Three-dimensional culture of single embryonic stem-derived neural/stem progenitor cells in fibrin hydrogels: neuronal network formation and matrix remodelling

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ABSTRACT

In an attempt to improve the efficacy of neural stem/progenitor cell (NSPC) based therapies, fibrin hydrogels are being explored to provide a favourable microenvironment for cell survival and differentiation following transplantation. In the present work, the ability of fibrin to support the survival, proliferation, and neuronal differentiation of NSPCs derived from embryonic stem (ES) cells under monolayer culture was explored. Single mouse ES-NSPCs were cultured within fibrin (fibrinogen concentration: 6 mg/ml) under neuronal differentiation conditions up to 14 days. The ES-NSPCs retained high cell viability and proliferated within small-sized spheroids. Neuronal differentiation was confirmed by an increase in the levels of βIII-tubulin and NF200 over time. At day 14, cell-matrix constructs mainly comprised NSPCs and neurons (46.5% βIII-tubulin+ cells). Gamma-
aminobutyric acid (GABA)ergic and dopaminergic/noradrenergic neurons were also observed, along with a network of synaptic proteins. The ES-NSPCs expressed matrissetase and secreted MMP-2/9, with MMP-2 activity increasing along time. Fibronectin, laminin, and collagen type IV deposition was also detected. Fibrin gels prepared with higher fibrinogen concentrations (8/10 mg/ml) were less permissive to neurite extension and neuronal differentiation, possibly owing to their smaller pore area and higher rigidity. Overall, it is shown that ES-NSPCs within fibrin are able to establish neuronal networks and to remodel fibrin through MMP secretion and extracellular matrix (ECM) deposition. This three-dimensional (3D) culture system was also shown to support cell viability, neuronal differentiation and ECM deposition of human ES-NSPCs. The settled 3D platform is expected to constitute a valuable tool to develop fibrin-based hydrogels for ES-NSPC delivery into the injured central nervous system.

**Keywords:** three-dimensional culture; neural stem cells; neuronal maturation; fibrin hydrogel; extracellular matrix; fibronectin; laminin; matrix metalloproteinase activity

## 1 INTRODUCTION

Nervous system disorders such as spinal cord injury (SCI), stroke, brain trauma and neurodegenerative diseases often result in life-long neurological deficits, with high burden to society and health systems. Regeneration of damaged central nervous system (CNS) tissue remains a challenging issue, as it comprises the regrowth of disrupted axons, the replacement of lost neural cells and the recovery of lost neural functions (Okano, 2010). As in the adult CNS the intrinsic capacity of endogenous neural progenitors to replace lost cells is insufficient, transplantation of exogenous neural stem/progenitor cells (NSPCs) has emerged as a possible therapy to compensate for injury-induced lost cells. In addition to secreting cytokines with neurotrophic and immunomodulatory effect, NSPCs are able to differentiate into neurons and glial cells, potentially contributing to the remyelination of spared axons and to the establishment of relay neuronal circuitries. In preclinical models of SCI and ischaemic stroke, transplanted NSPCs were shown to be capable of dividing, differentiating and integrating into the injured tissue, partly overcoming functional deficits (Lindvall and Kokaia, 2010; Okano, 2010; Lu et al., 2012). These findings led to the approval of clinical trials to assess the safety and preliminary efficacy of human fetal CNS-derived neural stem cell transplantation (https://clinicaltrials.gov).

Embryonic stem (ES) cells have been explored as an alternative source of NSPCs because of their pluripotency and unlimited capacity for self-renewal (McDonald et al., 2004, Colombo et al., 2006), particularly after the establishment of protocols to efficiently and reproducibly specify ES cells into pure populations of NSPCs (Ying et al., 2003; Incitti et al., 2014). Moreover, ES-derived NSPCs (ES-NSPCs) retain a higher proliferative capacity and neuronal differentiation efficiency compared with somatic NSPCs (Colombo et al., 2006), which can be of interest to counteract the high levels of cell death occurring following transplantation. Clinical trials involving ES-NSPC transplantation have also been approved (Geron Corporation, CA 94025, USA; Cell Cure Neurosciences Ltd, Jerusalem, Israel; Ocata Therapeutics Inc., MA 01752, USA; Chabiotech Co Ltd, Seoul, South Korea), as ES-NSPCs were shown to survive, divide, differentiate and promote functional recovery in rodent models of SCI (McDonald et al., 2004; Erceg et al., 2010).

Despite their functional benefits, NSPCs implanted as suspensions usually show poor cell survival, as well as limited differentiation along the neuronal lineage, as a result of the inhibitory milieu of the injured tissue (McDonald et al., 2004). Attempts are being made to improve NSPC transplantation
efficacy by combining NSPCs with biodegradable hydrogels (Pego et al., 2012). In addition to providing a structural support for axonal extension and allowing a homogenous distribution of transplanted cells, hydrogels contribute to sequester cell-secreted neurite-promoting factors and extracellular matrix (ECM) proteins, while supporting the exchange of nutrients within the surrounding tissue. Hydrogels may therefore contribute to the establishment of a more favourable microenvironment for NSPC survival and differentiation, and for regeneration (Aurand et al., 2012; Pego et al., 2012). Fibrin (Fb) is a Food and Drug Administration (FDA) approved hydrogel with widespread clinical use that emerged as a promising biomatrix for application in the CNS, either as an acellular graft (Petter-Puchner et al., 2007) or as a vehicle for cell and drug delivery (Amaral et al., 2011; Lu et al., 2012). Fibrin is a natural polymer formed during the blood coagulation cascade from its zymogen form, fibrinogen, with a central role in haemostasis. Owing to its mechanical properties, inherent biocompatibility and susceptibility to proteolytic degradation, Fb constitutes a suitable matrix for cells normally residing in soft tissues such as brain or spinal cord (Janmey et al., 2009) and, in this sense, has been successfully explored for the transplantation of NSPCs (Lu et al., 2012).

Commitment of ES cells into the neural fate is usually achieved by inducing embryoid body (EB) formation in suspension culture followed by treatment with retinoic acid (Ying and Smith, 2003). It has been shown that EBs containing neural progenitors proliferate and differentiate following embedment in fibrin hydrogels (Willerth et al., 2006). However, neural induction protocols relying on EB formation typically result in a heterogeneous cell population because of the establishment of concentration gradients of morphogens present from the outer edge of the multicellular aggregates to the core (Dhara and Stice, 2008; Wilson et al., 2014). This issue can be overcome using adherent monoculture systems to generate NSPCs from ES cells, which result in a more homogeneous population (>85% of cells expressing SOX1), because of uniform exposure of cells to soluble factors (Ying and Smith, 2003; Dhara and Stice, 2008; Chambers et al., 2009).

In this sense, the aim of this work was to establish the conditions for the three-dimensional (3D) culture of NSPCs derived from ES cells under adherent monoculture in fibrin hydrogels, when seeded as single cell suspension. For this purpose, mouse ES-NSPCs were embedded within Fb hydrogels as single cells, and the seeding density leading to the highest cell viability, after 14 days of culture under differentiation conditions, was established. NSPC-Fb constructs were then characterized in terms of cell morphology, proliferation, morphology, phenotype, distribution of cell-secreted ECM proteins, as well as in terms of matrix metalloproteinase (MMP) activity and matriptase expression. Moreover, the effect of fibrinogen concentration on NSPC viability, migration, proliferation, neuronal differentiation and MMP activity was assessed, as the density of bioactive domains and the structural and viscoelastic properties of hydrogels may have an impact on the behaviour of encapsulated cells (Aurand et al., 2012). Finally, to reveal whether or not this 3D culture system could be applied to human ES-derived NSPCs (hES-NSPCs), dissociated hES-NSPCs were cultured within Fb hydrogels for periods up to 21 days and cell viability, neuronal differentiation and ECM remodelling subsequently assessed.

