Hydrogel and neural progenitor cell delivery supports organotypic fetal spinal cord development in an ex vivo model of prenatal spina bifida repair

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Abstract

Studying how the fetal spinal cord regenerates in an ex vivo model of spina bifida repair may provide insights into the development of new tissue engineering treatment strategies to better optimize neurologic function in affected patients. Here, we developed hydrogel surgical patches designed for prenatal repair of myelomeningocele defects and demonstrated viability of both human and rat neural progenitor donor cells within this three-dimensional scaffold microenvironment. We then established an organotypic slice culture model using transverse lumbar spinal cord slices harvested from retinoic acid–exposed fetal rats to study the effect of fibrin hydrogel patches ex vivo. Based on histology, immunohistochemistry, gene expression, and enzyme-linked immunosorbent assays, these experiments demonstrate the biocompatibility of fibrin hydrogel patches on the fetal spinal cord and suggest this organotypic slice culture system as a useful platform for evaluating mechanisms of damage and repair in children with neural tube defects.

Keywords

Myelomeningocele, spina bifida, neural tube defects, organotypic, slice culture, hydrogels, fetal surgery

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Introduction

Myelomeningocele (MMC) is a severe, prenatally diagnosed form of spina bifida resulting from failed fusion of the caudal region of the neural tube during embryonic development. The diagnosis is synonymous with substantial and permanent disabilities secondary to leg weakness and paralysis, hydrocephalus, cognitive impairment, bladder and bowel dysfunction, and orthopedic abnormalities.1

Given the unfortunate prognosis of the MMC fetus, one option for pregnant mothers is to undergo in utero primary closure of the neural tube defect at 25 weeks’ gestation in an attempt to mitigate the secondary injury of the exposed spinal cord.2,3 Although the procedure has shown clinical benefit in a randomized trial,4 the operation is highly invasive, induces preterm labor, and has mixed long-term neurologic outcomes. Another drawback of current prenatal repair techniques is that they do not address the primary and chronic secondary spinal cord damage that has already occurred.5 The ability to provide MMC children with treatment options that can better enhance the regenerative capacity of already damaged spinal cord tissue is needed.

Tissue engineering–based techniques that enable complete tissue coverage of spina bifida defects while actively facilitating spinal cord regeneration have gained traction as an alternative treatment strategy in experimental models.5–8 Unfortunately, the testing of these approaches has been
challenged by traditional fetal models of MMC repair, all of which have a number of shortcomings. For example, in vivo repair of MMC in fetal rodents is technically difficult to perform due to their small size and fragility. There are also limitations in the number of fetuses that can be treated in vivo to avoid high risk of postoperative demise. Moreover, affected pups do not routinely survive into the postnatal period to adequately evaluate treatment effect. Fetal large animal models of MMC share many of the same challenges seen in rodent models and are very expensive to conduct. The establishment of an ex vivo model that combines some of the advantages of animal models with those inherent with dissociated two-dimensional (2D) cell cultures may be an ideal and complementary research platform to explore the molecular and cellular aspects of MMC disease mechanism and repair, thereby representing an advance in this multidisciplinary field.

In this study, we sought to develop a novel, injectable hydrogel-based patch for use during fetal MMC surgical repair. We then aimed to evaluate the effect of these hydrogels in an organotypic slice culture model of fetal MMC repair. Our hypothesis was that fibrin-based hydrogels would provide a supportive three-dimensional (3D) microenvironment for donor-derived neural progenitor cells of either human or rodent origin. Moreover, we speculated that hydrogel patches would be biocompatible with prenatal MMC spinal cord tissue and would facilitate ongoing neuronal differentiation and axonal regeneration in slice cultures.

Materials and methods

Fetal MMC rat model

This study was approved at the Johns Hopkins University and the University of Michigan under protocols RA19M88 and PRO00007385, respectively, in accordance with the National Institute of Health (NIH) Guidelines for the Care and Use of Laboratory Animals. To induce fetal MMC, timed-pregnant Sprague Dawley dams (n = 10, Charles River, Wilmington, MA) were gavage fed 60 mg/kg of retinoic acid (Sigma-Aldrich, St. Louis, MO) on E10.5 as previously described. On E21 (term = E22), the dams were deeply anesthetized, and all fetal pups (range, 8–12 per dam), regardless of sex, were harvested by a midline laparotomy (Figure 1(a) and (b)). Fetuses were euthanized by decapitation, and the spinal cords in those with isolated lumbosacral MMC (n = 60) were removed in toto in sterile manner and immersed in ice-cold Hanks’ balanced salt solution (HBSS) containing glucose (10 mM) and sucrose (75 mM; Figure 1(c)).

