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Engineering an integrated cellular interface in 3-dimensional hydrogel cultures permits monitoring of reciprocal astrocyte and neuronal responses

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Abstract

This study reports a new type of 3D tissue model for studying interactions between cell types in collagen hydrogels. The aim was to create a 3D cell culture model containing separate cell populations in close proximity without the presence of a mechanical barrier, and demonstrate its relevance to modelling the axon growth-inhibitory cellular interfaces that develop in the central nervous system (CNS) in response to damage. This provides a powerful new tool to determine which aspects of the astroglial scar response and subsequent neuronal regeneration inhibition are determined by the presence of the other cell types. Astrocytes (CNS glia) and dissociated dorsal root ganglia (DRG; containing neurons and peripheral nervous system [PNS] glia) were seeded within collagen solution at 4°C in adjacent chambers of a stainless steel mould, using cells cultured from wild type or green fluorescent protein expressing rats, in order to track specific populations. The divider between the chambers was removed using a protocol that allowed the gels to integrate without mixing of the cell populations. Following setting of the gels, they were maintained in culture for up to 15 days. Reciprocal astrocyte and neuronal responses were monitored using confocal microscopy and 3D image analysis. At DRG:astrocyte interfaces, by 5 days there was an increase in the number of astrocytes at the interface followed by hypertrophy and increased glial fibrillary acidic protein expression at 10 and 15 days, indicative of reactive gliosis. Neurons avoided crossing DRG:astrocyte interfaces, and neuronal growth was restricted to the DRG part of the gel. By contrast, neurons were able to grow freely across DRG:DRG interfaces, demonstrating the absence of a mechanical barrier. These results show that in a precisely controlled 3D environment, an interface between DRG and astrocyte cultures is sufficient to trigger reactive gliosis and inhibition of neuronal regeneration across the interface. Different aspects of the astrocyte response could be monitored independently, providing an insight into the formation of a glial scar. This technology has wide potential for researchers wishing to maintain and monitor interactions between adjacent cell populations in 3D culture.
Introduction

Tissue engineered cell culture models have the potential to provide powerful new tools for neuroscience research and the development of therapies. Recreating the 3-dimensional spatial environment of the central nervous system (CNS) allows cells in vitro to behave more like their in vivo counterparts, providing robust and controllable model systems that mimic the cell biology present in the nervous system (1-3). Here we report a new type of 3D tissue model for studying interactions between cell types, and demonstrate its relevance to modelling the inhibitory interfaces that develop in the CNS in response to damage.

Neuronal growth following CNS damage is limited, resulting in permanent paralysis and loss of function in many patients. CNS neurons have the capacity to regenerate if a permissive environment is created, but a common finding of strategies aimed at bridging CNS lesions, including our work with biomaterial conduits (4, 5), is that axons readily enter and traverse the bridging graft, but find difficulty in subsequently exiting the graft, or disassociating from engrafted cells, to re-enter host CNS parenchyma (6, 7). An important feature of many current bridging devices aimed at spinal cord injury (SCI) repair is that they contain or become populated by Schwann cells which accompany the neuronal regeneration. However, these peripheral nervous system (PNS) glia can trigger resident astrocytes to become reactive and growth inhibitory (forming a glial scar), preventing regenerating axons from crossing the Schwann cell - astrocyte transition zone (8, 9). This graft/host cellular interface has similarities to the PNS/CNS interface of the dorsal root entry zone (DREZ).

