Microextrusion printing cell-laden networks of type I collagen with patterned fiber alignment and geometry†

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Type I collagen self-assembles into three-dimensional (3D) fibrous networks. These dynamic viscoelastic materials can be remodeled in response to mechanical and chemical signals to form anisotropic networks, the structure of which influences tissue development, homeostasis, and disease progression. Conventional approaches for fabricating anisotropic networks of type I collagen are often limited to unidirectional fiber alignment over small areas. Here, we describe a new approach for engineering cell-laden networks of aligned type I collagen fibers using 3D microextrusion printing of a collagen-Matrigel ink. We demonstrate hierarchical control of 3D-printed collagen with the ability to spatially pattern collagen fiber alignment and geometry. Our data suggest that collagen alignment results from a combination of molecular crowding in the ink and shear and extensional flows present during 3D printing. We demonstrate that human breast cancer cells cultured on 3D-printed collagen constructs orient along the direction of collagen fiber alignment. We also demonstrate the ability to simultaneously bioprint epithelial cell clusters and control the alignment and geometry of collagen fibers surrounding cells in the bioink. The resulting cell-laden constructs consist of epithelial cell clusters fully embedded in aligned networks of collagen fibers. Such 3D-printed constructs can be used for studies of developmental biology, tissue engineering, and regenerative medicine.

Introduction

The extracellular matrix (ECM) consists of a heterogeneous mixture of macromolecules that form the non-cellular component of tissues.1 The fibrous structure of the ECM serves as a scaffold that provides chemical signals and mechanical support to its constituent cells. Reciprocal biochemical and biophysical interactions lead to continuous remodeling of the ECM, giving rise to a dynamic viscoelastic material with a rich diversity of structures and functions. One of the most common ECM structural motifs consists of aligned networks of type I collagen, which are associated with biological processes as diverse as collective cell migration,2 wound healing,3 metastasis,4 and tissue morphogenesis.5 These aligned patterns of collagen fibers have been challenging to reproduce ex vivo.

Collagen fibers are formed in vitro by the self-assembly of 300 nm-long tropocollagen monomers.6 Self-assembly is driven by a large positive entropy that results from the displacement of structured water surrounding tropocollagen monomers7,8 and depends on several parameters including concentration, pH, temperature, ionic strength, and molecular crowding.9–11 The resulting networks contain key biophysical and biochemical features observed in vivo and have been used extensively as models for native ECM.1

Nonetheless, in vitro networks of collagen are homogeneous and lack the anisotropy observed in vivo. In response to these limitations, several approaches have been described to induce collagen alignment either during or after self-assembly in vitro. During self-assembly, collagen fibers can be aligned using magnetic fields,12 flow fields,13–15 shear,16,17 mechanical instabilities,18 molding,19 or a combination of molecular crowding and spatial confinement,20 among others. After self-assembly, collagen fibers can be aligned by applying mechanical strain21,22 or shear.16 To the best of our knowledge, these approaches can only align collagen fibers uniaxially and are unable to spatially pattern collagen fiber orientation and geometry. As a result, existing approaches are unable to reproduce the complexity of collagen structures in native ECM. Moreover, it is challenging to fully embed cells or tissues into aligned networks of collagen using existing techniques. While strained elastomeric molds can be used to incorporate cells into networks of collagen with aligned fibers, it is unclear how to isolate the
effects of fiber alignment, compression, and collagen densification on cell behavior.  

