Functional Properties of Engineered Heart Slices Incorporating Human Induced Pluripotent Stem Cell-Derived Cardiomyocytes

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SUMMARY

Human induced pluripotent stem cell-derived cardiomyocytes (hiPSC-CMs) hold great promise for cardiac studies, but their structural and functional immaturity precludes their use as faithful models of adult myocardium. Here we describe engineered heart slices (EHS), preparations of decellularized porcine myocardium repopulated with hiPSC-CMs that exhibit structural and functional improvements over standard culture. EHS exhibited multicellular, aligned bundles of elongated CMs with organized sarcomeres, positive inotropic responses to isoproterenol, anisotropic conduction of action potentials, and electrophysiological functionality for more than 200 days. We developed a new drug assay, GRIDS, that serves as a “fingerprint” of cardiac drug sensitivity for a range of pacing rates and drug concentrations. GRIDS maps characterized differences in drug sensitivity between EHS and monolayers more clearly than changes in action potential durations or conduction velocities. EHS represent a tissue-like model for long-term culture, structural, and functional improvement, and higher fidelity drug response of hiPSC-CMs.

INTRODUCTION

The advancement in cardiac differentiation strategies for human pluripotent stem cells (hPSCs) (Burridge et al., 2012) has opened up opportunities for new in vitro studies of human cardiomyocytes (CMs). However, widespread and reliable use of hPSC-CMs requires the development of preparations that can recapitulate essential features of myocardial structure and function: e.g., elongated CMs in arrays that mimic myofiber bundles, coordinated contraction, fast and uniform conduction of action potentials (APs), and appropriate sensitivity to cardioactive drugs. To this end, a variety of strategies have been employed to make tissue-like constructs, including casting hPSC-CMs in hydrogels (Tzatzalos et al., 2015), seeding them onto synthetic matrices (Ma et al., 2014), and fabricating cell sheets (Matsuura et al., 2012). These efforts have resulted in structurally organized, multicellular preparations that promote more mature states of cardiac gene expression, conduction, cardiac handling, and conduction. However, their ultimate usefulness for in vitro studies may be hampered by the inability to maintain functionality during long-term culture and the absence of instructive cues typically present in the adult myocardium.

An emerging strategy is to use decellularized myocardial matrix as a source of biochemical, topographical, and biomechanical cues present in the heart to direct differentiation and maturation of PSC-CMs. Decellularized myocardial matrix decreases stem cell pluripotency and induces differentiation in iPSCs (Carvalho et al., 2012) and early cardiac progenitor cells (Lu et al., 2013). The idea that multicomponent extracellular matrix (ECM) can enhance cardiac differentiation has been demonstrated with hydrogels composed of solubilized acellular porcine myocardial matrix (Duan et al., 2011) or solubilized basement membrane Matrigel preparations (Zhang et al., 2012). Cell culture coatings made from these solubilized, acellular matrix sources or from decellularized sheets of supporting cells enhance the structural organization of CMs (Baharvand et al., 2005) and temporally advance the expression of cardiac genes and proteins in cardiac progenitor cells (French et al., 2012). Acellular matrix can improve the response of single hiPSC-CMs to cardiac drugs so that they more closely resemble that of adult CMs (Feaster et al., 2015). Although these findings suggest that decellularized matrix may be uniquely suited to guide cellular organization, promote CM lineage commitment, accelerate maturation, and promote better physiological responses to cardiac drugs, reseeding decellularized myocardium with hPSCs and differentiating these cells into a dense tissue-like network of CMs has proven difficult. Furthermore, an important step toward the creation of a truly tissue-like preparation of human CMs would be the demonstration of a high degree of electrophysiological and contractile function in preparations generated on decellularized matrices.

In this study, we have seeded CMs onto thin decellularized slices of ECM (dECM slices) derived from pig heart. Our goal was to create tissue-like constructs that would: support a functional network of hiPSC-CMs, organize and align CMs, exhibit coordinated contraction and uniform...
AP propagation, maintain functionality during prolonged culture, have a sufficiently large area to support reentrant arrhythmia, and exhibit a robust response to cardiac drugs for a wide range of concentrations and pacing rates. We find that CM-seeded tissues reliably recapitulate key structural, functional and electrophysiological features of native myocardium and proved more sensitive in drug screening assays where electrical pacing could be varied.

RESULTS

Thin Slices of Decellularized Myocardium Promote Growth and Global Alignment of hiPSC-CMs

We sectioned 12-mm diameter plugs of left ventricular myocardium from porcine hearts into 300-μm-thick slices using a vibratome (Figure 1A). Full decellularization was achieved after 3.5 h of exposure to detergents (Figure 1B). The resulting thin dECM slices did not maintain their shape when removed from liquid, a problem that was resolved by spreading each slice onto a plastic coverslip. Under these conditions, the perimeter of the dECM slice could adhere to the edges of the coverslip (Figure 1C). The dECM slice exhibited overall alignment of ECM and residual vasculature (Figure 1D).

We confirmed the removal of cells and nuclei by assaying slices before and after treatment with detergents. Native slices contained an abundance of cells, indicated by nuclear staining and intracellular F-actin (Figure S1A), which were absent in dECM slices (Figure S1E). Decellularization left intact the ECM components of collagen I (Figures S1B and S1F), collagen III (Figures S1C and S1G), and laminin (Figures S1D and S1H). The organization of collagen fibers in the dECM slice, as visualized by second-harmonic generation (SHG) imaging, was not altered by storing sectioned slices at −80°C before decellularization or by storing ventricular plugs at −80°C before slicing and decellularization (Figures S1I–S1L). Overall, each component analyzed in the

![Figure 1. Preparation of EHS](image)

(A) The workflow for preparing slices.
(B) Slices of myocardium before (left) and after decellularization (right).
(C) Slices attached to the edges of a plastic coverslip and placed in the wells of a standard culture dish before cell seeding.
(D) Light micrograph of decellularized slice showing ECM and portion of vasculature (arrow indicates vessel).
See also Figure S1.
dECM slices largely retained the structural alignment observed in native slices. Our decellularization method decreased the DNA content of slices more than 160-fold to approximately 0.12 μg/mg initial dry weight, as reported previously (Blazeski et al., 2015). Taken together, these data suggest that the detergent-treated slices are almost completely devoid of cells and nuclei but retain a mixture of ECM components that remains structurally organized.

Differentiated progeny from hiPSCs were seeded onto the dECM slices at d10–12 (Figure 1A) to form engineered heart slices (EHS). Our cardiac differentiation protocol yielded a mixture of, on average, ~83% cTnT-positive hiPSC-CMs and ~17% cTnT-negative non-CMs (Figures S2A and S2B). After 16 days of EHS culture (d26–28), these cells formed multicellular, aligned tissue layers (Figures 2A and 2B). CM alignment, however, decreased in the apical cell layers farthest from the surface of the matrix (Figure S3). Cells, which organized into multicellular strands, were made up mainly of cardiac troponin I (cTnI)-positive CMs, with vimentin-positive non-CMs located primarily in the center of the strands (Figure 2A). CMs also exhibited striations characteristic of sarcomere structures and stained positively for Cx43 (Figure 2C) localized along the periphery of the cells (Figure S3D). Transmission electron micrographs of hiPSC-CMs on slices showed the presence of z-lines in sarcomeres that were surrounded by mitochondria (Figure 2D).

Comparisons were then made between d55 hiPSC-CMs seeded at low density to make EHS and those on Geltrax-coated cell culture dishes to evaluate the effect of the matrix on cellular shape and organization. CMs grown on Geltrax were cobblestone-like and had randomly oriented sarcomeres (Figure 2E). Age-matched CMs on EHS were elongated, with sarcomeres arranged along the long axis of each cell (Figure 2F). Interestingly, the morphological trait of elongated cells and aligned sarcomeres became ingrained with time on the tissue slices. For example, when hiPSC-CMs were maintained for a prolonged period of time (139 days) as EHS were more elongated and more closely oriented at a pacing cycle length (PCL) of 666 ms (1.5 Hz, n = 8), the EHS area in the field of view decreased, on average, by 2.0% ± 1.3% from its value at rest (Figures 3A–3C; Video S1). The addition of 1 μM isoproterenol resulted in a 1.5 ± 0.6-fold larger area change, and subsequent washout of the drug brought the area change back down to 1.1 ± 0.4 times the baseline value (Figure 3). Increasing the pacing rate from 1.5 Hz (666 ms PCL) to 2 Hz (500 ms PCL) resulted in an area change of 0.7 ± 0.1 (negative force-frequency relationship) that was statistically significant, whereas decreasing the pacing rate from 1.5 to 1 Hz (1000 ms PCL) resulted in an area change (1.1 ± 0.1) that was not statistically significant (Figure 4B).

