Three-Dimensional Elastomeric Scaffolds Designed with Cardiac-Mimetic Structural and Mechanical Features

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Three-Dimensional Elastomeric Scaffolds Designed with Cardiac-Mimetic Structural and Mechanical Features

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Tissue-engineered constructs, at the interface of material science, biology, engineering, and medicine, have the capacity to improve outcomes for cardiac patients by providing living cells and degradable biomaterials that can regenerate the native myocardium. With an ultimate goal of both delivering cells and providing mechanical support to the healing heart, we designed three-dimensional (3D) elastomeric scaffolds with (1) stiffnesses and anisotropy mimicking explanted myocardial specimens as predicted by finite-element (FE) modeling, (2) systematically varied combinations of rectangular pore pattern, pore aspect ratio, and strut width, and (3) structural features approaching tissue scale. Based on predicted mechanical properties, three scaffold designs were selected from eight candidates for fabrication from poly(glycerol sebacate) by micromolding from silicon wafers. Large 20 × 20 mm scaffolds with high aspect ratio features (5:1 strut height: strut width) were reproducibly cast, cured, and demolded at a relatively high throughput. Empirically measured mechanical properties demonstrated that scaffolds were cardiac mimetic and validated FE model predictions. Two-layered scaffolds providing fully interconnected pore networks were fabricated by layer-by-layer assembly. C2C12 myoblasts cultured on one-layered scaffolds exhibited specific patterns of cell elongation and interconnectivity that appeared to be guided by the scaffold pore pattern. Neonatal rat heart cells cultured on two-layered scaffolds for 1 week were contractile, both spontaneously and in response to electrical stimulation, and expressed sarcomeric α-actinin, a cardiac biomarker. This work not only demonstrated several scaffold designs that promoted functional assembly of rat heart cells, but also provided the foundation for further computational and empirical investigations of 3D elastomeric scaffolds for cardiac tissue engineering.

Introduction

Cardiovascular disease remains the leading cause of death in the United States and other developed countries and is a tremendous public health burden, with $297.7 billion in direct and indirect costs in 2008 in the United States alone.1 In addition to myocardial infarction (MI) and other acquired cardiomyopathies, approximately one percent of newborns and an increasing number of adults around the world survive with congenital heart disease, a condition associated with extremely high morbidity.2 Many of these patients could benefit from more effective cardiac grafts to replace tissue damaged by infarct or malformed at birth. Current pharmacological and prosthetic-based interventions serve either to modulate or to replace dysfunctional tissue; cellularized scaffolds capable of mediating endogenous tissue regeneration have yet to be adopted clinically. In general, conventional cardiovascular prostheses such as synthetic vascular grafts,3 bioprosthetic heart valves,4 and cardiac support, patch, or plug devices5–7 are capable of ameliorating disease and routinely improve patient outcomes, quality of life, and life expectancy. However, without living cells, bioprosthetic devices have limited potential to recapitulate the full breadth of native tissue functions or to support new tissue growth, which is especially important for pediatric applications. Recent studies suggest that an actively contractile graft could perform better than a simpler, nonviable implant.8,9

Tissue-engineered constructs comprised of living cells cultivated within a wide variety of scaffolds, including biomaterials with three-dimensional (3D) pore architectures,10–12 3D gels,13–17 composite scaffolds,18–21 and de novo synthesized22–24

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or decellularized extracellular matrices, offer a potential alternative, and have shown promising results in both animal and human studies in helping to resolve diverse cardiovascular lesions. By varying the material of construction and the fabrication method, scaffold mechanical properties may be controllable over a broad range. Nevertheless, many of the scaffolds utilized in these constructs lack the specific structural and mechanical requirements necessary within the cardiac regeneration environment.

The design and implementation of a clinically effective and durable scaffold remain a critical unmet need in cardiovascular medicine, where combining elasticity, porosity, and strength have been particularly challenging. In particular, the optimal mechanical properties of a scaffold designed to repair infarcted myocardium remain an unresolved question. If the scaffold is intended solely as a vehicle for cells to allow more effective cell delivery than simple injection of dissociated cells, then scaffold mechanical properties should accommodate robust cell viability, proliferation, and differentiation before and after engraftment. Scaffolds designed with this intent could compromise relatively soft, naturally occurring or engineered biomaterials, with stiffness shown to optimally support maturation and synchronous, auxotonic contractility of cardiomyocytes, and capable of delivering cells in combination with bioactive factors. However, if the scaffold is further intended to provide myocardial reinforcement after infarction, then scaffold mechanical properties will likely derive from a balance of considerations, addressing not only effective cell delivery but also infarct location and timing of the intervention. In particular, distinct collagenous scar tissue anisotropies were shown to evolve in different infarct locations as part of a complex healing process, warranting customized anisotropic reinforcement strategies that have been explored by computational analysis and demonstrated in vivo. An early example of a scaffold designed with the dual intent of delivering cells and providing mechanical reinforcement explored the combination of fibrin gel and an anisotropic elastomeric knitted fabric made of hyuloronan benzyl ester. As the material of construction for scaffolds fabricated in the present study, we selected a unique synthetic bioresorbable elastomer poly(glycerol sebacate) (PGS), developed by Wang et al., and recently reviewed by Rai et al. PGS, the 1:1 molar polycondensate of naturally occurring metabolites glycerol and sebacic acid, reproduces synthetically the mechanical stiffness and elasticity of the collagen and elastin components of the extracellular matrix. PGS provokes minimal inflammatory response, can be relatively hemocompatible, and is supportive of cardiogenesis and vascularlogenesis. Moreover, PGS degrades primarily by surface hydrolysis, with relative preservation of scaffold form and function, thereby facilitating the computational design of scaffolds exhibiting predictable cell guidance and load-bearing capacities upon tissue evolution and PGS degradation.

We previously showed that microfabricated PGS scaffolds of low-to-moderate stiffnesses (~30–300 kPa) exhibited mechanical stability and supported the in vitro cultivation and synchronous contraction of cultured rat heart cells, although in a manner that was largely isometric under the conditions tested. With an ultimate goal of both delivering cells and providing mechanical support to the healing heart in the present study, we sought to design, develop, and demonstrate a set of PGS scaffolds with (1) elastomeric stiffness and mechanical anisotropy mimicking native myocardium, (2) systematically varied combinations of rectangular pore pattern, pore aspect ratio, and strut width, and (3) microfabricated structural features approaching tissue scale. To match scaffold mechanical properties to native myocardium entailed considering the directional dependence, or anisotropy. To achieve minimal resistance to cyclic contraction and stretch, we chose PGS because it maintained elastic behavior during cyclic loading tests and heart cell culture in vitro, and showed promise as a myocardial implant material in vivo. We undertook finite-element (FE) modeling of a systematically varied set of eight scaffold designs from which we selected three scaffolds, based on cardiac mimic mechanical properties, for fabrication and demonstration in a 3D in vitro model of engineered cardiac tissue. To enable fabrication of structural and geometrical features approaching tissue scale at a relatively high throughput, we chose replica molding as a method by which we could create precise, reproducible, microporous architectures in a variety of shapes and patterns.