Here, we describe a new approach to fabricate cell-laden networks of aligned type I collagen fibers using 3D microextrusion printing of collagen-Matrigel inks. We show that collagen can be 3D printed while simultaneously controlling the spatial deposition, geometry, and alignment of the resulting fibrous network. Whereas several studies have 3D printed collagen inks, we find that incorporating Matrigel into the ink allows us to print significantly lower concentrations of collagen (0.8 mg ml⁻¹). Our approach allows the collagen fiber geometry and orientation to be precisely analyzed throughout the volume of the printed construct. In addition, the low concentration of collagen more accurately reproduces the networks of collagen fibers used in cell culture experiments in studies of cancer cell migration and tissue-ECM interactions. We demonstrate that molecular crowding in the ink as well as substratum hydrophobicity can be used to tune collagen fiber alignment. We also demonstrate the ability to generate networks in which the collagen fibers are aligned in multiple directions. By combining 3D microextrusion printing and drop casting, we show that the geometry and alignment of collagen fibers can be patterned over mm-length scales, wherein distinct collagen fiber morphologies are separated by sharp interfaces with tunable geometry. We also show for the first time that microextrusion printing can be used to simultaneously bioprint epithelial cell clusters and control the alignment and geometry of collagen fibers surrounding them. Compared to other approaches, 3D microextrusion printing of collagen-Matrigel hydrogels is a simple, fast, and versatile technique that can generate large-scale cell-laden constructs with spatial control of collagen fiber alignment and geometry.

Materials and methods

Reagents and collagen gel preparation

Acid-solubilized bovine type I collagen (Advanced Biomatrix, Carlsbad, CA) was adjusted to a final concentration of 0.8 mg ml⁻¹ and neutralized to pH ~ 8 with the manufacturer-provided neutralizing solution. LAPONITE® XLG (BYK Additive and Instruments, Gonzales, TX), Pluronic F127 (BASF, Florham Park, NJ), or Matrigel (Corning, Corning, NY) were used as gelatinous additives at concentrations of 3 mg ml⁻¹, 250 mg ml⁻¹, and 4.2–10.1 mg ml⁻¹, respectively. The protein concentration of Matrigel was adjusted by dilution with 1:1 DMEM: F12 medium (Life Technologies, Carlsbad, CA). The molecular crowders, Ficoll 70 (70 kDa) and Ficoll 400 (400 kDa) (Sigma-Aldrich), were first dissolved in phosphate-buffered saline (PBS) and used at final concentrations of 6, 12, 18, and 24 mg ml⁻¹. The pH-adjusted collagen mixture was pipetted into a 3D-printing syringe and stored on ice for 1 h before printing.

3D microextrusion printing

Collagen inks were 3D printed using a microextrusion bioprinter (Inkredible+, CELLINK, Sweden) and conical polyethylene nozzles with diameters of 200 μm, 254 μm, or 400 μm (Nordson EFD, Robbinsville, NJ). All inks were 3D printed at room temperature (~20 °C) and were stored for ~5 min at room temperature prior to printing. The printing pressure and speed varied from 1–40 kPa and 20–80 mm s⁻¹, respectively. Printing paths were generated by writing G-code or by drawing 3D objects in Autodesk Inventor (Autodesk, San Rafael, CA) and importing the resulting STL file into Slic3r to generate G-code. Unless stated otherwise, samples were 3D printed onto a no. 1 glass coverslip at room temperature and cured in a 37 °C incubator at 5% CO₂ for 1 h before imaging. Prior to printing, coverslips were rinsed with 100% ethanol, air dried, and then cleaned with a UV/ozone cleaner (Jelight Company, Irvine, CA) for 7 min. Clean coverslips were silanized with exposure by trichloro(1H,1H,2H,2H-perfluorooctyl)silane (TCPFOS) (Sigma-Aldrich) vapors under vacuum for 24 h, 3,3,3-trifluoropropyltrichlorosilane (TFPTCS) (Alfa Aesar, Haverhill, MA) and methylcholrosilane (TMCS) (Sigma-Aldrich) vapors at atmospheric pressure for 20 min. Treatment with methoxy-poly(ethylene glycol)-silane (PEG-silane) (Laysan Bio Inc, Arab, AL) was achieved by coating clean coverslips with an ethanolic solution of 0.5% PEG-silane and 1% acetic acid for 30 min at 70 °C.