**EHS Contract Synchronously and Respond to Isoproterenol**

Spontaneous and asynchronous contractions were apparent in EHS within 24 h of cell seeding, but this transitioned to synchronous contractions in about 1 week. At d28 and d74, we evaluated contraction as the change in EHS area in a region where the slice edge was freed from the coverslip. The contracting hiPSC-CMs deformed the ECM, permitting us to monitor the change in area as an approximation of the degree of contraction. When stimulated at a pacing cycle length (PCL) of 666 ms (1.5 Hz, n = 8), the EHS area in the field of view decreased, on average, by 2.0% ± 1.3% from its value at rest (Figures 3A–3C; Video S1). The addition of 1 μM isoproterenol resulted in a 1.5 ± 0.6-fold larger area change, and subsequent washout of the drug brought the area change back down to 1.1 ± 0.4 times the baseline value (Figure 3). Increasing the pacing rate from 1.5 Hz (666 ms PCL) to 2 Hz (500 ms PCL) resulted in an area change of 0.7 ± 0.1 (negative force-frequency relationship) that was statistically significant, whereas decreasing the pacing rate from 1.5 to 1 Hz (1000 ms PCL) resulted in an area change (1.1 ± 0.1) that was not statistically significant (Figure 4B).

**EHS Exhibit Anisotropic Electrical Conduction and Retain Functionality during Long-Term Culture**

EHS at d54–58 could be pace-captured starting from a PCL of 100 ms to 425 ± 51 ms (n = 6) and exhibited propagation of APs throughout the entire preparation (Figure 4A; Video S2). Action potential durations (APDs) showed physiological rate dependence: APD$_{30}$ and APD$_{80}$ decreased during incremental increases in pacing rate (incremental decreases in PCL from 1000 to 400 ms, n = 4–6) (Figure 4C; Table S1). Both transverse and longitudinal conduction velocity (CV) showed physiological rate dependence, decreasing with increasing pacing rate (Figure 4D). The resultant CV anisotropy ratio remained relatively constant around 1.4 over the range of PCLs from 500 to 1000 ms (Figure 4E), suggesting that the increases in pacing rate primarily affected sodium channel availability (i.e., excitability) and did not cause significant changes in gap junctional conductance (de Diego et al., 2011).

The high degree of electrophysiological function in EHS was applicable to other hiPSC lines. This was demonstrated through the analyses of EHS made with hiPSC-CMs derived from a patient with confirmed LQT2. LQT2 is a cardiac disorder characterized by prolonged ventricular repolarization arising from mutations in the rapid delayed potassium channel (HERG) that increases the incidence of arrhythmias (Tester and Ackerman, 2014). As with wild-type (WT) EHS, LQT2 EHS cells were aligned and organized into multicellular strands (Figures SSA and SSB), in which non-CMs were in contact with the ECM, while the CMs...
were organized on top of these cells (Figures S5C and S5D). Like our previous observations in WT EHS, LQT2 EHS exhibited anisotropic propagation of APs throughout the slice (Figure S5E). APDs in LQT2 EHS were prolonged compared with those in WT EHS (Figure S5F), recapitulating the hallmark of LQT (Figure S5G, n = 3 LQT2 EHS and n = 5 WT EHS).

In addition to having structurally elongated and organized CMs (Figure 2G), EHS in long-term culture retained cell-connectivity and paceable, coordinated AP activity. Samples that were maintained in culture for more than 2 months (d62–82) exhibited isotropic propagation of APs, typical of thicker cell layers where cells become more randomly oriented (Figure S3), and had CVs ranging from...
13.9 ± 1.9 cm/s at 700 ms PCL to 18.4 ± 1.8 cm/s at 1900 ms PCL (Figure 5A; Table S2). The proportion of non-CMs in EHS at d68 increased from that at the time of seeding (Figure S2). EHS using the same batch of cells but cultured for more than 200 days, exhibited lower CVs, which at 1300 and 1500 ms PCL reached statistical significance when compared with d62–82 CVs at the same PCL (Figure 5A; Table S2). Notably, d201 EHS could be paced over a wider range of cycle lengths (CLs), from 2000 ms (now possible because of their lower spontaneous beating rates) to 400 ms (now possible because of their shorter APDs), than that for d62–82 EHS, from 1900 ms PCL to 700 ms. Mean APD_{80} and APD_{30} of d201 EHS were significantly shorter at all PCLs than those of d62–82, respectively (Figure 5C; Table S2), reflecting net increase in inward and/or net decrease in outward AP current in older EHS.

EHS Differ from Standard Monolayer Cultures in their Response to Ion Channel-Modulating Drugs, when Evaluated Using GRIDS Analysis

Based on the improved adult-like cellular morphology, more ventricular-like APs (Figures S6A and S6B) and increased expression levels of numerous ion channel transcripts of hiPSC-CMs after culture on EHS (Figure S6C), we hypothesized that cells on EHS would exhibit differences in drug responses in comparison with standard culture. To test the responsiveness of EHS to ion channel modulators, we developed an assay, called GRIDS (grid of responses indicating drug sensitivity), that characterizes the ability of a cardiac preparation to respond to pacing when exposed to a range of drug concentrations. We used it to compare EHS with age-matched monolayer cultures. APs of EHS and monolayers were evaluated at each drug concentration in the absence of pacing and when subjected to electrical stimulation at PCLs ranging from 2000 to 500 ms. By incorporating a range of pacing rates, GRIDS manifests the distinct sensitivities of different ion currents (Ravens and Wettwer, 1998) and their rate dependencies, and provides a “fingerprint” of drug sensitivity. We observed several types of responses: no spontaneous activity (Figure 6A, i), spontaneously generated APs when no pacing was applied (Figure 6A, ii), extra APs between pacing stimuli (Figure 6A, iii), one AP generated for each pacing stimulus—i.e., 1:1 pace-capture (Figure 6A, iv), and failure to maintain 1:1 pace-capture (lost beats) (Figure 6A, v). We evaluated the effects of each drug on augmenting paced activity by counting the proportion of EHS and monolayers that exhibited extra beats between paced beats (‘Fraction With Extra Beats’, exemplified in Figure 6A, ii–iii) and on the ability to retain capture by counting the proportion of EHS and monolayers that lost beats during pacing (‘Fraction With Lost Beats’ exemplified in Figure 6A, v). Further, if application of a drug resulted in spontaneous activity, the EHS or monolayers were counted as having extra beats and included in the ‘Fraction With Extra Beats’ visualized in the bottom row of each GRIDS analysis.
At 1 mM BaCl2, all of the EHS tested lost capture at 500 ms PCL. Taken together, these results suggest that EHS are less sensitive to BaCl2 than monolayers.

We then blocked the slow delayed rectifier K+ current (I_{KS}) with chromanol 293B to test its effects on EHS and monolayer cultures (Figure 6C). The fraction of monolayers exhibiting spontaneous activity when no pacing was applied was non-zero and constant at all concentrations except for 5 μM, when it was zero, and 60 μM, when all monolayers exhibited spontaneous activity (Figure 6C, bottom). However, at all concentrations tested, both EHS and monolayers could be pace-captured at all PCLs from 2000 to 500 ms (Figure 6C). Therefore, chromanol 293B had a limited effect on either preparation, suggesting that I_{KS} is poorly expressed or not functionally active in both monolayers and EHS. Aside from differences in potassium channels, we tested whether L-type calcium channels might also be differentially expressed in EHS and in monolayers, so the channel blocker nifedipine was applied (Figure 6D). As the drug concentration was increased, monolayers lost pace-capture over a wider range of PCL, whereas EHS never lost pace-capture (Figure 6D), indicating that EHS were insensitive to nifedipine in this regard.

