Structural and mechanical remodeling of the cytoskeleton studied in 3D microtissues under acute dynamic stretch

Condensed Running Title: The cytoskeleton in microtissues

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Abstract

When mechanically stretched, cells cultured on 2D substrates share a universal softening and fluidization response that arises from poorly understood remodeling of well-conserved cytoskeletal elements. It is known, however, that the structure and distribution of these cytoskeletal elements are profoundly influenced by the dimensionality of a cell’s environment (ie. on a 2D surface vs. within a 3D matrix). Therefore, in this study we aimed to determine whether cells cultured in a 3D extracellular matrix also follow the same softening response and to link this mechanical change to direct evidence of cytoskeletal remodeling. To achieve this, we developed a new high-throughput approach to measure the dynamic mechanical properties of cells and allow for sub-cellular imaging of physiologically relevant 3D microtissue cultures. We found that fibroblast, smooth muscle and skeletal muscle microtissues strain softened but did not fluidize, and upon loading cessation, they fully regained their initial mechanical properties. These responses required a filamentous actin cytoskeleton, and were mirrored by changes to actin remodeling rates, and direct visual evidence of actin depolymerization during stretching and repolymerization after stretch cessation. On the other hand, the response could not be attributed to either remodeling of microtubules or myosin motor activity. Our new approach for assessing cell mechanics has linked behaviors seen in 2D cultures to a soft 3D extracellular matrix, and connected visual remodeling of the cytoskeleton to changes in mechanical properties at the tissue-level.

Significance Statement

With every breath and movement, cells in our body are subjected to mechanical forces. These forces are key regulators of normal development and function, as well as disease progression. To understand how cells “feel” mechanical cues in their microenvironment, we have previously relied on two-dimensional experimental approaches and often assessed single cells in isolation. Here, we present a novel lab-on-chip device, which enables simultaneous mechanical stimulation and sub-cellular imaging of three-dimensional multi-cellular microtissues. In this article with this device, we quantitatively linked force-induced mechanical changes in microtissues to specific molecular remodelling pathways in the cytoskeleton. The approaches and insights presented in this study will deepen our understanding of the mechanobiological pathways governing tissue development and function in health and disease.
Introduction

With every breath, heartbeat and movement, cells in our body experience mechanical stretch, which in turn, creates continually unstable forces at focal adhesions, across the cell membrane, along cytoskeletal filaments and through the nucleus \textsuperscript{1,2}. In a cell, these forces direct functional and phenotypic behaviors by generating conformational changes, and thereby, altering ligand-receptor affinities \textsuperscript{1,2}. Importantly, this ability of cells to feel and adapt to mechanical forces has been linked to crucial events in normal development and function, as well as disease progression, including bone, muscle, heart and lung disorders, and cancer \textsuperscript{3,4}.

In particular, the well-conserved structural elements that make up the cytoskeleton of eukaryotic cells are in themselves mechanosensitive; in response to dynamic stretch, the cytoskeleton softens (decreased elasticity) and becomes more fluid-like \textsuperscript{5–8}. Then upon stretch cessation, it slowly regains its stiffness and resolidifies \textsuperscript{7}. Currently the molecular mechanism(s) behind strain softening remains unclear as there is limited visual-based evidence quantifying cytoskeletal remodeling following cyclic stretching \textsuperscript{9–11}. Furthermore, softening and fluidization has been observed in response to deformation at the subcellular \textsuperscript{5,6} and single cell levels \textsuperscript{7,8}, however the extent of the response remains poorly understood at the tissue-level \textsuperscript{12–15}. Nevertheless, in the body, this response has been linked to the maintenance of airway caliber \textsuperscript{12,13} and the regulation of blood pressure \textsuperscript{16}, but for unknown reasons, it is absent in certain pathological disorders. For example, unlike in healthy lungs, stretch from a deep inspiration does not dilate asthmatic airways \textsuperscript{17}.