Cell culture

MDA-MB-231 human breast cancer cells were cultured in 1:1 DMEM:F12 medium (Life Technologies) supplemented with 50 μg ml⁻¹ gentamicin (Sigma-Aldrich) and 10% fetal bovine serum (FBS) (Atlanta Biologicals, Flowery Branch, GA). Functionally normal EpH4 mouse mammary epithelial cells were cultured in 1:1 DMEM:F12 medium supplemented with 50 μg ml⁻¹ gentamicin, 2% FBS, and 5 μg ml⁻¹ insulin (Sigma-Aldrich). Both cell lines were cultured in an incubator maintained at 37 °C and 5% CO₂. Clusters of mammary epithelial cells were generated by suspending cells in culture medium supplemented with 0.1% (w/v) Pluronic F108 (BASF, Ludwigshafen, Germany) and incubating at 37 °C and 5% CO₂ overnight. The culture medium for mouse mammary epithelial cell clusters was supplemented with 5 ng ml⁻¹ hepatocyte growth factor (HGF) (Sigma-Aldrich).

Cell viability

The viability of mammary epithelial cell clusters within collagen-Matrigel constructs was evaluated using a live/dead viability kit (Fisher Scientific, Hampton, NH). Cell-laden constructs were washed with PBS and immersed in an aqueous solution containing 2 μM calcein AM and 4 μM ethidium homodimer-1 for 45 min at room temperature before acquiring fluorescence images. The resulting fluorescence images were converted to binary images in order to calculate the live and dead areas of each cell cluster.

Immunofluorescence staining

Samples were fixed in a 4% (w/v) solution of paraformaldehyde (Alfa Aesar) in PBS for 15 min and washed with PBS. To label F-actin, samples were permeabilized using a 0.3% (v/v) solution of Triton X-100 (Sigma-Aldrich) in PBS (PBST) for 15 min.
Samples were then blocked in a 10% (v/v) solution of goat serum (Sigma-Aldrich) in PBST for 1 h. Next, samples were incubated in a 1:200 (v/v) solution of Alexa Fluor 594 phalloidin (Thermo Fisher Scientific) in blocking solution for 2 h and washed with PBST. To label nuclei, samples were incubated in a 1:5000 (v/v) solution of Hoechst 33342 (Invitrogen, Carlsbad, CA) in PBS for 15 min and washed with PBS. All immunofluorescence staining was performed at room temperature.

**Microscopy**

Collagen fibers were imaged using a 10× or 20× air objective or a 40× oil-immersion objective on a Nikon A1 laser-scanning confocal microscope in reflection mode (488 nm argon laser with GaAsP detector). 30 μm z-stacks, scanned at 2 μm intervals, were acquired for each sample and the maximum-intensity z-projection was obtained using ImageJ (NIH). Confocal reflection microscopy (CRM) was also used to image collagen fiber orientation at 10 μm intervals throughout the depth of the sample. Cell-laden collagen inks were imaged using a Nikon Plan Fluor 2×/0.1 NA, 10×/0.30 NA, or 20×/0.45 NA objective and ORCA-03G digital CCD camera (Hamamatsu Photonics, Japan) or 40× oil-immersion objective and Nikon A1 laser-scanning confocal microscope.

**Quantifying collagen fiber alignment and geometry**

Collagen fiber alignment was quantified using the local gradient orientation method (created by Jean-Yves Tinevez) in ImageJ. Collagen fiber anisotropy was estimated using an alignment fraction, which represents the fraction of aligned intensity gradients with respect to the total number of intensity gradients identified in an image. We considered an intensity gradient oriented within 20° of the printing direction to be aligned. All fiber alignment calculations were performed using maximum-intensity z-projections of 30 μm z-stacks obtained using CRM. Lengths of collagen fiber bundles were measured manually using ImageJ, while fiber diameters were approximated using the BoneJ plug-in (created by Michael Doube) in ImageJ. Heat maps were generated using the heatmap function in MATLAB (R2015b; MathWorks, Natick, MA).