Organotypic slice cultures

The organotypic slice culture model was adapted from the interface method as described elsewhere. Briefly, fetal spinal cords in affected pups were aligned and placed on a block supported with 1% agarose. The caudal portion was then sliced into 400 µm sections using a vibrotome (Mcllwain Tissue Chopper; Ted Pella, Redding, CA; Figure 1(c)). Three to five transverse or longitudinal slices with intact topography were placed on top of the membrane inserts (PICM0RG50; Millipore, Burlington, MA) within 35 mm culture dishes (Figure 1(d)). The first group (n = 8) of slices were encapsulated in a fibrin hydrogel patch (250 mL droplet) without cells. The second group (n = 8) of slices were encapsulated in a 250-mL hydrogel droplet containing rat neural progenitors (1.25 × 10^6). The third group (n = 6) of slices, which served as a control, received no hydrogel or cell treatment. One milliliter of medium containing DMEM with HEPES (50%; Gibco, Gaithersburg, MD), horse serum (25%; Gibco), and HBSS (25%; Gibco) was added to ensure coverage of each slice. Organotypic cultures were maintained for up to 14 days under standard conditions (37°C, 5% CO2).

Hydrogel patches

Based on our prior studies and work from others demonstrating increased neural fiber formation and delayed astrocytosis, fibrin hydrogel patches (50–250 mL each) were generated by mixing human fibrinogen monomers (40 mg/mL; Calbiochem, Gibbstown, NJ), thrombin (50 U/mL; Sigma), and CaCl2 (3.75 mM; Merck, Kenilworth, NJ). In a subset of human neural progenitor experiments, self-assembling peptide nanofiber hydrogels (Puramatrix™ 0.15%; 3D Matrix Medical, Waltham, MA) were also used per the manufacturer’s instructions.

Neural progenitor cells

Approval for all human cell experiments was obtained from the Johns Hopkins University (IRB #202082) and the University of Michigan (IRB #38565). After written informed consent was obtained, human somatic cells were acquired from unused amniocentesis samples (n = 3, 8–10 mL) collected from MMC (n = 1) and normal (n = 2) fetuses (mean gestational age = 25 weeks, two males and one female). Specimen donations were anonymous and would have otherwise been discarded as medical waste. The amniocytes were isolated and expanded in culture. Integration-free induced pluripotent stem cells (iPSCs) were generated in our laboratory using a non-integrating, cytoplasmic Sendai virus (Cyto-Tune; Thermo Fisher, Waltham, MA) encoding for OCT4, SOX2, KLF4, and cMYC (Yamanaka factors), as previously described. Individual colonies were subsequently picked mechanically and propagated on Matrigel (BD Biosciences, San Jose, CA). Alkaline phosphatase staining was performed using the AP substrate kit (Millipore) according to the manufacturer’s instructions. Immunofluorescence staining
was used to assess pluripotency in established iPSC lines. Primary antibodies against NANOG (Abcam, Cambridge, MA), OCT4A (Santa Cruz, Dallas, TX), SOX2, SSEA3, SSEA4, TRA-1-60, and TRA-1-81 (Millipore) were used. Differentiation of iPSCs into neural progenitors was conducted as previously described.19 Briefly, neurospheres were created from iPSC clones derived from all three patients. Individual rosettes were re-plated into dishes containing neural stem cell media (Neurobasal media, B27 (Invitrogen, Carlsbad, CA), NEAA, GlutaMAX, and FGF-2). Neural progenitors were expanded and enriched using magnetic microbeads (Miltenyi Biotec, Gaithersburg, MD) conjugated with PSA-NCAM (Millipore) antibody. Karyotyping of neural progenitors was performed after five passages using the GTL-banding method on 20 metaphase preparations. To assess neural phenotype in 2D and 3D culture, the cells were evaluated by gene expression or fixed in 4% paraformaldehyde and stained with primary

