Mechanically Stimulated Contraction of Engineered Cardiac Constructs Using a Microcantilever

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Abstract—The beating heart undergoes cyclic mechanical and electrical activity during systole and diastole. The interaction between mechanical stimulation and propagation of the depolarization wavefront is important for understanding not just normal sinus rhythm, but also mechanically induced cardiac arrhythmia. This study presents a new platform to study mechanoelectrical coupling in a 3-D in vitro model of the myocardium. Cardiomyocytes and cardiac fibroblasts are seeded within extracellular matrix proteins and form constructs constrained by microfabricated tissue gauges that provide in situ measurement of contractile function. The microcantilever of an atomic force microscope is indented into the construct at varying magnitudes and frequencies to cause a coordinated contraction. The results indicate that changes in indentation depth and frequency do not significantly affect the magnitude of contraction, but increasing indentation frequency significantly increases the contractile velocity. Overall, this study demonstrates the validity of this platform as a means to study mechanoelectrical coupling in a 3-D setting, and to investigate the mechanism underlying mechanically stimulated contraction.

Index Terms—Atomic force microscopy, cardiomyocyte, mechanoelectrical coupling.

I. INTRODUCTION

T HE application of active mechanical stimulation like stretch to a cardiomyocyte can cause local depolarization, which leads to intracellular calcium release and contraction of the cell [1]–[3]. The process of mechanoelectrical coupling in cardiomyocytes has been extensively studied *in vitro* because of its implications for induction and termination of cardiac arrhythmias [4], [5]. To explore this phenomenon *in vitro*, previous studies have observed the electrical depolarization of single cells or monolayers plated on a 2-D substrate and exposed to either an applied force [6], [7] or fluid shear stress [8]. However, this configuration cannot mimic several aspects of the *in vivo*

Manuscript received January 2, 2014; revised July 3, 2014; accepted August 14, 2014. Date of publication September 15, 2014; date of current version January 16, 2015. The work of P. A. Galie was supported by National Institutes of Health under a T32 Interdisciplinary Cardiovascular Training Grant. *Asterisk indicates corresponding author*.

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This paper has supplementary downloadable material available at http://ieeexplore.ieee.org, provided by the authors (File size: 19MB).

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Digital Object Identifier 10.1109/TBME.2014.2357778

myocardium, not least of which is the three-dimensionality of native tissue.

Microscale tissue gauges provide a platform to study cardiomyocytes in a 3-D environment [9]–[11]. Mixtures of cardiomyocytes and fibroblasts isolated from neonatal rat hearts are combined with solubilized matrix and centrifuged into $800 \times 400 \ \mu$ m wells containing two flexible posts made from polydimethylsiloxane (PDMS) posts (referred to hereafter as microTUGs). After matrix polymerization and cell-mediated contraction, the mixtures form microscale constructs that are suspended between the PDMS posts. The formed constructs exhibit *in vivo*-like myofibrillar assembly and cell alignment. The force exerted by the contraction of the tissues can be measured *in situ* by observing the deflection of the PDMS posts. The system is a microscale adaptation of a previously described mesoscale cardiac tissue setup [12].

Traditionally, constructs within this system are stimulated by the application of an electric field to induce contraction. However, we have found that contraction of the constructs can also be instigated by indentation with the microcantilever of an atomic force microscope (AFM). AFM has been used extensively to measure the stiffness of cells and tissues in vitro [13]-[16], in addition to mechanically perturbing biological samples [17], [18]. Combining the tissue gauge platform with AFM creates a system capable of recapitulating the mechanoelectrical coupling of cardiomyocytes in a 3-D setting. The system also has the advantage of creating multiple samples for each condition since each substrate contains >100 wells, so that multiple tissues can be tested in each substrate with the rate limiting step being the use of AFM to perform the indentation. This study establishes this platform as a means to effectively study both mechanoelectrical coupling by measuring the effect of indentation magnitude and frequency on the response of the constructs.