**Rheological measurements**

Experiments were conducted using an MCR501 stress-controlled rheometer (Anton Paar, Ashland, VA). The temperature-dependent shear loss (\(G''\)) and storage (\(G'\)) moduli were measured using a 25 mm parallel-plate with a 600 μm gap. Before testing, samples (~300 μl) were loaded onto the bottom plate at a temperature of 4 °C. The temperature was raised to 37 °C at the beginning of the measurement to initiate gelation and samples were oscillated at 1 rad s⁻¹ and 0.5% strain for 10 min. Three independent measurements were acquired, and the average steady-state moduli were used for comparison.

**Contact angle measurements**

Advancing and receding contact angles were measured at room temperature using a Model 500-F1 contact angle goniometer (ramé-hart, Succasunna, NJ). All measurements were conducted on glass substrata and reported values represent an average of 10 measurements. Collagen and collagen-Matrigel samples were neutralized immediately before contact angle measurements.

**Statistical analysis**

Unless stated otherwise, data represent the mean of three independent replicates and error bars represent the standard error of the mean. Each independent replicate was conducted in triplicate, and two measurements were acquired for each sample. Statistical comparisons were performed using one-sided or two-sided \(p\)-values, which were calculated using Welch’s \(t\)-test or one-way analysis of variance. A \(p\)-value less than 0.05 was considered to be statistically significant.

**Results and discussion**

We began by evaluating the printability and shape retention of solutions of type I collagen during 3D microextrusion printing. “Shape retention” is a binary metric that describes the ability of a material to retain its shape after 3D printing. Materials with acceptable shape retention should demonstrate less than 10% change in length and width upon reaching equilibrium as compared to the programmed dimensions of the object. “Printability” is a qualitative description of the ability of a material to be continuously extruded at a constant printing speed and pressure and serves as a binary metric to assess the feasibility of an ink for microextrusion printing. Inadequate printability can be caused by a clogged nozzle or inconsistent flow.

A neutralized solution of collagen was pre-incubated on ice for 1 h prior to drop casting or 3D printing (Fig. 1a). Low-temperature pre-incubation allows collagen self-assembly to initiate at a reduced rate. Confocal reflection microscopy (CRM) revealed that both drop-cast (Fig. 1b) and 3D-printed (Fig. 1c) collagen inks form isotropic collagen fiber networks. Moreover, we found that the collagen ink had inadequate shape retention (Fig. S1, ESI†) and could not be accurately 3D printed into rectangular geometries. To enhance shape retention without increasing collagen concentration, rheological improvements are required.

Shear-thinning materials, including physically crosslinked gels, are commonly used to improve the rheological properties of 3D-printing inks.33,34 Shear and extensional flows generated during extrusion temporarily disrupt physical crosslinks, thus allowing a material to flow. After exiting the nozzle, the physical crosslinks reform and allow the printed shape to be retained. We therefore proceeded to mix collagen with shear-thinning gels to determine whether this would improve shape retention.

Aqueous LAPONITE® and Pluronic F127, which have well-characterized shear-thinning properties, have been used extensively as rheological modifiers for 3D printing and commercial applications.35,36 LAPONITE® is a synthetic hectorite clay that consists of nm-scale platelets. Pluronic F127 is a triblock copolymer of polyethylene oxide and polypropylene oxide, which forms micelles when dissolved in water above its critical micelle concentration (21% w/w) under ambient conditions.37
We combined LAPONITE® or Pluronic F127 with neutralized type I collagen and investigated their effects on shape retention, printability, and network structure. We found that collagen-LAPONITE® inks, which consisted of collagen mixed with a 3 mg ml⁻¹ LAPONITE® gel, frequently clogged the nozzle during printing. Similarly, collagen-Pluronic inks that consisted of collagen mixed with a 250 mg ml⁻¹ solution of Pluronic F127 frequently clogged the nozzle and did not retain their shape after printing (Fig. S1, ESI†).