2 Materials and methods

2.1 Generation of NSPCs from mouse ES cells

The mouse ES cell line (46C) established at the Institute for Stem Cell Research (Edinburgh University, Edinburgh, Scotland, UK) expressing green fluorescent protein (GFP) under the promoter
of the neural-specific Sox1 gene, was used (Aubert et al., 2003). The ES 46 cells were grown in KnockOut™ ESC/iPSC medium (Gibco; Thermo Fisher Scientific, MA 02451, USA) supplemented with 1% (v/v) penicillin/streptomycin (Pen/Strep), 2 μM GSK-3 inhibitor IX (Calbiochem; Merck, Darmstadt, Germany), and 2 ng/ml recombinant murine leukemia inhibitory factor (LIF, produced in bacteria; a gift from Dr Cláudia Lobato, IST, Lisbon, Portugal), in six-well tissue culture-grade plates coated with 0.1% (w/v) gelatin (Sigma-Aldrich, MO 63103, USA). Cell culture medium was refreshed every day. To start the monolayer culture protocol, ES cells were cultured at high density (1 × 10^5 cells/cm^2) for 24 h with LIF (Ying and Smith, 2003). To induce the neural phenotype, ES cells were cultured in monolayer in N2B27 medium, a 1:1 mixture of DMEM/F12 medium supplemented with modified N2 and Neurobasal medium supplemented with B27 and L-glutamine (Neurobasal/B27), containing 1% (v/v) penicillin/streptomycin (all from Gibco). Cell culture medium was refreshed every other day. At day 5 of the monolayer culture differentiation protocol, SOX1-GFP expression was analysed by fluorescence microscopy and flow cytometry, to assess the efficiency of neural conversion (see the Supplementary material online, Figure S1A,B). For the latter, cells were dissociated using StemPro® Accutase® (Gibco), counted, suspended in fluorescence-activated cell sorting (FACS) buffer [2% (v/v) fetal bovine serum (FBS) in phosphate-buffered saline (PBS)], and run on a flow cytometer (FACSCalibur™; BD Biosciences, CA 95131, USA). Live cells were gated based on forward scatter (size) and side scatter (cell complexity) criteria, and fluorescence gates set using undifferentiated 46C ES cells as a negative control. All analysis was performed using (FlowJo software, LLC; OR 97520, USA). The levels of SOX1-GFP expression of the ES-NSPCs used throughout this study are shown in the Supplementary material online (Figure S1B), together with a representative fluorescence histogram. The phenotypic characterization of ES-NSPCs at day 5 of monolayer culture neural commitment is provided in the Supplementary material online (Figure S1).

2.2 Culture of mES-NSPCs within Fb gels

Fibrinogen solution was prepared dissolving plasminogen-free fibrinogen from pooled human plasma containing factor XIII (Sigma) in ultrapure water, followed by dialysis against Tris-buffered saline (TBS, pH 7.4) for 24 h (Schense and Hubbell, 1999). The resulting fibrinogen solution was then sterile-filtered and its concentration determined by UV spectroscopy. Fibrinogen solution was then diluted to 12 mg/ml with sterile TBS. Fibrin gels (total volume 50 μl) were obtained by mixing equal volumes of the fibrinogen solution and a thrombin solution in TBS containing CaCl_2 and aprotinin. Fibrin gels were allowed to polymerize for 30 min at 37°C in a 5% CO_2 humidified incubator before the addition of cell culture media (3 ml/well). The cell–matrix constructs were cultured for periods up to 14 days in the presence of 5 μg/ml of aprotinin to delay Fb gel degradation, following a protocol for neuronal differentiation (Conti et al., 2005). Cells were initially cultured in N2B27 supplemented with 10 ng/ml basic fibroblast growth factor (bFGF; PeproTech) and, at day 2 of culture, the medium was switched to N2B27–Neurobasal/B27 (1:3) mix without growth factor. At day 8, half of the medium was replaced by the N2B27–Neurobasal/B27 (1:3) mix supplemented with 20 ng/ml brain-derived neurotrophic factor (BDNF) and 50 ng/ml nerve growth factor (NGF) (both from PeproTech EC Ltd.; London, United Kingdom). Half of the medium was changed every other day. To assess the effect of fibrinogen concentration on ES-NSPC behaviour within Fb, Fb gels with...
4, 6, 8 and 10 mg/ml fibrinogen concentrations were prepared. The ES-NSPCs were embedded within Fb hydrogels and cultured as described above.

2.3 Culture of human ES-NSPCs within Fb gels

H9-derived human (h) ES-NSPCs were purchased from Life Technologies and expanded according to the manufacturer’s protocol (Gibco; Thermo Fisher Scientific, MA 02451, USA). For cell embedment in Fb, hES-NSPCs were dissociated into single cells using StemPro Accutase and further suspended in Fb gel (1 x 10^6 NSPCs/ml) as described for mES-NSPCs. The cell–matrix constructs were cultured for periods up to 21 days in the presence of 5 μg/ml of aprotinin. Human ES-NSPCs were initially cultured in StemPro NSC SFM media (Gibco) containing bFGF and EGF and, at day 2 of culture, the medium was switched to the StemPro NSC SFM media–Neurobasal/B27 (1:1) mix, without growth factors. At day 8, half of the medium was replaced by the StemPro NSC SFM–Neurobasal/B27 (1:3) mix supplemented with 10 ng/ml of BDNF and 500 μm of dibutyryl cAMP (Sigma). Half of the medium was changed every other day.

2.4 Size of cellular spheroids formed within Fb gels

The size of the cellular spheroids present within Fb was assessed at day 14 of culture. Phase contrast images of the cell–Fb constructs were acquired (Axiovert 200; Carl Zeiss), and the diameter size of each spheroid measured using AxioVision software (Carl Zeiss; Oberkochen, Germany). Spheroids smaller than 25 μm diameter were not considered, as these mainly corresponded to single or paired cells (Mori et al., 2006).

2.5 Immunocytometry

Immunocytometric analysis was performed in six-pooled Fb drops, after cell isolation from the cell–Fb constructs. In brief, the constructs were washed twice with PBS and sequentially incubated with 1.25 mg/ml of collagenase type II (Gibco; 1 h at 37°C) and 1 x trypsin–EDTA (Gibco; 30 min at 37°C) under stirring (70 rpm). After trypsin inactivation with serum-containing media cells were gently dissociated, centrifuged and suspended in cell culture medium. For staining of intracellular markers, cells were fixed with 1% (w/v) paraformaldehyde (PFA; 20 min at 4°C), and permeabilized with 0.2% (v/v) Triton X-100 (10 min at 4°C). Cells were then incubated in blocking buffer for 20 min and incubated for 30 min with primary antibodies diluted in PBS containing 1% (v/v) normal goat serum (NGS; Biosource International Inc., CA 93012, USA) and 0.3% (w/v) saponin. Primary antibodies were detected by applying isotype-specific or species-specific secondary antibodies for 30 min. For immunolabelling of cell surface markers, the fixation and permeabilization steps were omitted, and the blocking/primary antibody solutions prepared without saponin. Cells were finally washed three times and suspended in FACS buffer containing sodium azide, for flow cytometry analysis (BD FACSCalibur). Primary and secondary antibodies are listed in the Supplementary material online, Table S1. Unlabelled cells were used to set fluorescence gates. Cells stained with secondary antibody only or with the correspondent isotype control were used to assess the background fluorescence associated with non-specific binding of the secondary antibody. Representative fluorescence histograms are shown in the Supplementary material online, Figure S3B.

2.6 Immunocytochemistry
Immunocytochemistry was performed in cell–Fb constructs previously fixed in cell culture media containing 2% (v/v) PFA (30 min at 37°C). All subsequent steps were performed under stirring (50 r.p.m.). When staining for intracellular markers cell–Fb constructs were permeabilized for 45 min with 0.2% (v/v) Triton X-100. Cell–Fb constructs were blocked for 1 h, incubated (overnight at 4°C) with the appropriate primary antibody, rinsed three times with PBS, and incubated for 1 h with the appropriate secondary antibody. When necessary, nuclei were counterstained with 4′,6-diamidino-2-phenylindole (DAPI, 0.1 μg/ml; Gibco). Samples were finally mounted in Fluoromount mounting media (Sigma-Aldrich) and observed under confocal laser scanning microscopy (CLSM) (TCS SP5 II; Leica Microsystems, Germany). Primary and secondary antibodies are listed in the Supplementary material online, Table S2.

2.7 Cell outward migration analysis

Neurite extension/cell outward migration from cellular spheroids was assessed at day 4 of ES-NSPC culture within Fb. Cell–Fb constructs were fixed and permeabilized as described for immunocytochemistry studies, incubated with PBS containing 1% (w/v) BSA for 1 h under stirring, and then for 1 h under stirring with Alexa Fluor 594-conjugated phalloidin (1:40; Invitrogen; Thermo Fisher Scientific, MA 02452, USA), for visualization of filamentous actin (F-actin). After rinsing with PBS, nuclei were counterstained with DAPI and samples mounted in Fluoromount. Fluorescence images were collected under CLSM. Spheroid and F-actin positive regions were obtained using automatic segmentation of the two-dimensional (2D) maximum intensity projections of DNA and F-actin CLSM image stacks, respectively. Given the segmented regions, the number of cell extensions protruding from the edge of each spheroid (sprouts), as well as the length of the longest sprout (maximal outgrowth), were analysed (Bessa et al., 2013). For each condition, 10 randomly selected spheroids from 2 different Fb gels were examined.