In addition to the GRIDS analysis, we evaluated the occurrence of drug-induced prolongations of repolarization, which is an index of liability for acquired long-QT syndrome (Wood and Roden, 2004), as well as conduction slowing, which can be an early sign of conduction block. At a PCL of 1500 ms, E-4031 prolonged APD_{80} (Figure 7A, i–iii) and slowed CV (Figure 7A, iv) in both EHS and monolayers in a concentration-dependent manner over a range of 50 nM to 10 μM. For concentrations up to 150 μM, BaCl2 increased APD_{80} and decreased CV in both preparations in a concentration-dependent manner (Figure 7B), although the decrease in CV reached statistical significance only in EHS. Chromanol 293B (Figure 7C) had no effect on APD_{80} or on CV at concentrations ranging from 1 to 60 μM in either EHS or monolayers. Nifedipine did not affect APD_{80} of EHS and monolayers (not shown), but shortened the plateau phase of the AP in EHS, an effect that was quantified by APD_{90}. At 1500 ms PCL, 0.1 and 0.3 μM nifedipine decreased APD_{90} to a greater extent in EHS than in monolayers (Figure 7D, i–iii), suggesting that EHS have more developed I_{Ca,L} current. Nifedipine does not cause conduction slowing in healthy adult myocardium (Mitchell et al., 1982) and did not significantly decrease CV in our preparations, except at a concentration of 0.01 μM in monolayers (Figure 7D, iv). We further modulated I_{Ca,L} in EHS by applying Bay K 8644, an L-type calcium channel activator, over a range of 0.3 μM–100 μM, and found that it increased APD and decreased CV in a concentration-dependent manner (Figures S7A–S7D). Finally, we found that EHS remained responsive to ion channel-modulating drugs even after long-term culture (d201), as in the case of Bay K

![Figure 4. Optical Mapping of EHS](image-url)
8644 and cromakalim, an activator of the ATP-dependent potassium current ($I_{K,ATP}$) (Figures S7E–S7G).

**DISCUSSION**

In this study, we repopulated dECM slices with hiPSC-CMs to make EHS with coordinated and syncytial contractile and electrophysiological function. On EHS, reseeded hiPSC-CMs organize in a similar manner to the ordered arrays of fibers found in the native myocardium (Veeraraghavan et al., 2014), becoming elongated and aligned in bundles, with elongated and oriented nuclei, and well-defined, aligned sarcomeres. This is an improvement from the morphology seen in standard cultures, whereby hiPSC-CMs appear more cobblestone-like and have randomly oriented sarcomeres. However, gap junction staining for Cx43 was observed around the periphery of cells in the EHS and does not localize at intercalated discs, reflecting a level of structural immaturity that has also been found in other studies of hPSC-CMs (Zhang et al., 2013). Nevertheless, our EHS are a well-connected and reproducible functional syncytium of CMs that exhibits uniform conduction and coordinated contraction over an area of about 1 cm². This overcomes the problem of variable and patchy conduction that occurs in other preparations using decellularized myocardium caused by non-uniform cell seeding (Guyette et al., 2016; Lu et al., 2013; Oberwallner et al., 2015).

The cellular organization of EHS and the resultant anisotropic conduction of APs is guided by the retention of the native oriented matrix in the dECM slice. This strategy to utilize the topographical cues of the ECM differs from those used in other studies to align CMs, including microcontact printing (Wang et al., 2014), hydrogel compaction in the presence of non-myocytes (Liu et al., 2011), and fabricated microgrooved (Rao et al., 2013), nanogrooved (Macadangdang et al., 2015), electrospun (Wanjare et al., 2017), and wrinkled (Wang et al., 2013) substrates. The use of decellularized matrix may confer benefits not present in the other approaches, as there is mounting evidence that the complex chemistry of the matrix can promote stem cell differentiation (Ng et al., 2011) and electrophysiological maturation of CMs (Herron et al., 2016). Our study of EHS demonstrates the suitability of this platform to study electrophysiological function over long-term culture.

Our experimental approach involves the routine production of batches of 10–20 thin tissue slices from ventricular plugs that are subsequently decellularized in parallel. This decellularization method leaves behind a scaffold that retains an organized and aligned structure, made up of multiple ECM components. While various decellularization methods have been developed (Badylak et al., 2011), we chose the method of Ott and colleagues because it preserves non-collagen proteins (particularly, fibrillin, heparin sulfate, and laminin; Guyette et al., 2016), with a trade-off of decreased retention of collagen (Akhyari et al., 2011). These non-collagen components may be particularly beneficial for promoting the differentiation and maintenance of hPSC-CMs (Nakayama et al., 2014). However, because the composition and mechanical properties of the ECM change during development of the heart from the postnatal to adult stage (Gershlak et al., 2013), further work needs to be done to identify the developmental stage that will yield ECM best suited for growth and maintenance of hPSC-CMs. Our method for decellularizing slices is amenable to the use of native myocardium from a variety of sources and can be used to compare the effect of different species, chamber locations, and developmental states of the ECM source on CM electrophysiology and contraction. Automation of the process of anchoring the dECM slice to a
support will be necessary if large numbers of ECM scaffolds are needed to make EHS for drug discovery and screening.

While EHS exhibited a positive inotropic response to isoproterenol, their fractional shortening (around 2%) was much lower than the 30% fractional shortening reported in adult hearts (Colan et al., 1984), and they also exhibited a negative force-frequency relationship. Further improvements, such as increasingly rapid pacing during culture, can be used to improve contractility and achieve the positive force-frequency relationship found in adult hearts.

Figure 6. Effect of Cardioactive Drugs on Paced Beats
(A) GRIDS maps of EHS and monolayer responses to E-4031. Sample traces of optical recordings from EHS (right, i–v, top black traces) and corresponding pacing stimuli (right, i–v, bottom blue traces) for examples of: (i) spontaneous activity, (ii) spontaneous activity when no pacing stimuli were applied, (iii) extra beats during pacing, (iv) capture of each paced beat, and (v) lost beats during pacing. PCL and concentration of E-4031 applied during the sample traces is indicated by the squares labeled i–v in the color grid (left). Color bars indicate the fraction of total monolayers or EHS which lost beats illustrated in (v) or gained extra beats illustrated in (iii). n = 5–10 independent EHS and n = 5–18 independent EHS.
(B) GRIDS maps for BaCl2. n = 3 independent monolayers and n = 3–6 independent EHS.
(C) GRIDS maps for chromanol 293B. n = 3 independent monolayers and n = 3–6 independent EHS.
(D) GRIDS maps for nifedipine. n = 3 independent preparations each of monolayers and EHS.
See also Figures S6 and S7.
myocardium (Ronaldson-Bouchard et al., 2018). EHS also exhibited rate-dependent decreases of APD and CV, as well as anisotropic conduction, as is found in the adult human heart (Yue et al., 2005). However, the CV in EHS was less than half of that measured in the adult ventricle (Durrer et al., 1970), and the anisotropy ratio of conduction was substantially less than that in the adult ventricle (Peters and Wit, 1998). The loss of CM alignment in layers of cells farther from the matrix surface likely contributed to a diminished anisotropy ratio.

Amenability to long-term culture is a powerful feature of EHS, because prolonged culture advances the structural organization, cardiac gene expression, and contractile and electrophysiological function of hPSC-CMs (Lundy et al.,

![Figure 7. Effect of Cardioactive Drugs on Action Potentials and Conduction Velocities](image)

Action potential recordings from EHS (i) and monolayers (ii) for increasing concentrations of (A) E-4031, (B) BaCl2, (C) chromanol 293B, and (D) nifedipine. Changes in APD90 (A–C, iii), APD50 (D, iii), and CV (A–D, iv) are plotted for each drug. EHS and monolayers were paced at 1500 ms cycle length for E-4031 and nifedipine and at 1000 ms cycle length for BaCl2 and chromanol 293B. Baseline values of APD and CV before the application of drug are indicated in red for EHS and blue for monolayers in (iii) and (iv) for each drug. Error bars denote standard deviation. Red *, p < 0.05 when comparing the percent change for EHS with 0. Blue *, p < 0.05 when comparing the percent change for monolayers with 0. #p < 0.05 when comparing baseline values for EHS with baseline values for monolayers. n values for EHS and monolayers exposed to each drug are the same as in Figure 6. See also Figure S7.
Maintaining multilayers of hiPSC-CMs in standard culture plates for periods of weeks is difficult, because they can detach from the underlying substrate, although individual hiPSC-CMs have been maintained for up to 120 days (Lundy et al., 2013), and multicellular embryoid bodies of hiPSC-CMs have been maintained for up to 360 days (Kamakura et al., 2013). In our study, EHS allowed for stable, long-term culture of a functional syncytium of hiPSC-CMs. Electrophysiological functionality was maintained for more than 200 days—EHS could be pace-captured at PCLs as short as 400 ms, and APs continued to propagate as before throughout the entirety of the preparation, although with some loss of CV that may occur as non-myocytes proliferate over time in EHS culture. Remarkably, hiPSC-CMs cultured long-term within EHS retained their elongated morphology and aligned sarcomeres even after removal from the dECM slice, suggesting that the ECM may have durable effects on cell phenotype. Aside from cues from the ECM, additional steps may be required to optimize the structural organization and function of EHS in long-term culture.