Some success has been achieved in overcoming these cellular interfaces (10-12). However, research in this area will be facilitated by the development of appropriate experimental models in which to develop a better understanding of the formation and persistence of inhibitory interfaces in the CNS, and to develop new therapies to prevent interface formation or overcome the glial scar barrier (13,
In order to understand the behaviour of specific cell populations, *in vitro* models offer several advantages over whole animal studies when control over cellular components is required, and where real-time measurement of responses that cannot be monitored *in vivo* would be of benefit (1, 3). Whilst many two dimensional culture models of the glial scar have been developed (14-21), three-dimensional systems permit a similar level of control, manipulation and monitoring, yet they maintain cells in a more relevant spatial arrangement (22). Three dimensional co-culture models that explore specific neuron-astrocyte interactions were initially developed more than 20 years ago (22), slice culture approaches have been used to investigate the role of extracellular matrix molecules (23), and 3D hydrogel systems have enabled the response of neural cells to exogenous forces (24, 25) and gradients of matrix stiffness (26) to be assessed. The inhibitory interfaces that develop at the edges of a CNS lesion or repair site or the DREZ are 3D structures so recreating them *in vitro* should logically require a 3D culture approach. Furthermore, it is clear that some of the limitations of previous culture models can be overcome using 3D matrices. Astrocytes on stiff matrices tend to adopt a reactive phenotype whereas in 3D hydrogels they are less reactive until stimulated (27), and neurons in culture also respond differently to 2D and 3D environments (28, 29).

However, the compliant mechanical environment of hydrogel matrices can be problematic when trying to reduce variables in culture systems, since cells in such an environment will respond to relatively small mechanical cues (30-32), including the stiff surfaces of the surrounding mould (27). This means that approaches such as embedding cell-seeded conduits or even DRG explants within astrocyte gels will provide mechanical cues that will introduce confounding factors. Making a 3D cell culture model that brings separate cell populations into close proximity without the presence of a mechanical interface is a challenge, but is necessary in order to isolate exactly which aspects of the glial response and the subsequent neuronal inhibition are determined by the presence of the other cell types.
The aim of this work was to develop novel methodology for engineering a 3D culture model that recreates the cellular interface which develops at the edge of a CNS lesion or the CNS/PNS boundary. Here we report that adjacent cell-seeded hydrogels integrated fully upon setting, providing no mechanical barrier to cells. This novel approach allowed astrocyte responses and neuronal behaviour to be quantified, providing useful new insight into how each cell population responds to the presence of the other over time. Although we have developed this model for nervous system research, it is equally applicable to modelling interactions between a wide variety of cell types and is therefore suited to many different areas of research.

Methods

Astrocyte cultures

All experiments were performed according to the UK Animals (Scientific Procedures) Act (1986) and approved by the Open University animal ethics advisory group. Sprague–Dawley rats (a β-actin-green fluorescent protein reporter line or wild type) were used from established in-house breeding colonies. Primary astrocyte cultures were prepared from postnatal 2-day-old rat cortices as reported previously (27). Cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM; Gibco) supplemented with penicillin and streptomycin (100 U/mL and 100 mg/mL, respectively; Sigma) and 10% v/v fetal calf serum in 75 cm² flasks (Greiner) coated with poly-D-lysine (Sigma). After 8 days in a humidified incubator at 37°C with 5% CO₂: 95 % air cells reached confluence, then flasks were shaken at 150 rpm for 4 h to detach microglia and less adherent cells. Resulting cultures were 95% astrocytes and 5% microglia (as determined by immunoreactivity for GFAP and IB4 lectin, respectively). Cells were trypsinized, washed, and counted so that the correct densities could be calculated prior to seeding into collagen gels.
Dorsal root ganglia preparation and culture

Dorsal root ganglia (DRG) were dissected from adult rats (200–300 g). Nerve roots were stripped and DRGs incubated in collagenase (0.125 %; Sigma) for 2 h at 37°C. Tissue was dissociated by trituration and washed twice by centrifugation with 25 ml of media to remove any remaining collagenase. Cell pellets were resuspended in DMEM and the resulting mixture of neurons and PNS glia was seeded within collagen gels.

