °C for 30 s, followed by 70 °C for 1 min, and lastly at 72 °C for 1 min. All the PCR cycles started in the hotstart mode at 95 °C for 3 min and terminated at 72 °C for 2 min.

It is worth mentioning that for each gene, three sets were prepared to validate the outcome. GAPDH was considered as the housekeeping gene, against which the gene expression is considered to be upregulated or downregulated. The gene expression was evaluated in the form of
2−ΔΔCT (CT, cycle threshold) values. ROX was the inbuilt reference dye (Sutter Instrument, Novato, CA, USA). The pipettes were filled with Germany). Prior to such measurements, patch pipettes (5–10 MΩ) were recording was carried out using the HEKA EPC10 amplifier (HEKA, 2.4.7.2. Whole-cell patch-clamp recordings

Whole cell patch clamp recordings

2.4.6. Fluorescence assisted cell sorter (FACS) analysis

To determine the cell fate processes quantitatively, FACS analysis for βIII-tubulin expression was conducted. The cells were enzymatically lifted from the channels and pooled as per groups (C2C12 monoculture with and without electric field, hMSc monoculture with and without electric field, hMSc/C2C12 coculture with and without electric field) to attain 10⁶ cell/ml. 1 ml of cell suspension was added to 500 μl of 4% paraformaldehyde for 1 h, while keeping the samples at 4 °C. The paraformaldehyde-cell mixture was pelleted by centrifugation and the pellet was added to 0.1% triton-100. Similarly, the pellet was recovered and dispersed into equal amount (50 μl) of nestin primary antibody (1:200). This was kept for 4 h. The pellet was washed and added with secondary alexafluor tagged antibody (50 μl of 1:1000) and DAPI (50 μl). After waiting for 1 h, the cell samples were washed and restored in 1 ml of PBS. The readings were assessed with flow cytometry runs for minimum of 1000 events with low speed flow. The recorded data were analyzed using BD FACS-Express.

2.4.7. Functional analysis of cells by fluorescence imaging and patch clamp technique

To assess the physiological property of cells in culture, KCl depolarization induced change in intracellular calcium by fluorescence imaging, and along with this, whole-cell patch-clamp recordings were performed. Cultured cells, adhered to Type 1 collagen coated cover-slips, were bath perfused with HEPES buffer of composition (in mM): 130 NaCl, 1.8 CaCl₂, 5 KCl, 20 HEPES, 6 glucose. The pH was adjusted to 7.4 with NaOH and osmolarity to 300–310 mOsm.

2.4.7.1. Intracellular calcium imaging

The cultured cells, plated over poly-L-lysine coated cover slip were incubated for 45 min at 37 °C with Ca²⁺ sensitive fluorescent dye, Fluo-4 AM (5 μM) with pluronic acid (0.01%) in HEPES buffer [34,35]. Cells were subsequently washed with fresh HEPES and kept in dark for 15 min for intracellular deesterification of Fluo-4 AM. The fluorescent dye loaded cells were further imaged under the fluorescent microscope (Olympus BX61, Japan). The bath volume of the chamber was 10 ml and the flow rate for perfusion of buffer was 1.5 ml per minute. During microscopic observations, the cells were excited at 495 nm and emission captured at 515 nm. The images were acquired using RETIGA A 2000DC camera. The change in [Ca²⁺], in single neurons was represented as change in Fluo-4 fluorescence intensity, as measured using by Image J software (NIH, USA).

2.4.7.2. Whole-cell patch-clamp recordings

Whole cell patch clamp recording was carried out using the HEKA EPC10 amplifier (HEKA, Germany). Prior to such measurements, patch pipettes (5–10 MΩ) were fabricated from borosilicate glass capillary (OD 1.5 mm; World Precision Instrument, WA, USA) using a P-2000 micropipette puller (Sutter Instrument, Novato, CA, USA). The pipettes were filled with internal buffer of composition (in mM): 130 K-gluconate, 4 Mg-ATP, 11 EGTA, 10 HEPES-KOH, and 1 CaCl₂. The buffer solution was characterised by pH 7.2 and Osmolarity 290–295 mOsm. Additionally, 2 mM QX314 was added to the internal solution to block sodium current. The holding potential of the cells was kept at −60 mV. During the measurements, access resistance was < 20 MΩ and the recorded currents were low-pass filtered at 2–4 kHz. The analysis was performed in Igor Pro and plotted in Origin 2016 software.

2.5. Statistical analysis

All the results were expressed as mean ± standard error (SE). Student’s t-test and one-way ANOVA with post-hoc Tukey’s test were performed to reveal the statistical differences among the different samples (electric field exposed and control, without the electric field stimulation). All the statistical analysis was performed using SPSS-16.0 software (SPSS Inc.@2010). The cut-off p-values were set at 0.05 and 0.01, lower than which were considered to be too low to accept the null hypothesis.

3. Results

Simao et al. reported the capability of perfusion based bioreactor to recapitulate the brain-like environment, which can facilitate neuronal differentiation of the iPSc-derived neural stem cells [36]. Such micro-environment plays critical role in cellular decision. Burdick et al. justified the need of the microenvironment adjustment to unlock the full pledged potential of the stem cells [37]. In order to create biophysically simulated microenvironment for the cellular differentiation, the present study was conducted using a custom-designed microfluidic device with embedded metal electrodes to apply mild electric field (8 ± 0.06 mV/mm) stimuli to hMSc/C2C12 coculture as well as monoculture. The cell viability together with gene and protein expression analysis allowed us to understand the cell fate in their proliferative/differentiative condition. The cell functionality and differentiation status of the cocultured cells are analyzed in the following sections.