The EHS preparation holds promise for preclinical cardiotoxicity testing, where accurate prediction of arrhythmia risk is essential to remove hazardous drugs from the development pipeline. Currently, drugs are tested for their ability to inhibit HERG and cause QT prolongation, a marker for risk of developing Torsades de Pointes, a tissue-level arrhythmia (Farkas and Nattel, 2010). Most studies of this kind are performed on heterologous expression systems that lack the full complement of cardiac ion channels (Ferrini et al., 2016). Such assays also do not account for off-setting mechanisms from non-HERG ion channels that may render a drug safe (Redfern et al., 2003). The EHS preparation addresses these shortcomings as a functional syncytium of human CMs and allows for a multitude of mechanisms by which drugs can affect excitability, including effects on ion currents and electrical coupling.

The GRIDS assay we developed provides a new tool for evaluating drug sensitivity in the context of cellular automaticity and excitability. Changes in spontaneous beating rates are often used to evaluate drug sensitivity of hiPSC-CMs (Gilchrist et al., 2015) and can be altered by drugs that act on ICa,L or IKr (Blazeski et al., 2012). However, electrophysiological measurements at variable spontaneous beat rates fail to control for the rate dependence of the various ion currents. On the other hand, GRIDS evaluates the effect of drugs during electrical pacing at different fixed rates. These periodic stimuli introduce controlled, dynamic changes into the system, and the resulting beating patterns are an integrated effect of automaticity, excitability, and refractoriness. The GRIDS map for a given drug is comprised of pace-capture responses across multiple dosing and electrical pacing regimes, and it can serve as a fingerprint of the drug sensitivity. The lowermost row of the GRIDS map reflects the effect of drug on spontaneous rate in the absence of electrical stimulation, while the remaining rows delineate the range of pace-capture across drug dosages. The leftmost column of the map delineates the range of pace-capture under drug-free conditions, while the remaining columns are at different drug dosages. The localization of red blocks in the upper right of the maps for E-4031 and nifedipine (monolayer only) indicates loss of capture of paced beats in the presence of high concentrations of the drug and short PCLs. Localization of blue blocks in the lower right of the maps for BaCl2 indicates a higher fraction of samples with spontaneous activity as drug concentration increased, and the large area of green for chromanol 293B and nifedipine (EHS only) indicates very little response to the drug. Further, across the four drugs tested, in the GRIDS maps the regions of red or blue blocks tended to be smaller, and the region of green tended to be larger, for EHS than for monolayers, revealing that EHS are comparatively less sensitive to these drugs when evaluated for effects on their excitability and ability to capture during pacing. This suggests that EHS express relatively more of the repolarizing currents IKr and IK1,j, which are responsible for maintaining the resting potential in adult ventricular cells (Doss et al., 2012), than monolayers. On the other hand, the absence of a chromanol 293B effect supports the notion that both EHS and monolayers have low levels of IKr, as has been previously described for hiPSC-CMs (Ma et al., 2011). The GRIDS maps also indicate that EHS remain excitable at all PCLs and concentrations of nifedipine, whereas monolayers are unable to be pace-captured for every beat for some combinations of PCL and nifedipine concentrations. Increased ICa,L in EHS would explain why, for the same level of ICa,L block at a given concentration of nifedipine, EHS would retain enough residual ICa,L to remain excitable while monolayers would not. Alternatively, the excitability of EHS may be governed more by IKs than by ICa,L (as in more mature ventricular tissue) compared with that of monolayers (either due to differences in ion channels or because hiPSC-CMs in EHS are less depolarized), so that block of ICa,L does not decrease excitability.

Our GRIDS maps were able to differentiate the responses of EHS and monolayers to the panel of drugs tested even though measurements of a single electrophysiological parameter, APD prolongation, did not provide a clear snapshot of relative drug sensitivity. In both EHS and monolayers, E-4031 and BaCl2 prolonged APD, chromanol 293B did not change APD, and nifedipine shortened APD. Further, we did not detect differences in drug responses between EHS and monolayers with respect to APD for E-4031 (Figure 7A, i–iii) and chromanol 293B (Figure 7C, i–iii), but we did detect them for high...
concentrations of BaCl$_2$ (Figure 7B, i–iii) and nifedipine (Figure 7D, i–iii). One limitation of the GRIDS assay is the spontaneous rate of the cells, which dictates the lower bound on the pacing rates that can be applied.

Differences in drug responses of EHS, which tend to be less sensitive but more robust in their ability to be electrically paced over a wider range of rates and drug concentrations when compared with monolayers, can be attributed to a variety of factors. One possibility is that cells in EHS are a more densely packed, thicker syncytium that experiences a lower effective drug concentration than cells in monolayers where diffusion is not limited. Also, differences in the mechanical and biochemical environment in EHS compared with monolayers can result in differences in cell phenotype and AP morphology seen in EHS (more elongated cells with organized sarcomeres), which can affect drug responses. In addition, EHS experience an increase in non-myocytes over time, and this modulation of cell-cell interactions in the preparation could also affect drug responses. Multicellular preparations with large areas, such as EHS, will be needed in future studies to assess the risk for reentrant arrhythmias, which require room for circuitous wavefront propagation to occur. Further, EHS are tissue-like models that can be used in studies aimed at treatment discovery and at creating clinically relevant disease models.

**Conclusion**

EHS take advantage of the complex biochemical and structural cues of the myocardial ECM to guide the alignment of CMs. Seeded hiPSC-CMs organize as multicellular, anisotropic bundles that contract the EHS and propagate APs uniformly throughout the preparation. EHS can be used for long-term culture of hiPSC-CMs to interrogate processes of cell maturation and response to drugs over time. Drug sensitivity can be evaluated for a range of concentrations and under different pacing rates using the GRIDS assay, which has revealed differences in drug sensitivity between EHS and cell monolayers. In conclusion, EHS are tissue-like models that can be used in long-term electrophysiology and drug studies.

**EXPERIMENTAL PROCEDURES**

An extended description of the methods can be found in the Supplemental Information.

**Preparation of EHS**

Slices of porcine myocardium 12 mm in diameter and 300 μm in thickness were sectioned and decellularized as described previously (Blazeski et al., 2015). The dECM slices were spread on plastic 12-mm-diameter coverslips, with the perimeter of each slice wrapped around the edges of the coverslip. Coverslips with slices were placed in wells of standard 24-well culture plates and kept in PBS with antibiotics for up to 2 weeks before reseeding.

**hiPSC Differentiation and Culture**

WT and LQT2 hiPSC lines with a heterozygous A422T mutation in the potassium voltage-gated, subfamily H, member 2 channel (KCNH2), which is commonly referred to as the HERG channel (Spencer et al., 2014) were gifts from Dr. Bruce Conklin. Both hiPSC lines were differentiated using a monolayer-based protocol (Boheler et al., 2014; Wang et al., 2015). The age of the EHS (d26–d201) is given as the time in days from the start of hiPSC-CM differentiation (d0). On d9, the medium was switched to RPMI 1640 containing B-27 with insulin, and this medium was used for the remainder of time both before and after seeding on the dECM slices.

On d10–12, hiPSC-CM monolayers were dissociated using 0.05% trypsin-EDTA and plated on dECM slices affixed to coverslips at a density of 0.8–1.3 million cells/cm$^2$. EHS were maintained in culture for 16–191 days before evaluation by optical mapping or contraction measurements.

**Imaging of ECM and EHS**

Standard fixation and immunostaining techniques were applied to slices before and after decellularization to label F-actin, nuclei, collagen I, collagen III, and laminin. Unstained dECM slices were also imaged by SHG. EHS were fixed and stained using standard techniques for cTnI, α-actinin, connexin 43 (Cx43), vimentin, filamentous actin (F-actin), and nuclei (DAPI). All samples were imaged by confocal microscopy. Nuclear elongation and orientation were analyzed in confocal images of EHS and monolayers using custom MATLAB scripts. Cellular structures in EHS were imaged by transmission electron microscopy.

**Contraction Measurements**

WT d24–78 EHS were placed in a 35-mm tissue culture dish filled with Tyrode’s solution and maintained at 31°C ± 0.1°C for the duration of the experiment. A section of each EHS was detached from the edge of the coverslip so that it could move freely. Each EHS was paced at 1, 1.5, and 2 Hz while the free region was imaged with a CCD camera. A custom MATLAB script was used to segment the image and calculate the change in EHS area over time, which was used as a measure of contraction.