2.8 Microstructure analysis of Fb gels

Fibrin gels microstructure was characterized using fluorescently-labelled fibrinogen. Fluorescent Fb gels were prepared as described above, using a 1:1.0 ratio of Alexa Fluor 488 human fibrinogen conjugate (Molecular Probes; Thermo Fisher Scientific, MA 02452, USA) to unlabelled fibrinogen. Acellular Fb gels were allowed to polymerize (30 min at 37°C), and immediately observed under CLSM, using a Plan-Apochromat ×63/1.4NA Oil objective. CLSM stacks of 10 optical sections covering a depth of 10 μm were acquired, and 2D projected. Pores were automatically segmented and their areas computed using (MATLAB software MathWorks®; MA 01760-2098, USA). Finally, the average pore area was determined. For each condition, four images from two different Fb gels were analysed.

2.9 Rheological analysis of Fb gels

A Kinexus Pro Rheometer (Malvern Instruments Ltd, Malvern, United Kingdom) was used to examine the viscoelastic properties of acellular Fb drops. Fibrin polymerizing solution (50 μl) was pipetted into a plate-plate geometry (4 mm in diameter with sandblasted surfaces) previously set at 37°C, and allowed to polymerize in situ for 1 h under a humidified atmosphere. A 10% compression was then applied to the gel to avoid slippage. For each condition, the linear viscoelastic region (LVR) was first determined performing stress sweeps (frequency 0.1 Hz, strain 1–100%). Frequency sweeps (frequency 0.01–100 Hz, strain 5%) were then performed within the LVR. The storage modulus (G'), a measure of the elastic energy stored during the deformation imposed by one oscillation of the
rheometer, was determined and used as a measure of Fb drop stiffness. The loss modulus (G″), which reflects the energy dissipated by Fb drops during deformation was also recorded. Six Fb gels were analysed per condition.

2.10 Data analysis and statistics

Each independent experiment was established from mES-NSPCs obtained from different neural commitments. In each flow cytometry analysis, 10,000 events were captured inside the gate. Sample distribution was tested for normality using the Kolmogorov-Smirnov test, and data subsequently analysed using the unpaired t-test. Unless mentioned otherwise, all data are presented as mean ± standard deviation (SD). The statistical significance level was set for p-values <0.05.

A detailed description of the methods used to culture mES-NSPCs on 2D substrates, as well as to assess cell viability, total cell number, proliferation, and the levels of MMP activity, matriptase, βIII-tubulin and NF200 is provided in the Supplementary material online (Methods S1).

3 Results

3.1 Optimization of mES-NSPC cell seeding density within Fb

To assess the optimal conditions to culture mES-NSPCs within Fb, different cell seeding densities were initially explored (1, 2 and 3 × 10⁶ cells/ml). Following entrapment within Fb, single ES-NSPCs were found well dispersed within the Fb gel, as shown by incubation of samples with calcein AM and Hoechst (see the Supplementary material online, Figure S2A). The lowest cell seeding density (1 × 10⁶ cells/ml) resulted in a lower number of dead cells and smaller-sized cellular spheroids after 14 days of culture, as shown by CLSM analysis of cell/Fb constructs previously incubated with calcein AM/propidium iodide (PI) (see the Supplementary material online, Figure S2B). These qualitative results were further supported by flow cytometry analysis of cell viability, which showed, for the lowest cell seeding density, the lowest percentage of PI-labelled cells (see the Supplementary material online, Figure S2C). As a result, this cell seeding density was selected for use in subsequent studies. The cell culture medium was supplemented with aprotinin, a cell protease inhibitor, as the mere addition of aprotinin to the Fb polymerizing solution was not sufficient to preserve cell–Fb construct integrity during the 14 days of culture (see the Supplementary material online, Figure S2D). The addition of 5 μg/ml of aprotinin to the medium was effective in preventing the overall Fb gel degradation, independently of the cell seeding density tested, while being permissive to Fb degradation by migrating cells (data not shown).

3.2 Cell viability and proliferation of mES-NSPCs within Fb gels

The distribution of live/dead cells within Fb gels was assessed after 7 days and 14 days of culture. Confocal analysis of cell–Fb constructs, previously incubated with calcein AM/PI, showed for both time-points the presence of live cells widely distributed throughout the Fb gel, often growing as cellular spheroids. Formation of cellular spheroids was associated with proliferative capacities and not with aggregation at the initial cell embedment, as only single cells were detected within Fb gels immediately after cell embedment (see the Supplementary material online, Figure S2A). Neurite-like extensions were observed at day 7, and at day 14 of culture; these formed a dense network of cellular processes (Figure 1a). Uptake of PI by cells was also observed at both time-points analysed. The
quantitative analysis of live/dead cells within Fb was assessed by flow cytometry, upon cell isolation from the Fb gels. Cell viability within Fb gels was found to be similar at both time-points and at levels greater than 85%, on average, as demonstrated by the percentage of cells excluding PI (Figure 1b). The percentage of cells remaining alive was found to be high for both time-points considered, as shown by the percentage of cells retaining ability to hydrolyse calcein AM (99.9% for day 7 and 94.3% for day 14; see the Supplementary material online, Figure S3A). The ES-NSPCs within Fb gels were able to proliferate, as demonstrated by the fivefold and twofold increase in cell number found over the first and second weeks of culture, respectively (Figure 1c). At day 14 of culture, the presence of small-sized cellular spheroids inside cell–Fb constructs, mostly frequently with diameters in the range of 50–75 μm (Figure 1d), also suggests cell proliferation of ES-NSPCs within Fb.

### 3.3 Phenotypic analysis of mES-NSPCs cultured within Fb gels

Differentiation into the neuronal lineage over the cell culture period was initially followed by western blot of βIII-tubulin (developing and mature neurons) and NF200 (mature neurons). As expected, expression of βIII-tubulin and NF200 was not detected before ES-NSPC embedment in Fb (data not shown). Throughout the culture period, the expression of these neuronal markers increased, attaining significantly higher levels at day 14 when compared with day 4 for both markers (Figure 2a,b). For this reason, a more detailed phenotypic analysis was performed at day 14 of cell culture by flow cytometry and immunocytochemistry (Figure 2c). At this time-point, a significant percentage of cells still expressed the NSPC marker nestin (53.1 ± 6.3%) and the neural stem cell (NSC) marker Sox1–GFP (37.7 ± 5.6%). The percentages found were always to be inferior to those of ES-NSPCs before embedment within Fb (87.7 ± 5.8% Sox1–GFP+ cells and 86.5 ± 7.2 nestin+ cells; see the Supplementary material online, Figure S1C). Under these conditions, ES-NSPCs differentiated mainly into βIII-tubulin immunoreactive neurons, as shown by the higher percentage of βIII-tubulin+ cells found (46.5 ± 12.5%) compared with those of cells staining positive for glial fibrillary acidic protein (GFAP) or O4, astrocytic and oligodendrocytic markers, respectively. After 14 days of culture, the percentage of cells expressing stage-specific embryonic antigen 1 (SSEA-1), a marker for undifferentiated mouse ES cells, dropped from 57.3 ± 17.6% (see the Supplementary material online, Figure S1C) to 20.6 ± 4.7%. CLSM analysis of samples processed for immunofluorescence at this same time-point (day 14) supported the previous results. Images depict cell–matrix constructs composed of a heterogeneous cell population encompassing NSPCs (nestin+ cells; Figure 2e) and developing and mature neurons (βIII-tubulin+ and NF200+ cells; Figure 2f,i). Neurites staining for βIII-tubulin were found mainly radially extending from cellular spheroids, forming a dense neuronal network throughout the whole Fb gel (Figure 2h). Astrocytes and oligodendrocytes were also detected but in lower amounts, and mostly located near the spheroids (Figure 2f,g). A reminiscent population marking for SSEA-1 could be also observed at this time (Figure 2d). The presence of synaptic vesicles was assessed through synapsin immunodetection. Cell–Fb constructs revealed an extensive network of synapsin (Figure 2j), and the presence of dendritic spines along the axonal processes (arrowheads in Figure 2j). Neuronal phenotype was further investigated. A ubiquitous expression of GAD67, a GABAergic phenotype marker, was observed throughout the cell–Fb constructs (Figure 2k). Furthermore, immunodetection of tyrosine hydroxylase (TH) revealed the presence of mature dopaminergic and/or noradrenergic neurons (Figure 2l); The TH-labelled neurons with long extensions are indicated by arrowheads.