3.1. Cell morphology analysis

The double lineage cocultured cells were channelized with the exogenous electric field. The influence of electric field on the cell morphology is illustrated in Fig. 2. Fig. 2a shows the cells, grown inside the channels with and without the electric field, on day 5 and day 9. The cells of different species/origin have adapted in the culture environment, without any biological discrimination or colonisation (Fig. 2b). The cells also exhibited regular morphological features, while physically communicating with each other. The cellular differentiation is regulated by the microenvironment, which modulates the genetic programming and alters the cellular fate. In order to assess any change in proliferation status quantitatively, we analyzed the cell samples at day 3, day 6 and day 9. The control sample was grown without electric field as a reference. The LOC device and chosen EF stimulation protocol supported cellular growth with a

Table 1: Primers used for qRT-PCR.

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Forward</th>
<th>Reverse</th>
</tr>
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<tbody>
<tr>
<td>nes (Nestin)</td>
<td>5’GGGAAAGAAGTTATGAACTCA-3’</td>
<td>5’AGCCTGTCTGGATGC3’</td>
</tr>
<tr>
<td>map2 (MAP2)</td>
<td>5’TGGTGGCGGAGTGGAGAAGA-3’</td>
<td>5’GGGATGGCATGGTTGATTA-3’</td>
</tr>
<tr>
<td>Tubβ3 (βIII-Tubulin)</td>
<td>5’GGGATTCAGCTTCTACACT-3’</td>
<td>5’CACATCCGGACCCAAATCCA-3’</td>
</tr>
<tr>
<td>GAPDH</td>
<td>5’GACCCGCTCTCATTGAGACCT-3’</td>
<td>5’TGATGGGATGGACCTTGGATGA-3’</td>
</tr>
</tbody>
</table>
considerable amount of viable cells. The signatures of the two inversely related processes (differentiation and proliferation) are reflected in the cell viability values (Fig. 2c). The maximum cell viability was found for the cells grown without electric field stimulation for up to day 9. The percentage of viable cells does not exhibit any statistical difference on day 3 and day 6, in both the cases. At day 9 in culture, the application of electric field decreased cell viability by ~11% than the initial culture (day 3). In other cases, we found the viability values to be statistically insignificant w.r.t. each other. The above results together denote the onset of cellular differentiation. Without the electric field, the cells have manifested the signature of slow proliferation.

3.3. Analysis of hMSC and C2C12 behaviour in monoculture in comparison to coculture

A distinct morphological comparison could be noticed among the three sets of the cultured cells (coculture, C2C12 monoculture, hMSC monoculture) (Fig. 3). The cocultured cells exhibited neurite-like projections, when stimulated with the electric field (Fig. 3a). The effect of the applied electric field on proliferation and differentiation of cultured muscle cells was studied in C2C12 monoculture (Fig. 3b). In the present study, the cells, exposed to electric field stimulation, exhibited syncytic structures (Fig. 3b). SEM images could reveal the formation of fused myotube structures on exposing them to electric field (see supplementary section, Fig. S3(a)). Interestingly, the electrical stimulation has facilitated alignment of C2C12 cells. On the other hand, the monoculture without the electric field shows normal proliferation with appreciable cellular spreading (Fig. 3b). The WST-1 assay of C2C12 monoculture on day 9 could be observed with increased proliferation, when compared with the day 3 culture (14%), for the cells, without exposure to electric field. On being stimulated with the electric field, the cells showed decreased WST-1 reading (14%), as they have undergone differentiation towards myotube-like structures (Fig. 2c). This has been confirmed with myogenin expression (Fig. S3(b)).

Upon the application of the electric field, hMSC monoculture started to grow in clustered structures (Fig. 3c, Fig. S4). To rule out the possibility that these structures were not merely formed with the debris of dead cells, we performed FDA/PI based live-dead assay (Fig. S4(b)) and WST-1 assay (Fig. 2c). The obtained staining pattern with FDA/PI showed sparsely dying core for most of the clusters together with a few of them with dying edge (Fig. S4(b)). The WST-1 assay was also not found to disregard the FDA/PI staining results, with well-established viability for the clusters.

We observed many of the clusters floating with a low degree of adhesion with LOC device substrate. Those were found to be smaller in size (≤100 μm in diameter). Some of the clusters with a large diameter (≥500 μm), were also observed and those mostly adhered firmly to the substrate (Fig. S4(a)). Consequently, the cells were seen to be grown with elongated processes. The clusters exhibited positive expression for neural stem cell marker, SOX2 (Fig. S4(c)) along with neural markers (section 3.4). The expression of these marker proteins has enabled to characterize cell colonies as the neurosphere-like clusters.
3.4. Expression of neural markers proteins: induction, commitment and maintenance of neural differentiation and architecture in the coculture

Next, we have validated the above observations using immunocytochemistry (Figs. 4–6). It is inevitable to validate the morphological change with expression of neural/neurogenic markers. We have emphasized on the expression of three protein markers, namely nestin (Fig. 4), an early neuronal marker; βIII-tubulin (Fig. 5), a marker expressed in immature neurons and MAP2 (Fig. 6), a marker expressed by all classes of immature and mature neurons. Nestin and βIII-tubulin were assessed for all the three experimental groups (coculture, C2C12 monoculture, hMSc monoculture). MAP2 has been characterised only for the cocultured cells as a confirmatory test along with PCR.

Fig. 4a is the representative image of immuno-stained cells from three experimental sets for nestin which has been performed on the day 5. Minimal immuno-reactivity of the nestin was detected in the control cells, grown without electric field. In contrast, a good amount of nestin is localized in the cytoskeleton of the electric field stimulated cells cocultured cells. Also, it is worthwhile to mention that the actin filaments were clearly observed in the control (un-or-less-differentiated

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**Fig. 3.** EF stimulation alters the cell morphological features of C2C12/hMSc coculture, C2C12 monoculture, hMSc monoculture. (a) Change of cellular morphology of the electric field stricken cells in coculture is associated with cellular elongated processes whereas unexposed cells show lack of altered cell phenotype. (b) In monoculture of C2C12, electric field stimulated cells show formation of syncytic structures. (c) hMSc monoculture, cells on application of electric field, forms clusters with growing outwardly proliferating edges. The hMSc cells growing without electric field are devoid of such cluster formation.
(intermediate filament protein) in case of the electric field-stricken cells. The neurosphere-like clusters, formed by monoculture of hMSc on exposure to electric field, has also expressed noticeable amount of nestin; whereas the electric field exposed C2C12 monoculture has failed to express the marker.