Most of our knowledge of cell mechanics and how cells respond to mechanical forces has been generated by growing cells on rigid, flat surfaces. Yet it is known that the physical environment in which a cell is grown alters its mechanical properties and behavior. For example, cells grown on stiff substrates tend to have their actin cytoskeleton arranged into dense stress fibers, and are stiffer, more solid-like and under greater pre-stress when compared to cells on softer substrates \textsuperscript{18–21}. In addition to matrix stiffness, it is suspected that the mechanical behavior of cells may be further altered by the dimensionality of their environment. In support of this growing hypothesis, culturing cells on a 2D substrate vs. within a more physiologically relevant 3D matrix
fundamentally changes the distribution and structure of the cytoskeleton by forcing unnatural apical-basal polarity of adhesion complexes. The difference between a rigid, flat, petri dish and a soft 3D extracellular matrix may also explain observed disparities in cellular behavior, and the loss of efficacy in costly clinical trials that often occurs when pharmaceutical treatments are developed using conventional 2D cell culture techniques. Thus, there exists a need for new high-throughput cell culture techniques capable of probing cell mechanical behavior while maintaining a physiologically relevant soft 3D environment.

To address this need, techniques that allow assessment of the mechanical behavior of cells within reconstituted 3D collagen gels have been a keen interest to the fields of mechanobiology, pharmacology, and tissue engineering. These methods have furthered our understanding of how tissue level forces are collectively generated by cells, and their mechanical behaviors, including strain softening. The centimeter scale of these bulk gels, however, limits throughput, causes imaging difficulties, and produces a high diffusive barrier for nutrients.

To overcome the limitations of bulk 3D cell cultures and to study the rapid dynamics and force generation during contractility, Legant et al. (2009) developed Microfabricated Tissue Gauges (microtissues). In the microtissue model, cells are cultured within a matrix composed of collagen and form around pairs of flexible vertical cantilevers into an array of dense, organized structures comparable to ex vivo tissue. High-throughput tensile force measurements can then be calculated from the visible deflection of the cantilevers. More recently, investigators fixed a magnetic microsphere to one of the cantilevers in each microtissue well, and with magnetic tweezers stretched one microtissue at a time for quasi-static stiffness measurements. The limitations in experimental throughput and actuation range of magnetically driven devices were addressed by our recently published Microtissue Vacuum-Actuated Stretcher (MVAS). In that publication, the MVAS allowed for high-throughput visualization of cellular remodeling during stretching due to a mostly planar deformation and following chronic (several days) conditioning.

We now present a new microtissue stretcher, the MVAS-force, which enables measurements of tensile force and dynamic stiffness. In contrast to our previous design,
only one of the cantilevers in the MVAS-force is actuated through a regulated vacuum pressure and forces are measured simultaneously from the passive bending in the other cantilever. In this article, this new approach allowed us for the first time to assess the mechanical properties of microtissues during dynamic loading and upon loading cessation, and to link the changes in mechanics to sub-cellular remodeling using responses to pharmacological treatments and by directly imaging the cytoskeleton. The findings that can be gained from our approach on a cell's ability, or impaired ability to sense mechanical forces are critical to understand pathways of development, normal function and disease progression in the body.

Results

Microtissue Morphology

Within the MVAS-Force, 3T3 fibroblast cells self-assembled around the cantilevers into dense, highly organized, three-dimensional constructs that morphologically resemble tissue. Top-down and cross sectional views of a fully compacted, representative microtissue are shown in fig. 1c. As been shown previously \(^{33,36,37}\), the cells compacted the collagen matrix away from the bottom and sides of the well into a tissue freely suspended around the tops of the cantilevers. The average microtissue thickness measured at its center after four days was \(97 \pm 2 \mu m\) (n=5) and was qualitatively uniform along the longitudinal axis.

Maximum intensity projections with orthogonal slices and centrally located magnified views of F-actin and cell nuclei within a representative microtissue at four days are shown in fig. 1d. Actin was highly polymerized into dense stress fibers that oriented with the longitudinal axis of the microtissue. The cell nuclei were also mostly aligned with the microtissue and evenly distributed in three dimensions.