Materials and Methods

Silicon mold fabrication

Silicon wafer molds were fabricated as previously described, with modifications. Mold designs were laid out using L-Edit (Tanner). CAD/ART Services (Brandon) generated Mylar mask transparencies with a laser photo-plotter (Orbotech 7008, 20,000 dpi). Four-inch-diameter silicon wafers were used as the base for generating scaffold molds (WRs Materials). Hexamethydisilazane, a photore sist adhesion promoter, was applied via spin-coating (4000 rpm, 30 s) and baked on a hotplate at 95°C for 60 min. Shipley 1822-positive photoresist was then applied via spin-coating (4000 rpm, 30 s) to achieve a 2.4-μm-thick coating, and baked in an oven at 92°C for 30 min. An MA-6 Photo Aligner (Suss Microtech, Inc.) applied hard-contact UV exposure for 9 s. Wafers underwent a 55 s developing step in Microposit MF319 (Microchem Corp.), and then were rinsed in diH2O and spun dry. Wafers were then cleaned briefly in a March PX-250 Plasma Asher (March Plasma Systems) at 55 W for 5 min, followed by an oven hard-bake step at 110°C for 30 min. Exposed silicon was etched in a Surface Technology Systems Inductively Coupled Plasma Etcher (STS-ICP), targeting 50-, 100-, and 150-μm depths. Photoresist was stripped by oxygen plasma (100 W, 5 min), followed by SVC-14 photoresist stripper (Rohn & Haas) for 20 min. Wafers were cleaned in a piranha bath (1:1 H2SO4:H2O2, 20 min), rinsed, and spun-dry. To improve release from the mold by reducing sidewall roughness and sharp corners, wafers with 100- and 150-μm etch depth were subjected to a 10-20-min isotropic silicon etch in an STS-Reactive Ion Etch (STS RIE). After oxygen plasma treatment (100 W, 20 min), another piranha cleaning was performed, after which the wafers were stored until use. Clean wafers were spin-coated with maltose (70% in dd-H2O) in a two-step coating process (1000 rpm, 10 s; 3000 rpm, 30 s) on a CEE100 spinner (Brewer...
before use in cell culture studies. To prepare two-layered (2L) scaffolds, they were soaked in a serum-containing culture medium for 5 to 8 days. Scaffolds were air-dried for use in characterization experiments.

3D ELASTOMERIC SCAFFOLDS WITH CARDIAC-MIMETIC PROPERTIES

PGS synthesis and scaffold fabrication

PGS prepolymer was synthesized as previously described. Briefly, prepolymer was synthesized by reacting glycerol and sebacic acid in a 1:1 molar ratio under heat and vacuum for 72 h. For curing, prepolymer was dissolved in 200 Proof ethanol at 50% (w/v), and cast volumetrically into maltose-coated silicon molds fitted with poly(dimethyl siloxane) (PDMS) gaskets. Volumes were adjusted to produce either open-through pores that extended from top surface to bottom surface or closed pores bounded by a thin PGS film. Wafers were heated to 110°C for 30 min to allow ethanol to evaporate, and then the prepolymer was cured under vacuum (10–15 mTorr) at 160°C for 6 h. Scaffolds and nonporous membranes were demolded by soaking in d1-H2O at 60°C–70°C for 1 to 3 days to dissolve the maltose sacrificial layer. Scaffolds were air-dried for use in characterization experiments or autoclave-sterilized (121°C, 30 min) and then soaked in a serum-containing culture medium for 5 to 8 days before use in cell culture studies. To prepare two-layered (2L) scaffolds, two one-layered (1L) scaffolds were stacked and bonded as previously described, with the top layer containing through pores and the bottom layer bounded by a thin PGS film on the lower surface. The long axes of the pores were aligned and offset by half the pore length to yield stacked scaffolds exhibiting two layers of interconnected pores. Manual alignment of the scaffolds was challenging due to the small size of the pores and deformation of the elastomeric PGS, especially evident with the 5:1 cubic rectangular (CR) scaffolds. To achieve bonding, interior surfaces were subjected to oxygen plasma (121°C, 30 min) and then soaked in a serum-containing culture medium for 5 to 8 days before use in cell culture studies. To prepare two-layered (2L) scaffolds, two one-layered (1L) scaffolds were stacked and bonded as previously described, with the top layer containing through pores and the bottom layer bounded by a thin PGS film on the lower surface. The long axes of the pores were aligned and offset by half the pore length to yield stacked scaffolds exhibiting two layers of interconnected pores. Manual alignment of the scaffolds was challenging due to the small size of the pores and deformation of the elastomeric PGS, especially evident with the 5:1 cubic rectangular (CR) scaffolds. To achieve bonding, interior surfaces were subjected to oxygen plasma (121°C, 30 min), aligned by eye under a stereomicroscope (Nikon SMZ1500; Nikon), and compressed under a weight (200 g) for 16 h at room temperature. To prepare 2L scaffolds with a PGS thin film across the bottom for cell retention, a 1L scaffold with film was placed film side down, treated with oxygen plasma along with a 1L scaffold with through pores, and the plasma-treated sides were bonded together using the method described. 2L scaffolds were examined dry to assess the successful layer-to-layer alignment, or autoclaved and prepared for cell culture as described for 1L scaffolds.