Fig. 5 depicts the expression level of βIII-tubulin on day 9, which is commonly found in the neural committed cells. Concomitantly, the cells stimulated with the electric field have a larger proportion of the cells showing βIII-tubulin positivity than the control. Furthermore, it could be noticed that βIII-tubulin proteins were localized in regions of the cells, mainly undergoing cytoplasmic extensions. This corroborates with the findings of Thrivikraman et al. who reported more positive βIII-tubulin expression for such neurite outgrowth [38]. On the other hand, the hMSc monoculture clusters, formed due to application of the electric field, has been stained positive for βIII-tubulin marker. Unlike the hMSc monoculture and coculture cells, the C2C12 monoculture has not expressed measurable βIII-tubulin, even after electric field...
MAP2 localization in the matured neurons mainly occurred in the neurites. The cell samples were collected on day 7 (image not included) and day 9 (Fig. 6) due to the late expression of MAP2. Such observation is in line with the previous observations, where the coculture cells, exposed to electric field, were able to express the markers in the neurite-like protruded regions (white arrows in Fig. 6a and b). On the contrary, the electric field unexposed cells have failed to express MAP2 protein.

The population of cells expressing these three vital neural specific markers (nestin/βIII-tubulin/MAP2) were estimated roughly by counting the positively expressed cells in randomly chosen 10 fields. Such analysis reveals that highest expressing marker is the nestin, which is followed by the βIII-tubulin, scoring around 33% lesser than the nestin. The MAP2 has a low expression in the cell population, only resulting about 25% positive cells.

Fig. 5. Differential marker expression (βIII-tubulin, actin, DAPI) on day 5, reveals signatures of onset of differentiation. (a) hMSc/C2C12 cocultured cells exhibited neurite-like projections with appreciable expression of βIII-tubulin, after the cells were exposed to electric field (absent in unexposed coculture cells). (b) Neural marker was absent in both electric field exposed and unexposed C2C12 monoculture. (c) Considerable amount of βIII-tubulin has been expressed for hMSc clusters, while the unexposed cells have failed to exhibit the same.
3.5. Gene expression analysis of neuron-specific markers (quantitative PCR for mRNA detection)

Quantitative PCR is widely regarded as the confirmatory test for the above discussed phenotypic characterization by immunostaining. Accordingly, mRNA expression of some of the specific genes were analyzed for day 3 and day 9 to assess the neural commitment (Fig. 6c). The bar graphs in Fig. 6c show fold change of the gene expression of neural-specific mRNA in terms of $2^{-\Delta \Delta CT}$ value with respect to the electric field unexposed cells of the same day (refer to table: S1, S2, S3 of annexure). An initial level of neural differentiation was confirmed by expression of nestin gene ($nes$). The fold change in $nes$ mRNA on day 9 with electric field, compared to that of the control of the same day, scored ~3.5 fold increase, which could be considered as a signature of neural induction. For the observation on day 3, the expression of $nes$ mRNA resulted quite low, which is only ~0.4 fold with respect to the unexposed cells.

The neural tubulin-III gene ($tubb3$) encodes for a class III member of the beta tubulin protein family. The heterodimer of the microtubule is the assemblage of beta-tubulins and alpha-tubulin. In case of neural cells, class-III member of the beta-tubulin serves as the subunit of the microtubules. This protein is primarily expressed in neurons and may be involved in neurogenesis. The mRNA expression corresponding to beta-tubulin of the coculture on day 3 with electric field stimulation has been observed with the noticeable increase. On day 3, the electric field stricken cells were able to express ~0.45 fold higher gene expression than the unexposed cells. This has been increased to as high as ~3.4 fold change on day 9, compared to that without electric field.

The third gene targeted was the $map2$, whose expression pattern commensurates with the trend of other neural genes. Similarly, the electric field exposed cells were able to elicit more expressed mRNA in comparison to the control or without electric field for both the observations. Importantly, the $map2$ expression was quite lower than other genes. This could be because of the fact that the appearance of this protein occurs once the neural cell starts to mature. In the first observation on day 3, the fold change in $map2$ mRNA manifested statistically insignificant expression for the electric field exposed cells, which is only ~0.4 fold higher with respect to the unexposed cells. The expression succeeded further with a drastic increase in mRNA amount on day 9 with ~2 fold change with respect to the unexposed cells.

3.6. FACS analysis for quantitative determination of cell differentiation

The monoculture of C2C12, upon exposure to the electric field, exhibited betaIII-tubulin expression very less, which has scored only ~27%. Also, the hMSc/C2C12 coculture cells exhibited ~46% betaIII-tubulin expression, under the electric field stimulation protocol (Fig. S5). Nevertheless, the hMSc monoculture also has achieved slightly lesser percentage of betaIII-tubulin, expressing cells of about ~37%. On applying electric field along with coculturing, the C2C12 cell achieved almost ~72% of increase in the neural marker expression. The multipotent stem cells also attained an efficacy of increased marker expression of ~21%, when they were cocultured with C2C12 under electric field stimulation. It is worthwhile to mention that the presently used culture protocol shows a good promise towards transdifferentiation of committed lineages.

