Mechanically patterned neuromuscular junctions-in-a-dish have improved functional maturation

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ABSTRACT Motor neuron (MN) diseases are progressive disorders resulting from degeneration of neuromuscular junctions (NMJs), which form the connection between MNs and muscle fibers. NMJ-in-a-dish models have been developed to examine human MN-associated dysfunction with disease; however such coculture models have randomly oriented myotubes with immature synapses that contract asynchronously. Mechanically patterned (MP) extracellular matrix with alternating soft and stiff stripes improves current NMJ-in-a-dish models by inducing both mouse and human myoblast durotaxis to stripes where they aligned, differentiated, and fused into patterned myotubes. Compared to conventional culture on rigid substrates or unpatterned hydrogels, MP substrates supported increased differentiation and fusion, significantly larger acetylcholine (ACh) receptor clusters, and increased expression of MuSK and Lrp4, two cell surface receptors required for NMJ formation. Robust contractions were observed when mouse myotubes were stimulated by ACh, with twitch duration and frequency most closely resembling those for mature muscle on MP substrates. Fused myotubes, when cocultured with MNs, were able to form even larger NMJs. Thus MP matrices produce more functionally active NMJs-in-a-dish, which could be used to elucidate disease pathology and facilitate drug discovery.

INTRODUCTION

Neuromuscular junctions (NMJs) are specialized synapses that form between motor neurons (MNs) and skeletal muscle fibers. On activation, presynaptic MNs release acetylcholine (ACh) into the synaptic cleft, which then binds to ACh receptors (AChRs) present on the postsynaptic muscle membrane; AChR ligation induces muscle fiber contraction. Diseases affecting MNs, such as amyotrophic lateral sclerosis (ALS), are characterized by the loss of NMJs as MNs degenerate during disease progression (Wijesekera and Leigh, 2009; Viollet and Melki, 2013). This class of diseases is particularly debilitating, as MN loss leads to muscle atrophy and paralysis and is often fatal. With limited treatment options that only extend lifespan by months (Engler et al., 2004), significant efforts have focused on refining in vivo disease models to test new therapeutic options. However, current animal models for many MN diseases replicate only a portion of the spectrum of phenotypes in human disease (Pioro and Mitsumoto, 1995; Picher-Martel et al., 2016). Coculture systems to create NMJs in vitro were developed to address concerns about animal models (Das et al., 2007, 2010). More recent versions use stem cell-derived MNs and cell line–derived (Umbach et al., 2013; Steinbeck et al., 2016), or stem cell–derived muscle (Demestre et al., 2015; Puttonen et al., 2015). Using human induced pluripotent stem cells in such systems has further enabled patient-specific MN disease modeling (Faravelli et al., 2014; Abujarour and Valamehr, 2015; Lenzi et al., 2016) and drug screening (Inoue et al., 2014). Although having human cells enables more complex NMJ-in-a-dish modeling, concerns about...
MN, skeletal muscle, and NMJ maturity have been raised (Siller et al., 2013). Moreover, NMJ development relies on the organization of embryonic muscle fibers, precise guidance of MN axons, and the interplay of signaling from both the presynaptic and postsynaptic cells (Witzemann, 2006; Wu et al., 2010; Singhal and Martin, 2011; Shi et al., 2012). Traditional unpatterned in vitro coculture models on rigid substrates do not faithfully recapitulate the complex architecture observed in vivo; myotubes exhibit poor fusion efficiency and are highly disorganized, leading to inefficient and irregular contraction (Das et al., 2007, 2010; Guo et al., 2011; Umbach et al., 2012). Unpatterned NMJ cultures also do not develop sufficiently large AChR clusters—postsynaptic membrane specializations that are a hallmark of functional, mature NMJs (Witzemann, 2006).