Scaffold characterization: scanning electron microscopy and mechanical testing

Samples were prepared for scanning electron microscopy (SEM) by sputter coating with gold–palladium (108auto sputter coater; Cressington Scientific Instruments) for 30 s. Samples were then mounted either flat, or at 45 degrees, within a Hitachi S3500 SEM (Hitachi High Technologies America), and imaged using 5-kV accelerating voltage. Measurements of scaffold dimensions were made using the measure tool in ImageJ. For uniaxial tensile mechanical testing, standardized specimens were prepared using a dog-bone punch (4 mm × 2 mm), and thicknesses were measured with a spring-loaded dial gauge (LS Starrett Co). Samples were cut from dry, 1L scaffolds with through pores, fitted with 5×5-mm paper apertures to set gauge length and assist in holding samples in mounts without tearing, and mounted on an Electroforce ELF 3200 mechanical tester (Bose) fitted with a 250-g load cell (Sensotech, Inc.), and controlled with WinTest software. Samples were strained to failure using a ramp speed of 0.1 mm/s (2% strain/s). Orthogonal material directions were tested with independent samples: parallel (PD) or perpendicular (XD) to the long axis of the pore structure. For each test direction, effective stiffnesses (EPD and EXD) were calculated from the slope of the stress–strain curve between 1% and 10% strain, and anisotropy, r, was calculated as the EPD/EXD ratio. Strain-to-failure (σfPD and σfXD) data were also collected from the stress–strain curve.

C2C12 cell culture

To examine the capacity of the three patterns to promote cell elongation and interconnectivity, C2C12 myoblasts were cultured on scaffolds for 5 days. C2C12 myoblasts elongate readily in culture, making this a good cellular model for preliminary studies of the effect of scaffold architecture on elongation. C2C12 cells were obtained from the American Type Culture Collection (ATCC) and maintained in a complete culture medium consisting of the Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and 1% penicillin/streptomycin (pen/strep). 1L scaffolds with through pores were punched to 5-mm-diameter discs and preincubated in a culture medium as described, dried gently by blotting with sterile gauze, and placed in sterile 12-well dishes with one scaffold per well. Cells were seeded onto scaffolds (50,000 cells/30 μL culture medium per scaffold) and allowed to attach for 45 min. Then, an additional 2 mL of medium was added, and constructs were cultured for 5 days. Media were completely replaced every 2 days, and constructs were harvested on culture day 5.

Heart cell isolation and culture

Heart cells were isolated from neonatal rats (aged 1 to 2 days) following an Institutional Animal Care and Use Committee-approved protocol, as previously described. Briefly, hearts were harvested, and ventricular tissue was isolated, minced, and digested overnight in a trypsin solution (1 mg/mL; Sigma) at 4°C. After trypsin digestion, cells were serially digested using type II collagenase solution (1 mg/mL; Worthington Biochemical Corp.) until dissociated into a single cell-suspension. Dissociated cells were preplated for 1 h in tissue culture flasks to enrich for cardiomyocytes, and cells that remained unattached to the flask were used to seed scaffolds. This preplating method has been characterized to yield a cell population containing 63%±2% cardiomyocytes, 33%±3% cardiac fibroblasts, 3% to 4% smooth muscle cells, and 2% to 3% endothelial cells, and has previously been successful in our hands at producing viable, contractile cultures. For cell seeding, 2L scaffolds bounded by a thin PGS film were punched into 5-mm discs, preincubated in a culture medium as described, dried gently by blotting, and placed within PDMS gaskets (6- to 7-mm inner diameter; 1- to 2-mm high) in 12-well tissue culture plates. Heart cells were seeded onto scaffolds (3.5 million cells/50 μL of culture medium per scaffold). Immediately thereafter, an additional
2 mL of medium was added slowly to avoid disturbing the heart cells within the gasket, and constructs were placed in a standard incubator. Gaskets were left in place for at least 24 h, to allow time for gravitational cell settling into the pores and initial cell binding to the scaffolds; gaskets were removed after 24 to 48 h to mitigate adverse effects on diffusion of a culture medium through the radial surfaces of the constructs. Media were completely replaced every 2 days, and constructs were harvested after 1 week.

**Electrophysiology**

Response to electrical field stimulation was assessed for 1-week cardiac constructs as previously described. Briefly, constructs were placed in a custom-designed, environmentally controlled test chamber. The chamber was fitted with two carbon electrodes (1/4-inch diameter; Ladd Research) separated by a distance of 1.5 cm. Constructs were intentionally positioned within the test chamber such that the scaffold PD was aligned in parallel to the stimulating electrodes and covered with 10 mL of 37°C culture medium. Platinum wire leads connected the electrodes to an electrical pulse generator (S88 × Dual Output Square Pulse Stimulator; Grass Technologies), which was used to apply monophasic square pulses with 2-ms duration and 1-Hz frequency. Initial voltage was set at 1 V and ramped in 0.5 V increments until each stimulus solicited a matching contraction in the construct. This voltage was recorded as the excitation threshold (ET). Contractions were observed and video imaged on a Zeiss Axiovert inverted microscope (Zeiss), at 10 × magnification.

**Histology**

After electrophysiological testing, and for samples not used for electrophysiology, constructs were rinsed gently with phosphate-buffered saline (PBS), fixed in 10% neutral buffered formalin overnight at 4°C, rinsed several times in PBS, and stored in fresh PBS at 4°C in preparation for histological assessments.

To assess cardiomyocyte presence and marker expression, constructs seeded with heart cells were permeabilized with 0.1% Triton X-100, blocked with 0.1% bovine serum albumin (BSA), stained with sarcomeric α-actinin primary antibody overnight (1:500; Sigma), followed by fluorescein-conjugated anti-mouse IgG secondary antibody (2 h, 1:200; Vector Lab), counterstained with DRAQ5 nuclear stain (1:200; Axxora LLC), and examined using an NIS-E Ultra-Fast Spectral Scanning Laser Confocal Microscope with NIS Elements Acquisition Software (Nikon). To assess muscle cell elongation and interconnectivity, constructs seeded with C2C12 cells were permeabilized and blocked as described, then stained for filamentous actin (f-actin) using fluorescein isothiocyanate (FITC)–phallolidin conjugate (1:40; Sigma), counterstained with DRAQ5 nuclear stain, and examined using the confocal microscope. To assess cell distribution within constructs seeded with heart cells, constructs were bisected, embedded in Tissue Tek O.C.T Compound (Sakura Finetek), and frozen for cross sectioning on a Thermo Shandon Cryotome SME (Thermo Scientific). Thirty-μm-thick sections were cut, stained with hematoxylin and eosin (H&E; Sigma), and imaged on a Zeiss Axiovert inverted microscope (Zeiss) to evaluate cellular infiltration into the scaffold.