3.7. Physiological analysis of cocultured cells

The confirmatory evidences of the neuron-like functionality of the cocultured cells were inferred by the intracellular calcium imaging and patch-clamp study.
3.7.1. Depolarization of the neural-like cells

3.7.1.1. Real time [Ca$^{2+}$]$_i$ fluorescence imaging. Localized puff stimulation of Fluo4 loaded cells with 60 mM KCl for 25 s using a glass micropipette of 1 MΩ resistance and 5 psi pressure showed an increase in intracellular calcium in electric field stimulated cells (Fig. 7a). However, it has failed to exhibit any change in fluorescence intensity in untreated control. This observation is consistent with the excitable nature of neurons due to the presence of voltage gated channels. As expected, non-neuronal cells mostly exhibited non-excitable behaviour and failed to show membrane depolarization upon high KCl application. Fig. 7b shows the real-time data plotted as intensity map, depicting depolarization induced increase in intracellular calcium in the cocultured cells (video file: Control_KCl and Experimental_KCl).

Supplementary video related to this article can be found at https://doi.org/10.1016/j.biomaterials.2019.119522

3.7.1.2. Whole cell voltage clamp recording. To further test the excitability of the electrical field stimulated cells, we measured KCl depolarization induced inward current. Previous reports have suggested that KCl depolarization is a measure of excitability [39]. Whole cell voltage clamp recordings showed strong inward current in electrical field stimulated cells with neurite-like protrusions upon puff application of 60 mM KCl. However, it has failed to exhibit any...
3.7.2. Calcium oscillation: Functionality analysis with calcium imaging

The electric field stricken cells were seen to have ruffle edges, along with a signature of neurite-like protrusions (Fig. 8a). The role of intracellular calcium oscillation has long been appreciated in regulating the cellular dynamics involved in the differentiation [40,41]. Interestingly, the calcium conduction between the intracellular to extracellular, subcellular to cellular or between two neighbouring cells are versatile and likely to depend on the type of biophysical cue.

The calcium dynamics in the hMSc and C2C12 cells were analyzed in cell samples on day 9. The profiles of calcium oscillation were recorded from the cells exposed to electric field, as shown in Fig. 8b and c (video file: Control_CalciumOscillation and Experimental_CalciumOscillation). The real-time recording shows the endogenous intracellular calcium oscillation in these cells, observed as change in pixel intensity. These changes in intracellular Ca\(^{2+}\) activity are attributed to the differentiation of hMSc/C2C12 cocultured cells to neural-like phenotype under electric field stimulation. Their minimal calcium activity has been seen to be increased due to the electric field stimulation.

4. Discussion

The present work demonstrates the impact of electric field stimulation and dynamic culture conditions on the transdifferentiation of coculture of C2C12 and hMScs. It is important to mention that hMScs are capable to form neuron-like cells through the neurogenic differentiation pathway, though it was claimed by Ma et al. that hMScs could not be differentiated to form fully functional neurons [42-44]. Hence, the term neuron-like has been used in the present context to signify the potential of immature differentiation phenomenon. The process commences with the transformation of multipotent stem cell to committed neuroprogenitor-like cells. This requires the interplay of multiple factors, which regulates the shape of the stem cells for neural differentiation. Any change in the shape of the cells can develop intracellular tension. The application of electric field generates cellular tension, mediated by the cytoskeleton organization. This results in the forceful alteration of the cellular decision, leading to the induction of mechanosensitive signal pathway towards neural differentiation [45]. The above observations establish the apparent changes in cell morphology, characterised by thin extended axon-like protrusions. In this context, we first analyse the relevance of the electric field stimulation and examine the novelty of our culture protocol in the backdrop of earlier studies involving such biophysical stimulation.

4.1. Rationale of electric field stimulation-based culture protocol

The application of electric field stimulation can modulate cell functionality to different extent or through different pathways. Depending on the cell type and culture condition, the electric field could be applied intermittently as well as continuously. For example, Jain et al. demonstrated that continuous application of the low electric field of < 25 mV/mm for 6 h to the neuroblastoma cell line (N2a cells) could elongate the neurite effectively [46]. The relationship between the specific electric field and to the stem cell response cannot be generalized as the effect is quite diverse. The impact of such stimulation would depend on the cellular microenvironment, effect of cocultured cells and mode of application. Usually, the intermittent electric field application can be programmed with relatively lower electric field intensity. Thrivikraman et al. followed the protocol of exposing the cells in the electric field of 100 mV/cm for 15 min/day in combination with intracellular and extracellular nano-manipulators in the form of Au nanoparticles to trigger neurogenic/cardiomycogenic differentiation in hMScs [38]. In another study, it has been noticed that upon exposure of the hMSc to DC electric field of 100 mV/cm for 10 min/day, the cells exhibit neurogenic differentiation on PANI substrates [47]. Another mode of application is to give a short duration pulse of the relatively higher electric field. For example, Mobini et al. used DC electrical stimulation of 10 V/cm electric field for 6 h to the adipose-derived mesenchymal stem cell cultures. This culture protocol guided them towards osteogenic differentiation. Zhao et al. reported that a 115 V/m (DC) electric field for 2h/day can induce directional migration and differentiation of neural precursor cells (NPCs) after culturing for 3 days [48]. Hamerick et al. showed that the application of electrical stimulation of physiological nature (1 V/mm) and supra-physiological nature (6 V/mm and 10 V/mm) for short duration of 6 h facilitated galvanotaxis of the murine adipose-derived stromal cells [49]. In this context, the physiological range of electric fields occurring endogenously in the animal tissues has a strength as large as 2 V/cm. This also represents the upper limit of the electrical activities found at a wound site [50]. Fröhlich et al. reported that a low endogenous electric field of 2.29 ± 0.27 mV/mm can guide the neocortical network formation [51]. In the above backdrop, the present experiments were
conducted with electric field strength of 8 ± 0.06 mV/mm.