II. MATERIALS AND METHODS

MicroTUG substrates were constructed using soft lithography. Briefly, SU-8 photoresist (MicroChem) was spun onto silicon wafers, exposed, and developed to create the geometry of the posts within the wells. PDMS (Dow-Corning) stamps were cast off the silicon masters, and then used to generate the PDMS microTUGs used for the experiments. A complete description of microTUG fabrication can be found in [9]. To generate the constructs, 750 000 cells of the suspension isolated from the ventricles of 2–3 day old Sprague-Daley rats were seeded in a 1 mg/mL salmon fibrin (Reagent Proteins, Pfenix, Inc.) gel supplemented with 0.25 mg/mL bovine collagen type I. Constructs were fed daily with M199 medium supplemented with 10% chick embryo extract and 0.1% antibiotic antimycotic solution.

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The solution contains 75 μ M streptomycin, a known inhibitor of stretch-activated channels. However, we found no significant difference in construct response to 1 Hz indentation with and without the presence of streptomycin.

At day 5, the microTUG substrates were transferred to the AFM setup, consisting of a Bioscope AFM (Bruker) probe fitted to an inverted microscope with a heated stage. A 3.5- μ m diameter bead was fixed to the probe tip and lowered into contact with 1 mL of medium bathing the microTUG substrate, and the stage was kept at 37 °C. Measurements were made within one hour after removing the substrate from the incubator, since CO₂ level was not controlled. Only nonspontaneously contracting constructs were targeted for mechanical stimulation (about 10-20% of the constructs spontaneously beat at 5 days in culture). Once contact was made with the construct, the indentation depth and frequency were set and the construct contractions were measured using a camera attached to the inverted microscope. The frame rate was set to 11 frames/s, and at least one full contraction was required for a successful measurement. To quantify the deflection of the microTUG posts, ImageJ was used to measure the displacement of the post during construct beating. Contraction velocity was calculated by dividing the post displacement during contraction by the time to peak.

After testing, the microTUG substrates were fixed in 3.7% paraformaldehyde, permeabilized in 0.1% Triton X-100, and stained with Texas Red phalloidin (1:50), DAPI (1:1000), and a primary antibody for alpha-actinin (1:100). Data are presented as mean \pm standard deviation for n = 3 tissues, and contraction magnitude and speed values were compared using one-way ANOVA and Tukey's post-hoc test with statistical significance accepted for p < 0.05.

III. RESULTS

A schematic of the testing configuration can be found in Fig. 1(a). The cell-seeded hydrogel (pink) is suspended between the two posts (blue) within the 400 \times 800 μ m microTUG substrate wells (gray), and the AFM probe (black) is lowered to the top surface of the construct. The dimensions of the post are shown in Fig. 1(b). The post has a 190 μ m head that the contracting fibroblasts wrap around to anchor the construct. A bright-field image from the high-speed camera is shown in Fig. 1(c). The AFM tip (triangular shape) is placed over the center of the construct, which is attached to the opposing TUG posts. Fig. 1(d) shows a construct stained for nuclei (blue), actin (red), and alpha-actinin (green). The stain demonstrates self-organization of the cell-seeded constructs. Alpha-actinin concentrated in the z-disks of the sarcomeres distinguishes the cardiomyocytes from fibroblasts (since actin is present in both cell types). Fibroblasts are present in the outer edges of the construct, while cardiomyocytes are located primarily in the center. Therefore, the AFM probe makes contact on or near the centerline of the construct for all measurements to assure that the tip is not indenting an area devoid of cardiomyocytes.