3D Interface culture construction

Interface cultures were prepared by simultaneously casting 400 µl collagen gels (32) containing 3 million astrocytes per ml or 16 dissociated DRGs per ml in adjacent compartments of a stainless steel mould (Figure 1). Gels were composed of 10% cell suspension in DMEM, 10% 10x minimum essential medium (MEM; Sigma), and 80% type I rat tail collagen (2mg/ml in 0.6 % acetic acid; First Link). The MEM and collagen were mixed together and neutralized using sodium hydroxide, assessed by colour change of phenol red pH indicator; then the mixture was added to the cell suspension, mixed to ensure even distribution of cells throughout the suspension, and transferred to the mould. The two culture populations were separated by the presence of a 0.1 mm thick stainless steel divider which was removed after 2 min to allow gels to integrate. During the casting process the mould was maintained at 4°C until removal of the divider, whereupon it was incubated at room temperature for 10 min, then 37°C for 20 min. Controlling the temperature in this way ensured that the adjacent gels integrated prior to setting. In each integrated model, either the astrocyte or the DRG culture was generated using cells from GFP animals. After setting, interface cultures were carefully removed from the mould and maintained in a petri dish with 8 ml DMEM and incubated as before. Cultures were maintained for up to 15 days before fixing in 4% paraformaldehyde. In some experiments, control interface cultures were included which contained DRG cultures in both compartments.
Immunofluorescence staining, microscopy and image analysis

Antibody sources, dilutions, and immunofluorescence staining were carried out as previously described (27). Gels were stained for glial fibrillary acidic protein (GFAP), CS56 and βIII tubulin, for astrocytic reactivity, chondroitin sulphate proteoglycans and neurites, respectively. Hoechst 33258 was used to label cell nuclei. Confocal microscopy was performed using a Leica TCS SP5 confocal microscope (Leica Microsystems) to sample designated fields within the astrocyte and DRG compartments of the interface cultures.

Quantification of astrocyte cellular responses

Confocal fields for quantification of astrocytes were 400 x 400 x 40 µm (XYZ) with 1 image per 1 µm in the Z direction, 3 fields were sampled at the interface and 3 were sampled from a control region of the astrocyte compartment 5.5 mm away from the interface, according to a predefined protocol. The volumes of GFP labelling and GFAP and CS56 immunoreactivity were measured using Volocity image analysis software (Improvision, Perkin-Elmer). Automated analysis protocols were developed and cell number was determined by counting Hoechst stained nuclei, so that cell density could be assessed and GFP/GFAP volumes could be expressed per cell. For astrocyte:DRG interfaces, four independent interface gels were analysed, from at least 2 separate cell preparations.

Quantification of neurite outgrowth

Confocal fields for quantification of neurons were 400 x 400 x Z µm (XYZ) with 1 image per 1 µm in the Z direction. The Z displacement varied for neuronal quantification to include the full length of all neurites in each field. Five regions at the interface, and 5 regions 5.5 mm away from the interface were sampled according to a predefined protocol, ensuring that the orientation of the interface was aligned parallel to the edge of the field. The length of βIII-tubulin immunostained neurites was measured with Openlab Software (Improvision), the number of neurites per neuronal cell body was counted, and the angle of deviation of each neurite from growth perpendicular to the interface was
calculated. The number of neurites which crossed the interface in each field was also assessed. For DRG:DRG interfaces 3 independent interface gels were analysed from 3 separate cell preparations.

Statistical analysis

Data were analysed using GraphPad Prism software (Version 4). Normality and quality of variance tests were performed on all data to determine which test was appropriate. A paired t-test was used with significance level 95% for comparison between 'at the interface' and 'away from the interface' data at comparable time points. For comparison between time points, an ANOVA with Tukey's post-hoc test was performed with significance level 95% for comparison of neuronal length, number of neurites per neuron and crossing behaviour of neurites at the interface. If variances of data sets were significantly different, then Welch's correction was applied. All values are indicated as mean ± standard error of the mean. p-values were taken as an indicator of statistical significance as follows: *p < 0.05, **p < 0.01, and ***p < 0.001 and where significant differences were present between two means in a figure these were indicated using a line.