**FE modeling**

Extending our recently described method, an FE periodic model was used to predict the effective stiffnesses and anisotropy ratio of scaffolds with varying pore pattern, pore length, and strut width. The FE model was developed using Z-set software (ZSet Resources, Mines ParisTech, ONERA, NWNumerics, http://zebulon.onera.fr/English/index.html). Briefly, scaffolds were designed as periodic tessellations defined by the translation of a unit cell along vectors of periodicity. The unit cell comprised void space in the shape of the pore surrounded by PGS with width equal to one-half the scaffold strut width, with axes of symmetry along the PD and XD directions. Periodic, homogeneous boundary conditions were prescribed on the unit cell to compute the effective stiffnesses of the heterogeneous structure (i.e., structure comprising PGS struts and void pore space). For the PGS phase, the elastic modulus ($E_{PGS}$) was assigned based on mechanical test data, assuming that the PGS material exhibited isotropic linear elastic behavior for strain < 1%. For the void space, an elastic modulus near zero (0.0001 MPa) was assigned to obtain a condition number amenable to stiffness matrix inversion during the FE code execution; for both phases, a Poisson’s ratio equal to 0.45 was assumed. The FE model was also used to model degradation of the scaffold. Since PGS degrades by surface hydrolysis, strut width is expected to decrease while the pore area is expected to increase concomitantly over time. To ascertain the effects of this surface hydrolysis-dominated degradation on $E_{PD}$, $E_{XD}$, and $r$, the model was applied over a decreasing range of strut widths with concomitant increase in pore area and a constant PGS-effective stiffness; PGS bulk hydrolysis was assumed to be negligible. The FE model was also used to simulate macroscopic loading of the scaffold, as previously described.

**Statistical analysis**

For mechanical testing, statistical significance was assessed in Minitab 15 Statistical Software (Minitab Inc.) using a general linear model analysis of variance (ANOVA) with two fixed, crossed factors (pore pattern and test direction) with Tukey’s post hoc testing to establish among- and between-group significance. Significance was asserted at $p < 0.05$. Statistical differences in ET were assessed using Minitab 15 Statistical Software with a one-way ANOVA and Tukey’s post hoc testing with significance asserted at $p < 0.05$.

**Results**

**Scaffold design**

Two pore patterns were considered, CR and hexagonal rectangular (HR), indicating four or six first-neighbor pores respectively (Fig. 1). FE modeling of the tensile mechanical properties of two patterns (CR and HR), two aspect ratios (2:1 and 5:1 pore length: pore width, L/l), and two strut widths ($w$, 30 and 50 μm) provided a blueprint for selecting three scaffold designs for fabrication (Fig. 1) from a total of eight possible combinations of pore pattern, pore aspect ratio, and strut width (Table 1). With an ultimate goal of both delivering cells and providing mechanical support to the healing heart, our scaffold design goals were to meet dual objectives of microfabricating structural features approaching tissue scale and matching scaffold mechanical properties.
to physiological values for elastomeric stiffness and anisotropy, properties that depend on cardiac structure and strongly influence pumping function.

As a starting point to select target scaffold structural features, we modeled pore inner dimensions \( L \), \( l \) and minimum strut widths \( w \) deemed to be technically achievable. Rectangular pore shape was selected based on empirical evidence that this pore shape was mechanically anisotropic and was capable of aligning fibroblasts and their collagenous matrix and muscle cells. For modeling purposes, the bulk PGS stiffness, \( E_{\text{PGS}} \), was prescribed as \( 791 \pm 58 \text{ kPa} \) (mean \( \pm \text{SEM} \), \( n = 6 \)), as empirically measured for nonporous, 112-21-mm-thick PGS membranes cured at 10 to 15 mTorr and 160°C for 6 h. FE simulations of PGS scaffold mechanical properties were run with minimum, maximum, and mean values of \( E_{\text{PGS}} \) to predict scaffold-effective stiffnesses in two orthogonal directions, \( E_{\text{PD}} \) and \( E_{\text{XD}} \). These values were used to predict anisotropy, \( r = E_{\text{PD}}/E_{\text{XD}} \). For selection of target scaffold mechanical properties, we used ranges measured for explanted rectangular specimens of normal adult rat left ventricular myocardium (LV) harvested in two orthogonal directions (circumferential and longitudinal). We reasoned scaffold mechanical properties (\( E_{\text{PD}}, E_{\text{XD}}, \) and \( r \)) should recapitulate LV stiffnesses in the two orthogonal directions (157 \( \pm \) 43 kPa and 84 \( \pm \) 23 kPa, mean \( \pm \text{SD} \) of \( n = 9 \) specimens) and the corresponding LV anisotropy ratio \( (r = 1.87) \). Of note, this anisotropy ratio offers only a partial measure of anisotropy of the scaffold, considering only the directions relevant to direct comparison between the scaffold and LV.

Three scaffold designs (Fig. 1) were selected for fabrication, 2:1 cubic rectangular (2:1 CR), 5:1 cubic rectangular (5:1 CR), and 2:1 hexagonal rectangular (2:1 HR). The cubic and hexagonal pore patterns, respectively, refer to the existence of four or six first-neighbor pores. The 2:1 and 5:1 aspect ratio pores refer to the ratio of long-pore length:short-pore length, \( L/l \), while \( w \) refers to the strut width. The three scaffold designs were chosen for best comparison between structural features and matching of effective stiffnesses predicted by the finite-element (FE) model to those previously measured for normal adult rat left ventricular (LV) myocardium.

**Table 1. Mechanical Properties Were Predicted for Candidate Scaffold Designs**

<table>
<thead>
<tr>
<th>Scaffold Design</th>
<th>Predicted Mechanical Properties</th>
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<tr>
<td>Symmetry</td>
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<tr>
<td>Cubic</td>
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Of eight possible combinations of two pore patterns (cubic and hexagonal), two pore aspect ratios (2:1 and 5:1) and two strut widths (50 and 30 \( \mu \text{m} \)), three scaffold designs (shaded and bold-face) were selected based on best comparison between structural features and matching of predicted mechanical properties \( (E_{\text{PD}}, E_{\text{XD}}, \text{and} r) \) to those of normal adult rat left ventricular (LV) myocardium.
50-μm struts was selected (Table 1, row 5), while the other candidates were discounted for anisotropy ratios far higher than the physiological range (Table 1). Comparing 2:1 CR and 2:1 HR, $E_{PD}$ was decreased, and $r$ was increased by more than four-fold, but $E_{PD}$, and the feature size ($L, l, w$) were maintained constant (Fig. 1, column 2 vs. 4).