**Electrophysiological Studies**

Each EHS was placed in Tyrode’s solution and stained with 10 μM di-4-ANEPPS for 10 min at 37°C. The EHS was rinsed several times in a dish with warm Tyrode’s solution, and then immersed in Tyrode’s solution containing 10 μM blebbistatin to suppress contraction. The dish was placed on a 37°C heated stage for the duration of the experiment. At least 5 min after adding blebbistatin, the EHS was stimulated with a point electrode and optically mapped using a CMOS camera (MiCAM Ultima-L, SciMedia). The EHS was paced by 5 ms monophasic rectangular pulses at stepwise increasing rates starting at 0.5 Hz. For some samples, E-4031, chromanol 293B, nifedipine, Bay K 8644, BaCl$_2$, or cromakalim were added for 7 min before mapping. Mapping data were analyzed using custom MATLAB scripts (details provided in
Supplemental Information). APDs at 30% and 80% repolarization (APD30 and APD80) were calculated from the optical voltage signal. For drug studies, APD and CV measurements at each concentration were plotted as a percentage of APD and CV measured at baseline, with no drug present.

**Statistics**
All data are presented as mean ± SD. A Wilcoxon rank-sum test was used to determine statistical significance between control and drug groups for WT EHS contraction experiments, and between WT and LQT2 EHS. Paired, unequal variance, two-tailed t tests were performed to determine the statistical significance between experimental measurements of d62–82 and d201 EHS. Differences were considered statistically significant at p < 0.05.

**SUPPLEMENTAL INFORMATION**
Supplemental Information can be found online at https://doi.org/10.1016/j.stemcr.2019.04.002.

**AUTHOR CONTRIBUTIONS**
A.B. and L.T. designed the experiments, reviewed and interpreted the data, and wrote the manuscript. A.B. and R.Z. performed cell culture and differentiation under the guidance of K.R.B. J.L. performed the flow cytometry analysis. J.E. performed the slice orientation analysis. A.B. collected and analyzed the rest of the data. All authors edited the manuscript.

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Supplemental Information

Functional Properties of Engineered Heart Slices Incorporating Human Induced Pluripotent Stem Cell-Derived Cardiomyocytes

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Figure S1. Characterization of dECM slices. Related to Figure 1. Immunostaining of slices for F-actin (green) and DAPI (blue) indicates the presence of (A) cellular content before but (E) not after decellularization. The extracellular matrix components for (B,F) collagen I, (C,G) collagen III, and (D,H) laminin in the slice both before (top row) and after (bottom row) decellularization. Second harmonic generation imaging showing the collagen fibers in (I) a slice sectioned and decellularized from a fresh porcine heart, (J) a slice sectioned from a porcine heart frozen for > 48 hours at -80°C and subsequently decellularized, and (K) a slice originating from a plug frozen for > 48 hours at -80°C and subsequently sectioned and decellularized. (L) The degree of orientation of the collagen fibers was similar in all three groups, n = 8 for each group. Error bars denote standard deviation. NS indicates means are not significantly different.
Figure S2. Characterization of cell populations. Related to Figure 2. Cardiomyocytes were analyzed via flow cytometry at several stages: on day 17 (d17) after initiating cardiomyocyte differentiation according to (A) standard protocols, (B) after re-plating into 2D tissue culture plates at d17 and long-term maintenance in culture for an additional 61 days, and (C) after re-plating onto dECM slices at d17 to form EHS and maintained in culture for an additional 61 days. Data shows percentage of non-cardiomyocyte (cTnT-, blue) cell and percentage of cardiomyocytes (cTnT+, red) in each panel. Histogram counts were extracted by gating on unstained cells and cells stained only with a secondary antibody control (to remove autofluorescence and background nonspecific secondary staining).
Figure S3. Morphology of hiPSC-CMs on EHS. Related to Figure 2. EHS seeded with hiPSC-CMs had multiple layers of cells, with (A,C) CMs in contact with the ECM of the slice exhibiting greater alignment than (B,D) CMs in a second layer sitting on top of the first CM layer, farther from the slice and forming the surface of the EHS. (D) EHS also exhibited localization of Cx43 around the perimeters of the cells (white arrows). Top row images are from d26 EHS and bottom row images are from d69 EHS.
**Figure S4.** Contraction of EHS in response to different pacing rates. Related to Figure 3. (A) Contraction traces of EHS exposed to 1μM isoproterenol plotted in Fig. 3C. Each cycle of contraction and relaxation was fit by a polynomial of degree 5 using a least squares fitting method. The data for each cycle are plotted in a different color, and the fitted curve for that cycle plotted on top using a dashed black line. The difference in the minimum (representing maximum EHS contraction during one cycle) calculated by our data compared to that calculated by the best fit line was 0.02±0.01% for the control, 0.06±0.04% for the isoproterenol, and 0.05±0.03% for the washout traces plotted in Fig. 3C. (B) Contraction amplitudes at 500 ms and 1000 ms pacing rates were normalized by the contraction amplitudes at a pacing rate of 666 ms. Error bars denote standard deviation. NS indicates mean is not significantly different from 1. n ≥ 6 for each group.
Figure S5. **EHS comprised of LQT2 hiPSC-CMs.** Related to Figure 4. EHS were seeded with LQT2 hiPSC-CMs 10 days after the start of differentiation and maintained in culture for 13 days before evaluation (at d23). (A) Immunostaining for cardiac troponin I (green), DAPI (blue) and vimentin (magenta) reveals alignment of hiPSC-CMs. (B) Cardiomyocytes on LQT2 EHS stained with α-actinin (green) and F-actin (red) had striations typical of sarcomeric structures (inset scale bar = 10 µm). (C) Confocal images of the EHS in (A) display the distribution of cardiomyocytes (stained for cardiac troponin I, green) and non-myocytes (stained for vimentin, magenta). Images were taken at intervals of 1.7 µm from the surface of the EHS. Cell nuclei are indicated by DAPI staining in blue. (D) An orthogonal slice of the entire volume reveals multiple layers of cells, with cardiomyocytes on the surface (top arrow) and non-myocytes underneath, closest to the ECM of the slice (bottom arrow). (E) Activation map of EHS with LQT2 hiPSC-CMs paced at 1000 ms cycle length shows anisotropic conduction. Black lines are isochrones at 40 ms intervals. Rectangular symbol indicates pacing site. (F) Average traces for wild type (WT, black) and LQT2 (blue) EHS paced at 1000 ms cycle length reveal differences in action potential morphology. (G) APD_{80} and APD_{30} were greater in LQT2 EHS versus WT EHS at 1000 ms and 700 ms paced cycle lengths. Error bars denote standard deviation. n = 5 for WT EHS and n = 3 for LQT2 EHS.
Figure S6. Action potential morphologies and qRT-PCR analysis of ion channel transcripts. Related to Figure 6. Sample action potential traces (A) illustrate the differences in action potential shape of EHS and monolayers (d40-70). Action potential durations (B, APD$_{80}$ values are connected using solid lines and APD$_{30}$ values are connected by dashed lines) plotted for pacing cycle lengths ranging from 2000 ms to 500 ms. Error bars denote standard deviation. * indicates p<0.05 when comparing mean APD$_{80}$ for monolayers and mean APD$_{80}$ for EHS at the specified cycle length. & indicates p<0.05 when comparing mean APD$_{10}$ for monolayers and mean APD$_{30}$ for EHS at the specified cycle length. n = 28-31 for EHS and n = 15-16 for monolayers. (C) Expressions of ion channel transcripts were altered in d69 EHS compared to control d22 hiPSC-CMs prior to seeding on EHS. n = 3 for EHS and n = 4 for controls. Error bars indicate standard deviation.
Figure S7. Application of cardioactive drugs to EHS after prolonged culture. Related to Figures 6 and 7. (A) Sample traces of EHS paced at 700 ms indicate the changes in action potential shape in the presence of increasing concentrations of Bay K 8644. (B) Bay K 8644 prolonged APD₈₀, and (C) decreased conduction velocity in a concentration-dependent manner. (D) Bay K 8644 also decreased the maximum capture rate. (E) Sample traces of d201 EHS paced at 1900 ms at baseline (black trace), after the application of 1 μM cromakalim (red trace), and after the additional application of 1 μM Bay K 8644 (green trace) indicate changes in action potential shape. (F) Cromakalim and Bay K 8644 both decreased APD₈₀ and (G) slowed conduction velocity compared to baseline values. Baseline values of APD₈₀ and conduction velocity before the application of drug are indicated in (B,C,F,G). Error bars denote standard deviation. n = 3 in (B,C,D) and n = 4 in (F,G). * indicates p<0.05 when comparing the percent change to 0 using paired, unequal variance t-test.
Video S1. Contracting EHS. Related to Figure 3A-C. (Top) Free edge of EHS shortens in response to pacing at a cycle length of 666ms. (Bottom) The degree of contraction is indicated by the continuous plot of EHS area, which decreases from 100% (the maximum area at rest) with each paced beat.