Microtissues strain soften

It has been widely reported that acute dynamic stretching changes the mechanical properties of cells grown on 2D surfaces; they become softer (decreased elasticity) and more fluid-like (increased phase lag) \(^{7,11,38}\). In that regard, we started by investigating
whether or not 3D microtissues composed of 3T3 fibroblasts share this behavior by assessing dynamic mechanical properties under progressively larger strains at 0.25Hz.

Microtissue storage stiffness decreased in a strain-dependent manner (N=22, linear regression: $R^2 = 0.97$, $p<0.001$), (fig. 2a). Comparing 9% to 1% strain, the average storage stiffness decreased by 26 ± 2% (repeated measures t-test: $p<0.001$).

Unlike previously published findings on cells in 2D culture, where softening is accompanied with a more fluid-like behavior (or fluidization) $^7$, the phase lag of microtissues decreased with strain amplitude (linear regression: $R^2 = 0.95$, $p<0.01$) (fig. 2b), indicating a greater amount of energy stored for a given amount dissipated at higher strains. Therefore in contrast to cells in 2D, microtissues become more elastic-like as they soften.

The softening response in microtissues was also accompanied with a decrease in prestress (linear regression: $R^2=0.98$, $p<0.001$) (fig. 2c). Comparing 9% strain to 1% strain, the average prestress decreased by 2.0 ± 0.4µN (repeated measures t-test: $p<0.001$).

The strain softening response was reversible upon returning to small amplitude oscillations (fig. 2d). The stiffness and prestress recoveries are shown in fig. 2e,f, respectively (N=8). After 160 seconds, microtissue stiffness recovered to 99± 3% of its initial value. The prestress also agreed well with initial measurements. The recovery curves followed stretched exponential functions (equation 1) with agreeing time ($\tau$) (38 ± 1 vs. 35 ± 6 seconds) and power law constants ($\beta$) (0.87 ± 0.01 vs. 0.92 ± 0.08) (SI 1).

$$F(t) = a * e^{-t/\tau^\beta} + d$$ (1)

We have shown that, as with cells in 2D culture, 3D microtissue cultures strain soften. Depolymerization of actin filaments $^{7,9-11}$ and perturbing of myosin motor binding $^{12,13,39-41}$ are previously hypothesized mechanisms for cells in 2D culture. On the other hand, the involvement of microtubules has been largely overlooked despite their contributions to overall cell mechanics $^{42-45}$ and their dynamic instability $^{46}$. Accordingly, we investigated the roles of these three cytoskeletal proteins to microtissue strain softening. We begin with examining the involvement of actin microfilaments.
**Softening is actin dependent**

To assess the role of the actin cytoskeleton in strain softening, f-actin was depolymerized with Cytochalasin D (CytoD). As expected, CytoD treatment reduced the resting tension, stiffness and phase lag (SI 2). Importantly, CytoD treatment also muted the softening response (N=16 repeated measures t-test, p<0.001) (fig. 3b). In fact the stiffness of CytoD treated microtissues did not change with strain amplitude (repeated measures t-test, P>0.05). Upon stretch cessation, CytoD treatment also prevented tension recovery; further demonstrating that CytoD treated microtissues do not soften (fig. 3c). These results indicate that strain softening is dependent upon changes to a densely polymerized actin cytoskeleton.

**Stretch remodels and depolymerizes actin**

We have shown that f-actin is required for strain softening in microtissues. To further link the softening response to changes in the cytoskeleton, we labeled actin filaments with a live-cell stain. Then by comparing images taken before and following various durations of static resting or oscillatory stretching, we assessed whether stretching increases the rate of actin remodeling. Heat maps of correlation coefficients showing actin remodeling (areas with low correlation coefficients indicate high remodeling) within a centrally located region of the microtissue are in fig. 4a. Actin remodeling was spatially heterogeneous throughout the tissue and increased with time for both static and loading conditions. Importantly, compared to the static condition, oscillatory loading increased remodeling of actin filaments (decreased the average correlation coefficient) after one and five minutes (N=6, repeated measures t-test, P<0.05) (fig. 4b).