To improve cell and NMJ maturation in coculture systems, some studies used microcontact printing (μCP), topography, and three-dimensional (3D) molds to organize myotubes in vitro. μCP guides cell growth by constraining cells to areas where extracellular matrix (ECM) proteins have been selectively deposited (Chen et al., 2017). Myoblasts attach and fuse on patterned ECM, creating aligned myotubes (Engler et al., 2004; Altomare et al., 2010). Over time, however, proteins from culture medium may deposit on the substrate, obscure the pattern, and eliminate cell alignment (Choi et al., 2012b). Physical stimuli, including topographical cues, have also been used to align myotubes in vitro (Wang et al., 2012), but these systems use substrates orders of magnitude more rigid than muscle (Choi et al., 2012a). They often result in muscle that is thinner and less mature than their in vivo counterparts (Gaudel et al., 2008; Zhu et al., 2011; Lawlor et al., 2014; Gibbons et al., 2016). Both of these approaches rely on two-dimensional (2D) methods, so 3D models have been developed that create aligned myobundles, which persist for several weeks in culture. Generally, these constructs are produced by using micromolds in which myoblasts are cultured (Morimoto et al., 2013; Madden et al., 2015; Bettadapura et al., 2016) or via self-assembly around T-shaped flexible posts (Sakar et al., 2012; Cvetkovic et al., 2014; Raman et al., 2016); however, both are often fabricated with complicated lithography processes, and resultant systems are difficult to process and image based on their 3D structure.

Methods that produce mature muscle but also rely on simple fabrication methods could be optimal for widespread use in NMJ-in-a-dish models. In this article, we describe a simple mechanically patterned hydrogel that can be used as a MN and myotube coculture platform. The system is based on a micropatterned hydrogel cell culture substrate that aligns cultured myotubes by juxtaposing soft and stiff regions similar to the alternating pattern of aligned myotubes with adjacent softer neurons to innervate the firm muscle in vivo (Choi et al., 2012b). Such patterns result in improved fusion, differentiation, and AChR formation for cells of either mouse or human origin that are responsive to pharmacological stimuli and, on coculture with MNs, form functional NMJs, thus providing a significant improvement over standard cell culture vessels without the requirement of expensive specialized equipment.

RESULTS
Design and fabrication of mechanically patterned hydrogels
We developed a two-step photopolymerization method that produces polyacrylamide (PA) hydrogels containing a grating pattern with alternating elasticity profiles (Figure 1A), resulting in a hydrogel with 100-μm-wide “stiff” stripes interspersed with 200-μm-wide “soft” stripes, that is, alternating regions of 12.6 ± 0.63 and 0.72 ± 0.05 kPa, respectively, as measured by atomic force microscopy (Figure 1B). Using this method, we successfully fabricated a wide range of geometrical patterns with micrometer-sized features. A sharp change in stiffness was observed at the interface of soft and stiff regions, creating a steep gradient of >100 kPa/mm, consistent with prior observation (Vincent et al., 2013) and indicating that there was limited diffusion during photopolymerization. For this NMJ-in-a-dish system, we chose to model myogenic and neurogenic stiffness (Engler et al., 2006), although elastic properties can be easily modified by changing acrylamide concentrations and micropattern dimensions for alternate applications.

Improved myotube fusion and differentiation on mechanically patterned hydrogels
Mouse C2C12 and human embryonic stem cell (hESC)–derived myoblasts (Caron et al., 2016) were cultured on mechanically patterned hydrogels to assess the influence of mechanical patterning on myotube fusion and differentiation. To decouple the effect of the compliant hydrogel from the mechanical patterning, myoblasts were also cultured on an unpatterned, myogenic hydrogel and on glass. Within 24 h, myoblasts preferentially migrated to and aligned with the

![Figure 1: Fabrication of mechanically patterned hydrogel. (A) Patterning process by which sequential exposures of the hydrogel to UV light—the second exposure including a photomask—creates the patterned hydrogel. (B) Hydrogel pattern and plot of elastic modulus vs. position orthogonal to the pattern direction. Average modulus values for “soft” (teal) and “stiff” (blue) regions.](image-url)
Data were normalized to glass substrates (M-cadherin (left) and N-cadherin (right) for the indicated conditions. Scale bar, 200 μm. (B) mRNA transcript levels for bright-field image of mouse myoblasts on the mechanically patterned substrate. Scale bar, 200 μm. (B) mRNA transcript levels for M-cadherin (left) and N-cadherin (right) for the indicated conditions. Data were normalized to glass substrates (n = 5).

FIGURE 2: Durotactic pattern recognition and fusion marker expression. (A) Bright-field images of mouse myoblasts on the indicated substrates. Scale bar, 100 μm. Inset, low-magnification bright-field image of mouse myoblasts on the mechanically patterned substrate. Scale bar, 200 μm. (B) mRNA transcript levels for M-cadherin (left) and N-cadherin (right) for the indicated conditions. Data were normalized to glass substrates (n = 5).