Results

Methodology was optimised for the generation of 3D interface models. Specialised moulds were engineered from stainless steel which incorporated chambers separated by 0.1 mm thick dividers (Fig 1). By optimising the temperature and timing of gel setting and removal of the dividers, a protocol was developed that allowed the collagen in adjacent gels to integrate without the cell populations mixing. The result was that adjacent gels were physically contiguous, thus facilitating the investigation of subsequent cell-cell interactions without the presence of confounding mechanical cues.
Astrocytes undergo reactive gliosis at the interface

Reactive gliosis was monitored using confocal microscopy to detect astrocyte population density, volume and GFAP expression after 5, 10 and 15 days in 3D culture (Fig 2, A & B). Astrocytes at the interface with the DRG culture were compared to those 5.5 mm away from the interface in the same gels. There was double the number of astrocytes at the interface compared to the control cells away from the interface at 5 days. At 10 and 15 days the numbers of astrocytes in the interface regions remained higher than those in the corresponding control regions (Fig 2C). Over the course of the 15 days there was an increase in both astrocyte volume (based on cytoplasmic GFP measurement per cell) and GFAP expression (based on the amount of GFAP immunoreactivity per cell) indicative of both hypertrophy and activation. At 5 days there was a slight increase in astrocyte volume at the interface, then this increased to be significantly greater than control areas at 10 and 15 days (Fig 2D).

There was no difference in GFAP expression at 5 days but at 10 and 15 days there was considerably greater GFAP expression in the astrocytes at the interface compared to cells in the control regions 5.5 mm away from the interface (Fig 2E). CSPG immunoreactivity was associated with cells and extracellular matrix (Fig 3A) and there was a trend towards greater CSPG immunoreactivity at the interface than in control areas at all three time points (Fig 3A and 3B), although this difference was not statistically significant.

Further experiments in models with astrocyte:astrocyte interfaces showed no changes in astrocyte reactivity between the interface and control regions. In addition, incubation of astrocyte gels with DRG-conditioned media did not increase reactivity (data not shown).

Neurons are inhibited from regenerating across an astrocyte interface

Neurite growth was assessed at day 5 at the interface between DRG:astrocyte gels, at the interface between DRG:DRG gels, and 5.5 mm away from these interfaces in control areas (Fig 4A). There was a slight reduction in the number of neurites detected per 3D field in all regions within the DRG gels
that were in contact with astrocyte gels compared to those in the DRG:DRG interface group although this was not statistically significant (Fig 4B). Figure 4C shows that there was a similar amount of neurite outgrowth in all regions of all DRG gels, with mean neurite length in the range 133.3 – 159.1 µm. Interestingly, despite the overall amount of neurite growth being equivalent in the different groups, there was a marked difference in the orientation of neurites and their ability to penetrate the interfaces. In the DRG:DRG cultures the neurites projected equally in all directions at the interface, whereas in the DRG:astrocyte cultures neurites at the interface grew parallel to it (Fig 4D). When the ability of neurites to cross the interface was analysed there was a highly significant reduction ($P<0.001$) in the proportion of neurites that crossed DRG:astrocyte interfaces compared to those that crossed DRG:DRG interfaces (Fig 4E).

**Discussion**

Here we report for the first time a method for making a physically integrated 3D interface between adjacent collagen hydrogels and its utilization to generate a model of the glial scar interface that demonstrates the development of reactive gliosis and neurite growth inhibition. Fluorescent tracking of different cell populations, combined with immunofluorescence, confocal microscopy and 3D image analysis provided a method for monitoring neuronal and glial cell responses. Astrocytes at the interface responded to the presence of DRG cultures by increasing in number, hypertrophy and increased reactive gliosis. The presence of the astrocyte interface caused no reduction in neurite growth within the DRG compartment, but significantly inhibited growth towards or penetration through the astrocyte interface by regenerating neurites. This was in contrast to similar DRG:DRG 3D cultures in which neurites grew in all directions and freely crossed the DRG:DRG interface, thus confirming the physical integration of the two gels.