We have shown that oscillatory stretch increases remodeling of actin filaments in living cells in 3D cultures. To investigate whether remodeling arose simply from organizational changes or depolymerization/repolymerization of filaments, microtissues were fixed and stained immediately following various durations of stretching at 9% strain. Representative images, average heat maps and average f-actin expression per cell (fig. 5a,b,c, respectively) all indicated that f-actin rapidly depolymerized with oscillatory stretching (N>14).
To show that f-actin also repolymerizes following stretch cessation, microtissues were fixed and stained after various durations of recovery following five minutes of stretching. Average heat maps and f-actin expression per cell (fig. 5b,d) show complete recovery to initial expression values (t-test P>0.05) (N>11). Although our time resolution of f-actin expression was poor and the uncertainties are large, the rate of f-actin recovery appeared to be within the same order of magnitude (tens of seconds) as the rate of tension and stiffness recovery, suggesting that the mechanical measurements reflect actin repolymerization.

**Myosin and microtubules do not contribute to strain softening**

We have identified that actin filaments play a major role in the strain softening response of 3D microtissues, however, the mechanical behavior of cells \(^\text{42-45,47}\) and microtissues (SI 2) is also highly dependent upon myosin activity and microtubules. To assess the contribution of myosin and microtubules to strain softening, we examined the response following myosin inhibition with blebbistatin (Bleb) and microtubule depolymerization with nocodazole (Noco).

Myosin inhibition with blebbistatin decreased microtissue stiffness (N=10, repeated measures t-tests, P<0.05) and prestress (p<0.01) (SI 2). Myosin inhibition, however, did not affect strain softening (N=10, repeated measures t-test, P>0.05). As expected, it did reduce the tension recovery (fig. 6c) because of the decrease in prestress that unsurprisingly accompanied myosin inhibition. However, as there was no change in the rate of recovery, myosin was not likely responsible for the recovery following softening (SI 1). Although it is possible that there was incomplete inhibition of myosin motors, it is unlikely according to blebbistatin’s measured dose-response curve (SI 3). Moreover, even with incomplete inhibition, one would still expect a decrease in the softening response if myosin were responsible.

In keeping with the hypothesis that microtubules are mainly compressive elements that oppose acto-myosin activity \(^\text{42-45}\), microtubule depolymerization increased microtissue stiffness (N=15, repeated measures t-tests, P<0.001), and prestress (P<0.001) (SI 2). Microtubule depolymerization, however, had no effect on strain softening in terms of the percent change in storage stiffness (P>0.05) (fig. 6b). It did increase the absolute
tension recovery (fig. 6c) as expected, because of the increase in prestress that accompanied microtubule depolymerization. However, again the molecular mechanism for softening was not likely affected by microtubule depolymerization, as there was no change to the rate of recovery (SI 1). Interestingly, oscillatory stretching did increase remodeling in microtubules with significant differences from static conditions after one and five minutes of stretching (SI 4) (N=7, repeated measures t-tests, p<0.01 and p<0.05, respectively). However, stretching did not change the degree of microtubule polymerization per cell (SI 4) (N>14, 1-way ANOVA).

**Strain softening is a conserved response for microtissues**

We have shown that microtissues composed of 3T3 fibroblasts, mimicking connective tissue, strain soften through actin depolymerization. To assess whether this behavior is unique to fibroblasts or is shared among other cell types, we assessed strain-softening responses in microtissues composed of human airway smooth muscle cells (HASM) or skeletal muscle cells (C2C12). Both smooth and skeletal muscle microtissues strain softened (fig. 7a) (linear regression: $R^2=0.99$, $0.98$ P<0.001, 0.001, respectively) and shared strain-dependent decreases in prestress (fig. 7c) (linear regression: $R^2=0.98$, $0.94$ P<0.001, 0.01, respectively). The phase lag of skeletal muscle microtissues decreased with strain (linear regression: $R^2=0.99$, P< 0.001), whereas smooth muscle microtissues did not ($R^2=0.43$, P=0.14) (fig. 7b).