Stiffer myogenic regions on the mechanically patterned hydrogel, whereas myoblasts on the unpatterned hydrogels and glass exhibited no organization (Figure 2A), consistent with myocytes and stem cells (Choi et al., 2012b; Vincent et al., 2013; Wen et al., 2015). Before differentiation, myoblasts on patterned hydrogels also expressed more mRNA of M- and N-cadherin, markers indicative of cell fusion (Figure 2B).

After myogenic differentiation was initiated, elongated and multinucleated myotubes were present on all three substrates. However, the fusion index and average myotube width were larger on the mechanically patterned substrates than for both the unpatterned myogenic hydrogel and glass (Figure 3, A–C); only myotubes on patterned hydrogels approach sizes consistent with in vivo muscle, independent of species (Gaudel et al., 2008; Zhu et al., 2011; Lawlor et al., 2014; Gibbons et al., 2016). In addition to fusion indices, mouse myoblasts expressed MyoD and myogenin transcription factors on mechanically patterned hydrogels in a temporally regulated manner (Figure 3D), consistent with their sequential expression during differentiation (Choi et al., 2012a). Mechanically patterned human myotubes also expressed significantly more myosin heavy chain transcript, whereas mouse myotubes tended toward higher expression, suggesting that they are more mature than myotubes cultured on unpatterned myogenic hydrogels or glass (Figure 3, E and F). Together these data suggest that alignment, in addition to substrate elastic properties, could drive maturation and fusion, which may also contribute to deficiencies previously observed in hESC-derived myotube fusion (Demestre et al., 2015; Puttonen et al., 2015).

Mechanical patterning drives formation of functional AChR clusters

AChRs are critical components of NMJs and must be clustered in fused myotubes to function (Huh and Fuhrer, 2002). MNs secrete agrin to induce clustering, and so, on addition of exogenous agrin, we determined whether mechanical patterning could enhance AChR clustering. For both humans and mice, we found the largest AChR cluster sizes in myotubes cultured on mechanically patterned hydrogels (Figure 4, A and B). During NMJ development, agrin is released from approaching MNs and binds a receptor complex on the myotube consisting of dimers of muscle-specific receptor tyrosine kinase (MuSK) and low-density lipoprotein receptor-associated protein (Lrp4), initiating a signaling cascade that induces AChR clustering (Figure 4C; Wu et al., 2010; Shi et al., 2012). Lrp4 transcript was most highly expressed in myotubes in mechanically patterned myotubes, independent of species (Figure 4D). MuSK transcript was similarly highest with mechanically patterned hydrogels, but only in human myotubes (Figure 4E). To determine whether endogenous agrin from MN coculture could drive cluster formation, we cocultured primary MNs after myoblast fusion into myotubes. Cocultures were initiated after 4 d of myotube differentiation and maintained for 7 d. We found that mechanically patterned hydrogels drove formation of the largest AChR clusters, whereas AChR clusters on both the glass and unpatterned myogenic substrates remained small and punctuate (Figure 5, A and B). In the coculture system, MNs generally reside on top of the existing regions of mechanically patterned myotubes, avoiding the softer areas, and extend neurite outgrowths that terminate on the myotubes, with some instances of colocalization with AChR clusters, an indicator of putative NMJ formation (Figure 5C). Together these data suggest that NMJ-in-a-dish systems should support robust myotube alignment to facilitate AChR clustering, but these data do not yet establish the propagation of signals across the forming NMJ.

Spontaneous contractions were observed for myotube cultures only with mechanically patterned hydrogels (Figure 6, A and B, left); spontaneous activity increases during development, and thus our data are consistent with mechanically patterned myotubes being more functionally mature than unpatterned myotubes (Altimare et al., 2010). Although these data suggest functional clusters in the NMJ-in-a-dish system, they do not establish that the junctions are indeed functional. To determine whether AChR clusters can stimulate contraction, we exogenously added a bolus of 1 mM ACh. On stimulation, myotubes on both glass and mechanically patterned hydrogels contracted, although mechanically patterned myotubes underwent larger displacements during contraction, indicating a more intense contraction (Figure 6, A and B, right). Mechanically patterned myotube contractions were also shorter in duration and occurred at a higher frequency than myotube contractions on glass (Figure 6, C and D), again suggesting that the AChR clusters, which are a critical component of NMJs, are more functionally mature.