Mechanical stimulation was implemented by displacing the base on the AFM cantilever at a constant rate for a specified distance and then immediately retracting the probe at the same



Fig. 1. (a) TUG schematic indicating the position of the AFM probe as it contacts the microtissue. (b) Postdimensions. (c) Bright-field image of the construct just prior to contact with the AFM probe (scale bar = 50 μ m). (d) Construct stained with Texas Red phalloidin (red), DAPI (blue), and alpha-actinin (green). Scale bar = 20 μ m.

rate to form an indentation ramp. Each indentation cycle had a period of 100 ms. Indentations of the sample by the bead attached to the AFM probe using a ramp size greater than 0.5 μ m were found to induce a net contraction of the construct. Assuming the construct has a resting elastic modulus of 5 kPa [19], a 0.5 μ m indentation corresponds to a normal force of approximately 3.9 nN. Having discovered that indenting with the AFM tip causes a coordinated contraction of the construct, we sought to determine if the indentation depth and frequency affected the resultant contraction force of the tissue. We modulated the indentation depth by setting the ramp size of the AFM tip to 1, 2, and 3 μ m. The ramp size does not exactly match the indentation depth of the probe because the construct causes deflection of the probe tip relative to the probe head, and actual indentation depths were not measured due to the pacing of the AFM probe during testing. Using the spring constant of the cantilever and a Hertz model for contact with a substrate of elastic moduli ranging from 1 to 5 kPa, we calculated the indentation depth for a given ramp size (Supplementary Fig. 1) and found that at least 80% of the ramp size translated to construct indentation. Three AFM indentation frequencies were chosen (0.5, 1, and 2 Hz), resulting in a total of nine testing conditions. For control experiments, nonstimulated spontaneously contracting tissues were analyzed.

Fig. 2(a) shows the averaged peak force exerted by the constructs, calculated from measurements of the TUG post displacement and a spring constant of 0.45 μ N/ μ m [10] for the three indentation lengths and frequencies. The data indicate that



Fig. 2. (a) Peak force as a function of indentation ramp size of the AFM probe. (b) Contraction speed as a function of ramp size. (* denotes p < 0.05). (c) Example traces of constructs stimulated by 2, 1, and 0.5 Hz. The timing of indentation is represented by a dotted gray line.

ramp size has no significant effect on the resultant contraction force magnitude. The varying ramp sizes and frequencies all induced an approximately 10- μ m displacement of both of the TUG posts, corresponding to a peak force of between 4 and 5 μ N. These values are within the range of force values reported using electrical stimulation (4–8 μ N) [10]. We also measured the contractile velocity to determine if indentation depth and frequency had any effect on the velocity at which the constructs contract. Fig. 2(b) shows that ramp size did not significantly affect contraction velocity. However, an indentation frequency of 2 Hz induced a significantly higher contraction velocity compared to constructs stimulated at 0.5 Hz. Interestingly, the frequency of the AFM-stimulated contraction events did not mirror the frequency of the indentation, but rather remained approximately at 1 Hz. Therefore, after the initial contraction, the indentation was not synchronous with the initiation of a contraction event. This is evident in Fig. 2(c), which exhibits the force exerted by tissues stimulated with three indentation frequencies.

Fig. 3(a) compares averaged force–time traces of constructs stimulated by a 1 μ m ramp size at 0.5, 1, and 2 Hz with constructs undergoing spontaneous contractions. The traces were averaged over several periods and at least three different tissues to create a



Fig. 3. (a) Averaged traces of force generated by constructs stimulated by 1 μ m indentations at different frequencies and by spontaneously contracting constructs. (* denotes p < 0.05 compared to constructs stimulated by 2 Hz frequency.) (b) Time to peak, 20% relaxation, and 80% relaxation (relative to time to peak) for the average traces plotted in (a). (* denotes p < 0.05 compared to 1 and 0.5 Hz frequency, ** denotes p < 0.05 compared to AFM-stimulated constructs.)

representative waveform of the construct contraction. The figure illustrates that indentation frequency has a significant effect on the contraction and relaxation dynamics. Constructs stimulated by 2 Hz indentations display a shorter contraction period than 0.5 and 1 Hz. Interestingly, spontaneously contracting constructs display the shortest contraction period. Although in contrast to the indentation-driven contractions, the spontaneous beating did not occur at a consistent frequency. The waveforms are quantified in Fig. 3(b) by plotting the time to peak, time to 20% relaxation, and time to 80% relaxation (relative to time to peak). The time to peak is significantly shorter in constructs stimulated by 2 Hz indentations compared to 1 and 0.5 Hz. The spontaneously contracting constructs exhibit significantly different times to peak compared to all the indentation-induced contracting constructs, suggesting a quicker depolarization.