To understand why LAPONITE® and Pluronic F127 adversely impact collagen shape retention and printability, we imaged gels using CRM, which revealed that collagen did not self-assemble into fibers when mixed with either additive in drop-cast (Fig. S2a and b, ESI†) or 3D-printed (Fig. S2c and d, ESI†) gels. Instead, large fragments of reflective material, which were likely aggregates of collagen and the additives, were observed. Similar aggregation has been reported for mixtures of collagen with polyethylene glycol or hyaluronic acid.²⁰ These results suggest that collagen self-assembly is adversely affected by the charged LAPONITE® particles and the Pluronic micelles. These possibilities are further supported by measurements of the shear storage moduli for the inks: collagen-LAPONITE® and collagen-Pluronic moduli differ significantly from pure collagen, LAPONITE®, or Pluronic moduli (Fig. S3, ESI†).

We therefore searched for an alternative material to improve shape retention and printability without disrupting collagen self-assembly. Matrigel is a gelatinous mixture of basement membrane proteins,³⁸,³⁹ primarily laminin (~60%), type IV collagen (~30%), and entactin (~8%). Matrigel is compatible
with collagen fibrillogenesis and also gels at similar temperatures to collagen (~37 °C), as shown by measurements of its temperature-dependent $G'$ and $G''$ values (Fig. S3, ESI†). We therefore mixed Matrigel with collagen and examined the shape retention and printability of the resulting inks. CRM images of drop-cast (Fig. 1d) and 3D-printed (Fig. 1e) collagen-Matrigel samples revealed intact collagen fiber networks. In addition, collagen-Matrigel inks were found to have good printability and shape retention (Fig. S1, ESI†) and could be 3D printed into narrow (~600 μm) lines (Fig. 1f and g). We therefore further explored the use of collagen-Matrigel inks for 3D printing.

We evaluated collagen fiber anisotropy in drop-cast and 3D-printed inks by quantifying the fraction of aligned collagen fibers. We found that the alignment fraction of drop-cast (27.9 ± 0.79%) and 3D-printed (27.9 ± 1.4%) pure collagen inks was identical, indicating that 3D printing did not alter the alignment of collagen fibers (Fig. 1h). Increasing collagen concentration (1.6 mg ml$^{-1}$; 27.9 ± 1.1% or 2.4 mg ml$^{-1}$; 28.2 ± 1.1%) had no effect on collagen fiber alignment (Fig. S4, ESI†). Similar results were observed with collagen-LAPONITE® (Fig. S2e, ESI†) and collagen-Pluronic (Fig. S2f, ESI†) inks. In collagen-Matrigel samples, however, fiber alignment was considerably higher in 3D-printed rectangles (31.5 ± 1.1%) and lines (38.1 ± 0.87%) than in drop-cast (27.0 ± 0.18%) configurations (Fig. 1i). We found that constraining drop-cast collagen-Matrigel inks to microfluidic channels (1 mm width and height) did not affect collagen fiber alignment (Fig. S5, ESI†). These data suggest that alignment in 3D-printed collagen-Matrigel rectangles is lower than in 3D-printed lines because the printing nozzle disrupts the alignment of existing fibers while printing new ones. To evaluate the spatial distribution of collagen alignment in 3D-printed samples, we acquired CRM images at varying $x$ and $y$ positions and generated alignment fraction heat maps (Fig. S6, ESI†). These heat maps revealed that 3D-printed collagen-Matrigel samples had regions of higher alignment than 3D-printed collagen samples and were more uniformly aligned than 3D-printed collagen-LAPONITE® or collagen-Pluronic samples. Alignment fraction did not vary as a function of sample depth (Fig. S6, ESI†).