Figure 1. Organotypic slice culture rat model of fetal myelomeningocele (MMC) hydrogel patch treatment. (a) Gross inspection of representative lumbosacral defect (dotted oval) in a fetal MMC rat after maternal retinoic acid exposure. (b) H&E sagittal section through fetal MMC rat demonstrating lumbosacral defect (dotted oval, magnification: 4×). (c) Gross appearance of intact rat MMC spinal cords adjacent to an agarose block (asterisk). (d) Schematic view of MMC organotypic system showing spinal cord slices encapsulated within a hydrogel patch. Membrane inserts allow for nutrient absorption to ensure viability. No donor cells are depicted. (e) Representative transverse section (400 µm) of MMC lumbar spinal cord embedded in fibrin hydrogel on brightfield microscopy (day 0, magnification: 4×). Note the preservation of gross topography including median fissures. (f) Representative longitudinal section (400 µm) of MMC lumbar spinal cord embedded in fibrin hydrogel (day 0, magnification: 4×), white arrow = caudal end.
antibodies at 96 h. Confocal imaging was performed using a Nikon A-1 confocal microscope (Melville, NY). The cells (5 × 10^3/mL) in hydrogel suspension were also assessed by live/dead assay (Molecular Probes, Thermo Fisher) at 8 and 24 h. Percent viability was calculated in a blinded fashion from six random images at each time point using ImageJ (NIH, Bethesda, MD).

Rat fetal neural progenitors were purchased from a commercial vendor (GSC-8010; GlobalStem, Rockville, MD) or isolated from fetal Sprague Dawley rat central nervous tissue. At passage 4, the cells were labeled with mKate red fluorescence protein (RFP) based on a lentivirus driven by a synapsin promoter (IncuCyte; Essen Biosciences, Ann Arbor, MI; multiplicity of infection = 1). Immunohistochemical detection of donor neural cells within slice cultures was assessed using an RFP antibody (Thermo Fisher).

**Immunostaining**

All spinal cord tissue was fixed in 4% paraformaldehyde prior to immunostaining using the following primary antibodies: nestin (1:800, Cat. No. MAB5326; Millipore), musashi-1 (MSI1; Millipore), b-III tubulin (TuJ1, 1:800, Cat. No. Ab107216; Abcam), glial fibrillary acidic protein (GFAP, 1:500; Cell Signaling, Danvers, MA), caspase 3 (Cas3, 1:100; Millipore), Ki67 (1:200; Millipore), oligodendrocyte transcription factor (OLIG2; Thermo Fisher), NeuN (Millipore), microtubule-associated protein-2 (MAP2, 1:1000; Thermo Fisher), and NF200 (Sigma). Three random areas of tissue were imaged using a multiphoton microscope (Olympus FV1000, Waltham, MA) and converted into an RGB stack after tracing of total tissue area. Degree of staining was quantified in a blinded fashion based on the optical density method as described elsewhere.20,21 The densities were calculated based on minimum and maximum color thresholds for each section using ImageJ.

**Quantitative gene expression**

Relative gene expression of spinal cord tissue was analyzed by quantitative reverse transcription polymerase chain reaction (qPCR). Total RNA was extracted from explants using a MagMAX-96 Total RNA Isolation Kit (Life Technologies, Carlsbad, CA) and MAG Max Express (Applied Biosystems, Foster City, CA). RNA quantity and quality were determined spectrophotometrically using a NanoDrop 2000 spectrophotometer (Thermo Fisher). Reverse transcription was conducted using the SuperScript VILO kit (Invitrogen) according to the manufacturer’s protocol. Finally, qPCR was performed using Fast SYBR Green Master Mix (Applied Biosystems) and an AB-Quant Studio3 real-time PCR machine (Thermo Fisher). PCR primers were designed with Primer-BLAST (NIH; see Supplemental Table), and GAPDH was used as a reference gene for the normalization of target gene expression using the 2−ΔΔCt method.

**Neurotrophic factor analysis**

Enzyme-linked immunosorbent assays (ELISA; Merck Millipore) were used to measure brain-derived neurotrophic factor (BDNF), neurotrophin-3 (NTF3), and neurotrophin-4/5 (NTF4) levels in slice cultures. These specific neurotrophic factors were measured given their association with the promotion of corticospinal axon growth and immunomodulatory properties.22–24 ELISAs were performed twice on pooled slice cultures within a single dish according to the manufacturer’s instructions, and concentrations were normalized to total protein for each sample.