Video S2. Optical mapping video of EHS. Related to Figure 4A. The EHS was paced in the center at a cycle length of 500ms, and the sample action potential recordings are from the site denoted by the magenta dot on the optical map. The bottom trace indicates the pacing pulses, and the vertical dashed line spanning both the upper and lower traces indicates the time point in the action potential corresponding to the displayed optical map.
Table S1. Electrophysiological measurements of d54-58 EHS. APD30, APD80, longitudinal and transverse conduction velocities, and anisotropy ratios correspond to plots in Figure 4. Data represent mean±SD.

| Paced Cycle Length (ms) | n  | APD30 (ms) | APD80 (ms) | Longitudinal Conduction Velocity (cm/s) | Transverse Conduction Velocity (cm/s) | Anisotropy Ratio |
|-------------------------|----|------------|------------|----------------------------------------|---------------------------------------|-----------------|
| 400                     | 4  | 129±3      | 183±6      | 8.9±1.5                                | 7.4±1.0                               | 1.2±0.1         |
| 500                     | 6  | 152±15     | 214±20     | 11.6±2.1                               | 8.3±1.7                               | 1.4±0.2         |
| 600                     | 6  | 165±19     | 235±23     | 13.6±2.1                               | 9.7±1.7                               | 1.4±0.1         |
| 700                     | 6  | 178±24     | 253±29     | 15.4±2.4                               | 10.9±1.8                              | 1.4±0.1         |
| 800                     | 6  | 186±28     | 267±32     | 16.6±1.8                               | 11.4±0.9                              | 1.5±0.1         |
| 900                     | 6  | 198±24     | 283±34     | 18.9±2.3                               | 13.0±1.3                              | 1.5±0.2         |
| 1000                    | 6  | 211±21     | 295±32     | 18.8±1.3                               | 13.4±1.4                              | 1.4±0.1         |

Table S2. Electrophysiological measurements of d62-82 and d201 EHS. APD30, APD80, and conduction velocities correspond to plots in Figure 5. Data represent mean±SD.

| Paced Cycle Length (ms) | APD30 (ms) d62-82 (n=3) | APD30 (ms) d201 (n=5) | APD80 (ms) d62-82 (n=3) | APD80 (ms) d201 (n=5) | Conduction Velocity (cm/s) d62-82 (n=3) | Conduction Velocity (cm/s) d201 (n=5) |
|-------------------------|--------------------------|-----------------------|--------------------------|-----------------------|-----------------------------------------|---------------------------------------|
| 400                     | 93±14                    | 139±19                | 139±19                   | 9.5±1.5               | 11.1±1.5                                | 12.3±1.7                              |
| 500                     | 111±11                   | 162±16                | 162±16                   | 13.2±1.8              | 13.8±1.9                                | 14.4±2.0                              |
| 700                     | 128±16                   | 184±21                | 184±21                   | 15.9±1.7              | 16.6±1.8                                | 14.6±2.0                              |
| 900                     | 137±15                   | 194±20                | 194±20                   | 17.8±1.6              | 20.2±21                                 | 15.1±1.7                              |
| 1000                    | 143±15                   | 202±21                | 202±21                   | 18.1±1.5              | 213±23                                 | 15.4±1.3                              |
| 1200                    | 153±18                   | 215±22                | 215±22                   | 18.4±1.8              | 212±22                                 | 16.0±1.3                              |

\[72x579\]
Supplemental Experimental Procedures

Unless otherwise stated, reagents were acquired from Thermo Fisher Scientific, Waltham, MA.

1. Preparation of EHS

Hearts obtained from slaughterhouse pigs were rinsed in distilled and deionized water to remove blood and stored overnight at -20°C. The following day, the hearts were allowed to thaw at room temperature for 1 hour. A metal 12 mm diameter punch was sterilized using 70% ethanol and used to punch out plugs of tissue from the left ventricle. Plugs were trimmed in order to fit into 35 mm culture dishes and stored at -80°C until slicing, a minimum of 16 hours. Plugs were allowed to thaw in room temperature distilled water supplemented with 100 U/mL Penicillin-Streptomycin and 0.1X antibiotic-antimycotic. After thawing, plugs were blotted dry, placed in a 35 mm culture dish with the epicardium pressed against the bottom of the dish, and embedded in 4% w/v low gelling temperature agarose (Sigma-Aldrich Corp., St. Louis, MO) dissolved into distilled water with penicillin-streptomycin and antibiotic-antimycotic. The agarose was allowed to solidify at 4°C for 15 minutes, and then the agarose disc containing the plug was removed from the culture dish and attached to the cutting stage of a vibratome (7000smz, Campden Instruments, Lafayette, IN) using cyanoacrylate glue (3M, Maplewood, MN) with the epicardium positioned at the top of the plug. The plug was sectioned into 300μm-thick slices parallel to the epicardium using a ceramic blade oscillating at a frequency of 100 Hz with an amplitude of 1 mm and advancing at a speed of 0.01 to 0.03 mm/second. The cutting solution in which the plug was immersed (phosphate buffered saline (PBS) supplemented with antibiotics) was kept at 4-8°C during slicing. Slices were stored in PBS supplemented with antibiotics at 4°C overnight.

Slices were decellularized using a procedure modified from Ott et al. (Ott et al., 2008). All detergents for decellularization were diluted in distilled water supplemented with antibiotics. PBS was also supplemented with the same antibiotics. Slices were each placed in a well of a 12-well plate, and 1 mL of each of the decellularization solutions was added to each well. The plate was placed on a rotator (Orbit 1000 Digital Shaker, Labnet International Inc., Edison, NJ) and agitated 180 rpm in the presence of the following solutions: 1% sodium dodecyl sulfate (SDS) for 3 hours (replaced with fresh solution after 1.5 hours), water for 15 minutes, 1% Triton-X 100 (Sigma-Aldrich Corp.) for 7 minutes, and PBS for 45 minutes (replaced with fresh solution every 15 minutes). Samples were left in PBS on rotator at 160 rpm overnight to rinse out any remaining detergents.

Plastic 12 mm coverslips were immersed in 70% ethanol and wiped dry. After an overnight wash in PBS, slices were carefully handled with forceps, spread over the coverslips and wrapped around the edges of the coverslips. Slices attached to the coverslips were placed into wells of a 24 well plate, immersed in PBS with antibiotics, and stored at 4°C until seeding (up to 2 weeks).

2. hiPSC Differentiation and Culture

Wild type and LQT2 hiPSCs were plated into wells of 6-well plates coated with 1:200 Geltrex:DMEM/F-12. For the first 22 hours, hiPSCs were maintained in Essential 8 medium (E8) with 10 μM Y-27632 dihydrochloride (Tocris Bioscience, Bristol, UK). Afterwards, hiPSCs were rinsed with DMEM/F-12 and fed with E8 medium every day. On the fourth day, when cells had reached about 80% confluence, the medium was replaced with RPMI 1640 supplemented with B-27, minus insulin and 6 μM CHIR-99021 (Selleck Chemicals, Houston, TX) to initiate differentiation (d0 of differentiation). Over the course of the next week, medium was changed as follows: RPMI 1640 with B-27 without insulin (B-27 minus) on d2, B-27 minus and 5 μM IWR-1 (Sigma-Aldrich Corp.) on d3, B-27 minus on d5 and d7, and RPMI 1640 with B-27 with insulin (B-27 plus) on d9 and every other day afterwards. Spontaneous beating in the monolayers was observed starting at d7 to d10. On d10 to d12, the hiPSC-CM monolayers were washed with 0.5 mM EDTA (Mediatech, Inc., Manassas, VA) and then incubated in EDTA for 5 minutes at 37°C. Afterwards, the EDTA was aspirated off, and 0.05% Trypsin-EDTA was added for 3 minutes at 37°C. Cells were trypared before Defined Trypsin Inhibitor was added to stop the digestion. The resultant suspension was centrifuged at 200 g for 5 minutes (Centrifuge 5702, Eppendorf AG, Hamburg, Germany). After aspirating off the supematant, the cell pellet was resuspended in B-27 plus. The PBS was removed from dECM slices, and the suspension of hiPSC-CMs was pipetted on top of the slices at a density of 0.8-1.3 million cells per cm². Slices were maintained in culture for 16 to 190 days, and the B-27 plus medium was replaced every other day for the duration of culture.