### 3.4 Distribution of cell-secreted ECM proteins, MMP activity and matriptase expression in 3D cultures of mES-NSPCs within Fb gels
The distribution of cell-secreted ECM proteins was observed after 14 days of culture through immunodetection of fibronectin, laminin and collagen IV. As fibronectin is a common contaminant in commercially available fibrinogen, acellular Fb gels were also assessed for the presence of fibronectin, but no positive staining was detected in Fb gels (data not shown). Cell–Fb constructs revealed the presence of cell-secreted fibronectin fibrils following a similar deposition pattern to that described for neurites, and often forming bundles near the cellular spheroids (Figure 3a). This deposition pattern is in accord with the role of fibronectin as a mediator of neuronal migration and outgrowth during development (Sheppard et al., 1993). A network of laminin was also observed, mainly inside and near cellular aggregates (Figure 3b). Collagen type IV was also detected at this time-point, whether in the form of a tight network inside cell spheroids or as fibrils deposited within the Fb gel, as depicted by arrowheads in Figure 3c. In 2D cultures of ES-NSPCs on poly-d-lysine/laminin-coated glass coverslips carried out in parallel, the deposition of these three ECM proteins was also observed (Figure 3d–f). The activity of MMPs was studied by performing gelatin zymography in conditioned media of ES-NSPC cultures within Fb gel and by loading 20 μg of total protein from conditioned media. No MMP activity was detected in Fb gels cultured without cells (data not shown). Zymograms showed that MMP-2 and MMP-9 were secreted by ES-NSPCs within Fb, the latter being detected at a later period of culture (Figure 3g). The MMP-2 activity levels increased along the culture period, attaining levels significantly higher than those of day 4 of culture at day 14 (Figure 3g). The MMP-9 activity was only detected from day 8 of culture onwards, and no significant differences were found among the different time-points analysed (Figure 3f). Matriptase expression of ES-NSPCs cultured within Fb was subsequently investigated, as ES-46C cells were reported to express this protease when cultured on a collagen-coated surface under neuronal differentiation conditions (Fang et al., 2011). Western blot was performed for three independent experiments and matriptase levels normalized to GAPDH expression. As expected, matriptase expression was detected in ES-NSPCs before embedment in Fb (matriptase levels 1.32, data not shown). Significant differences in matriptase expression were only found at day 14 of cell culture, for which 1.3-fold higher levels were found compared with day 8 (Figure 3h).

### 3.5 Effect of fibrinogen concentration on mES-NSPC behaviour within Fb gels

To assess the effect of fibrinogen concentration on NSPC behaviour within Fb, Fb gels were prepared with 4, 6, 8 or 10 mg/ml of fibrinogen and their structural and viscoelastic properties characterized. Fibrin gels prepared with 4 mg/ml or 6 mg/ml of fibrinogen revealed a similar microstructure, while those prepared with higher concentrations of fibrinogen (8 mg/ml and 10 mg/ml) showed a denser network of Fb fibre bundles and smaller pore size (Figure 4a). Image analysis of 2D projected CLSM stack images further confirmed the qualitative results obtained, revealing a decrease in the average pore area with increasing fibrinogen concentration (0.71 ± 0.02, 0.65 ± 0.05, 0.53 ± 0.01 and 0.49 ± 0.02 μm² for fibrinogen concentrations of 4, 6, 8 and 10 mg/ml, respectively), which was statistically significant when comparing Fb gels prepared with 8 mg/ml or 10 mg/ml of fibrinogen with those prepared with 4 mg/ml or 6 mg/ml of fibrinogen (Figure 4b). Rheological analysis of Fb gels consistently showed a significant increase of the storage (G’) and loss modulus (G’’), with increasing fibrinogen concentrations, with G’ always exceeding G’’ (Figure 4c).

The ES-NSPCs were then embedded within Fb gels and cultured for periods up 14 days. The impact of fibrinogen concentration on neurite extension/cell outward migration was assessed at day 4 of culture. Fibrin gels prepared with 8 mg/ml or 10 mg/ml of fibrinogen were found to be less permissive.
to neurite extension/cell outward migration compared with gels prepared with 4 mg/ml and 6 mg/ml of fibrinogen, as shown by the significant decrease in the number of protruding cellular processes (sprouts) as well as in the maximal outgrowth from cellular spheroids found after image analysis (Figure 5a). The effect of fibrinogen concentration on neuronal differentiation was investigated at day 14 of culture. Compared with Fb gels with 4 mg/ml or 6 mg/ml of fibrinogen, Fb gels prepared with 8 mg/ml or 10 mg/ml showed a less dense network of cellular processes labelled for βIII-tubulin (as shown by CLSM; Figure 5b). After cell isolation from the constructs, flow cytometry analysis also revealed a lower percentage of βIII-tubulin immunoreactive cells within Fb gels prepared with 10 mg/ml of fibrinogen, comparing to those found in Fb gels prepared with 4 or 6 mg/ml of fibrinogen. (Figure 5c). The analysis of the mean fluorescence intensity ratio of the histograms (mean fluorescence intensity of cells stained for βIII-tubulin normalized to that of cells incubated only with secondary antibody), also suggests lower expression of βIII-tubulin at the single level in Fb gels prepared with 8 mg/ml and 10 mg/ml of fibrinogen (Figure 5d), further supporting immunocytochemistry data. Despite hampering neuronal differentiation of ES-NSPCs within Fb, fibrinogen concentration did not affect cell viability, regardless of the fibrinogen concentration considered, as shown by a Trypan Blue dye exclusion assay, performed at day 14 of culture (Figure 5e). When analysing the size of cellular spheroids present in cell–Fb constructs at this same time-point, large-diameter spheroids (diameter ≥ 125 μm) were found within Fb gels prepared with 8 mg/ml or 10 mg/ml of fibrinogen (see the Supplementary material online, Figure S4A). These findings suggest that Fb gels with fibrinogen concentrations ≥8 mg/ml possibly constitute less permissive microenvironments for neurite extension and cell migration, which is in line with the inhibitory effect on cell outward migration observed for fibrinogen concentrations of 8 mg/ml and 10 mg/ml, at day 4 of culture (Figure 5a). To assess the effect of fibrinogen concentration on cell proliferation, total cell number and the percentage of proliferative cells were determined at day 14 of culture, counting the average number of viable cells per Fb drop and determining the fraction of cells incorporating BrdU (5-bromo-2’-deoxyuridine), respectively. Results obtained showed no significant effect on either the total cell number or the percentage of proliferative cells at this time-point, when varying the fibrinogen concentration (see the Supplementary material online, Figure S4B,C). Finally, the effect of fibrinogen concentration on MMP activity present in conditioned media of ES-NSPCs cultured for 14 days within Fb was also investigated. Except for an increase in MMP-2 activity levels found when increasing the fibrinogen concentration from 8 mg/ml to 10 mg/ml, no significant differences were found among MMP-9 activity levels when varying the fibrinogen concentration (see the Supplementary material online, Figure S4D).

3.6 Culture of hES-NSPCs within Fb gels

It was then investigated whether this 3D culture model could support the neuronal differentiation of single hES-NSPCs. Confocal analysis of cell–matrix constructs incubated with calcein AM/PI after 14 days of cell culture, showed the presence of live cells widely distributed throughout the Fb gel growing as cellular spheroids, similarly to mES-NSPCs in Fb (Figure 6a). At the same time, the quantitative analysis of live/dead cells isolated from Fb gels revealed a high percentage of viable cells (98.4%) and an average of 6.7% PI+ cells (dead cells; Figure 6b). Immunocytochemistry analysis showed the presence of nestin+ cells (Figure 6c), as well as a population of neurons expressing βIII-tubulin with neurites radially extending from cellular spheroids (Figure 6d). Synaptic proteins (synapsin) with a distribution pattern resembling that of neurites were also observed (Figure 6e). Similarly to mES-NSPCs, hES-NSPCs were found to deposit laminin and fibronectin (Figure 6f,g).
Fibronectin deposition was found to be closely associated with neurites expressing βIII-tubulin, as shown by double immunocytochemistry performed at day 21 (Figure 6h).