A few attempts are also being made to explore differentiation of stem cells in conditioned media, without biophysical cues. The adult human brain harbours neural stem cell (NSC) at the anterior sub-ventricular zone (SVZ) of the forebrain and hippocampus, which shows oligopotenency towards neural lineages, like glial cells and functional neurons [52]. The stem cells migrate to the injured brain area to provide repairing functions. Comparing the hNSC and hESC, the hMScs are most widely investigated, because it has enormous potential to recreate the multiple lineages. MSCs have been transplanted in rodent models of stroke, brain, and spinal cord trauma and Parkinson’s disease [53–55]. These treated animals have survived, with functional recovery. However, there are certain controversies regarding differentiation potential of stem cells with mesenchymal origin into neurons and glia, as they are ectoderm originated [56]. But it has been reported that the mesenchymal cells have been transformed into neurons. For example, Snyder et al. reported that on transplantsing them into mouse brain, hMScs retained their capability to differentiate into neuronal and glial cells, in vivo [57]. Thrivikraman et al. have established the ability of the hMSc to differentiate into neuron-like lineage with upgradation of nestin, βIII-tubulin, like early neuron-specific markers [58]. It is worthwhile to mention that several other studies have established the event of neural-like differentiation of hMSc.

Some biomaterials substances have proven to be a good choice for differentiation of hMScs into neurons. For example, Yim et al. demonstrated the differentiation of hMScs towards neurogenesis by regulation of the morphology using micropatterned surfaces [59]. Her et al. created a 3D-environment with varying substrate stiffness to differentiate the hMSc into the neuronal lineage [60]. Some studies are also been conducted to reveal the capacity of the hMScs to differentiate towards a specific type of neuron. For example, Singh et al. transformed the hMSc cells into dopaminergic neurons [61]. Danielyan et al. reported the cholinergic neural-like differentiation of hMScs under the influence of erythropoietin [62]. The manipulation of cell-surface interaction by silicon nanowires facilitates the differentiation of hMScs into neural-like lineage, are reported by Kim et al. [63].

In the present context, it should be emphasized that the specific use of in-built electrodes within a microfluidic device allows us to create a uniform electric field distribution in the microfluidic culture (Fig. 1c and d). This has not been achieved before in earlier studies in the context of transdifferentiation. Also, the magnitude of electric field stimulation is fairly low and such a field is capable of guiding transdifferentiation of the volume of C2C12 and hMScs.

4.2. Establishing the hypothesis for the coculture differentiation

4.2.1. Exogenous and endogenous electric field interaction crossing the capacitive cell membrane

It is important to know the magnitude of electric field experienced by cell coculture as the cells are shielded by the phospholipid membrane. This will help us to deduce the molecular pathway taken by the cells to respond to the applied stimuli. King et al. described the effective relationship between the endogenous and the exogenous field, while estimating the effective electric field reaching the cell interior [64]. The cell membrane, which has a very low conductivity, may or may not cause a drop in the electric field reaching inside the cells. Thus, it is very crucial to estimate the effective strength of electric field inside the cells, overcoming the capacitive barrier of the lipid membrane. The ratio of the electric field experienced inside the cell to that of the outside is,

\[ \frac{E_1}{E_1} = \frac{9\eta}{(2 + 5\eta + 2\eta^2) - 2(1 - 2\eta + \eta^2)(a/b)^3} \]  

(2)

where \( \eta \) and \( a/b \) can be expanded as,

\[ \eta = \frac{\sigma_m}{\sigma_1}, \]

\[ \left( \frac{a}{b} \right)^3 \approx 1 + \frac{8\delta}{b} \]  

(4)

where, \( \delta \) is the thickness of the cell membrane, \( a \) is the outer radius, \( b \) is the inner radius of the cells, \( \sigma_m \) and \( \sigma_1 \) are the conductivity of the cell membrane and the extracellular saline fluid, respectively, the extracellular field is \( E_1 \), and \( E_2 \) is the internal electric field.

In case of conducting tissue, like heart, muscle, brain, the conductivity is about \( \sigma_1 \sim 150 \text{ S/m}. Also, on the basis of the above formulation, it could be inferred that during the exposure of low-frequency electric field, the spherical cells experience a large drop in the electric field reaching inside the cell (\( E_2/E_1 = 6 \times 10^{-3} \)). In contrast, the elongated cells (length 500 μm) with cylindrical shape, experience almost the same electric field as of the extracellular matrix (\( E_2/E_1 = 0.99 \)).

The initial phase of cell differentiation starts with lack of electric field effect on the internal core of the cells. During this period, the surface receptors play a key role. Previous evidence suggests the presence of a membrane-mediated electric field response [65]. Once the cells are elongated, the core electric field experienced by the cells is close to the value of the extracellular electric field. This leads to direct effect of electric field on the intracellular region.

In the present case, initially, hMSc and C2C12 have a length of around 100–300 μm and 60–90 μm, respectively. These cells experience negligible core electric field. Once the cells elongate to more than 500 μm along the axial direction, they started experiencing the core electric field, leading to intracellular electrocoupling activities. Moreover, C2C12, which are known to be multipotent/oligopotent stromal cells, receive paracrine signalling from the hMSc. This is elaborated further in the following discussion.

4.2.2. Neural differentiation by electric field mediated changes in cellular microenvironment

It is important to reiterate that the cells communicate initially with the electric field through the membrane receptors only. For example, Bezanilla et al. proposed that the electric field is sensed by the membrane protein by translocating the charges or by the movement of dipoles within the membrane [66]. The dislocation of charges and/or dipoles in an electric field produces electric current. In the case of the protein voltage sensors, the charges remain confined inside the cell membrane and keep moving within the protein [66]. Hence, the polarised proteins can modulate the cell behaviour. There is strong evidence, to suggest that a DC electric field can polarise a number of proteins. These include VEGF/EGF, acetyl choline receptors, receptors, and integrins [67–70]. These polarised surface receptors initiate an intracellular signalling cascade, mediated through src kinase (src), small GTPases, Phosphatase and tensin homolog (PTEN) and phosphoinositol kinase pathways [70,71].