Substrate compliancy mediates clustering of ACh receptors on mechanically patterned hydrogels

To examine the roles of culture substrate stiffness versus myotube fusion and assembly in AChR clustering during NMJ development on mechanically patterned hydrogels, we cultured mouse myoblasts on patterned hydrogels with pathological stiff regions (Figure 7). After denervation—a hallmark of MN diseases—muscle fiber atrophy is accompanied by increased fibrosis, which increases the elastic modulus of muscle tissue (Engler et al., 2004; Carlson, 2014). Thus we fabricated mechanically patterned hydrogels with alternating soft (0.8 kPa) and pathological (58.1 kPa) stripes (Figure 7A). Mouse myoblasts differentiated on the pathological patterns showed no difference in fusion or myotube width from those cultured on patterned substrates with myogenic stiff regions (Figure 7, B and C). AChR cluster area was sensitive to elastic modulus, with myotubes cultured on the pathological patterns expressing smaller
AChR clusters than those cultured on myogenic mechanically patterned substrates (Figure 7, D and E), indicating that a previously unknown mechanotransduction pathway mediates AChR clustering. However, there was no correlation between myotube width and the size of the AChR clusters it expressed (Figure 7F), indicating that the mechanism of AChR clustering is independent of the extent of myotube fusion.

**DISCUSSION**

Despite the proliferation of culture methods to assemble NMJs (Das et al., 2007; Umbach et al., 2012; Morimoto et al., 2013; Chipman et al., 2014; Faravelli et al., 2014; Inoue et al., 2014; Abujarour and Valamehr, 2015; Demestre et al., 2015; Puttonen et al., 2015; Lenzi et al., 2016; Steinbeck et al., 2016; Uzel et al., 2016) and probe their mechanics (Tay et al., 2016), the possibility of functional neuron-to-muscle connectivity is insufficiently validated due to technical limitations with in vitro coculture systems. The mechanically patterned hydrogel system described here creates more mature (and wider) myotubes with larger AChR clusters that are more responsive to ACh than conventional or topographically patterned substrates. This is likely the result of agrin secretion by MNs, which signals through MuSK and Lrp4 to induce NMJ assembly (Wu et al., 2010). Forced clustering of myotubes on stiffer stripes also likely drives alignment by overlapping their traction forces, as well as increasing the probability that M- and N-cadherin-rich regions will make contact between cells, a step that is required for fusion (Choi et al., 2012a,b). Alignment of hESC-derived myotubes also illustrates that mechanical patterns may help to specify muscle during development and provides a platform for drug discovery.

During NMJ development, AChR clustering initiated by agrin signaling is a hallmark of the maturing NMJ synapse (Huh and Fuhrer, 2002; Witzemann, 2006). On the mechanically patterned culture system, AChR clustering significantly improves compared with both the glass and unpatterned myogenic culture substrates (Figure 4). In addition, we found that under coculture conditions in which the myotubes are exposed to endogenous agrin released from MNs, mechanically patterned myotubes exhibit AChR clusters that are >20× larger than those in myotube monocultures treated with rat recombinant agrin; however, myotubes cultured on glass or unpatterned myogenic substrates do not respond in such a robust manner (Figures 4B and 5B). We attribute the improved clustering to the fact that the MNs likely continuously release agrin onto the myotubes at a physiologically relevant level, whereas the monoculture myotubes were exposed to only one bolus dose of agrin added to the culture medium at day 1 after the onset of differentiation. Myotubes cultured on the mechanically patterned substrate, but not on glass or unpatterned myogenic substrates, express up-regulated levels of Lrp4 transcript (Figure 4D). Lrp4 acts as a receptor for agrin and, through its interactions with MuSK, initiates an intracellular cascade that results in AChR clustering on the muscle surface (Kim et al., 2008; Zhang et al., 2008). Thus lower
expression levels of Lrp4 in myotubes grown on glass and unpatterned myogenic substrates may preclude their ability to respond to the improved agrin signaling that occurs in the co-culture system.