**Statistical analyses**

Quantitative data were presented as mean ± standard error. Data were analyzed by the Mann–Whitney test or one-way analysis of variance with post hoc testing by Kruskal–Wallis correction for multiple comparisons, as appropriate, using Prism 8 (GraphPad, La Jolla, CA). The results were considered to be statistically significant if p < 0.05.

**Results**

**3D hydrogel surgical patches support human neural progenitors**

Human amniotic fluid cells were isolated from three fetuses, including one with MMC, and were successfully reprogrammed into iPSCs using non-integrating Sendai virus. Under standard stem cell media conditions, amniocytes uniformly began to aggregate while reducing cytoplasmic volume and losing spindle-shaped morphology. After 28 days, there were multiple candidate iPSC colonies with well-defined borders and high nuclear-to-cytoplasmic ratios from all patients (Figure 2(a)). Morphologically distinct colonies expressed high levels of alkaline phosphatase (Figure 2(a), inset), and immunofluorescence staining of iPSCs revealed uniformly high expression of several pluripotency markers, including NANOG, OCT4, SOX2, SSEA3, SSEA4, TRA-1-60, and TRA-1-81.

Human iPSCs were then exposed to neural differentiation culture conditions. Neurospheres attached and formed neural rosettes typical of neuroepithelial cells within 7 days in adherent culture. All tested neural progenitor cell lines maintained a normal karyotype without clonal aberrations (Figure 2(b)). Directed differentiation in the neural lineage was confirmed by positive staining for multiple neural progenitor markers, including SOX2, nestin, and MSI1 (Figure 2(c)), and by the presence of axonal projections over time by light microscopy. The cells expressed moderate...
levels of TuJ1 but no detectable levels of GFAP, OLIG2, NANOG, and OCT4. Following resuspension of neural progenitors in hydrogel patches, gelation occurred within 15–30 min at 37°C. Neural progenitors within this 3D microenvironment demonstrated >60% viability when evaluated at 24 h based on intracellular esterase activity and plasma membrane integrity (Figure 2(d)). Immunofluorescent staining revealed predominantly nestin-positive cells, consistent with an immature neural progenitor cell phenotype.

Quantitative expression of various genes associated with neural development and differentiation showed that neural progenitors in hydrogel suspension did not alter RNA transcripts, including levels of *NES*, the gene encoding for nestin, which were comparable to those expressed in fetal brain positive controls (Figure 3(a)). The intermediate neural progenitor cell phenotype was confirmed by low *PAX6* expression in combination with negligible *MAP2*, *OLIG2*, and *GFAP* transcripts. Evidence of continued neurotrophic activity by neural progenitors resuspended in hydrogels was supported by significantly increased expression of *BDNF* and *NTF3* transcripts when compared with those in standard 2D culture (Figure 3(b)). Collectively, these data suggest that human neural progenitors derived from amniotic fluid iPSCs can survive and maintain neurogenic properties within 3D hydrogel matrices.

**3D hydrogel patches are a vehicle for rat neural progenitors**

Rat neural progenitors from all cell sources were successfully expanded and characterized prior to resuspension in
hydrogels (Figure 4(a)–(c)). Neural progenitors expressed both nestin and GFAP, and maintained >75% viability when resuspended in 3D culture at 8 and 24 h based on intracellular esterase activity and plasma membrane integrity. mKate2-labeled cells were visualized at 24 h (Figure 4(d)–(f)).

Slice culture model recapitulates development of the MMC spinal cord

Retinoic acid was given to pregnant rat dams to induce a posterior lumbosacral tissue defect in 66 (82.5%) of 80 term fetal pups (Figure 1(a) and (b)). The mean defect size was 7.9 mm × 5.4 mm. Lumbar spinal cord tissue harvested from affected rats was characterized by varying degrees of dorsal element injury. Nevertheless, representative immunohistochemistry showed a predominance of neuronal phenotypes in the expected topographical distribution within the gray matter of transverse sections (Figure 5(a)) and a relative paucity of astrocytes based on GFAP staining that were localized to the spinal cord periphery. The evaluation of slice cultures on day 1 demonstrated similar topography, which quantitatively revealed high levels of TuJ1, NeuN, and MAP2 and relatively low levels of GFAP and nestin at baseline (Figure 5(b)). There was no oligodendrocyte expression based on OLIG2. Evidence of baseline neurotrophic activity within slice cultures on day 1 was suggested by significant expression of Bdnf and Ntf3 transcripts when compared with age-matched rat dorsal root ganglion positive controls (Figure 5(c)).