To identify the mechanism by which collagen is aligned during 3D microextrusion printing, we investigated why anisotropy is generated in collagen-Matrigel inks but not pure collagen. One possible explanation is that molecular crowding in collagen-Matrigel inks improves alignment in 3D-printed

![Fig. 2 Identifying the mechanism for collagen fiber alignment in 3D-printed collagen inks. CRM images of 3D-printed collagen inks containing Ficoll 400 concentrations of (a) 6 mg ml$^{-1}$, (b) 12 mg ml$^{-1}$, (c) 18 mg ml$^{-1}$, or (d) 24 mg ml$^{-1}$. All CRM images represent the maximum-intensity z-projection of a 30 μm z-stack. (e) Alignment fraction of 3D-printed collagen-Ficoll 400 inks. Collagen-Ficoll samples were printed using a 254 μm-diameter conical nozzle at a printing speed of 40 mm s$^{-1}$. Scale bars = 50 μm. (f) Schematic of conical nozzle during 3D printing and (g) 3D-printing path used to identify different printing regimes shown in panel (f). (h) Normalized width of 3D-printed line as a function of printing speed and nozzle diameter. *$p < 0.05$; **$p < 0.01$.](https://example.com/fig2.png)
samples by increasing the molecular weight of collagen assemblies prior to extrusion. To test this hypothesis, we incorporated different concentrations of the molecular crowders Ficoll 70 or Ficoll 400 into collagen inks that were then drop cast or 3D printed. CRM images of 3D-printed collagen-Ficoll 400 hydrogels revealed isotropic networks at Ficoll concentrations of 6 mg ml⁻¹ (27.3 ± 0.27%) and 12 mg ml⁻¹ (27.7 ± 1.4%), aligned networks at 18 mg ml⁻¹ (37.5 ± 1.7%), and entangled networks at 24 mg ml⁻¹ (28.8 ± 2.3%) (Fig. 2a–e). Ficoll 70 gave similar results (Fig. S7a–e, ESI†), consistent with previous theoretical⁴⁰ and experimental⁴¹ studies, which have suggested that molecular crowding may not strongly depend on the size of the crowding agent. Consistently, we did not observe differences in the rate of collagen self-assembly in collagen-Ficoll 70 and collagen-Ficoll 400 inks (Fig. S8, ESI†). Adding Ficoll did not increase collagen alignment in drop-cast samples (Fig. S7f–n, ESI†). These results suggest that there is an optimal range for the molecular weight of collagen assemblies required to fabricate aligned networks of collagen. Below this range, collagen assemblies are either too small to be aligned or not stable enough to be extruded without being disrupted. Above this range, collagen assemblies are either too large to print without clogging the nozzle or are extruded as entangled networks.
During 3D microextrusion printing, hydrogels are subjected to shear and extensional flows in the conical printing nozzle, which may be sufficient to align collagen assemblies in the printing direction. We found that classical lubrication theory applied to the 254 μm-diameter conical nozzle yields shear and strain rate estimates on the order of 100 s⁻¹. This order of magnitude is consistent with shear rates previously reported to align type I collagen.16,42 Additionally, we investigated three magnitude is consistent with shear rates previously reported to align collagen fiber morphology.44 We found that collagen alignment remained constant as Matrigel protein concentration varied from 3 mg ml⁻¹ (28.36 ± 0.76%) to 6 mg ml⁻¹ (26.68 ± 0.46%) (Fig. 3d).

To investigate the effects of nozzle diameter and printing speed on collagen fiber alignment, we 3D printed collagen-Matrigel inks at different speeds using conical nozzles with different exit diameters. Collagen alignment increased as nozzle diameter decreased (Fig. 3e–h), which is consistent with the suspected role of shear and extensional flows in aligning collagen during printing. Similarly, we found that the alignment fraction increased at faster printing speeds (Fig. 3i–l).

We also observed that collagen-Matrigel inks had reduced surface wetting as compared to pure collagen inks (Fig. S10a, ESI†), which suggested that surface chemistry might play a role in dictating the alignment of collagen fibers. To test this hypothesis, we used different silane treatments to modify the hydrophobicity of the glass substratum on which the collagen inks were printed. Hydrophobicity was evaluated using advancing contact angle measurements and receding contact angle measurements. Collagen-Matrigel inks were 3D printed onto hydrophobic trimethylchlorosilane (TMCS) (θ_R/θ_W = 84°/79°), 3,3,3-trifluoropropyl-trichlorosilane (TFPTCS) (θ_W/θ_R = 88°/73°), or trichloro(1H,1H,2H,2H-perfluorooctyl)silane (TCPFOS) (θ_W/θ_R = 104°/83°)-treated glass, while methoxy-poly(ethylene glycol)-silane (PEG-silane) (θ_W/θ_R = 35°/21°) was used as a hydrophilic control (Fig. S10b, ESI†). Advancing contact angle measurements showed a decrease in contact angle with increasing silane concentration (Fig. S10c–e, ESI†), indicating a reduction in hydrophobicity.