To probe the role of substrate stiffness versus myotube alignment and fusion in the formation of AChR clusters, we differentiated mouse myotubes on patterned substrates of pathological stiffness. Cluster size was responsive to substrate stiffness, with smaller clusters expressed on pathological patterns, although there was no change in the extent of myotube fusion (Figure 7). These data suggest that AChR cluster size is regulated independently of myotube fusion, that is, clusters do not increase in size with increased myotube fusion, thus indicating a novel role for mechanotransduction in the control of AChR clustering. This pathway may be mediated by Wnt signaling, which is responsive to mechanical cues and involved in the Lrp4/MuSK signal cascade leading to AChR clustering (Hernandez and Salinas, 2012; Du et al., 2016). Mechanosensitivity of the NMJ may have implications for MN or muscle diseases in which muscle fibrosis occurs, such as ALS and Duchenne muscular dystrophy, and may contribute insight into the mechanisms of muscle denervation and atrophy (Klingler et al., 2012; Carlson, 2014).

Although this system is relatively straightforward to assemble, several important caveats should be considered relative to existing systems. Most notably, we find that this system enables imaging for days in coculture, which is sufficient for nascent NMJs to form; beyond that time, our system exhibits similar delamination issues to other 2D technologies (Wang et al., 2012), which could limit long-term utility with disease modeling. The 3D systems, despite complex fabrication, could provide long-term monitoring of patient-derived NMJs, although in vitro efforts in three dimensions have primarily focused on muscle (Sakar et al., 2012; Cvetkovic et al., 2014; Madden et al., 2015; Bettadapura et al., 2016; Raman et al., 2016). A second consideration is the relative maturity of the junctions themselves; AChR clusters resemble more plaque-like morphology (Marques et al., 2000; Huh and Fuhrer, 2002), and muscle twitch duration is long (Schiaffino and Reggiani, 2011; Umbach et al., 2012), even when improved by mechanical patterning, suggesting formation of functional but fetal-like synapses and immature slow fiber-type muscles. Although only spontaneous contractions induced by MNs were examined here, improved optogenetic control could further tease out NMJ maturity by examining signal propagation across junctions. Indeed, optogenetic control of muscle in these types of systems has shown that human cells can be optically controlled to examine more complex behaviors in-a-dish (Sakar et al., 2012; Raman et al., 2016).

Despite these caveats, benefits specific to this NMJ-in-a-dish model are its relative ease of fabrication and the observation that mechanical patterning improves the maturation of hESC-derived skeletal muscle (Caron et al., 2016). Other studies illustrated that human progenitors can be used to create functional NMJs (Demes-tre et al., 2015; Puttonen et al., 2015), but, to the best of our knowledge, this is the first demonstration that human myoblasts can be matured, fuse into patterned myotubes, and form functional NMJs in vitro. They appear to express more Lrp4 and MuSK and are more ACh sensitive than these other efforts, and so NMJs in these cells may represent a more mature state; however, side-by-side comparisons of these 2D methods is required to draw a better-supported conclusion. In addition, this system reveals that a mechanotransductive pathway that is not dependent on myotube fusion influences the formation of AChR clusters and may mediate the cellular response to pathological conditions, such as the fibrosis that occurs in muscle diseases. Thus we believe that this system represents the most straightforward and useful 2D NMJ-in-a-dish model for studying NMJ development and MN or muscle disease pathophysiology.

FIGURE 4: Patterning drives clustering of ACh receptors. (A) Representative fluorescence images of AChR clustering on agrin-treated myotubes cultured on glass, myogenic (Myo), or mechanically patterned (MP) hydrogels at 5 d (mouse) or 7 d (human) after induction of differentiation. AChRs were labeled with BTX. Scale bars, 25 μm. (B) Average AChR cluster size for mouse (left; n = 5) and human (right; n = 20). (C) Agrin-driven AChR clustering involving Lrp4 and MuSK. (D) Lrp4 and (E) MuSK expression was assessed by qPCR, and data were normalized to transcript expression on glass and plotted for the indicated species (n = 6). *p < 0.05 and ***p < 0.001 based on ANOVA comparisons of the indicated groups.
Mechanically patterned gel fabrication