**Mold and scaffold fabrication**

The chosen designs, 2:1 CR, 5:1 CR, and 2:1 HR, were etched into silicon molds to allow for easily reproducible micromolding, as the same mold may be used indefinitely with appropriate care and cleaning. Pore and strut dimensions of the molds for 2:1 CR and 2:1 HR were identical ($L = 250 \mu m$, $l = 125 \mu m$, $w = 50 \mu m$), while for 5:1 CR, the pore and strut dimensions were varied ($L = 250 \mu m$, $l = 50 \mu m$, $w = 30 \mu m$). Wafer sets were etched to three different depths (50, 100, and 150 μm), to extend the range of strut width to strut height that could be reproducibly fabricated.

As compared to previous PGS micromolding efforts, three methodological improvements were made. First, wafer fabrication included a final RIE step to smooth sharp corners in the molds etched to 100 and 150 μm to facilitate demolding of the corresponding PGS scaffolds. Second, a maltose-coating method developed by Wang et al. replaced sucrose as the sacrificial layer to reduce concerns of crystallization leading to unintended microarchitecture on the scaffold surfaces. Third, the PGS pre-polymer was dissolved in ethanol before casting to provide more reproducible control of scaffold thickness.

Large 20×20 mm PGS scaffolds were cast, cured, and demolded from molds with all three scaffold designs and three thicknesses of 50, 100, and 150 μm (Fig. 2). Scaffolds faithfully matched the intended design and retained consistent dimensions after multiple uses of each mold ($L = 126.4 \pm 1.6$, $56.5 \pm 1.2$, and $123.5 \pm 2.4 \mu m$; $L = 250.9 \pm 1.3$, $247.0 \pm 1.1$, and $246.5 \pm 2.5 \mu m$; and $w = 41.2 \pm 2.8$, $22.7 \pm 2.5$, and $46.1 \pm 1.0 \mu m$ for 2:1 CR, 5:1 CR, and 2:1 HR molds etched to a depth of 100 μm, respectively). Scaffolds for heart cell culture were fabricated with an ~50-μm-thick PGS film on one surface to assist in cell retention during seeding and culture (Fig. 2H, K). The 100-μm layer height was selected for subsequent studies of scaffold mechanical properties and cell compatibility demonstration, since these 1L scaffolds were thick enough to allow for handling, but thin enough to minimize diffusion limitations.

**Scaffold mechanical properties**

Average effective stiffnesses of PGS scaffolds tested dry ranged from 273 to 293 kPa for $E_{PD}$ and 85 to 179 kPa for $E_{XD}$, while lower values were obtained for PGS scaffolds tested after 1 week of immersion in 37°C PBS (131 to 206 kPa for $E_{PD}$ and 37 to 138 kPa for $E_{XD}$). When compared with stiffness values predicted by the FE model, experimentally measured stiffnesses obtained for PGS scaffolds matched closely, and trends were maintained (Fig. 3A). Specifically, model predictions of $E_{PD}$ were 11.0%, 33.8%, and 3.98% greater than empirical values and for 2:1 CR, 5:1 CR, and 2:1 HR, respectively. Predictions of $E_{XD}$ were 1.06% greater, and

![FIG. 2. One-layered (1L) poly(glycerol sebacate) (PGS) scaffolds with precisely defined, anisotropic pores were rapidly and reproducibly micromolded from silicon masters. Representative light micrographs (A–C) of etched silicon molds demonstrating three scaffold designs: 2:1 CR (A, D, G, H, I), 5:1 CR (B, E, J, K, L), and 2:1 HR (C, F). Scanning electron micrographs of PGS scaffolds delaminated from silicon molds illustrate successful pattern replication (D–F) and full-thickness scaffolds from 50-μm deep (G, J), 100-μm deep (H, K), and 150-μm deep molds (I, L), viewed from above (D–F) or 45 degrees (G–L). Scaffolds were fabricated either with through pores (G, J, I, L), or with an ~50-μm-thick PGS film at the base for enhanced cell retention (H, K), indicated by black arrowheads. Scale bars: 200 μm.](image-url)
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ANOVA with Tukey’s post hoc testing.

Empirical data represent the mean ± standard error of at least three independent measurements. * indicates significant difference between corresponding measurements (i.e., EpD vs. ExD, p < 0.05, and EpD vs. ExD, p = 0.011), as determined by ANOVA with Tukey’s post hoc testing.

27.1% and 53.2% lesser than empirical values for the same patterns. As predicted, anisotropy was demonstrated with a significant difference between experimental EpD and ExD as a whole (all scaffold EpD compared to all scaffold ExD, p < 0.001) and for both the 2:1 CR (p < 0.007) and 2:1 HR (p = 0.021) groups individually. Scaffold pattern alone did not yield significant differences in EpD or ExD across the groups (i.e., ExD measured for 2:1 CR was not significantly different from ExD measured for 5:1 CR). Experimental anisotropy ratios (r = 1.5, 1.8, and 3.4 for 2:1 CR, 5:1 CR, and 2:1 HR, respectively) matched the trends and values predicted by the model (r = 1.7, 3.4, 7.5, respectively). As compared to values previously measured for the LV myocardium, scaffold EpD was higher than physiologic in all three groups; scaffold ExD was higher in the two CR groups and physiologic in the HR group; and scaffold r was lower than physiologic in the 2:1 CR group, physiologic in the 5:1 CR group, and higher than physiologic in the 2:1 HR group. Strain to failure (σf) of all three scaffold patterns was greater than one and showed a significant difference between experimental σf in the PD and XD directions (p = 0.0014) (Fig. 3B). No significant differences were found among groups for σfEpD or σfExD. As compared to previously measured rat LV data (σfEpD = 0.9 ± 0.1; σfExD = 1.1 ± 0.2),26 scaffold σfEpD and scaffold σfExD were either similar or slightly higher for all three groups.

The FE model predicted that all the three scaffold designs would exhibit gradual decreases in EpD and ExD as a function of simulated degradation-mediated decreases in PGS strut width (Fig. 4A–C). Moreover, the model predicted that the anisotropy ratio would increase markedly during degradation of the HR design, particularly as the strut width decreased below 40 μm (Fig. 4C), but would be maintained relatively constant, although with slight increases, during degradation of the two CR designs. Simulations further suggested that the 2:1 HR scaffold would respond to mechanical loading in the XD direction principally by flexural deformation of struts oriented in the PD direction (Fig. 4E), while the two CR scaffolds would instead respond to mechanical loading by tensile extension of struts oriented in the XD direction (2:1 CR; Fig. 4D).