It is interesting that in our system, adjacent compartments of astrocytes and DRG cells were necessary and sufficient to elicit the development of the neurite growth-inhibitory interface. This is
the first time that this has been demonstrated using a 3D culture model and provides a reproducible and high-throughput method for detailed investigations of the dynamics of interface formation and regenerative failure. By maintaining the astrocytes in a 3D hydrogel environment it was possible to detect the events of reactive gliosis from a much earlier stage than would be achieved using a monolayer culture in which the astrocytes would exhibit a more reactive phenotype in the absence of a stimulus (27). When mature astrocytes and Schwann cells come into contact on 2D surfaces they react to each other’s presence by producing migration-inhibitory molecules that concentrate at the cellular interface (33-36). Previous in vitro models have been used to explore this phenomenon further, for example studying the growth of axons across boundaries between monolayer cultures of astrocytes and Schwann cells (8), or across the DREZ in slice cultures from embryonic and mature spinal cord (37). Various studies have demonstrated the advantages of modelling astrocytes and neurons in 3D culture (3, 22, 27-29), but it is technically challenging to achieve a cellular interface in 3D culture without also generating a mechanical interface. Indeed, Yu and Bellamkonda showed that a mechanical barrier, formed by the elasticity mismatch between adjacent agarose gels, greatly influenced the ability of neurites to cross the resulting 3D interface (38). A key advantage of a culture model over an in vivo model is being able to limit the number of variables. It was therefore our aim to generate a culture system in which cell-cell interactions could be studied in isolation without the additional mechanical interface that can be present in simpler 3D culture systems or in vivo (particularly where an implanted material has mechanical properties that are mismatched in relation to the host tissue).

The behaviour of astrocytes and neurons in our 3D system mimicked key aspects of the reactive gliosis and inhibition of neuronal regeneration that occurs in vivo at the site of a CNS lesion or at the mature PNS:CNS boundary (7, 39). It is interesting to note that in our model we were able to explore the timing of three separate aspects of the astrocyte response: at day 5 there was an increase in cell numbers at the interface and a slight increase in cell volume with no apparent increase in GFAP
volume/cell; while at subsequent time points there was little further change in the number of cells but a steady increase in both cell volume and GFAP volume/cell. It would be interesting to conduct further experiments on this system to unravel the relative importance of these three aspects of astrocyte behaviour in the development of neurite growth-inhibitory interfaces. There was no change in astrocyte reactivity in the control regions away from the interface.

In our previous study we showed that primary astrocytes in 3D culture exhibited relatively low reactivity and could then react to a stimulus (in that case TGFβ1) to become reactive (27). Here the astrocytes became reactive in response to contact with dissociated DRG cultures, which contained similar cellular components to the mature DREZ or the permissive environment within a CNS repair graft (4, 34, 40). In this case the astrocyte response was localised to the region of the gel immediately adjacent to the interface, and the increase in cell density, hypertrophy and increased GFAP expression resulted in a 3D meshwork of reactive astrocyte processes resembling the inhibitory glial environment that forms at the boundary of a CNS lesion (6).

The culture system reported here permits the simultaneous analysis of neuronal and glial responses in 3D cultures, and could be amenable to supplementation of the culture media or incorporation of different cell types (e.g. transfected cells or primary cells from genetically altered animals) to allow investigation of neuro-glial biology or testing of CNS injury therapies. Furthermore, it is straightforward to monitor the same culture over time (time-lapse) to allow detailed studies of cell-cell dynamics as the inhibitory interface develops and matures. In a wider context, this technology might be useful to researchers interested in modelling interfaces in non-neural tissues or but at subsequent time points there was little further change in the number of cells.
tissue implants, where the ability to maintain and monitor interactions between adjacent cell populations in 3D would be of interest (41-44).

In summary we have demonstrated a method for generating 3D cell-cell interfaces and used this to simulate the cellular interactions that occur during the formation of an inhibitory interface in the CNS. The results show that an interface between a DRG culture and an astrocyte culture is sufficient to trigger reactive gliosis in the astrocyte population and inhibition of neuronal regeneration across the interface after 5 days. Our method permitted different aspects of the astrocyte response to be monitored independently, providing an insight into the progression of events that leads to the formation of a glial scar.