Upon decreasing the strain amplitude, the stiffness and tension of HASM and C2C12 microtissues recovered as previously observed with the 3T3 fibroblasts (fig. 7d,e). The time constant for stiffness recovery in HASM cells was statistically greater than either 3T3 or C2C12 microtissues (1-way ANOVA, P<0.05) (SI 1). On the other hand, the time constants for tension recovery agreed well with each.

**Discussion**

We aimed at assessing the response to acute oscillatory stretch in cells grown in conditions that mechanically and biologically recapitulate the 3D environment that a cell would experience were it in the body. In order to fulfill this goal, we developed a novel
approach to allow high throughput measurements of dynamic mechanical properties and direct visualization of the cytoskeleton in 3D microtissue cell cultures. Our approach consists of an array of vacuum-driven actuators to stretch microtissues and optically tracked force-sensors to measure their mechanical behavior. Advantages and limitations of our approach are summarized in SI 5. In using our approach, we showed that microtissues soften under dynamic stretching through actin depolymerization. These results are further discussed the sections that follow.

**Microtissues soften under dynamic loading**

We showed that the prestress and storage modulus of living 3D microtissue cultures composed with three different cell types decrease with dynamic strain amplitude. This finding agrees well with previous reports in cells in 2D culture $^{7,38,48}$ and ex vivo tissue strips $^{12,13}$. Strain softening has been identified as an important mechanism for maintaining homeostasis throughout the body. For example, it may explain how a large tidal stretch from a deep inspiration can open contracted airways in healthy lungs $^{12,13}$ and could contribute to the regulation of blood pressure in arteries $^{16}$.

In addition to softening, cells in 2D have long been reported to exhibit a more fluid-like behavior when stretched $^{49-51}$. More recently this fluidization response of cells has been associated with that of a class of inert materials called soft glasses (eg. foams, dense emulsions, pastes and slurries) $^7$. In contrast to these reports and the soft glassy rheology hypothesis, skeletal muscle and fibroblasts microtissues actually became more elastic-like with greater strain amplitudes, and smooth muscle microtissues did not phase transition. Although this apparent contradiction could arise from environment differences between growing cells our 3D microtissues and past reports done in 2D culture, it is more likely that the fluidization response was hidden by the elasticity of extracellular matrix.

After removing most of the cellular component to the mechanical behavior of the microtissue with CytoD treatment, the remaining behavior, describing the contribution of the matrix, is primarily elastic and contributes little to energy dissipation (SI 2). In turn, a more elastic-like behavior could be perceived as the cells soften (and perhaps fluidize) because the cellular contribution to the overall mechanical behavior becomes less important compared to the elasticity of the matrix. It would, however, still be of interest
in future work to assess whether or not microtissues follow the same time-scale invariance \(^{52}\) that has given traction to the hypothesis that the mechanical behavior of cells follow Sollich’s (1997) theory on soft glassy rheology \(^{53}\).

Upon strain cessation, we showed that microtissue prestress and stiffness return to pre-intervention values along well-conserved trajectories that could be modeled over three magnitudes of time with a stretched exponential. Stretched exponentials have previously been used to describe relaxation processes in disordered systems \(^{54,55}\) and can appear from a linear superposition of simple exponential functions with a nontrivial distribution of relaxation times \(^{56}\). It, therefore, should not be overly surprising that such a function can appear from the complex nature of the cytoskeleton and the added intricacies that arise when considering an aggregate of cells. In spite of this, we found that recovery time constants for prestress and stiffness were between 35 and 43 seconds for all tested treatments that showed a recovery response and cell types (except for stiffness recovery in HASM microtissues). This time constant appears to be within the same order of magnitude as previously reported recoveries in cortical actin stiffness following transient stretching cells in 2D culture \(^{7}\). Unfortunately the authors of that work did not fit their curves to stretched exponentials. They did, however, concede that recovery occurs with timescales that grow with the elapsed time since stretch cessation and is slower than an exponential process. These are characteristic features of stretched exponentials.