Hydrogel treatment supports neuronal maturation within slice cultures

Spinal cord slices cultured for 4, 7, or 14 days within hydrogels maintained cytoarchitectural integrity as shown by hematoxylin and eosin (H&E) staining (Figure 6(a)) with strong expression of both nestin and TuJ1 throughout

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**Figure 3.** Gene expression and neurotrophic activity of human neural progenitors in 3D hydrogels at 96 h. (a) Representative quantitative gene expression data demonstrating relative specificity of neural progenitors derived from human iPSCs when compared to fetal brain positive controls. Placement of neural progenitors from 2D plates into 3D fibrin hydrogel suspensions did not alter gene expression, characterized by upregulation of NES, the gene encoding for nestin. Data were normalized relative to housekeeping gene (GAPDH) and are presented as mean ± SEM, *p ≤ 0.05 and **p ≤ 0.01 when compared to the other groups (Kruskal–Wallis), n = 4 independent biological replicates. (b) Quantitative gene expression showing relative neurotrophic activity of neural progenitors within 2D and 3D fibrin hydrogel microenvironments. There was upregulation in BDNF and NTF3, the genes encoding for brain-derived neurotrophic factor and neurotrophin-3, respectively. Human fetal brain RNA was used as a positive control, and data were normalized relative to housekeeping gene (GAPDH) and are presented as mean ± SEM, *p ≤ 0.05 (Kruskal–Wallis), n = 4 independent biological replicates.
Figure 4. Characterization of donor rat neural progenitor cells. (a) Brightfield microscopy of rat neural progenitors in 2D culture (magnification: 10×). (b) Positive immunofluorescence staining of neural progenitors using antibodies against nestin (Cy3 secondary, magnification: 40×) at 24 h. (c) Positive immunofluorescence staining of neural progenitors using antibodies against GFAP (FITC secondary, magnification: 40×) at 24 h. Nuclei were counterstained with DAPI. (d) Representative appearance of mKate2-labeled neural progenitors within 3D hydrogel patch (magnification: 40×) at 24 h. Scale bar represents 50 μm. (e) Low-power light microscopy image showing rat neural cells contained within hydrogel patch (white arrow = patch edge, magnification: 10×). Scale bar represents 500 μm. (f) High-power light microscopy images of hydrogel patch edge (white arrow, magnification: 40×), n=8–10 independent biological replicates for each experiment. Scale bar represents 100 μm.

the slice culture tissue (Figure 6(b)). Active outgrowth of axonal projections >500 mm from the tissue periphery was visualized in all groups by phase microscopy and by positive NF200 staining (Figure 6(c)). Although there was evidence of progressive hydrogel degradation over time by microscopy, approximately 25% of the remaining hydrogel was grossly visible on day 14.

Quantitative histopathologic alterations in neural cell phenotypes within transverse sections of slice cultures after 7 days (E21 + 7) or 14 days (E21 + 14) are summarized in Figure 7. Based on blinded analyses of immunohistochemistry images (Figure 7(a) and (b)), there were no significant changes in Ki67 and Cas3 expression over time (Figure 7(c) and (d)). There was increased expression of neuronal phenotypes at E21 + 14, including a statistically significant increase in TuJ1 after hydrogel treatment without cells (E21 + 14: 0.98 ± 0.11 vs E21 + 7: 0.51 ± 0.11, p=0.0125, Figure 7(g)). Based on MAP2 and NeuN expression, neuronal maturation scores increased over time in all groups and was significantly increased in the hydrogel group compared to time-matched controls (Figure 7(h) and (i)). Conversely, GFAP expression declined by E21 + 14 and was statistically significant after hydrogel treatment without cells (E21 + 14: 0.32 ± 0.06 vs E21 + 7: 0.54 ± 0.11, p=0.0418; Figure 7(j)). There were no statistical differences in GFAP expression among the three groups. Oligodendrocyte transcripts were undetectable in all cultures, and no mKate2-positive cells were visualized within slice cultures treated with hydrogel plus cells. Taken together, these data suggest acceptable biocompatibility of 3D fibrin hydrogels with MMC spinal cord slices in culture. Hydrogel treatment also correlated with enhancement of mature neuronal phenotypes when compared to controls without hydrogel.