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Given the suspected role of shear and extensional flows in aligning collagen assemblies during printing, we next investigated the range over which collagen fiber alignment and geometry can be tuned. We 3D printed collagen-Matrigel inks that had different concentrations of Matrigel protein (Fig. 3a–c). The average collagen fiber diameter was unaffected by changing Matrigel concentration (Fig. S9a, ESI†). However, the average length of collagen fiber bundles was significantly higher at a low concentration of Matrigel (3 mg ml⁻¹; 117 ± 11 μm) than at higher concentrations (4.5 mg ml⁻¹, 68.4 ± 5.8 μm; 6.0 mg ml⁻¹, 47.5 ± 2.4 μm) (Fig. S9b, ESI†). These results are consistent with findings that the concentration of Matrigel can impact collagen fiber morphology.44

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measurements showed that collagen-Matrigel inks became increasingly dewetting as substratum hydrophobicity increased [Fig. S10c, ESI†]. We quantified a normalized alignment fraction, which represents the alignment fraction on silane-treated glass relative to a control from the same ink printed onto untreated glass. We found that collagen alignment is maximized at intermediate substratum hydrophobicity (TF P T C S ) (Fig. 3m–p). We conclude that there is an ideal range for substratum hydrophobicity that maximizes alignment. Below this range the ink spreads after exiting the nozzle (Fig. 3m) and above this range the ink coalesces (Fig. 3o), both of which reduce alignment. Taken together, these data reveal that alignment can be tuned by modulating shear and extensional flows during printing as well as the hydrophobicity of the underlying substratum.

Controlling these parameters allowed us to spatially pattern collagen fiber alignment and geometry. By 3D printing...
collagen-Ficoll and collagen-Matrigel inks into a fan-shaped pattern (Fig. 4a), we aligned collagen fibers in multiple directions (Fig. 4b–d). A lower concentration of Matrigel (3 mg ml$^{-1}$) was needed to obtain alignment in the fan-shaped pattern, which suggests that the optimal Matrigel concentration depends on the geometry of the 3D-printed construct. To demonstrate control of shape retention and printability, we 3D printed an outline of a shield using collagen-Matrigel (Fig. 4e). We found that we could spatially control collagen fiber geometry by printing lines of collagen-Matrigel with a high Matrigel concentration (7.2 mg ml$^{-1}$) and drop casting low Matrigel concentrations (3 mg ml$^{-1}$) in between (Fig. 4f). CRM images revealed distinct regions with different collagen fiber geometries that were separated by sharp interfaces (Fig. 4f). To demonstrate the flexibility of this approach, we simultaneously controlled collagen fiber alignment and geometry by drop casting pure collagen in between 3D-printed lines of collagen-Matrigel (Fig. 4g). We observed that collagen fibers were aligned at the interface between these regions, which is consistent with previous studies of collagen-collagen and collagen-Matrigel interfaces. Moreover, we were able to generate complex alignment patterns by 3D printing interfaces with both concave (Fig. 4h) and convex (Fig. 4i) curvature. Single confocal slices of the convex interface are shown in Fig. S11 (ESI†). These results demonstrate that 3D printing can be used to generate a variety of complex patterns of aligned collagen fibers that cannot be achieved with other approaches.