In the literature strain softening exists in a paradox with a large number of studies reporting strain stiffening and actin reinforcement in response to stretch \(^{57}\). Under a sustained stretch or examining different locations along the stress-strain curve, reconstituted crosslinked actin gels \(^{58,59}\), cells \(^{60}\), and 3D cell cultures \(^{28,61}\) have all been shown to strain stiffen. This nonlinear effect arises from reorganization of actin filaments \(^{58,59}\) and, for tissues, reorientation of cells and percolation of these local effects across the cell culture \(^{61}\). Although our loading loops were mostly linear, strain stiffening can be observed to a degree at large strain amplitudes (the total harmonic distortion from nonlinearities increased from 0.041 ± 0.002 at 1% strain to 0.152 ± 0.004 at 9% strain (P<0.0001, repeated measures t-test)). However, this effect was still much smaller than the softening behavior produced by dynamic loading. On the other hand, downstream
signaling cascades have been reported to cause mechanical stiffening through actin microfilament reinforcement as a sort of negative feedback to maintain localized mechanical stress. Microtissues have previously been shown to stiffen following 15 minutes of dynamic stretching and we have also reported actin reinforcement in microtissues under chronic (2 days) conditioning. Importantly in those investigations quasi-static stiffness measurements and f-actin expression were evaluated following loading and after a period much greater than the time-scale of stiffness recovery and actin repolymerization we report here. That said, over our relatively short experimental time, we did not observe any differences between initial and fully recovered stiffness, prestress and f-actin expression measurements.

**Stretch depolymerizes actin in microtissues**

Although strain softening of cells and tissues has been widely reported, the molecular mechanism(s) behind this response remains unclear. Here we investigated the contributions of actin microfilaments, myosin motors and microtubules.

Firstly, in a cell, the actin cytoskeleton is a filamentous network that gives the cell its shape and opposes tensile forces. In 2D culture, stretching of cells has been reported to depolymerize actin filaments. Our results in 3D cultured microtissues agree with those observations. We found that 1) f-actin was necessary for strain softening and the recovery response; 2) actin remodeling in living cells increased with stretch; 3) short-term stretch lowered f-actin expression; and 4) upon stretch cessation, f-actin expression recovered along the same timescale as tension and stiffness recovery. These findings strongly suggest that strain softening, at least in part, arises from actin depolymerization.

Secondly, myosin motors regulate the mechanical behavior of cells by generating tension through crosslinking and actively pulling on actin filaments. Furthermore, strain softening in reconstituted actin-myosin networks has been attributed to disruption of myosin crosslinks. Perturbing of the binding of myosin has also been implicated in the softening response in airway tissue strips. In contrast, we found softening in microtissues was invariant on myosin activity and that there was no change to the rate of the recovery response following strain cessation. This strongly suggests that myosin has no role in the softening response of cells in 3D culture.
Lastly, although our understanding of the role of microtubules in cell mechanics is still being refined \(^{70}\), it is thought that they act as compressive struts to oppose actin-myosin contractility, as in tensegrity architecture \(^{42-45}\). Accordingly, and in keeping with several other investigations measuring cell traction forces and stiffness in 2D \(^{42,71}\), we found that depolymerization of microtubules with nocodazole increased microtissue stiffness and prestress (SI 2). Comparably fewer studies have assessed how stretching cells and tissues affects microtubule remodeling and polymerization. Microtubules have visually been seen to buckled while cells deform \(^{44}\), and in axons grown in 2D culture, microtubules have been shown to disassemble under large (75\%) dynamic loading \(^{72}\). In contrast, we did not observe any changes to microtubule polymerization and nocodazole treatment had no affect on the softening response. Dynamic stretching did, however, increase microtubule remodeling. Whether stretching directly caused disassembly/assembly of microtubules or they simply remodeled in accordance with actin depolymerization, and whether or not the observed microtubule remodeling contributes to changes in the mechanical behavior of cells are interesting questions for future investigations.