4 Discussion

The present study established the 3D culture of NSPCs derived from mouse ES cells under adherent monoculture, in fibrin hydrogels. The ES-NSPCs were seeded within Fb in the form of single cells and cultured under neuronal differentiation conditions. Cell viability, morphology, proliferation, phenotype, distribution of cell-secreted ECM proteins, and proteolytic activity were subsequently characterized. Furthermore, the behaviour of ES-NSPCs within Fb in response to fibrinogen concentration was investigated.

The cell seeding density leading to the highest retention of cell viability in an Fb gel with fixed composition (fibrinogen concentration in the polymerizing gel: 6 mg/ml) was initially established. When seeded at $1 \times 10^6$ cells/ml, ES-NSPCs were able to retain in average more than 85% of cell viability throughout the culture period and to proliferate, as shown by the nine-fold increase in total cell number observed at day 14 of culture (Figure 1a–c). The high levels of cell viability observed may be associated with the presence of bFGF during the first 2 days of culture, owing to its mitogenic effect on SOX1+ cells (Ying et al., 2003), as well as to the presence of BDNF and NGF during the last period of culture. These two neurotrophins are important for neuronal survival (Levi-Montalcini and Angeletti, 1963; Shetty and Turner, 1998) which, apart from being constitutively-secreted by NSPCs (Lu et al., 2003), were added to the media. Moreover, the ability of Fb to sequester bFGF and BDNF through its heparin-binding domain (Martino et al., 2013) possibly contributed to their retention within fibrin and to prolong their bioavailability to cells. The addition of aprotinin to cell culture media, a serine protease inhibitor herein used to delay fibrin degradation, may have also contributed to cell viability, as aprotinin was shown to exert neuroprotective effects in vitro (Davis et al., 1975) and to increase cell viability of dissociated embryoid bodies (EBs)-containing neural progenitors in Fb (Willerth et al., 2006). ES-NSPCs entrapped within Fb as single cells rapidly divided forming spherical-shaped clone-like multicellular clusters. The formation of large-sized cellular spheroids (diameter greater than 300 μm) is often a concern, as this can compromise the delivery of oxygen/nutrients/growth factors to the inner part of the spheroids, and ultimately lead to the formation of necrotic cores (Sen et al., 2001). However, ES-NSPCs seeded at $1 \times 10^6$ cells/ml resulted in the formation of small-sized spheroids with diameters below values that could compromise cell viability in the core of the spheroids (Figure 1d). Fibrin was able to sustain a pool of NSPCs, as shown by the high percentage of total cell population expressing the NSC marker SOX1-GFP and the NSPC marker nestin at the end of culture, regardless of bFGF withdrawal from the cell culture medium at day 2 of culture (Figure 2c,e). This is an important feature, as hydrogels for delivery of ES-NSPCs should provide an adequate microenvironment for the maintenance of a pool of NSPCs following transplantation, not only because of their potential to differentiate into mature neural cells but also because functional recovery is thought to be mainly mediated by neurotrophic factors secreted by NSPCs (Lu et al., 2003; Ying et al., 2003). Single ES-NSPCs seeded within Fb gels efficiently generated neurons, giving rise to a dense neuronal network (46% of βIII-tubulin+ cells; Figure 2) and, to a lesser extent, to astrocytes and oligodendrocytes. The capacity of maturation of these neuronal cells was shown by the increase in NF200 expression (Figure 2b), and detection of GABAergic (GAD67+ cells) and dopaminergic/noradrenergic neurons (TH+ cells), as well as by the presence of a network of synaptic vesicles and dendritic spines (Figure 2j–l). The ability of Fb to support ES-NSPC differentiation into such neuronal populations (namely, into GABAergic neurons) is valuable, as
several therapeutic strategies could profit in the transplantation of cells allowing a long-term supply of GABA (Kim et al., 2010). When differentiated on gelatin-coated surfaces, ES-NSPCs were found to express markers for glutamatergic, serotonergic and motor neuronal subtypes, in addition to GABAergic and dopaminergic markers (Incitti et al., 2014). Still, the ability of Fb to sustain ES-NSPC differentiation into glutamatergic, serotonergic and motor neurons remains an issue to be addressed.

As Fb acts as a provisional ECM that can be remodelled by invasive cells through cell-secreted proteases and cell-secreted ECM proteins, the ECM proteins distribution and the proteolytic activity in cell/Fb constructs were further analysed. Laminin and fibronectin expression by NSPCs was initially detected in floating aggregates of NSPCs (neurospheres) (Campos, 2004), and, more recently, collagen IV and vitronectin were also shown to be present in EB-containing neural progenitors (Sart et al., 2014). It was found that, when cultured within Fb, ES-NSPCs secrete endogenous ECM proteins, namely laminin, fibronectin and collagen type IV, contributing to Fb remodelling (Figure 3a–c). In comparison with laminin or collagen IV, fibronectin showed a wider distribution inside cell–Fb constructs, which can be related to the ability of Fb to sequester fibronectin through its fibronectin high-affinity binding domains (Brown and Barker, 2014). MMPs are key players in ECM remodelling because of their ability to cleave ECM components such as collagen, laminin and fibronectin (Tonti et al., 2009), as well as to proteolytically degrade fibrinogen and fibrin (Ahmed et al., 2007). Increasing evidence is supporting MMP involvement in neurogenesis, neurite elongation and axonal growth, as well as in synaptogenesis along homeostasis and pathologic events (Tonti et al., 2009; Fujioka et al., 2012). Interestingly, it was observed that ES-NSPCs within Fb secrete MMPs (Figure 3g) which, by locally contributing to ECM and Fb degradation, possibly facilitated neurite extension and cell migration, leading to Fb remodelling and to the formation of a more mature engineered tissue. In line with this hypothesis, Man et al. (2011) showed that neurite extension from dorsal root ganglia (DRG) explants entrapped in Fb is dependent on fibrinolysis and on MMPs and serine protease secretion. Interestingly, an increase of MMP-2 activity was found throughout the culture period, suggesting that this protease may play an important role in mediating ES-NSPC neurite extension and cell infiltration within Fb. MMP-9 activity was detected at a later stage of ES-NSPC culture within Fb, in accordance with the progress of neuronal differentiation and maturation observed (Barkho et al., 2008). The expression of matriptase, a type II transmembrane serine protease that acts upstream of MMP-2, promoting ES-NSPC migration and neuronal differentiation (Fang et al., 2011), was also assessed. ES-NSPCs within Fb retained the ability to express matriptase, showing an increase in matriptase expression during the second week of cell culture (Figure 2h), in accordance to its role in neuronal differentiation. The present study did not assess the expression of plasmin – one of the major fibrinolytic serine proteases – because ES-NSPCs were cultured within Fb in the presence of aprotinin.

As the structural and viscoelastic properties of hydrogels affect stem cell behaviour and fate (Banerjee et al., 2009; Man et al., 2011), the effect of fibrinogen concentration on NSPC behaviour was assessed. In accord with the literature, changes in fibrinogen concentration are expected to affect Fb structural and viscoelastic properties to a greater degree when compared to other Fb components, such as thrombin and calcium (Brown and Barker, 2014). Increasing the fibrinogen concentration resulted in increased Fb stiffness and in a denser Fb network with the presence of smaller pores (Figure 4), in accordance with the literature (Brown and Barker, 2014). Still, significantly changes in average pore diameter were only observed when increasing fibrinogen concentration to 8 mg/ml or 10 mg/ml. In the present study, ES-NSPCs within Fb were found to sense the changes in Fb viscoelastic and structural properties, as shown by the decrease in both neurite extension/cell...
outgrowth migration and neuronal differentiation in Fb gels with small-sized pores and stiffer matrices (Figure 5a–c), corresponding to 8 mg/ml or 10 mg/ml of fibrinogen. To sustain neuronal differentiation, neurite outgrowth and cell infiltration, Fb gel mechanical properties should ideally match those of native neural tissues while Fb structural and biological properties should ensure support to cells and provide an adequate density of bioactive domains for integrin/growth factor/ECM binding (Man et al., 2011). The fibrinogen concentrations that induced a better performance of ES-NSPCs within Fb (in terms of neurite extension, neuronal differentiation, and formation of smaller-size spheroids) were those of 4 mg/ml and 6 mg/ml, which resulted in Fb gels with a storage modulus comparable to that of neural tissue in the adult brain (400–1000 Pa) (Elkin et al., 2011). The findings are in agreement with previously published studies that reported a decrease in βIII-tubulin expression levels for adult NSCs cultured in alginate gels when increasing the elastic modulus of these gels (Banerjee et al., 2009). In contrast, no decrease in cell proliferation was observed when increasing the elastic modulus of Fb hydrogels (Figure S4c) in the present study. This result may be related to the lower percentage of BrdU-incorporating cells found at this time-point (<5%), which may be too low to allow the detection of significant differences among the fibrinogen concentrations tested. Nevertheless, when considering an in vivo scenario, Fb gels produced with fibrinogen concentrations of 8 mg/ml and 10 mg/ml can also be of interest to reduce the polymerization time and the degradation rate (Aurand et al., 2012; Brown and Barker, 2014), as in vitro these Fb hydrogels also supported neuronal differentiation.