To determine the response of cells to 3D-printed collagen networks, we cultured human breast cancer cells on top of 3D-printed collagen-Matrigel constructs. Based on previous reports that breast cancer cells orient along collagen fibers, we hypothesized that cells would align themselves in the printing direction. Cells were suspended in culture medium and subsequently seeded on top of polymerized collagen-Matrigel. We observed that cells oriented in the direction of collagen fiber alignment for both 3D-printed lines and curved printing paths (Fig. 5a). We found that the cell alignment fraction, which represents the fraction of cells oriented within 20° of the fiber alignment direction, was significantly higher for cells cultured on 3D-printed collagen-Matrigel networks (44.81 ± 1.93%) than for those on isotropic (27.27 ± 3.16%) networks (Fig. 5b). In addition, we incorporated human breast cancer cells into unpolymerized collagen, which was subsequently drop-cast on top of and in between 3D-printed lines of collagen-Matrigel (as in Fig. 4g). We observed that cells oriented along collagen fibers that were aligned perpendicular to the interface between 3D-printed collagen-Matrigel and drop-cast collagen (Fig. 5c). We found that the cell alignment fraction within 100 μm of the interface (33.25 ± 5.52%), where collagen fibers are aligned, was higher than in isotropic collagen networks (19.60 ± 1.79%) (Fig. 5d). We also 3D printed collagen-Matrigel inks that contained mouse mammary epithelial cell clusters. After printing, the cell-laden collagen-Matrigel construct was gelled at 37 °C, submerged in culture medium supplemented with 5 ng ml$^{-1}$ hepatocyte growth factor (HGF) in order to induce branching morphogenesis, and cultured for two days. Live/dead staining indicated that cell clusters remained viable after 3D printing (Fig. S12, ESI†). Fluorescence and CRM images of the cell clusters were acquired immediately after gelation as well as after one and two days in culture (Fig. 5e).

We found that the clusters extended actin-rich protrusions in the direction of collagen fiber alignment. As the number of days in culture increased, the circularity of clusters decreased and the orientation of the clusters approached the orientation of the printing direction (0°) (Fig. 5f). These 3D-printed collagen networks can be used to study a wide variety of interactions between cells and aligned networks of collagen fibers.

While collagen-Matrigel inks enable the fabrication of aligned networks of collagen fibers, the batch-to-batch variability and high cost of Matrigel preclude its use in clinical settings. This motivates the need for a similar gelatinous material that is compatible with collagen fibrillogenesis and that has viscoelastic properties conducive to 3D microextrusion printing. A synthetic formulation of Matrigel might be useful for improving printability and shape retention, which would allow for the fabrication of 3D cell-laden constructs while preserving native collagen fiber structure.

**Conclusions**

Here, we showed that 3D microextrusion printing of collagen-Matrigel inks can be used to fabricate cell-laden networks of aligned fibers of type I collagen. By tuning collagen self-assembly conditions and printing parameters, we demonstrated that collagen fiber geometry and alignment can be spatially controlled. Our results suggest that shear and extensional flows generated during 3D printing are responsible for aligning collagen and that the size of collagen assemblies during printing, which is dictated by molecular crowding in the ink, can be used to modulate alignment in 3D-printed networks of collagen. Moreover, we showed that cells cultured on top of 3D-printed collagen networks orient in the direction of collagen fiber alignment. We also demonstrated that collagen-Matrigel inks could be used to bioprint cell-laden constructs, wherein aligned networks of collagen surround fully embedded epithelial cell clusters. Aligned networks of collagen fibers that are generated by 3D printing will be useful in fields ranging from developmental biology to tissue engineering and regenerative medicine.

**Author contributions**

B. A. N., P.-T. B., and C. M. N. designed experiments; B. A. N. and P.-T. B. performed modeling calculations; B. A. N. performed experiments, completed data analysis, and prepared figures; B. A. N., P.-T. B., and C. M. N. wrote the manuscript.

**Abbreviations**

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<td>2D</td>
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<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>CRM</td>
<td>Confocal reflection microscopy</td>
</tr>
<tr>
<td>ECM</td>
<td>Extracellular matrix</td>
</tr>
</tbody>
</table>
Conflicts of interest

There are no conflicts to declare.

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