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Disclosure Statement

The authors have no competing financial interests

References


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Figure legends

Figure 1. Interface experimental set up. A Photograph of stainless steel interface mould; one 0.1 mm thick stainless steel divider is shown; however, up to three dividers could be inserted into the frame. Two culture populations in collagen gels were seeded either side of a steel divider, which was then removed after 2 min to allow gels to integrate. B Diagrammatic representation of one astrocyte:DRG interface gel.

Figure 2. Qualitative assessment of astrocyte reactivity at the interface with dissociated DRG cells. Astrocyte morphology and immunoreactivity was assessed using immunostaining and confocal microscopy for GFP (green), GFAP (red) and Hoeschst (blue) at days 5, 10 and 15 in culture. Representative confocal projections are shown of astrocytes at the interface with dissociated DRG
cells, and away from the interface, in an astrocyte only region at (A) low and (B) high magnification.

Scale bar in (A) = 150 µm and scale bar in (B) = 40 µm. Over time, astrocytes in contact with / in close
proximity to dissociated DRG cells become ramified and hypertrophic, which corresponds to an
increase in staining for GFAP.

Quantification of astrocyte reactivity at the interface with dissociated DRG cells: C GFP positive
astrocytic cell nuclei (stained with Hoechst) were quantified at and away from the interface with
dissociated DRG cells. Significantly more astrocytes were observed at the interface with dissociated
DRG cells than away from the interface at day 5 and 10 in culture. D Quantification of GFP staining
per cell revealed that astrocytes were becoming hypertrophic over time at the interface with
dissociated DRG cells. No differences were observed in astrocyte size away from the interface. E
Quantification of GFAP staining per cell revealed that astrocytes were becoming reactive at the
interface with dissociated DRG cells, with significantly greater GFAP expression at day 15 in culture
compared to away from the interface. * P < 0.05, ** P < 0.01.

Figure 3. CSPG expression at the interface with dissociated DRG cells. A Astrocyte CSPG
immunoreactivity was assessed using immunostaining and confocal microscopy, for CS56 (red) and
Hoechst (blue) at days 5, 10 and 15 in culture. Representative confocal projections are shown of
astrocytes at the interface with dissociated DRG cells, and away from the interface, in an astrocyte-
only region at high magnification. Scale bar = 100 µm. B Quantification of the volume of CSPG
staining revealed a trend towards greater CSPG staining at the interface with DRG cells although
these differences were not significantly different at any time point.

Figure 4. Assessment of neuronal growth in interface gels. A Neuronal growth was investigated using
immunostaining and confocal microscopy, for GFP (green), βIII tubulin (red) and Hoechst (blue) at
days 5, 10 and 15 in culture. Representative confocal projections are shown of control gels (an
interface gel of dissociated GFP DRG cells with dissociated WT DRG cells) and test gels (an interface
gel of GFP astrocytes with dissociated WT DRG cells). Arrowheads indicate neurites growing parallel to the interface. Scale bar = 150 µm.

Neuronal growth was assessed in control gels (an interface gel of dissociated GFP DRG cells with dissociated WT DRG cells) and test gels (an interface gel of GFP astrocytes with dissociated WT DRG cells) at days 5 in culture. B The number of neurites was quantified in confocal projections. Whilst there was a trend for a reduced number of neurites in the DRG:astrocyte interface gels, this did not reach statistical significance (p = 0.0547 at the interface vs control DRG:DRG interfaces). C Mean neurite length did not differ significantly between the different interfaces, at or away from the interface. D Analysis of the angle of neurite growth revealed differences between the control and test interfaces. Neurites were present at all angles in control interfaces, whereas in test interfaces, neurites were orientated predominantly parallel to the interface with astrocytes. E Significantly fewer neurites in the test gels crossed the interface compared to control gels in which more neurites (GFP positive) crossed the interface. *** p < 0.001.
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