C2C12 cells on one-layer scaffolds

C2C12 cells displayed distinct behaviors when cultured on each of the three pore arrangements (Fig. 5). Cell elongation along the PD-axis was qualitatively visible in the 5:1 CR design, at the top surface of the struts and within the pores (Fig. 5B, E, H), while elongation along the XD-axis was relatively more pronounced in the 2:1 CR design (Fig. 5A, D, G) and elongation along both PD- and XD-axes as well as diagonally between pores was observed in the 2:1 HR design (Fig. 5C, F, I). The direction of cell elongation appeared to correlate with cellular bridging across struts and connecting with cells in neighboring pores.

Heart cells on two-layer scaffolds

Replica-molded 1L scaffolds were assembled into 2L scaffolds with an underlying thin PGS film by manual stacking with offset pores. The 2L scaffolds exhibited combined thickness of the porous region of the scaffolds measuring ~200-μm thick, twice the thickness of 1L scaffolds of the same pattern, while the underlying PGS film was ~50-μm thick (Fig. 6).

Heart cells began to contract spontaneously after 3 days in culture and continued to do so throughout the 1-week study. After 1 week, scaffolds were subjected to electrical field stimulation to determine the ET, which describes the stimulating voltage required to capture the resultant contraction. Constructs based on all scaffold designs contracted in response to electrical field stimulation (Fig. 7). Although the 5:1 CR design showed lower mean ET than the 2:1 CR design, the difference was not significant (p > 0.05); however, ET was significantly lower for the 2:1 HR design than for the 2:1 CR design (p = 0.042).
Cross sections of fixed 1-week samples showed heart cells seeded throughout the full depth of 2L scaffolds of all three scaffold designs (Fig. 8A–C), suggesting that the static seeding method used was sufficient for cells to reach the porous areas of at least 2L scaffolds with interconnected pores up to 200-μm deep. The 5:1 CR pattern qualitatively yielded the thickest layer of cells across the top strut surfaces, possibly due to the smaller pore size and greater difficulties in manual layer alignment, preventing some pore filling (Fig. 8B inset). Because the heart cell isolation retained cardiac fibroblasts along with the cardiomyocytes, sarcomeric α-actinin was used as a specific marker to verify the presence of cardiomyocytes after 1 week in culture. In all three designs, α-actinin was prevalent (Fig. 8D–F), and there was some degree of sarcomeric banding (Fig. 8G–I, arrowheads), consistent with the observation of coordinated contractions. While the neonatal heart cells remained predominantly immature and rounded at 1 week, positive immunostaining for α-actinin provided evidence of early-stage cardiomyocyte differentiation, and functional responses to electrical stimulation provided evidence of heart cell viability, interconnectivity, and contractility.

**Discussion**

In this study, we designed, developed, and demonstrated a set of 2L elastomeric PGS scaffolds with offset and fully interconnected pores that provided a platform for cell delivery while recapitulating the geometry and mechanical properties of normal heart muscle.

Utilizing a structured engineering design plan allowed interrogation of scaffold mechanical properties in a manner directly relevant to the objective of designing scaffolds with biomimetic mechanical properties; others48,66 have used a similar approach to scaffold design. Specifically, we developed an FE model describing periodic, tessellated scaffolds with rectangular pores, for probing changes in scaffold stiffness and anisotropy while varying three factors: pore pattern (CR and HR), rectangular aspect ratio (2:1 and 5:1), and strut width (50 μm and 30 μm). This computational approach helped pare down the scaffolds for fabrication from eight candidates to three cardiac-mimetic options, thereby increasing efficiency and yield while reducing time and expense. While our previous work showed that FE modeling could retrospectively predict empirically determined scaffold mechanical properties,52 including the influence of tissue filling the pores,35 the present work extended use of FE modeling to intentionally choosing scaffolds that provide the desired architecture, stiffness, and anisotropy, for fabrication and demonstration in a 3D in vitro model of engineered cardiac tissue. As metrics, we selected (1) mechanical matching of the scaffolds to normal rat myocardium (Table 1; Figs. 1, 3 and 4), (2) structural evidence of cell-to-cell connectivity in 1L constructs (Fig. 5), (3) contractile response to electric stimulation in 2L constructs (Fig. 7), (4) heart cell delivery while recapitulating the geometry and mechanical properties of normal heart muscle.
penetration and seeding throughout 2L scaffolds (Fig. 8), and (5) maintenance of neonatal heart cell phenotype in 2L constructs (Fig. 8).

An interest in replicating tissue-scale architecture motivated the choice of the two CR designs. Moving from 2:1 CR to 5:1 CR, $l$ and $w$ were intentionally reduced toward the tissue scale. As such, comparison of 2:1 vs. 5:1 CR designs offered a potential ability to tease out differences due to scaffold structural features while maintaining scaffold mechanical properties relatively close to each other and to those

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**FIG. 5.** Scaffold design influenced elongation and interconnectivity of C2C12 cells. Schematic diagrams (A–C) depict effects of scaffold feature size and pore pattern on C2C12 cell elongation and interconnectivity, where black lines indicate predominant direction of cell elongation with respect to a central pore. Note that similar effects were seen for each pore, but for clarity were shown only for a central pore. Confocal micrographs of 5-day constructs stained for filamentous actin using FITC-phalloidin and for nuclei using DRAQ5, showing cells on the surface of the uppermost strut (D–F) and within the pores of 2:1 CR (D, G), 5:1 CR (E, H), and 2:1 HR (F, I) scaffold designs. In the 5:1 CR design, cell guidance in the PD was provided by 30-μm struts (E) and 250 × 50 μm pores (H), while in the 2:1 HR design, cell-to-cell connections between pores were observed diagonally as well as in the PD and XD directions (F). Scale bars: 20 μm. Color images available online at www.liebertpub.com/tea

**FIG. 6.** Two-layer (2L) scaffolds with three-dimensional (3D) pore microarchitecture were fabricated by offset stacking of 1L scaffolds. Representative scanning electron micrographs of three PGS scaffold designs: 2:1 CR (A, D), 5:1 CR (B, E), and 2:1 HR (C, F), viewed from above (A–C) or 45 degrees (D–F), show offset stacking along the long pore axis. Scale bars: 200 μm.
of LV. An interest in studying effects of scaffold mechanical properties motivated the comparison of 2:1 CR and 2:1 HR designs. Moving from 2:1 CR to 2:1 HR reduced EXD and increased r, offering a potential ability to discover effects due to changes in pore pattern and certain scaffold mechanical properties, since feature sizes were maintained.