We next determined whether sustained indentations were necessary for continuous contractions. Because the contraction frequency is steady at approximately 1 Hz, the indentations are not necessarily in phase with the contractions after the initial stimulation. Moreover, we sought to determine whether a given ramp size could continue to stimulate a coordinated contraction, or if the construct would develop tolerance and require higher indentation depths over time. Fig. 4 shows an example waveform of a tissue stimulated with 1- μ m ramp sizes at a frequency of 1 Hz. AFM indentation is started, stopped, and then restarted. The figure shows that once the AFM stimulation is turned off (between 4 and 8 s), coordinated contractions abruptly terminate. Supplemental Video 1 illustrates the effect of pausing AFM



Fig. 4. Force trace of a construct stimulated by $1-\mu m$, 1 Hz indentations for two distinct cycles. Indentations are applied between 0.5 and 3.5 s, and 8 and 10 s.

indentation. In between indentation cycles, several cells within the construct contracted (as evidenced by the 1 μ N force peaks). However, there was no coordinated contraction during this time. For all the constructs tested, continuous AFM stimulation was required to sustain the contractile events. The longest period tested was 3 min.

IV. DISCUSSION

The method described in this study provides a novel means to elucidate the mechanoelectrical coupling of cardiomyocytes in a 3-D environment. Previous studies have shown that both active [6], [7] and passive [20], [21] mechanical stimulation can modulate the contractile dynamics of cardiomyocytes. However, previous studies were conducted on 2-D surfaces coated with extracellular matrix proteins, which is not as representative of the *in vivo* composition of the myocardium. The microTUG system provides a platform to study cardiomyocytes in a 3-D environment, and incorporation of AFM allows for controlled mechanical stimulation of these constructs. Additionally, the microTUG system allows for direct quantification of contraction velocity and force magnitude, which are defining properties of striated muscle.

AFM ramp sizes more than 0.5 μ m were found to induce coordinated contraction events at approximately 1 Hz, regardless of the indentation frequency (within a range from 0.5 to 2 Hz). Modulating indentation depth and frequency did not significantly affect the peak force exerted by the constructs. These results imply that the mechanism underlying the mechanically induced contraction is not affected by the extent of the probe indentation, once a certain threshold required for contraction has been surpassed. The contraction velocity was also unaffected by indentation depth, but increased significantly with indentation frequency. Therefore, even though the resultant contraction force is unaffected by ramp size or frequency, the time to peak of the contraction can be controlled by the frequency of the microcantilever.

Beyond providing a platform for exploring mechanoelectrical coupling of cardiomyocytes in 3-D, the capability for mechanical point stimulation has additional benefits. Current stimulation techniques for microTUGs involve globally applied electric [9], [10] or magnetic [11] fields. Point stimulation provides the means to track the propagation of depolarization wavefronts in anisotropic conditions like ectopic foci. Although the setup used for this study was limited to frame rates of about 10 Hz, combining a higher frame rate camera with a voltage sensitive or calcium dye would allow for tracking of the depolarization and repolarization wavefronts. Changes could be made to the construct composition, stiffness, cell alignment, and other parameters to study their effect on how the depolarization wavefront propagates in response to changes in preload, afterload, and inotropy.

V. CONCLUSION

This study demonstrates the ability of AFM indentations to stimulate coordinated contractions of *in vitro* cardiac microtissues. Further research is required to determine the mechanism underlying this effect. Importantly, the system provides a means to investigate clinically relevant conditions such as the induction and termination of ventricular arrhythmias by acute mechanical stimulation, precordial impact, and mechanical pacing.

ACKNOWLEDGMENT

P. A. Galie thanks R. Alvarez and A. Mu for their assistance in isolating the cardiomyocytes and cardiac fibroblasts.

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