Finally, the present study assessed whether or not this 3D culture model could support human ES-NSPC viability, neuronal differentiation and ECM remodelling. Although the culture of human fetal/adult NSPCs within hydrogels has been extensively explored (either seeded as single cells or as cellular aggregates), only a few studies report the 3D culture of dissociated NSPCs derived from human pluripotent stem cells within hydrogels (Sun et al., 2015; Alessandri et al., 2016). Similar to mES-NSPCs, hES-NSPCs within Fb were found to retain cell viability and sustain a pool of NSPCs, while expressing neuronal and synaptic proteins. These results are in good agreement with the studies of Alessandri et al. (2016), in which NSPCs derived from human induced pluripotent stem cells encapsulated as single cells within Matrigel revealed high retention of cell viability (97.8% of viable cells), with expression of NSPC and neuronal markers. In addition, the present study verifies that hES-NSPCs cultured within a 3D Fb hydrogel were able to remodel Fb through cell-secreted fibronectin and laminin – two ECM proteins present in the embryonic neural stem cell niche (Solozobova et al., 2012).

5 Conclusions

In the present study, it was shown that mES-NSPCs obtained in adherent monoculture, when seeded within a Fb hydrogel as dissociated cells remain viable, while retaining the ability to proliferate, differentiate into mature neurons and establish neuronal networks. In this 3D system, ES-NSPCs were also able to secrete ECM proteins and MMPs, therefore contributing to Fb remodelling and to the formation of a more mature engineered neural tissue. The ES-NSPCs were found to respond to changes in Fb viscoelastic and structural properties, with neurite extension/cell outgrowth migration and neuronal differentiation being hindered in Fb gels with stiffer matrices and small-sized pores (herein corresponding to gels prepared with 8 mg/ml or 10 mg/ml of fibrinogen). This 3D culture model was also shown to support cell viability, neuronal differentiation, expression of synaptic proteins and ECM deposition of human ES-NSPCs seeded as single cells within Fb. The settled 3D platform is expected to constitute a valuable tool for the development of Fb-based hydrogels...
specifically designed for ES-NSPC delivery into the injured CNS. Moreover, it could be of interest for pharmacological screening and to unravel molecular mechanisms implicated in neurogenesis in an in vitro 3D microenvironment, which better mimics in vivo conditions.

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Conflict of interest

The authors have declared that there is no conflict of interest.
REFERENCES


Figure 1 Cell viability, proliferation, and cellular spheroid diameter of mouse embryonic stem–neural stem/progenitor cell (mES-NSPCs) within fibrin (Fb) hydrogels. (a) Distribution of live and/or dead cells at day 7 and day 14 of culture; cell–Fb constructs were incubated with calcein AM and propidium iodide (PI) for detection of live and dead cells, respectively, and imaged by confocal laser scanning microscopy (CLSM); representative two-dimensional projections of CLSM stack images covering a depth of 300 μm are shown. (b) Quantitative analysis of cells incorporating PI (PI⁺, non-viable cells) at day 7 (D7) and day 14 (D14) of culture, as determined by flow cytometry (n = 3 independent experiments). (c) Total number of viable cells per Fb drop at day 0 (D0; upon cell embedment within Fb), day 7 and day 14 of culture (n = 3 independent experiments). (d) Histogram of cellular spheroid (neurosphere) diameter distribution at day 14 of culture; analysis was performed in four Fb drops from two independent experiments. [Colour figure can be viewed at wileyonlinelibrary.com]
Figure 2. Phenotypic characterization of mouse embryonic stem–neural stem/progenitor cell (mES-NSPCs) within fibrin (Fb) hydrogels. Western blot analysis of (a) βIII-tubulin (developing and mature neurons) and (b) NF200 (mature neurons) expression over the culture period (n = 3 independent experiments). (c) Quantitative expression of stage-specific embryonic antigen 1 (SSEA-1; mouse embryonic stem cells), SOX1-green fluorescent protein (GFP) (early neuroectodermal marker), nestin (neural stem/progenitor cells), βIII-tubulin, glial fibrillary acidic protein (GFAP; astrocytes) and O4 (oligodendrocytes) at day 14 (D14) as determined by flow cytometry (n = 2 independent experiments). Representative two-dimensional (2D) projections of confocal laser scanning microscopy (CLSM) stack images of cell/Fb constructs at day 14 processed for immunofluorescence labelling of (d) SSEA-1, (e) nestin, (f) GFAP, (g) O4, (h) βIII-tubulin, (i) NF200, (j) synapsin (synaptic vesicles), (k) GAD 67 (GABAergic phenotype) and (l) tyrosine hydroxylase (TH, dopaminergic and noradrenergic phenotype). GAPDH, glyceraldehyde-3-phosphate-dehydrogenase. [Colour figure can be viewed at wileyonlinelibrary.com]
Figure 3. Characterization of mouse embryonic stem–neural stem/progenitor cell (mES-NSPC) culture in terms of deposition of extracellular matrix (ECM) proteins and proteolytic activity. Representative two-dimensional projections of confocal laser scanning microscopy stack images of cell–fibrin (Fb) constructs processed for immunofluorescence labelling of (a) fibronectin, (b) laminin and (c) collagen type IV, at day 14 of culture. (d–f) Immunofluorescent labelling of ES-NSPCs cultured on poly-d-lysine/laminin-coated glass coverslips, at the same time-point. (g) Activity of secreted matrix metalloproteinases (MMPs) as determined by gelatin zymography (n = 3 independent experiments; *p < 0.05); ES-NSPCs embedded within Fb secrete matrix metalloproteinase (MMP)-2 and MMP-9, with MMP-9 being detected at a later stage of culture; MMP-2 activity increased at day 14 (n = 3 independent experiments; *p < 0.05). (h) Western blot analysis of matriptase expression over the culture period, showing that ES-NSPCs within Fb endogenously express matriptase (n = 3 independent experiments; *p < 0.05).

[Colour figure can be viewed at wileyonlinelibrary.com]
Figure 4. Effect of fibrinogen concentration on fibrin (Fb) hydrogels microstructure and stiffness. (a) Microstructure of Fb gels prepared with 4, 6, 8 or 10 mg/ml of fibrinogen spiked with Alexa Fluor 488-labelled fibrinogen; representative two-dimensional (2D) projections of confocal laser scanning microscopy (CLSM) stack images covering a depth of 10 μm are shown. (b) Average pore area of Fb hydrogels, as determined by image analysis of the 2D projected CLSM stacks (n = 4; *p < 0.05). (c) Storage modulus (G’) and loss modulus (G’’) of Fb hydrogels, as assessed by rheological analysis (n = 6 independent experiments; *p < 0.05; **p < 0.01; ***p < 0.005). [Colour figure can be viewed at wileyonlinelibrary.com]
Figure 5. Effect of fibrinogen concentration on mouse embryonic stem–neural stem/progenitor cell (mES-NSPC) outward migration, viability and neuronal differentiation within fibrin (Fb) hydrogels. (a) Representative two-dimensional (2D) projections of confocal laser scanning microscopy (CLSM) stack images of cell–Fb constructs processed for F-actin/DNA staining at day 4 of culture; images for Fb hydrogels prepared with 4 mg/ml or 10 mg/ml of fibrinogen are shown. Neurite extension/outward migration from cellular spheroids was quantified by image analysis; bar graphs depict the number of sprouts (above) and the maximal outgrowth (below) per spheroid ($n=10$ spheroids from two replicate cultures; *$p<0.05$; **$p<0.01$; ***$p<0.005$). (b) Neuronal network formed within Fb hydrogels at day 14 of culture, as assessed by immunofluorescence labelling of βIII-tubulin; representative 2D projections of CLSM stack images of cell–Fb constructs covering a depth of 100 μm are shown. (c–d) Flow cytometry analysis of βIII-tubulin expression at day 14 of culture. (c) Percentage of positive cells, showing a decrease in the percentage of βIII-tubulin$^+$ cells when increasing the fibrinogen concentration from 4/6 mg/ml to 10 mg/mL ($n=4$–6 independent experiments; **$p<0.01$); (d) the analysis of the mean fluorescence intensity ratio of the histograms also suggest a lower expression of βIII-tubulin in these Fb gels. (e) Percentage of viable cells at day 14 of culture using Trypan Blue dye exclusion method ($n=4$–6 independent experiments). [Colour figure can be viewed at wileyonlinelibrary.com]
Figure 6. Cell viability, phenotype and extracellular matrix (ECM) deposition of human embryonic stem–neural stem/progenitor cells (ES-NSPCs) cultured within fibrin (Fb) hydrogels. (a) Distribution of live and/or dead cells at day 14 of cell culture; cell–Fb constructs were incubated with calcein AM and propidium iodide (PI) for detection of live and dead cells, respectively, and imaged by confocal laser scanning microscopy (CLSM); representative two-dimensional (2D) projections of CLSM stack images covering a depth of 150 μm are shown. (b) Quantitative analysis of live and dead cells at day 14, as determined by flow cytometry (n = 3 independent experiments). (c–h) Representative 2D projections of CLSM stack images of cell/Fb constructs processed for immunofluorescence labelling of (c) nestin, (d) βIII-tubulin, (e) synapsin (synaptic vesicles), (f) laminin, (g) fibronectin, and (h) fibronectin and βIII-tubulin; inset shows a single confocal image (0.17 μm in depth) demonstrating close association of fibronectin fibrils with neurites. Images (c–g) correspond to day 14 of cell culture and image (h) to day 21. [Colour figure can be viewed at wileyonlinelibrary.com]
Supplementary Materials and Methods