3. Imaging of Extracellular Matrix and EHS

dECM slices or EHS were fixed in 4% paraformaldehyde solution (Affymetrix, Inc., Cleveland, OH) for 10 minutes and rinsed twice with PBS. Samples were stored in PBS at 4°C until immunostaining. To immunostain for ECM proteins, slices were immersed in Target Retrieval Solution (Dako North America, Inc., Carpinteria, CA) for
were of the orientation FFT was calculated by converting from Cartesian to polar coordinates. The magnitude of the 2D subregion continuously shifted to the were fiber degree of alignment. SHG acquired from the same hearts that were stored at decellularized slices (native (in cacodylate buffer. After undergoing two rounds of rinsing with water for 5 minutes each, the samples were stained shielded from light, and left for one hour in a solution of 1% osmium tetroxide in 3 mM MgCl₂ times (15 minutes per rinse) in a 0.1 M sodium cacodylate buffer supplemented with 4',6-Diamidino-2 Phenylindole, Dihydrochloride (DAPI) for 25 minutes at room temperature. Samples were subjected to three more rounds of TBS-T washing before the addition of a drop of Prolong Gold Antifade Mountant and a glass slide on top of each sample. Alternatively, some samples were permeabilized as previously described and stained with Alexa Fluor 488 Phalloidin and DAPI at room temperature for 25 minutes. Samples were washed in TBS-T and mounted for imaging as described above. Images of stained samples were acquired using a confocal microscope (LSM 510 Meta, Zeiss, Oberkochen, Germany). Fibrillar collagen in unstained dECM slices was also imaged by second harmonic generation (SHG) using a multiphoton microscope (710NLO, Zeiss) with excitation at 880 nm, and emission acceptance at 415-450 nm.

To immunostain for cellular proteins, EHS were permeabilized with cold 0.5% Triton-X 100 (Sigma-Aldrich Corp.) in PBS for 20 minutes, followed by blocking with 10% goat serum (Life Technologies, Carlsbad, CA) for 1 hour at room temperature. Primary antibodies against cardiac troponin I (T8665-13F, United States Biological, Pittsburg, PA), α-actinin (A7811, Sigma-Aldrich Corp.), connexin 43 (C6219, Sigma-Aldrich Corp.), or vimentin (M0725, Dako North America, Inc.) in antibody diluent (Dako North America, Inc.) were added overnight at 4°C. The next day, EHS were washed with TBS-T as described above. Afterwards, samples were stained with 4',6-Diamidino-2 Phenylindole, Dihydrochloride (DAPI) for 25 minutes at room temperature. Samples were subjected to three more rounds of TBS-T washing before the addition of a drop of Prolong Gold Antifade Mountant and a glass slide on top of each sample. Alternatively, some samples were permeabilized as previously described and stained with Alexa Fluor 488 Phalloidin and DAPI at room temperature for 25 minutes. Samples were washed in TBS-T and mounted for imaging as described above. Images of stained samples were acquired using a confocal microscope (LSM 510 Meta, Zeiss).

To prepare samples for transmission electron microscopy, EHS were fixed in 2.5% glutaraldehyde and 3 mM MgCl₂ in 0.1 M sodium cacodylate buffer (pH 7.2) overnight at 4°C. Samples were subsequently rinsed three times (15 minutes per rinse) in a 0.1 M sodium cacodylate buffer supplemented with 3 mM MgCl₂ and 3% sucrose, shielded from light, and left for one hour in a solution of 1% osmium tetroxide in 3 mM MgCl₂ and 0.1 M sodium cacodylate buffer. After undergoing two rounds of rinsing with water for 5 minutes each, the samples were stained in 2% aqueous uranyl acetate in the dark for 1 hour. Samples were then dehydrated in a graded series of ethanol washes (30%, 50%, 70%, 90%, and 100% of ethanol), followed by two washes with propylene oxide (for 5 minutes each) and an overnight incubation in 1:1 propylene oxide/epon. The next day, samples were incubated in epo with catalyst three times for 2 hours each at room temperature and for 2 days at 60°C. Samples were examined using a transmission electron microscope (Philips/FEI BioTwin CM120 TEM, Hillsboro, OR).

4. Calculation of Collagen Fiber Orientation

SHG images were acquired of three types of decellularized slices (dECM slices) after decellularizing: fresh slices (native slices from sections of fresh plugs prepared right after the hearts were acquired that were decellularized), frozen slices (fresh slices that were stored at -80°C for over 48 hours), and frozen plus (fresh slices that were stored at -80°C for over 48 hours, thawed, and subsequently sectioned). Both fresh and frozen slices were acquired from the same hearts, and fresh and frozen slices were derived from adjacent sections of the same plug. SHG z-stack images had an area of 1.875 mm² and were processed in MATLAB to quantify fiber orientation and fiber degree of alignment. After compressing z-stack images in ImageJ using a maximum intensity z-axis projection, each compressed image was converted to an RGB image using ImageJ and segmented in MATLAB. SHG images were then segmented into 6889 subregions. The sectioning frame originated at the top left corner of the image and continuously shifted to the right or downwards by 0.0133 mm to create a new subregion with an area of 0.071 mm².

The absolute logarithmic magnitude of the 2D Fast-Fourier transform (2D-FFT) was calculated for each subregion in the segmented image. The direct current frequency was shifted to the center of the 2D-FFT plot, and the magnitude of the 2D-FFT was normalized so that it had a magnitude of 1. Since the 2D-FFT is symmetric, only half of it was used in the analysis. The angle of orientation from the horizontal axis of each coordinate in the 2D-FFT was calculated by converting from Cartesian to polar coordinates.

The radial sum of the 2D-FFT, normalized to the number of pixels along each radius, was plotted as a function of the orientation for each subregion. The orientation of the maximum 2D-FFT sum determined the angle perpendicular to the fiber orientation in that subregion. The mean and standard deviation of the fiber orientation were calculated for all subregions in each image. Mean values of fiber orientation were plotted for 8 slices within
each group. The fiber degree of alignment was calculated as the maximum 2D-FFT sum divided by the total sum of the 2D-FFTs of all subregions within an image.

5. Calculation of Nuclear Elongation and Alignment

Time-matched EHS and monolayers were immunostained with DAPI and imaged using a confocal microscope, as described above. Images of nuclei were thresholded above a baseline noise level and segmented, removing overlapping areas as necessary, by using a previously described method (Plissiti et al., 2014) with some modifications. A recursive search for concavity points was performed; when a concavity point was found, the outline between it and the convex hull was searched for more concavities. Further, the case where non-adjointing sections of an outline are part of the same nucleus was permitted. An ellipse was fitted to each nucleus using a method previously described (Fitzgibbon et al., 1999). The nuclear elongation ratio, mean angular orientation, and standard deviation of the angular orientations were calculated, as previously described (Bray et al., 2010).

6. Flow Cytometry Analysis

Flow cytometry analysis was performed on d17 (prior to monolayer seeding), and on monolayers and EHS that had been seeded at d17 and cultured for 61 days (d68). Monolayers and d17 cells were washed with 5 mM EDTA and subsequently incubated in 5 mM EDTA at 37°C for 5 minutes. The EDTA solution was aspirated off and replaced with 0.05% Trypsin-EDTA for 4-6 minutes at 37°C, until the cells easily detached from the bottom of the wells when the culture dish was agitated. The Trypsin-EDTA solution containing cells was neutralized using Defined Trypsin Inhibitor, and the cells were centrifuged at 200 g for 5 minutes before fixation. EHS were unhooked from their underlying plastic coverslip, transferred to a 12-well plate and rinsed with DPBS containing calcium and magnesium (DPBS +/+). After removing the DPBS, EHS were incubated in 10 mg/mL of collagenase IV with 10% fetal bovine serum in DPBS +/+ and 50 ug/mL of DNAse I in 0.15 M NaCl (Sigma-Aldrich Corp.) for 30 minutes on a shaker at 37°C. The collagenase and DNase solution were subsequently removed, and the EHS were rinsed twice with PBS (without calcium or magnesium). After removing the PBS, EHS were incubated in 0.05% Trypsin-EDTA for 5 minutes on a shaker at 37°C and broken up by pipetting up and down. The Trypsin-EDTA solution containing the recovered cells was neutralized and centrifuged as described above prior to fixation.

Dissociated cells were fixed in 4% paraformaldehyde (Affymetrix) for 10 min at room temperature. Cells were then simultaneously and permeabilized and blocked in PBS containing 0.1% BSA (Sigma-Aldrich Corp.), 5% goat serum, and 0.1% Triton-X 100 (Sigma-Aldrich Corp.) for 30 min. Cells were incubated with mouse anti-cTnT antibody diluted 1:200 in FACS buffer (PBS with 0.1% BSA and 0.1% Triton-X 100) for 1 hr at 4°C, washed three times with FACS buffer, and incubated with anti-mouse Alexa Fluor® 488 antibody in FACS buffer (1:200, Invitrogen) for 30 min at 22°C in the dark. After washing and re-suspending in PBS with 0.1% BSA, cells were strained through a 30 µm filter and run on a FACSCalibur cytomter (BD Biosciences, Woburn, MA). Secondary controls consisted of cells incubated only with anti-mouse Alexa Fluor® 488 secondary antibody. Single cells were identified and gated based on their forward and side scatter, and cardiomyocytes were gated based on their cTnT expression. Data were analyzed using FlowJo X Software. Histogram counts were extracted by gating on unstained cells and cells stained only with secondary antibody control (to remove autofluorescence and background nonspecific secondary staining).