In relation to the present study involving coculture with biophysical stimulations, the external field can also have electrophoretic and electroosmotic effects on the negatively charged proteins [72,73]. A DC electric field induces electrophoretic clustering of the membrane structures. For example, Lin et al. observed that mesenchymal stem cells experience the polarisation of lipid rafts towards the cathode after forming clusters [74]. The β1-integrin molecules of the cell membrane along with some other known/unknown protein members form subsequent clusters or lipid rafts. The integrin/lipid raft complex is stabilized by Cav1, which is an integral inner membrane protein. This subsequently interacts with intracellular proteins, like RhoA, to activate signal transduction mechanism, through inactivation of p190 Rho-GTPase [74].

Lim et al. mentioned that upon application of electric field, the hMScs can be guided along several differentiation pathways. They found that mesenchymal stem cells of human alveolar bone origin can
undergo neurogenesis on reduced graphene oxide with the mediation of a pulsatile electric field [75]. They proposed that the cell-to-substratum adhesion machinery confers the regulatory effect on the neural differentiation, which is increased on graphene-coated surfaces. The neural processes, like axon and dendrons formation, are initiated as a result of regulated substrate adhesion. The focal adhesion kinase has a bidirectional role in neural commitment. Focal adhesion kinase (p125fak) has specific interaction with NCAM and the SRC-related tyrosine kinase, which promotes the growth of neural processes [76]. In contrast, FAK has also been described as the negative regulator of axon outgrowth by exerting pruning actions on the axons. This plays a sophisticated role in the axon dynamics [77]. In relation to electric field application, FAK exhibits altered expression and cellular distribution, which guides stem cell differentiation.

Another hypothesis is that electric field stimulation introduces conformational changes of the membrane proteins as the field-induced configuration gets trapped in local potential well. This may be less of thermodynamically favourable conformation than the native stable three-dimensional structure. For example, Aaron et al. suggested subsequent changes in the membrane protein conformation with downstream signal transduction, leading to genetic reprogramming [78]. In addition, the electrical stimulations are found to regulate the voltage-sensitive ion channel activities with higher activation towards the cathode [67,79]. As the cells began to differentiate due to the cell surface mediated genetic reprogramming under the influence of extracellular electric field, the differentiated cells acquired elongated morphology. Such elongated morphology enables the cells to experience the intracellular electric field, which can influence cytoskeletal reorientation. For example, Li et al. provided evidence that the galvanic stress generates superoxides, which cause cytoskeletal reorganization through the activation of AKT/ERKs pathways [80].

The above pathway can be proposed for hMSCs towards the neural differentiation under electric field stimulation. Nevertheless, the presence of the specialized media with B-27 and astrocytic conditioned medium has caused neural induction in the presence of electric field. This can be due to superficial membrane interaction through the membrane receptor. Alternatively, the selective electroporation of the membrane in the presence of electric field stimulation has allowed the signalling molecules to enter and to modulate the cell behaviour epigenetically.

4.2.3. Induced neural differentiation of C2C12 cells by the paracrine effect

Several researchers reported the phenomenon of dedifferentiation of adult somatic cells (C2C12) into induced multipotent stem cells after treating them with transforming growth factor-β(1) (TGF-β(1)). For example, Mu et al. demonstrated that a lower concentration of TGF-β(1) could promote C2C12 myoblasts to express stem cell markers as well as to repress myogenic proteins, which stimulate dedifferentiation [81]. Watanabe et al. successfully modulated the C2C12 myoblasts into active neural phenotype by activation of REST/NRFS target genes. This allows the activation of terminal neuronal differentiation genes by overriding muscular differentiation [82]. Other than invasive genetic engineering procedures, small chemo-molecules can act as inducers of neuronal differentiation for C2C12. For example, Williams et al. developed a protocol for C2C12 (cocultured with PC12) to differentiate into neurons using neurodazene [83].

Williams et al. have reported the recycling of synaptic vesicle stained with FM1-43 dye at the exposure of high K⁺ ions [84]. Zhao et al. found that an electric field strength of 150 mV/mm for 20 h has a stimulatory effect on the cell division by altering the cell division orientation. Tituskin et al. reported that the exposure of the cells for 60 min at 2 V/mm electric field caused actin remodelling [85,86]. The intermittent resting time is important as the cells use that as a recovery period.

There is strong evidence that the hMSCs were characterised as a good vehicle to deliver growth factor and micro-RNAs. For example, Suzuki et al. used the transformed hMSCs to modulate neuroprotection and neural proliferation of the rat adult muscle, in vivo. [87]. Also, hMSCs are reported to be capable of modulating other cell lines by efficient exosome secretion [88]. Aaron et al. reported the autocrine and paracrine effect on the cells through the secretary growth factors [78]. Dabrowska et al. established the biological aspect of paracrine signalling of the hMSC to other cell types. They showed that hMSCs, being multipotent cells are capable of influencing the organotypic hippocampal culture. The putative regenerative interactions among Wharton-jelly-isoalted hMSC and organotropic-hippocampal cells mediate the molecular mechanisms mainly through mesenchymal cell paracrine activity [89]. This helps in creating the neuroprotective effect towards the primary cells of hippocampal culture. The secretion profiling of the spent media revealed that the hMSCs are capable of releasing TGFβ (tumor necrosis growth factor β), IL-2 (interleukin-2), IL-10 (interleukin-10), FGF (fibroblast growth factor), and IGF (insulin-like growth factor) and VEGF (vascular endothelial growth factor) [89].

In the present work, it is hypothesized that the hMSCs have mediated the C2C12 based neurogenesis through secreted exosomes containing neurotrophic factors, endocytosed by the C2C12 cells. The C2C12 cells were influenced and redirected towards neurogenesis, probably through activation of B-cell translocation gene 2 (bcl2) [83]. Through the intracellular delivery of the exosomes, the cells receive the intact and functional miRNA and growth factors without any perturbation. Likewise, the low intensity constant electric field for a prolonged duration time mimics the physiological neurogenic condition. This commits the hMSCs into neural differentiation, which in turn have cross-talk with the C2C12 through paracrine signalling. This study also implies that C2C12, being the stromal origin multipotent cells, can be transdifferentiated into neuron-like cells. As experimental evidence, the pattern bound cell culture of hMSC and C2C12 (supplementary information) shows induction of neurogenesis in the C2C12 cells, which were adjacent to the hMSC clusters (Fig. 5).