New micromolding methods were established for rapid, precise, reproducible production of PGS scaffolds with high aspect ratio of strut height: strut width (5:1) and thin (30 μm) strut width. The silicon-based replica molding method developed and optimized for PGS scaffold production was more precise and high throughput than our previously described laser microablation method.11,20,51 Specifically, fabrication by micromolding yielded more precise in-plane scaffold features than could be achieved by current laser microablation techniques, in part due to negligible pore tapering from top to bottom of 150-μm-thick scaffolds. Micromolding allowed strut widths of 30 μm that were not achievable with laser microablation, due to depth-dependent changes in laser intensity. Additionally, minimal pore tapering assisted in maintaining consistent mechanical properties and pore geometry, and enabled a closer match with the numerical predictions on idealized structures. The addition of an RIE step at the end of the silicon mold fabrication process smoothed both the corners and walls of the etched silicon, allowing for more consistent demolding of the PGS scaffolds, which in turn provided the opportunity to demold larger areas and to fabricate higher aspect ratios of feature height versus feature width than previously achieved.51,58,64,65 Selecting maltose as the sacrificial layer to coat etched silicon wafers enabled more reliable demolding of PGS scaffolds than a sacrificial layer of sucrose, which was impaired by uncontrollable sucrose crystallization (a result of moisture and humidity or exposure to high temperature). Finally, volumetric dispensing of PGS dissolved in ethanol provided more reliable control over mold filling, scaffold thickness, and thin-film thickness as compared to the previously used method of PGS prepolymer melt casting.51

The ability to demold larger areas of scaffolds than are required for in vitro studies is critical as we consider scale-up

FIG. 7. Scaffold design affected the contractility of heart cells. Excitation threshold (ET, V) shown for heart cells cultured on 2L scaffolds for 1 week. Data are mean ± standard error of at least five independent samples. Significance was asserted at p < 0.05 (indicated by *) using a one-way analysis of variance (ANOVA) with Tukey’s post hoc testing.

FIG. 8. Heart cells distributed throughout depth of 2L constructs and expressed a cardiac-specific marker. Full-construct cross-sections, cut to a thickness of 60 μm and stained with hematoxylin and eosin (* indicates polymer), show cells throughout the depth of scaffolds of all three designs (A–C, cut in parallel to the long-pore axis; and B, inset, cut in parallel to the short-pore axis). Cells formed a multicellular layer across top struts of 5:1 CR 2L scaffold (B, inset). Representative 3D renderings of 50-μm-thick stacks taken on a confocal microscope after staining with α-actinin (arrowheads indicate sarcomeric banding) indicate cardiomyocyte phenotype in all three scaffold designs: 2:1 CR (D, G), 5:1 CR (H), and 2:1 HR (F, I). Scale bars: 100 μm (A–C), 50 μm (D–F), and 10 μm (G–I). Color images available online at www.liebertpub.com/tea
of scaffold fabrication toward anatomical dimensions for future clinical applications. Micromolding of PGS scaffolds yielded very consistent scaffold dimensions and pore shapes over relatively short time periods. The molds used in this study were 20 × 20 mm, and more than 20 molds could be cast and cured simultaneously, resulting in much greater and faster yield than laser microablation could produce. In addition, molds could be cleaned and reused, facilitating consistent scaffold geometry. While PGS scaffolds fabricated in the present study were purposefully restricted to planar dimensions of 20 × 20 mm, suitable for in vitro cell culture and mechanical testing, the ability to demold larger areas of scaffolds is critical. One advantage of our replica molding approach is that maximum planar dimensions are limited only by the dimensions of the silicon substrate and mask aligner. In the current study, we used a mask aligner capable of accommodating four- or six-inch-diameter silicon wafers; even larger (eight or twelve inch) wafers are routinely used in semiconductor fabrication. Thickness limitations are also a consideration for scale-up to human myocardial applications. Our design for multilaminar scaffolds was developed to overcome these limitations, and ongoing work will further enhance our capabilities of generating thick cardiac grafts. Importantly, engineered cardiac constructs with thicknesses on the length scale of native myocardium will require microvascular-like networks to sustain the viability of highly metabolically active heart cells. As such, recent studies have explored a variety of biomaterials and techniques to produce prevascularized grafts with promising results.

The aspect ratio of molded feature height versus feature width for scaffolds was as high as 5:1 (150-µm strut height; 30-µm strut width), which exceeded previous reports for replica-molded PGS scaffolds in which the feature height-versus-width ratios were 2:1 or lower. High-aspect ratio struts in combination with stacking of multiple layers may help to achieve tissue-scale thicknesses for tissue engineering applications, as many more layers would otherwise be needed to achieve the same overall thickness, increasing the chances of misalignment or pore closure resulting from multiple rounds of stacking and lamination. We produced 2L scaffolds with a fully interconnected pore network by a manual stacking process as previously described, but this was difficult due to the flexibility and deformability of PGS scaffolds with cardiomimetic mechanical properties. In ongoing work, we are exploring more rigorous stacking methodologies to improve pore and strut alignment in multilayered scaffolds.

In consideration of our objective to match scaffold stiffness, anisotropy, and strain-to-failure to cardiac-mimetic values, we undertook uniaxial tensile testing of the fabricated scaffolds. Although $E_{PD}$ was higher than physiologic in all three scaffold designs, and $E_{XD}$ higher in both CR designs, we previously showed decreases in accordant-like honeycomb scaffold $E_{PD}$ and $E_{XD}$ both with cyclic mechanical fatigue and with in vitro culture of cell hearts. It is reasonable to assume that our rectangular pore scaffolds will show qualitatively similar responses to fatigue testing or cell culture as our accordant-like honeycomb scaffolds, resulting in stiffnesses decreasing into a more physiologic range.

Tensile mechanical testing of scaffolds in orthogonal directions justified FE model predictions as a means to select only candidate scaffolds with desired mechanical properties for fabrication. Although FE model predictions overestimated actual values in the PD direction and underestimated empirical values in the XD direction, model trends were consistent. Possible explanations for the minor differences include error during experimental testing, particularly for the 2:1 HR pattern tested in the XD direction, and differences between stiffnesses of PGS scaffolds and stiffnesses of nonporous PGS membranes utilized in our FE simulations, which could in turn be due to potential influences of the mold on local temperatures during curing. Despite the minor discrepancy, we considered the model validated for use with the scaffold pore designs we chose.