Mechanically patterned hydrogels were fabricated using a two-step PA photopolymerization method (Figure 1A) modified from established methods (Marklein and Burdick, 2010; Tse and Engler, 2010) but that differs from them (Choi et al., 2012b). First, single-modulus PA hydrogels were polymerized onto methacrylated 18-mm glass coverslips from a prepolymer solution containing acrylamide (3%), bis-acrylamide (0.4%), and 2-hydroxy-4’-(2-hydroxyethoxy)-2-methylpropiophenone (0.5%). The prepolymer solution was sandwiched between the methacrylated glass and a chlorosilanated glass slide, dehydrated overnight, and then rehydrated with a secondary solution consisting of acrylamide (6.25%), bis-acrylamide (0.4%), and 2-hydroxy-4’-(2-hydroxyethoxy)-2-methylpropiophenone (0.5%) to minimize the amount of differential swelling that might occur in a hydrogel. The resultant hydrated PA gel was exposed to UV light (350 nm) for 5 min. After polymerization, the hydrogels were functionalized with a photoactivating cross-linker, sulfo-succinimidyl 6-(4’-azido-2’-nitrophenylamino)hexanoate (Sulfo-SANPAH; Pierce). The hydrogels were immersed in 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid buffer (pH 8.4, 50 mM) containing Sulfo-SANPAH (0.2 mg/ml) and exposed to UV light (350 nm) for 10 min. After several washes in PBS, the hydrogels were incubated in collagen IV (100 μg/ml) overnight at 37°C.

Mechanical characterization

Substrate stiffness was measured via indentation using an MFP-3D-Bio (Asylum Research, Santa Barbara, CA) atomic force microscope (AFM). We used chromium/gold-coated, silicon nitride (SiN) cantilevers with pyramid tips (PNP-TR; NanoWorld) with a nominal spring constant of ∼30 pN/nm, as determined from the MFP-3Ds built-in thermal calibration function. Samples were mounted on glass slides using vacuum grease and immersed in PBS. The probe was indented into the sample with an approach velocity of 2 μm/s and a force trigger of 2 nN. AFM data were analyzed using custom code in Igor Pro (Wavemetrics); the substrate spring constant, that is, Young’s modulus, was determined using a linearized Sneddon model (Kaushik et al., 2011).

Cell line and human embryonic stem cell–derived myotube culture

C2C12 mouse myoblast cell line, which was validated by the American Type Culture Collection (CRL-1772), was seeded on the surface of mechanically patterned and myogenic hydrogels or glass coverslips coated with collagen IV (100 μg/ml) at a density of 4000 cells/cm² and maintained in growth medium consisting of DMEM (with 4.5 g/l glucose, with glutamine, without sodium pyruvate), fetal bovine serum (FBS; 20%), and penicillin/streptomycin (1%). After 2 d in growth medium, the cultures were switched to differentiation medium consisting of DMEM, horse serum (2%), insulin (2 μg/ml), and penicillin/streptomycin (1%). At 1 d after the onset of differentiation, cultures were treated with agrin (100 ng/ml) to induce AChR clustering, and at 2 d after the onset of differentiation, cultures were treated with cytosine β-d-arabinofuranoside (10 μM) to inhibit myoblast proliferation. At 5 d after the onset of differentiation, cultures were either assayed for functional activity or terminated for immunofluorescence and RNA analysis.

The hESC-derived myoblasts (GENEA002) were obtained and validated by Genea Biocells (La Jolla, CA) through prior publication (Caron et al., 2016). Cells were seeded onto hydrogels or glass coverslips coated with collagen I (100 μg/ml) at a density of 15,000 cells/cm² and maintained in growth medium (SKM-02; Genea Biocells) for 3 d. Cultures were then changed to differentiation medium (SKM-03; Genea Biocells) and maintained for 7 d before functional analysis or fixation for immunofluorescence and RNA analysis.