Culture of mES-NSPCs on laminin-coated glass coverslips under neuronal differentiation conditions

Glass coverslips previously coated with 20 µg/mL of poly-D-lysine (P7280, Sigma) were incubated overnight with 5 µg/mL of laminin (L2020, Sigma). After cell dissociation, cell suspension was transferred to poly-D-lysine/laminin-coated glass coverslips at 2×10⁴ cells/cm². ES-NSPCs were cultured for 14 days under neuronal differentiation conditions, following the same protocol described for neuronal differentiation of ES-NSPCs within Fb gels (for detailed description see section 2.2 of the manuscript).

Cell viability and total cell number

For qualitative cell viability analysis, cell-Fb constructs were sequentially incubated with 4 µM calcein AM (Invitrogen; 30 min at 37°C) and 6 µM propidium iodide (PI; Sigma; 10 min at 37°C), for detection of live and dead cells, respectively. After incubation, samples were washed twice with PBS, transferred to cell culture media, and immediately observed under the confocal laser scanning microscope (CLSM; Leica TCS SP5II). Quantitative analysis of cell viability was performed in a pool of six Fb drops after cell isolation from the cell-Fb constructs, as described in Materials and Methods section. Cell number per drop was determined using a Neubauer chamber and the percentage of viable cells determined using Trypan Blue dye exclusion method or by flow cytometry. For flow cytometry analysis single cell suspensions were incubated with 67 nM calcein AM (20 min at 37°C) or with 6 µM PI (10 min at 37°C) to label live and dead cells, respectively. Cells were finally washed 3×, suspended in FACS buffer and analysed by flow cytometry, unlabeled cells were used as negative control. Representative histograms are represented in Figure S3A. The average cell number per Fb drop was determined dividing the total number of viable cells counted using the Neubauer chamber by the number of Fb drops.

5'-Bromo-2'-deoxyuridine (BrdU) assay

The percentage of proliferative cells was assessed by immunocytometry detection of BrdU incorporated into cells synthesizing DNA. BrdU (Sigma) was added to the cell culture medium of six replicate Fb drops at the final concentration of 50 µM, and ES-NSPCs kept in culture for additional 18 h. Cells were isolated from the gels as described, fixed in 1% (w/v) paraformaldehyde (PFA; 20 min), permeabilized with 0.2% (v/v) Triton X-100 (10 min at 4°C) and further exposed to 2 M HCl for 20 min. Cells were subsequently incubated in blocking buffer (PBS buffer containing 5% normal goat serum (NGS, Biosource) and 0.1% (w/v) saponin) for 20 min followed by incubation with anti-BrdU (V450 conjugated anti-BrdU antibody, 560810, BD Horizon; 8 µg/mL) for 30 min. Cells were finally washed 3× and suspended in FACS buffer containing sodium azide for immunocytometry analysis (FACS Canto™ II, BD Biosciences). Samples not exposed to BrdU but processed in parallel for immunodetection of BrdU were used as negative control. Representative histograms are shown in Figure S4C.

Gelatin zymography

MMP activity in conditioned media of mES-NSPC cultures before and after embedment within Fb was analysed by gelatin zymography, as previously described (Cardoso et al., 2014). For this purpose, media was collected at day 5 of the neural commitment protocol and after 4, 8, and 14 days of ES-NSPC culture within Fb. Total protein was determined and 20 µg of total protein loaded per lane. MMP activity was analysed using Quantity One software (BioRad). Data presented correspond to three independent experiments. MMP activity was analysed using Quantity One software (BioRad), and interpreted based on the literature (Frankowski et al., 2012).

Western Blot analysis

Western blot analysis was performed on protein extracts of mES-NSPCs, before and after cell embedment within Fb. For this purpose, cold RIPA buffer was added to dissociated ES-NSCs at day 5 of neural commitment or added directly to a pool of six cell/Fb constructs collected at day 4, 8, and 14 of culture. The constructs were sonicated, centrifuged at 3000 x g (10 min at 4°C), and protein concentration in the supernatant fractions determined by the DC protein assay (BioRad). Protein extracts (5 µg for matriptase expression, and 20 µg for βIII-tubulin and NF200 expression) were incubated at 95°C for 5 min and subjected to a 10% or 4-12% SDS-PAGE. After SDS/PAGE separation the proteins were transferred to a nitrocellulose membrane (GE Healthcare) and blocked in TBS-T
buffer containing 5% (w/v) skim milk or 5% (w/v) bovine serum albumin for 1 h. The membranes were then incubated with rabbit polyclonal anti-matriptase (sc-48830, Santa Cruz Technologies; 1:2000), rabbit monoclonal anti-βIII-tubulin (MRB-435P; Covance; 1:5000), rabbit polyclonal anti-NF200 (AB8135, Abcam; 1:2000), and mouse monoclonal anti-GAPDH (5G4-6C5, HyTest; 1:40000) overnight at 4°C. After rinsing with TBS-T buffer membranes were incubated for 1 h with horseradish peroxidase-conjugated secondary antibodies and the blots developed with horseradish peroxidase-reactive chemiluminescence reagents (GE Healthcare). After exposure to autoradiographic film, the relative density of the protein band was further analysed with Quantity One software (BioRad).
### Table S1 – Primary and secondary antibodies as well as isotype controls used for immunocytochemistry studies.

<table>
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<th>Isotype control</th>
<th>Dilution</th>
<th>Origin and product number</th>
<th>Secondary Antibody</th>
<th>Dilution</th>
<th>Origin and product number</th>
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<td>Santa Cruz Biotechnology sc-21702</td>
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<td>-</td>
<td>1:100</td>
<td>Chemicon MAB353</td>
<td>Goat anti-mouse IgG1-PE</td>
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<td>Developing and mature neurons</td>
<td>5% (v/v) NGS in NIS</td>
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<td>-</td>
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<td>BD BioSciences 558713 (Isotype)</td>
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</table>

NGS, normal goat serum; SSEA-1, stage-specific embryonic antigen 1; GFAP, glial fibrillary acidic protein.
### Table S2 – Primary and secondary antibodies used for immunocytochemistry studies.