The FE model was then used to predict how PGS degradation and loading would be expected to impact scaffold stiffness and anisotropy. All three scaffold designs exhibited decreases in $E_{PD}$ and $E_{XD}$ as a function of decreasing strut width and increasing pore area, which were covaried to approximate the degradation by surface hydrolysis that occurs in vitro and in vivo. Gradual reduction in scaffold stiffness during scaffold degradation is a primary objective of tissue engineering applications where it is envisioned that regeneration of a mechanically functional, fully biological tissue can gradually replace the mechanical support temporarily contributed by a biodegradable scaffold.

The marked increase in anisotropy predicted during simulated degradation of the HR scaffold can be attributed to its architecture, comprising discontinuous struts in the XD direction and continuous struts in the PD direction. In particular, while CR scaffolds were predicted to respond to XD loading by tensile extension of XD-oriented struts (Fig. 4D), HR scaffolds were predicted to respond to XD loading primarily by flexural deformation (i.e., bending) of PD-oriented struts (Fig. 4E). These distinct modes of deformation exhibited by the HR-versus-CR scaffolds can account for their distinct changes in effective stiffness predicted in association with simulated degradation. Flexural deformation of the PD-oriented struts is theorized to contribute nonlinearly to the effective stiffnesses and the associated anisotropy ratio of the HR scaffold, due to the cubic (i.e., third power) dependence of the strut segment’s cross-sectional area moment of inertia on the strut width. By contrast, tensile deformation of the XD-oriented struts is theorized to contribute only linearly to the effective stiffnesses and associated anisotropy ratio of the CR scaffolds, due to the linear dependence of the strut segment cross-sectional area on the strut width. Experimental studies of how these two scaffold designs will respond to cyclic fatigue is the subject of ongoing work, not only in vitro but also in vivo where PGS degradation has been shown to occur even more quickly.

In consideration of our objective to provide scaffold pore patterns that enable the formation of a syncytium of muscle cells, we examined C2C12 cells, which have a propensity to form connections with one another and align along scaffold microstructures. C2C12 cells formed connections with cells in adjacent pores by proliferating and elongating across intervening struts in all three scaffolds in a design-dependent manner (Fig. 5). Taking scaffold feature size into account, we compared 2:1 CR and 5:1 CR designs, with the latter presenting architectural features closer to tissue scale. The 2:1 CR pores directed cell elongation and interconnectivity in both PD and XD directions, while the 5:1 CR pores
encouraged relatively more elongation and interconnectivity in the PD direction, possibly due to the smaller pore dimensions and strut widths. Regarding effects of pore pattern and mechanical properties, we compared 2:1 CR and 2:1 HR designs, with the latter providing a higher number of neighboring pores, lower $E_{XD}$ and higher $r$. The HR pores promoted C2C12 cell elongation and interconnectivity in multiple directions, including PD, XD, and even diagonally, suggesting that this elongation outward from the center of the pore was likely directed by the presence of six neighboring pores and cell-to-cell connections formed among them.

We also examined neonatal rat heart cells, which have a propensity to form a syncytium characterized by coordinated, synchronous contraction. Heart cells were cultured on 2L scaffolds bounded by a thin PGS film, thereby increasing the capacity for cell retention and delivery and building scaffold thickness toward the eventual goal of reconstructing full-thickness myocardial defects. ET was significantly lower in the 2:1 HR group as compared to the 2:1 CR group (Fig. 7), suggesting a dependence of construct contractile function on scaffold design. One explanation for this finding is that six neighbor pores enhanced formation of a contractile syncytium; another plausible alternative is that the lower ET in the neighbor pores enhanced formation of a contractile syncytium. Heart cells were cultured on 2L scaffolds bounded by a thin PGS film, thereby increasing the capacity for cell retention and delivery and building scaffold thickness toward the eventual goal of reconstructing full-thickness myocardial defects. ET was significantly lower in the 2:1 HR group as compared to the 2:1 CR group (Fig. 7), suggesting a dependence of construct contractile function on scaffold design. One explanation for this finding is that six neighbor pores enhanced formation of a contractile syncytium; another plausible alternative is that the lower ET in the neighbor pores enhanced formation of a contractile syncytium.

The next step for bringing this construct closer to application is validation in a preclinical infarct model. In vivo validation of our approach is being addressed in ongoing studies. One potential limitation of the current study is that target mechanical properties of the PGS scaffolds were chosen as those of normal, not infarcted, myocardium, whereas the optimal mechanical properties of a scaffold for myocardial repair remain unknown. Likewise, an in vivo study by Stuckey et al. matched PGS membrane mechanical properties to those of normal rat myocardium, and demonstrated that these PGS membranes reduced post-MI hypertrophy in a rodent model in contrast to a stiffer material. In Stuckey et al., the PGS membranes had a modulus of 300 kPa, corresponding to a membrane stiffness of 0.117 N/mm (0.67 lbs/inch). Clinical studies have shown that a cardiac support device (Acorn CorCap™ Cardiovascular, Inc.) can limit post-MI remodeling associated with deterioration of cardiac function in humans. These CorCap™ devices had an estimated membrane stiffness of 0.315 N/mm (1.8 lbs/inch), based on compliance data in a cardiac-relevant strain range of 10% to 25%; a modulus was not estimated because material thickness was not provided. In a recent in vivo study by Fomovsky et al., anisotropic reinforcement post-MI in a canine model yielded improved systolic function, while isotropic reinforcement did not yield functional improvement, consistent with the predictions of a computational model. While Fomovsky et al. studied acellular poly(ethylene terephthalate) grafts in vitro, their promising results highlight the importance of combining computational modeling and experimentation to address anisotropy in the setting of myocardial regeneration and repair, and support the utility of an inherently flexible technique for the production of anisotropic cardiac grafts.

**Conclusion**

2L PGS scaffolds designed based on FE simulations and exhibiting offset, anisotropic, and interconnected pores provided a platform for cell delivery while simultaneously recapitulating the geometry and mechanical properties of normal heart muscle. Herein, a 2L scaffold with a network of fully interconnected pores supported the in vitro development of contractile muscle constructs from neonatal rat heart cells over 1 week of culture. The modular nature of the scaffold and its layer-by-layer assembly provided the ability to extend the scaffold in three dimensions, but especially provided a method to achieve physiologically relevant tissue thickness. The FE model developed in this study may be further manipulated to optimize the beneficial qualities of the scaffolds studied herein. As such, this work not only...
demonstrated several scaffold designs that promoted neonatal rat heart cell culture and contraction in vitro, but also provided the foundation for further computational and empirical investigations of 3D elastomeric scaffolds for cardiac tissue engineering.

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Disclosure Statement

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