FIGURE 5: Patterning improves AChR clusters induced by MN coculture. (A) Representative fluorescence images of AChR clustering on mouse myotubes cocultured with mouse MN for 7 d. Myotubes were differentiated for 4 d before the start of the coculture. AChRs were labeled with BTX. Scale bar, 20 μm. Arrows indicate BTX immunoreactivity. (B) Average AChR cluster size (n = 20) for glass (Gl), soft myogenic substrates (Myo), and mechanically patterned hydrogels (MP). ***p < 0.001 based on ANOVA comparisons between substrate conditions. (C) Representative fluorescence image of patterned mouse myotube/mouse MN coculture at 7 d. Cells were immunostained with β-III-tubulin (TuJ1, green) and BTX (red) and counterstained for nuclei with Hoechst 33352 (blue). Scale bar, 30 μm. Arrows indicate BTX immunoreactivity that colocalizes with β-III-tubulin immunoreactivity, suggestive of a putative NMJ.
incubated in PBS containing 5% goat serum for 20 min at room temperature. Samples were incubated in primary antibodies at 4°C overnight, washed in PBS, and then incubated in secondary antibodies for 45 min at room temperature, followed by a 10-min incubation in Hoechst 33352 (1:1000 in PBS) to counterstain nuclei. Primary antibodies used in this study were hybridoma-derived anti–myosin heavy chain (1:50, MF20; Developmental Studies Hybridoma Bank) and rabbit polyclonal anti–β-III-tubulin (1:500, ab18207; Abcam). Alexa Fluor 488 secondary antibody was used (1:1000; Fisher Scientific). To label AChR clusters, samples were incubated with α-bungarotoxin (BTX) conjugated with Alexa Fluor 555 (1:500; Molecular Probes) for 45 min at room temperature.

Myoblast fusion index was calculated as the percentage of Hoechst-positive cell nuclei residing within MF20-positive myotubes. Average myotube width was calculated by measuring the width at the widest point on the myotube using the ImageJ measure function. For each metric, data were collected from 20 representative 20× images. BTX-immunoreactivity on 5 (mouse) or 20 (human) representative 60× images was used to evaluate AChR clustering. Average cluster size was quantified by manual tracing using ImageJ software. The width of each myotube that expressed AChR clusters was recorded and plotted against AChR cluster area to evaluate the correlation between cluster size and myotube width.

Quantitative PCR
RNA was collected from cultures using TRIzol isolation, and cDNA was synthesized from 2 μg of RNA using reverse transcriptase PCR (37°C for 60 min, 99°C for 5 min, and 5°C for 5 min) with Superscript II Reverse Transcriptase (Invitrogen) and 2.5 mM random hexamer mix (Invitrogen). Quantitative PCR (qPCR) was carried out using a CFX96 Real-Time PCR Detection system using iQ SYBR Green Supermix (2 min at 50°C followed by 10 min at 95°C for one cycle, then 15 s at 95°C followed by 1 min at 60°C for 40 cycles) and custom designed primers (Supplemental Table S1). Expression levels were calculated based on a standard curve generated by fibronectin plasmid. All data were normalized to glyceraldehyde-3-phosphate dehydrogenase (GAPDH), and data from the myogenic and MP hydrogels are presented as the fold change from the glass control.

Functional assessment of myotubes
Bright-field video images were acquired with a CoolSnap HQ camera (Photometrics Scientific) through a Nikon Eclipse Ti inverted microscope with a 10× objective.

Myotube monocultures were assessed for ACh responsiveness after 5 d in culture (mouse) or 7 d in culture (human). Videos were collected at 9.2 frames/s and captured before and immediately after

Isolation of rat primary MNs and their coculture with myotubes
The use of rats in this study was approved by UC San Diego’s Institutional Animal Care and Use Committee review board (Protocol # S11032). Mouse embryonic motor neurons were isolated from the spinal cords of embryonic day 12 CD1 mouse embryos using a discontinuous density gradient centrifugation purification technique (Gingras et al., 2007). Purified motor neurons were cocultured with established C2C12 myotube cultures at 4 d after the onset of myotube differentiation. The cocultures were maintained in motor neuron medium consisting of Neurobasal medium, B27 supplement (2%), glutamine (0.5 mM), hydrocortisone (400 ng/ml), insulin (2 μg/ml), FBS (10%), and penicillin/streptomycin (1%).

Immunofluorescence staining, fusion characterization, and AChR characterization
Cultures were fixed with 3.7% formaldehyde for 25 min at room temperature, permeabilized with 1% Triton-X for 5 min, and then
stimulation with 1 mM ACh chloride (Sigma-Aldrich) in a 10-μl volume. Kymographs of myotube contraction were generated using ImageJ software.

Statistical analysis
All data are presented as the mean ± SEM. Statistical analysis was carried out with GraphPad Prism software using one-way analysis of variance (ANOVA) tests, unpaired t test, or linear regression analysis, as appropriate. Statistical significance was considered at p < 0.05.

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