4.3. Compatibility of heterogenic murine and human origin cells in a coculture

Heterogenic coculture is noncanonical and difficult to establish, compared to homogeneous culture. It is necessary to have cell-to-cell cross-talk and healthy growth pattern of the cells to have a synergistic output. Mesenchymal stromal cells, being the most potent cell type of regenerative cells, were chosen to exploit the extensive paracrine capability to modulate the functionality of other cell types. C2C12 cells were chosen to be modulated by the hMSCs under the influence of the electric field.

Sassoli et al. mentioned that coculture of these two cells (hMSC and C2C12) under canonical culture system can potentially improve the ability for skeletal muscle repair/regeneration [90]. The crosstalk between the bone-marrow hMSCs and C2C12 myoblast was established by analyzing the chemical communication between cells of two different origins. The secretion of vascular endothelial growth factor (VEGF) by the hMSCs was thought to be the molecules responsible for such an outcome [90]. Ryu et al. made attempts to show that the C2C12 and hMSCs, when layered together onto chitosan substrates, promote myogenesis [91]. The mouse embryonic stem cells were successfully transformed to hepatocyte-like cells by the liver nonparenchymal cell lines in the coculture [92]. Such cocultured cells can successfully exhibit the rare noncanonical differentiations.

In the above context, Fig. 2 reveals that hMSCs and C2C12 cells were well maintained in the close vicinity without imposing any detrimental mutual exclusion effects. The cells could communicate extensively and there was no trace of colonization of same cells.
4.4. Expression of neural markers by cocultured cells under the influence of electric field

Nestin is a neural progenitor-specific member of type IV intermediate filament (IF) proteins. It is known that nestin is specifically expressed in neural progenitors, but never in matured neurons and glial cells [93]. This plays a pivotal role in neural transdifferentiation in the present context. As a consequence of neural differentiation, the hMSc/C2C12 coculture has been characterised with the expression of nestin. Unlike most other IF proteins, nestin cannot form stable filaments by itself, but can form co-polymer in vitro and in vivo by association with type III IF proteins, such as vimentin [94]. Chang et al. similarly found elevated expression of nestin in DC electric field stimulated cells [95]. Also, the same research group reported the associated expression of βIII-tubulin along with GFAP and O4. In corroborartion with the reported marker expression study, it has been found that the cocultured cells have expressed βIII-tubulin as well, in the present case. The FACS analysis quantitatively establishes that a substantial percentage of cells was expressing βIII-tubulin (see the supplementary section, Fig. 54). Importantly, MAP2 is expressed as a compulsory neural marker after nestin expression. The electric field has been reported to have enhanced fibroblast growth factor receptor (FGF-R) and its ligand modulated signalling pathway [85]. As a consequence of electric field application, MAP2 may have been expressed by the action of FGF signalling pathway. Further signalling analysis by Chang et al. also revealed that FGF-2-induced nestin expression is mediated through FGFR-MAPK-ERK signalling axis and the transcriptional factor Sp1 [96]. In the present study, we found the NCAM silencing results in lack of neural morphology of the electric field exposed cells, when compared to the NCAM expressing cells (see the supplementary section, Fig. 57).

4.5. Physiological changes provide confirmatory evidence towards neurogenesis in coculture

4.5.1. KCl mediated membrane depolarisation

A higher amount of extracellular potassium destabilizes the membrane potential of excitatory cells, thereby causing opening of voltage gated sodium and calcium ion channels. In addition to the calcium current observed in our patch clamp experiments, KCl depolarization of cocultured cells also showed bolus of calcium entry in electric field stimulated cultures (Fig. 7). This is a characteristic property of the excitable cells, thereby strengthening our claim of similarity to neurons.

4.5.2. Calcium oscillation

When neural stem cells are exposed to electromagnetic stimulation, spontaneous Ca\(^{2+}\) transition occurs due to an increase in Cav\(_{1.2}\) (L type) channel activity [97]. This stimulates neurogenesis by Ca\(^{2+}\) influx via L-type Ca\(^{2+}\) channel, consisting mainly of CREB phosphorylated transcription factor-mediated pathway. In particular, in the case of stimulated bone marrow-derived hMSc, increased levels of neuronal marker gene expression are associated with CREB-mediated phosphorylation [98]. It has been speculated that electric field exposure increases CREB to DNA binding activity, particularly for classes of nucleotides called CAMP [99].

In the present study, we have monitored the calcium oscillatory activity in 30% of the cells, which were specifically in physical contact with each other (Fig. 8). This process is important for many cellular phenomena, including cell proliferation, differentiation, signalling and other adaptive functions [100,101]. Electric field stimuli caused a dominant activity of L-type calcium channel along with the activation of N-type calcium channels in neuronal gene expression. Nonetheless, the neurons can detect voltages with the help of two different classes of voltage-activated calcium channel and recruitment of certain intracellular signalling pathways. Other than the calcium channels, the formation of gap junctions as the representative of electric synapse can also mediate the movement of Ca\(^{2+}\). This is perhaps the possible reason for calcium oscillation in the physically communicating cells.

4.6. hMSc/C2C12 monoculture functionality under electric field stimulation

The differences in cellular functionality are more evident when we analyse the baseline results of monocultures. Murine C2C12 myoblasts, being multipotent, have the ability to retain their differentiation potential, when grown under biophysical stimulation. The myoblasts fuse with neighbouring cells to form multinucleated myotubes [102]. Similar findings were also reported by Park et al. and they demonstrated that with electric field stimulation, the C2C12 cells started to express the myotube maturation markers, such as myoD [25,103]. Thelen et al. reported that electrical stimulation at 3 V/cm with 2 Hz enhances contractile properties in the C2C12 myotubes [104]. They even found expression of SERC Ca\(^{2+}\) type 1 proteins, which provide strong evidence for myogenic differentiation of the C2C12 cells, when exposed to electrical stimuli. Tanaka et al. found the C2C12 cells align on application of electric field [103]. In summary, the electrical cues cause the fusion and differentiation of the C2C12 cells into multinucleated structure formation.