Temporal increase in neurotrophic activity after hydrogel treatment

To evaluate the neurotrophic activity within slice cultures, ELISA was performed, which demonstrated a trend toward increased neurotrophic factor secretion within all slice cultures by E21 + 14. There was a statistically significant increase in NTF4 in slice cultures after hydrogel treatment with rat neural progenitors (E21 + 14: 107.6 ± 9.2 vs
Figure 5. Baseline characteristics of cytoarchitecture and neural cell phenotypes within the myelomeningocele (MMC) lumbar spinal cord and slice cultures at E21. (a) Representative native MMC lumbar spinal (transverse sections) using TuJ1, NeuN, MAP2, and GFAP antibodies demonstrate a reduction in dorsal elements and predominance of neuronal phenotypes (magnification: 4×). (b) Quantification of neural phenotypes within transverse sections of fetal MMC organotypic slice cultures (OSCs) based on optical density (y-axis) calculated from immunohistochemistry data (n=5/group). Values are presented as mean ± SEM, n=4 independent biological replicates. (c) Quantitative gene expression showing robust neurotrophic activity of fetal MMC OSCs at 24h based on brain-derived neurotrophic factor (Bdnf) and neurotrophin-4/5 (Ntf4) upregulation. Age-matched rat dorsal root ganglion (DRG) RNA was used as a positive control. Values are presented as mean ± SEM, *p ≤ 0.01 (Mann–Whitney), n=5 independent biological replicates.

Figure 6. Maintenance of cytoarchitecture and progressive neurite growth of organotypic slices of MMC spinal cord in 3D hydrogels. (a) H&E staining of representative transverse slice after 7 days in vitro (E21 + 7) with hydrogel patch (magnification: 10×). Scale bar represents 100 mm. (b) Positive immunofluorescence staining of typical transverse E21 + 7 MMC slice using antibodies against nestin (left) and TuJ1 (right, Cy3 secondary, magnification: 20×). Scale bar represents 500 mm. (c) Representative light microscopy at the periphery of MMC slice (asterisk, left) with robust neurofilament (NFH)-positive staining of neurite extensions at E21 + 14 (right, FITC secondary, magnification: 40×). Scale bar represents 500 mm.
Figure 7. Quantitative histopathologic analysis of myelomeningocele organotypic slice cultures after hydrogel patch treatment (with or without neural progenitors) for 7 days (E21+7) or 14 days (E21+14). (a) and (b) Representative appearance of slice culture sections after staining using antibodies against TuJ1 and MAP2, respectively. Scale bar represents 500 mm. (c) and (d) Ki67 and Cas3 as markers of cell proliferation and apoptosis, respectively. Control slice cultures in media alone. (e) and (f) Nestin and PSA-NCAM to evaluate neural progenitor phenotypes. (g)–(i) TuJ1, MAP2, and NeuN as markers of neuronal cells. TuJ1 was significantly increased in E21+14 hydrogels without cells. (j) GFAP to evaluate astrocytes. GFAP was significantly decreased in E21+14 hydrogels without cells. Optical density values are presented as mean ± SEM, *p ≤ 0.05 (Kruskal–Wallis), n = 4–13 independent biological replicates.
cannot ambulate independently despite in utero repair. These children, and more than half with spina bifida still continue to be ongoing challenges for clinical results associated with prenatal repair, bowel and bladder control remain issues.

E21 + 4: 58.5 ± 12.0, p = 0.0152; Figure 8). These neurotrophic factor levels were comparable to those from positive control dorsal root ganglion cells grown in 2D culture.

**Discussion**

In 2011, a landmark multi-institutional randomized trial was published demonstrating the efficacy of open surgical repair in human MMC fetuses. Despite the promising clinical results associated with prenatal repair, bowel and bladder control continue to be ongoing challenges for these children, and more than half with spina bifida still cannot ambulate independently despite in utero repair. Accordingly, novel experimental approaches aimed at further improving neurologic outcomes in these children have been pursued by multiple laboratories worldwide.