7. Contraction Measurements

Tyrode’s solution was prepared by combining 1.8 mM CaCl₂, 5 mM glucose, 5 mM HEPES, 1 mM MgCl₂, 5.4 mM KCl, 135 mM NaCl, and 0.33 mM NaH₂PO₄ in ddH₂O and adding NaOH to raise the pH to 7.4 (all chemicals from Sigma-Aldrich Corp.). EHS were placed in a 35 mm dish filled with Tyrode’s solution and set on a stage heated to 31±0.1°C (Warner Instruments, Hamden, CT). A section of each EHS was unhooked and allowed to move freely throughout the duration of the experiment. The EHS were allowed to equilibrate in Tyrode’s for 5 minutes before the start of pacing with a point electrode. Each sample was paced for 1 minute at each of 3 cycle lengths (1000 ms, 666 ms, 500 ms), while the freely moving region was kept approximately vertical in the field of view and imaged at 4x magnification using a CCD camera (Swiftcam, Swift, Schertz, TX) at a rate of 14-17 fps with 320x256 pixel resolution. Samples were also imaged during pacing at 666 ms after the application of 1 µM isoproterenol (Sigma-Aldrich Corp.) for 2 minutes and after washout of the drug for 2 minutes. We applied a pacing cycle length shorter than 1000 ms in the presence of isoproterenol to overcome the increase in spontaneous rate in response to the drug. Custom MATLAB scripts were used to segment the images by applying a user-defined threshold to assign each pixel in the image as either belonging to the EHS, which was darker, or the background, which was lighter. From this, the area of EHS was calculated in each image. The change in area from a fully relaxed state (reference frame, designated as 100% EHS area) was determined for each frame in the time series. The minimum area (maximum change in area) was averaged over time (multiple cycles of contraction) and relaxation for
each pacing condition. The change in area in the presence of isoproterenol and after washout was compared to the change in area prior to the application of drug for each EHS.

8. Electrophysiological studies

EHS were stained with 10 μM of the voltage-sensitive dye di-4-ANEPPS (Sigma-Aldrich Corp.) in Tyrode’s solution for 10 minutes at 37°C. Afterwards, EHS were rinsed twice with Tyrode’s and placed in a 35 mm dish filled with Tyrode’s and 10 μM of the contraction inhibitor blebbistatin (Sigma-Aldrich Corp.). This dish was set on a stage heated to 37°C and allowed to equilibrate for at least 5 minutes. Samples were point paced with at least 30 stimulus pulses at a range of cycle lengths, starting from 2000 ms and decreasing until the they lost capture. EHS were optically mapped during pacing using a 100x100 pixel CMOS camera (MiCAM Ultima-L, SciMedia, Costa Mesa, CA).

During drug studies, recordings at a range of pacing rates were taken at baseline (prior to the addition of drug) before replacing the solution in the dish with Tyrode’s supplemented with blebbistatin and the lowest concentration of drug studied. The sample was paced at the same range of rates and mapped 7 minutes after the addition of drug. Afterwards, the bath solution was replaced with Tyrode’s with blebbistatin and the next lowest drug concentration and the procedure repeated. This method was applied for all concentrations of each drug and, except where noted in the figures, only one drug was tested per sample. In these studies, we superfused EHS or monolayers with the following drugs: E-4031 (Tocris Bioscience), BaCl₂ (Sigma-Aldrich Corp.), Chromanol 293B (Tocris Bioscience), Nifedipine (Tocris Bioscience), Bay K 8644 (Tocris Bioscience), and Cromakalim (Sigma-Aldrich Corp.).

Optical mapping data was analyzed using custom MATLAB scripts. Recordings at each pixel were de-noised using a previously described method (Little and Jones, 2010) to regulate total signal variance and convolved with a 5x5 spatial Gaussian filter. Activation times were defined as the maximum of the derivative of membrane potential (dV/dt), which was calculated as previously described (Chartrand, 2011). Histograms of local conduction velocities for each EHS were fitted to a Gaussian curve and the mean of the curve was defined as the average conduction velocity (CV). To determine longitudinal and transverse CVs, a bimodal Gaussian curve was fitted to the local CVs. Action potential durations at 30 and 80 percent repolarization (APD₃₀ and APD₈₀) were determined for all local traces over the recording region for each EHS and fit with Gaussian curves to determine the mean value for each EHS, as described for CV measurements. For drug studies, average APD and CV measurements for each concentration were normalized by average APD and CV measurements at baseline (without the drug).

9. Quantitative RT-PCR

mRNA was isolated from EHS (d69) and monolayers (d22) using the following procedure: incubation in TRIzol Reagent for 5 minutes at room temperature, incubation in chloroform for 3 minutes, centrifugation at 12,000 g for 15 minutes at 4°C, collection of colorless phase that separated at the top of the centrifuged sample, addition of isopropyl alcohol and incubation at room temperature for 10 minutes, centrifugation at 12,000g for 10 minutes at 4°C, solubilization of RNA pellet in 75% ethanol, centrifugation at 7,500g for 5 minutes at 4°C, air drying of sample, and resuspension of the RNA pellet in DEPC-treated water. mRNA from two EHS were combined for each EHS replicate.

Reverse transcription was performed to create cDNA with the PCR Master Mix kit, using the MyGo Mini PCR system (IT-IS Life Science Ltd., Republic of Ireland). RT-PCR was performed on each target in triplicate, using the following primers:

| Gene Target | Associated Protein                      | Primer         |
|-------------|----------------------------------------|----------------|
| CACNA1C     | L-type calcium channel                  | Forward CACGGCTTCCCTCGAATCTTG  |
|             | L-type calcium channel                  | Reverse CTGGAGATGGTGCTGCAATTG     |
| KCNH2       | Voltage-gated potassium channel         | Forward AGGAGCGAACCCACAATGTC    |
|             | Voltage-gated potassium channel         | Reverse AGGTGGTGGCGGAAAGTGTG    |
| TNNI1       | Troponin I, slow skeletal type          | Forward GGTGATGAGGAGCGATACG   |
|             | Troponin I, slow skeletal type          | Reverse CAGGCTGGAGGGAAGAATGTG   |
| TNNI3       | Troponin I, cardiac                     | Forward CCAACTACCGCGTCTATGCC  |
|             | Troponin I, cardiac                     | Reverse CTTCTCGGTCCTCCCTTC     |

The PCR program run for each sample consisted of 120 seconds hold at 95°C, 40 cycles of amplification that alternated between 90°C and 65°C, 10 seconds pre-melt hold at 95°C, and a melting step that increased from 60°C to 97°C at 0.1°C/second. CT values were obtained using MyGo Mini PCR Software (IT-IS Life Science Ltd.)
and CT was calculated for each gene of interest (GOI) in EHS to determine fold change over transcript levels expressed monolayers according to the formula:

\[ \Delta C_T = C_{EHS,GOI} - C_{EHS,ACTN2} - C_{Mn,GOI} + C_{Mn,ACTN2} \]

where \( C_{EHS,GOI} \) refers to CT of EHS for the GOI, \( C_{EHS,ACTN2} \) refers to CT of EHS for ACTN2 (α-actinin, which was used as a normalizing gene), \( C_{Mn,GOI} \) refers to CT of monolayers for the GOI, and \( C_{Mn,ACTN2} \) refers to CT of monolayers for ACTN2.

10. Statistics

All data are presented as mean±SD. Measurements of nuclear elongation were log transformed and reported as the interval of the log-transformed mean ± SD after inverse transformation into linear space (Bland and Altman, 1996). For contraction experiments, a one-tailed Wilcoxon test was used to determine statistically significant differences from 1 for isoproterenol-treated and washout groups. For studies on rate-dependence of contraction, a two-tailed Wilcoxon test was used to determine statistically significant differences from 1 for groups paced at 500 ms and 1000 ms cycle lengths. A two-tailed Wilcoxon test was also used for statistical significance between APD measurements for WT and LQT2 EHS. Paired, unequal variance, two-tailed t-tests were performed for all other drug studies and unpaired, unequal variance, two-tailed t-tests were performed for optical mapping studies not involving drugs, studies comparing d62-82 EHS to d201 EHS, and for orientation analysis of fresh slices, frozen slices, and frozen plugs.

11. Supplemental References

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