<table>
<thead>
<tr>
<th>Target</th>
<th>Blocking solution</th>
<th>Primary Antibody</th>
<th>Dilution</th>
<th>Origin and product number</th>
<th>Secondary Antibody</th>
<th>Dilution</th>
<th>Origin and product number</th>
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</thead>
<tbody>
<tr>
<td><strong>Cellular Markers</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Mouse ES cells</td>
<td>5% (v/v) NGS in PBS</td>
<td>Mouse monoclonal anti-SSEA-1</td>
<td>1:100</td>
<td>Santa Cruz Biotechnology sc-21702</td>
<td>Alexa Fluor 594 goat anti-mouse</td>
<td>1:100</td>
<td>Invitrogen A11020</td>
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<td>NSPCs</td>
<td>5% (v/v) NGS in PBS containing 0.05% (v/v) Tween-20</td>
<td>Mouse monoclonal anti-nestin</td>
<td>1:100</td>
<td>Chemicon MAB353</td>
<td>IgG</td>
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<td></td>
<td></td>
<td>Mouse monoclonal anti-nestin (human)</td>
<td>1:200</td>
<td>Chemicon AB22035</td>
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<td></td>
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<tr>
<td>Developing and mature neurons</td>
<td>5% (v/v) HS and 3% (v/v) BSA in PBS containing 0.05% (v/v) Tween-20</td>
<td>Rabbit monoclonal anti-tubulin</td>
<td>1:500</td>
<td>Chemicon AB15207</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
<td>1:100</td>
<td>Invitrogen A11070</td>
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<tr>
<td></td>
<td></td>
<td>Mouse monoclonal anti-tubulin</td>
<td>1:500</td>
<td>Covance MMS-435P</td>
<td>Alexa Fluor 488 donkey anti-mouse IgG</td>
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<td>Invitrogen A21202</td>
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<td><strong>Astrocytes</strong></td>
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<td>Rabbit polyclonal anti-GFAP</td>
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<td>DAKO Z0334</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
<td>1:100</td>
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<td><strong>Oligodendrocytes</strong></td>
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<td>Mouse monoclonal anti-O4</td>
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<td>Chemicon MAB345</td>
<td>Alexa Fluor 594 goat anti-mouse IgG</td>
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<td><strong>Mature neurons</strong></td>
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<td>Rabbit polyclonal anti-NF200</td>
<td>1:1000</td>
<td>Chemicon AB8135</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
<td>1:100</td>
<td>Invitrogen A11070</td>
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<tr>
<td>Synapt 1, phosphoproteins associated to synaptic vesicles</td>
<td>10% (v/v) NGS in PBS</td>
<td>Rabbit polyclonal anti-synapsin</td>
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<td>Chemicon AB1543P</td>
<td>Alexa Fluor 660 donkey anti-goat IgG</td>
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<td>GABAergic neurons</td>
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<td>Goat polyclonal anti-GAD67</td>
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<td>Santa Cruz Biotechnology sc-7512</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
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<td>Invitrogen A11070</td>
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<td>Dopaminergic and noradrenergic neurons</td>
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<td>Rabbit polyclonal anti-tyrosine hydroxylase</td>
<td>1:500</td>
<td>Chemicon AB152</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
<td>1:100</td>
<td>Invitrogen A11070</td>
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Table 1. ECM markers

<table>
<thead>
<tr>
<th>ECM marker</th>
<th>Source and Dilution</th>
<th>Primary Antibody</th>
<th>Secondary Antibody</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibronectin</td>
<td>1% (v/v) BSA in PBS</td>
<td>Rabbit polyclonal anti-fibronectin</td>
<td>Alexa Fluor 647 donkey anti-rabbit IgG</td>
<td>1:100</td>
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<tr>
<td>Laminin</td>
<td>1% (v/v) BSA and 4% (v/v) FBS in PBS</td>
<td>Rabbit polyclonal anti-laminin</td>
<td>Sigma-Aldrich L0303</td>
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<td>Collagen IV</td>
<td>5% (v/v) BSA in PBS</td>
<td>Goat polyclonal anti-collagen IV</td>
<td>Chemicon AB169</td>
<td>1:50</td>
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</tbody>
</table>

Supplementary References


Frankowski H, Gu YH, Heo JH, Milner R, Del Zoppo GJ. 2012, Use of gel zymography to examine matrix metalloproteinase (gelatinase) expression in brain tissue or in primary glial cultures, Methods Mol Biol, 834: 221-33.
Figure S1. Phenotypic characterization of mESC-NSPCs at day 5 of monolayer neural commitment. (A) Representative fluorescence image of the monolayer culture showing cells expressing Sox1-GFP (early neuroectodermal marker) after neural phenotypic induction. (B) Percentages of Sox1-GFP+ cells attained in the different neural commitments performed as assessed by flow cytometry: live cells were gated using forward scatter (FSC, reflecting size) and side scatter (SSC, reflecting complexity) criteria, and fluorescence gates set using undifferentiated 4iC ES cells; a representative dot plot and fluorescence intensity histogram is shown. (C) Immunocytochemical analysis of SSEA-1 (mouse embryonic stem cells), nestin (neural stem/progenitor cells), βIII-tubulin (developing and mature neurons) and GFAP (astrocytes) expression (N = 3 independent experiments); fluorescence intensity histograms from a representative experiment are shown; the blue histogram represents the fluorescence of cells incubated only with the isotype-specific secondary antibody.
Figure S2. Distribution of viable cells within Fb. (A) Distribution of viable cells following embedding in the hydrogel: fibrin gels containing mES-NSPCs (1×10⁶ cells/mL of Fb) were incubated with 4 μg/mL calcein AM for 30 min and 2.8 μg/mL Hoechst 33342 dye for 30 min at 37°C (both from Invitrogen), for detection of live cells (in green) and DNA (in blue), respectively, and immediately imaged by CLSM; representative 2-D projections of CLSM stack images covering a depth of 140 μm are shown. (B) Distribution of live/dead cells at day 14 of culture, as a function of cell seeding density (1, 2 or 3×10⁶ mES-NSPCs/mL of Fb); cell-Fb constructs were incubated with calcein AM and PI for detection of live and dead cells, respectively, and imaged by CLSM; representative 2-D projections of CLSM stack images covering a depth of 200 μm are shown. (C) Quantitative analysis of cells incorporating PI (PI⁺; dead cells) at day 14 of culture as determined by flow cytometry (N = 2-3 independent experiments). (D) Degradation of cell-Fb constructs, as a function of aprotinin (AP) concentration within Fb gel; mES-NSPCs were embedded within Fb gels prepared with 5, 10 and 25 μg/mL of AP, and cultured in medium without AP; representative images of whole cell-Fb constructs acquired using a stereomicroscope (SZX10, Olympus) at day 0 of culture are shown; the incorporation of AP in Fb gel was not sufficient to delay the degradation of cell/Fb constructs, independently of the concentration used. Fb gels prepared with 25 μg/mL of AP degraded at a lower rate, as suggested by the presence of residual Fb gel at this time point (arrows). Scale bars, 2 mm.
Figure S3. Representative flow cytometry fluorescence histograms correspondent to: (A) Live/dead assay performed at day 7 and 14 of mES-NSPC culture within Fb; cells were isolated from the cell/Fb constructs, incubated with calcine AM and PI (for detection of live and dead cells, respectively), and analysed by flow cytometry. (B) Phenotypic characterization of mES-NSPCs cultured within Fb for 14 days: SSEA-1 (mouse embryonic stem cells), Sox1-GFP (early neuroectodermal cells), nestin (neural stem/progenitor cells), βIII-tubulin (developing and mature neurons), GFAP (astrocytes), and O4 (oligodendrocytes); the grey line represents the fluorescence of cells incubated with secondary antibody only or with the correspondent isotype control.
Figure S4. Effect of fibrinogen concentration on cellular spheroid diameter, total cell number, percentage of proliferative cells, and MMP activity, at day 14 of mES-NPC culture within Fb. (A) Histogram of cellular spheroids diameter distribution as a relative frequency (N = 252-373 spheroids); large cellular spheroids (diameter > 125 μm) were only observed in Fb gels formed with the two highest fibrinogen concentrations tested (6 and 8 mg/mL). (B) Total cell number per Fb drop, after cell isolation from cell/Fb constructs (N = 3-6 independent experiments). (C) Percentage of proliferative cells assessed by BrdU incorporation and flow cytometry (N = 3-4 independent experiments); representative fluorescence histograms are shown; the fluorescence of cells not exposed to BrdU but processed in parallel for immunodetection of BrdU is also shown; varying the fibrinogen concentration did not significantly alter the total cell number per drop neither the percentage of proliferative cells (BrdU+) cells in the range of fibrinogen concentrations tested. (D) MMP-2 and MMP-9 activity present in cell culture conditioned media, as determined by gelatin zymography; MMP-activity levels were normalized to those of the lowest fibrinogen concentration tested (N = 3 independent experiments. *, p < 0.05); apart from the difference found between MMP-2 activity levels obtained for fibrinogen concentrations of 8 and 10 mg/mL, no significant differences were among the fibrinogen concentrations tested, independently of the MMP considered.