In this study, we developed an injectable hydrogel patch containing human neural progenitor cells as proof of concept for a cell-based tissue engineering strategy for fetal MMC repair. The immature human neural progenitors generated using transgene-free reprogramming methods stained positive for nestin, remained mostly viable within 3D scaffolds for over 24 h, and expressed genes associated with neurotrophic activity. Although neural progenitors derived from human amniotic fluid could potentially be used as an “off-the-shelf” cell therapy product for affected fetuses at the time of repair, autologous use of these patches may also be possible given that amniotic fluid itself could easily be obtained upon prenatal diagnosis, which usually occurs by ultrasound at 18–20 weeks of gestation. An autologous approach would have the additional advantage of alleviating concerns regarding the allogeneic immune rejection of donor cells. Currently, the time required for the preparation of our human neural progenitors took 10–12 weeks from amniotic fluid, but more rapid differentiation protocols from pluripotent stem cells have been described. In addition, transdifferentiation of amniocytes into the neurogenic lineage and isolation of specific neurogenic cell populations that may be uniquely accessible in the MMC fetus may be viable alternatives for use in hydrogel patches.

Our rationale for selecting neural progenitors as the ideal cell type for the MMC patch was based on previous work showing that neural progenitors can improve axonal regeneration in both animal and slice culture models of adult spinal cord injury. In addition to neural progenitor differentiation into neurons, the functional multipotency of these cells, including host cell integration, re-myelination of host neuronal circuitry, and constitutive secretion of neurotrophic and neuroprotective factors responsible for regrowth and axonal regeneration, has been demonstrated. However, in models of MMC, the use of cells derived from neural lineages to enhance spinal cord regeneration has been much more limited. Unique aspects of the MMC spinal cord compared to those in other types of spinal cord injuries include the potential ability to intervene with treatment in a more permissive fetal environment and more chronic nature of the neural tube injury. The latter is thought to be secondary to a primary neural tube defect combined with ongoing mechanical and chemical trauma incurred with the absence of overlying soft tissue. Due to their potential to engraft and to support endogenous neural cells within the host through immunomodulatory and trophic factors, the utility of mesenchymal stem cells has been more thoroughly investigated and has been beneficial in some MMC models of repair.

The second major finding from our study is the successful establishment of an organotypic model specific to the affected lumbar spinal cord in fetal MMC. Specifically, slice cultures from MMC pups demonstrated robust survival of endogenous cells within fibrin hydrogels for up to 14 days and showed evidence of continued differentiation into mature neuronal phenotypes with robust neurite outgrowth along its periphery. Ex vivo slice culture models of spinal cord regeneration have been widely embraced by neuroscientists as an informative research tool for over a decade. In our study, we also found minimal glial responses after hydrogel treatment, which is not a trivial

**Figure 8.** Neurotrophic activity as measured by enzyme-linked immunosorbent assay within myelomeningocele organotypic slice cultures after hydrogel patch treatment (with or without neural cells) for 4 days (E21 + 4) or 14 days (E21 + 14). (a) There was a trend toward increased brain-derived neurotrophic factor (BDNF) over time and a significant increase at E21 + 14 in organotypic slice cultures treated with hydrogel and neural cells when compared to E21 + 4. Age-matched rat dorsal root ganglion (DRG) cells were used as a positive control, n = 4 independent biological replicates. (b) There was a similar trend toward increased neurotrophin-4/5 (NTF4) levels over time. Values are presented as mean ± SEM, *p ≤ 0.05 and **p ≤ 0.01 (Mann–Whitney), n = 4 independent biological replicates.
finding given that previous histopathologic studies of the MMC placode have shown a paucity of mature neurons but with more extensive gliosis and fibrosis consistent with secondary injury.  

The MMC slice culture model is relatively simple to establish, easy to reproduce, and convenient for observing real-time changes in fetal spinal cord repair and regeneration. Further investigations in this model should allow us to test the effect of different neuroregenerative biomaterials and cell populations on different stages of development of the MMC spinal cord in a more controlled setting than what has been previously attempted. In contrast, in vivo fetal surgical manipulations in both rodents and sheep are often tedious to perform and are associated with high fetal morbidity and mortality. Ex utero murine embryo culture has also been described as a model of fetal treatment, but this technique is limited by the short time window (e.g., less than 24 h) between intervention and demise. Although our data suggest that fibrin hydrogels can support neuronal maturation over time, the addition of neural progenitors to the model did not consistently improve the regeneration of slice culture tissue. These results are in contrast to previous in vivo work in a prenatal ovine model, suggesting that the addition of bioactive fetal-derived cells may have benefit when compared to scaffold alone. Given our cell viability data and relatively short time frame of ex vivo culture, it is also possible that the number of neural progenitors delivered was insufficient to demonstrate functional multipotency within the MMC spinal cord. Whether there may be a pruning effect of fibrin hydrogels on donor neural progenitors is unknown. However, we should also acknowledge that the MMC spinal cord used in our experiments was harvested near term (E21), which does not take advantage of the likely enhanced regenerative capacity of the spinal cord earlier in gestation. Some investigators have suggested that donor cells may be more likely to differentiate into neural lineages within an embryonic spinal cord niche (E16).  