In monoculture, hMSCs, which are known to have altered adhesion due to electric field application, form clusters (see the supplementary section, Fig. S4(a)). These neurosphere-like structures were the outcome of less adhesiveness of the hMSCs, in comparison to the electrically stimulated C2C12 which are capable of secreting collagen I (adhesion protein) [25]. Moreover, the neurosphere formation is influenced in the presence of EGF [105]. The electric field stricken hMSCs are known to secrete EGF and to create polarity [67]. The EGF is known to have positive influences in the formation of neurosphere [106]. The combinatorial effect of the above factors contributes to the formation of neurosphere-like clusters for electric field stimulated hMSC monocultures.

In particular, hMSCs are capable of exerting a trophic effect on the neighbouring cell, which can alter the cellular behaviour. For example, in coculture pellets of hMSC and chondrocytes, it has been observed that due to the extended trophism of bone marrow stem cells, the chondrocytic differentiation was aggravated, irrespective of culture condition or cellular origin [107]. This implies that the paratrophic effect of the hMSC in the form of chemical cue or contact cue in cocultures is a general phenomenon with potential for redirecting the cells into the differentiative path along with increased cell growth.

The human mesenchymal stem cells, isolated from the umbilical cord and Wharton jelly, secrete activin A into the extracellular milieu, when they are cocultured with the neural stem cells isolated from subventricular zone (SVZ). It is also reported that the differentiating stem cell in neural path secretes exosome containing miRNA (miR-125b), which can be endocytosed by other cells [108]. This further recruits the cells in the neurogenic path. For instance, the coculturing of neural stem cells (NSCs) with hMSC can drive the former along the path of neurogenesis [109]. The electric field directed exocytosis by the stem cells plays the role of neighbouring cell recruitment. It is worth mentioning that the rapid loss of stemness of in vitro-isolated NSCs possess a serious challenge for neural regeneration. During any injury of the nervous system, a limited population of the newly generated cells is recruited to the damage sites, and also takes part in neural repair [1]. Mesenchymal stem cells are promising cell sources for CNS regeneration due to their greater abundance and easier accessibility, as compared to neural stem cells and embryonic stem cells.

4.7. Relevance in the context of cellular reprogramming

There are no effective treatments for neurodegenerative diseases such as Alzheimer’s disease (AD), Parkinson’s disease (PD), Huntington’s disease and amyotrophic lateral sclerosis. Such diseases could be well
managed by introducing regenerative engineering approaches. This concept has been proposed to be more effective than the conventional drug therapy as well as invasive clinical approaches like deep brain stimulation [110,111]. The choice of the cells to perform such neural transdifferentiation for cell replacement therapy is unfortunately restricted to embryonic stem cell and induced pluripotent stem cell (iPSc). It is known that iPScs can be converted to DA neurons which restore phenotypes in animal models of Parkinson’s disease [112]. iPScs respond well to a pattern of the interacting surfaces, which could divert the cells towards neurogenesis, along with the possibility of generating iPSc-derived nucleus pulposus (NP) cells in a mouse model in hypoxic condition [113,114]. Nevertheless, apart from the successful implementation of iPSc for reprogramming of cells towards neurogenesis, the present protocol reveals a new approach for transdifferentiation of non-neuronal cell (C2C12 (pre-myoblast) cells) to neural-like phenotype, when cocultured with hMScs under electric field stimulation. This
essentially supports the hypothesis for transdifferentiation of somatic cells with stem-like properties (example C2C12) into neuron-like by an optimal microenvironment. In the backdrop of this hypothesis, it should be mentioned that hMScs, while undergoing differentiation in culture, can regulate the functionality of other cell types. In the present context, the dedifferentiating and transdifferentiating potential of C2C12 along with exploiting the stemness of hMSc, allows us to induce a sub-population of cells with characteristics of muscle stem cells into non-muscle lineage, through application of intermittent electric field (Fig. 9). This could be used for transmuting cellular functionality into generating autologous stem cells for regenerative medicine.

5. Conclusions

The following important conclusions emerge from this work.

(a) The present study demonstrates the unique ability of low-intensity electric field stimulation (8 ± 0.06 mV/mm) towards transdifferentiation of C2C12 myoblast cells to neural-like cells, when co-cultured together with hMScs, in conditioned media within custom-made PMMA microfluidic device. The novel protocol has facilitated the transdifferentiation of a non-stem cell type towards neuron-like cells under the influence of differentiating hMScs by electric field stimulation.

(b) The monocultured C2C12 cells have manifested cell fusion under the influence of similar culture conditions. On the other hand, the hMSc monocultures formed neurosphere-like clusters. Both the cell types have exhibited an altered phenotype, but failed to exhibit the level of neural-like morphology, in isolation.

(c) Statistically significant upregulation of neural markers in cocultured cells was recorded at protein and mRNA level. The calcium oscillation is clearly exhibited in the cocultured cell population. The paracrine signalling or the physical communication of the differentiated hMScs drives the C2C12 cells to undergo transdifferentiation towards neural-like morphology.

(d) The electrophysiological analysis of Ca2+ depolarization strengthens our finding of differentiation of C2C12 cells to neural-like cells upon electrical stimulation. Further, our patch clamp study demonstrates an important characteristic of neural-like cells, significant intracellular calcium-signalling, in the electric field stimulated cocultured cells.

To conclude, biomicrofluidics based devices with tailored electric field stimulation allow us to study the aspect of transdifferentiation of stromal cells, cocultured with the mesenchymal stem cells. The co-culturing of multiple cell lines can make difficult differentiation pathways realisable with the intervention of electric field stimulation.

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Appendix A. Supplementary data

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