To date, most biomaterials described in animal models of MMC have comprised acellular, biodegradable sponges, sheets, or films whose primary aim is to provide tissue coverage over the spinal cord. The choice of polymerizing hydrogels in our study was based on prior reports of their use in other neuroregeneration models. We and others have also shown that fibrin supports neural progenitor survival in vivo and can provide physical and trophic support for nervous tissue ingrowth in neural tissue engineering models. The safety and efficacy of fibrin glue (Tisseel; Baxter, Deerfield, IL) as a watertight sealant in neurosurgical applications have already been established, and the potential of fibrin hydrogels as a delivery vehicle for donor neural cells and growth factors represents another advantage as a substrate for MMC repair. Desirable properties for any hydrogel suitable for fetal MMC surgical repair include rapid transition from liquid to solid phase, impermeability in the aqueous in utero environment, favorable attachment to surrounding tissues, and biocompatibility with fetal neural elements. In addition, the ideal hydrogel product should be injectable and capable of filling any cavity defects specific to the fetus, thereby providing a scaffolding to bridge damaged axons within the spinal cord.

One drawback of fibrin hydrogels has been the rapid degradation by native plasmin over the first several weeks after application. Although modifying the chemical properties of our hydrogels may extend its degradation until there is complete ingrowth of adjacent epithelium, another strategy to circumvent short degradation times would be to employ a biodegradable sheet on top of the hydrogel patch. This bi-layered approach could also serve an important barrier function to further minimize mechanical trauma. Continued advances in the evaluation of these biomaterials may eventually enable more widespread adoption of fetoscopic repair of MMC defects, which remains technically demanding compared to open surgery but has been shown to induce less maternal–fetal morbidity.

Moving forward, we propose that our slice culture model could be used as a unique “disease-in-dish” approach to understand mechanisms of mechanical and chemical spinal cord injury in MMC. The direct and paracrine effects of neural stem cells on the fetal spinal cord remain essentially unknown, and techniques for long-term experiments using spinal cord slice cultures have only recently been described. Since there remains a paucity of well-established biomimetic models that permit effective in vitro screening of fetal spinal cord regeneration, our approach represents a novel, high-throughput platform for evaluating next-generation MMC repair strategies, including stem cells and bioactive molecules, in both the prenatal and postnatal setting. Other stem cell populations, neurotrophins, and growth factors could be screened to better identify the optimal microenvironment to facilitate neural regeneration in this condition. Despite the results presented herein, there are several important caveats and limitations that should be acknowledged. First, as with most animal models produced using teratogenic compounds, there is inherent heterogeneity in the degree of spinal cord damage in the retinoic acid model of MMC. Such differences may explain some of the variability in our slice culture assays. Second, until we accumulate long-term data from slice cultures, we have yet to quantify neurite extension as a marker for degree of axonal regeneration and neural networking capacity. Such data may also allow us to measure functional aspects within these slice culture models as described by others. Third, we did not utilize baseline control groups that include direct exposure of our constructs ex vivo to culture media containing amniotic fluid. While not essential to demonstrate the biocompatibility of hydrogels in our model, inclusion of these groups would be ideal to better mimic the in vivo environment.
environment where the neural tube defect remains in contact with amniotic fluid. Such studies may also allow us to assess neural inflammation and the role of microglia during tissue regeneration. Finally, findings within this small animal model may be species-specific and therefore would likely require subsequent validation in scale-up studies on human tissues prior to clinical translation.

Conclusion

In summary, we developed hydrogel surgical patches for use during prenatal spina bifida repair and demonstrated viability of both human and rat neural progenitor donor cells within this 3D microenvironment. The patches were then tested in a 14-day organotypic slice culture model, which demonstrated ongoing neuronal development, neurite extension, limited astrocytosis, and robust neurotrophin activity. Thus, these experiments reveal the biocompatibility of fibrin hydrogel patches within the fetal MMC spinal cord and suggest an ex vivo organotypic system as a useful tool for evaluating mechanisms of damage and repair in children with neural tube defects.

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Supplemental material

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