**Conclusions**

In this article, we presented a new high-throughput approach for both assessing dynamic cell mechanics and for visualization of remodeling at the sub-cellular level in response to stretch within physiologically relevant 3D microtissue cultures. Our approach offers the ability to link behaviors observed in 2D culture to cells within a soft 3D matrix comparable to human tissue, and to connect visual remodeling of the cytoskeleton to changes in mechanical properties. In that regard, we found that fibroblast, smooth muscle, and skeletal microtissue cultures all share a conserved softening response when dynamically stretched and recovery following stretch cessation. Furthermore, by directly quantifying cytoskeletal remodeling, softening of microtissues appeared to arise from rapid actin depolymerization. This suggests that actin microfilaments are sensors of mechanical stretch in cells, and in turn, form a feedback loop to control the mechanical behavior of tissues. The ability of cells to feel and react to mechanical stimuli from their
environment is an important mechanism for maintaining homeostasis in the body and is a critical aspect to fully understand many pathological disorders.

**Methods**

**Device design**

Our original MVAS device was modified to allow *in situ* measurements of microtissue tension and dynamic stiffness. The MVAS-Force consists of six independently controllable rows of ten microtissue wells (fig. 1). Vacuum chambers border one side of each row of wells. Within each well, there are two cantilevers spaced apart by 500µm. One cantilever is secured on a flexible membrane that deforms to stretch the microtissue when a vacuum is applied through an external electronic regulator (SMC ITV0010) controlled via Labview software (movie 1). The other cantilever acts as a passive force sensor. Its deflection is optically tracked and converted into a force measurement using a spring constant of $k_{\text{cantilever}}=0.834\text{N/m}$ estimated with Euler-Bernoulli beam theory and verified with atomic force microscopy (SI 6).

**Device Fabrication**

Devices were fabricated as previously described with slight modifications. Detailed assembly steps are illustrated in SI 7. Briefly, the MVAS-Force consists of three layers fabricated through mold replication from SU-8 (Microchem) masters made with standard photolithographic methods on polished silicon wafers (Universitywafers.com). All photomasks were ordered from CAD Art Services Inc. The top layer of the MVAS-Force comprises the open-top microtissue wells and enclosed vacuum chambers. The thin middle membrane is fabricated with the cantilevers around which the microtissues compact. Finally the bottom layer contains vacuum chambers that match the top layer, and bottom chambers that equalize the pressure on either side of the membrane to minimize out of plane motion. All three layers were cast in polydimethylsiloxane (PDMS) from the SU-8 negatives with a 10:1 monomer to curing agent ratio and then plasma bonded together. To aid in tracking the bottom of the cantilevers, the middle membrane was fluorescently dyed with Rhodamine B (RhoB).
Cell culture

NIH3T3 fibroblast (ATCC) and C2C12 skeletal muscle (ATCC) cells were cultured in Dulbecco's Modified Eagle's Medium (DMEM) (Hyclone Laboratories Inc.). Human ASM cells (Donor 12) (previously characterized by Gosens et al. 73) immortalized by stable transfection with human telomerase reverse transcriptase were obtained as a generous gift from Dr. William Gerthoffer (University of South Alabama) and maintained in DMEM/F12 (Invitrogen 11330). All culture media was supplemented with 10% fetal bovine serum (FBS), 50mg/ml streptomycin and 50U/ml penicillin antibiotics (all from Hyclone Laboratories Inc.). Cells were grown at 37°C with 5% CO₂ on 100mm tissue culture dishes (Fisher) until 80-90% confluent.

Microtissue fabrication

Microtissue fabrication was performed as described previously33,37, with modifications. Briefly, the device was sterilized with 70% ethanol, and treated with 0.2% Pluronic F-127 (P6866, Invitrogen) for two minutes to reduce cell adhesion. 250,000 cells were resuspended in 1.5mg/ml rat tail collagen type I (354249, Corning) solution containing 1x DMEM (SH30003.02, Hyclone), 44 mM NaHCO₃, 15 mM d-ribose (R9629, Sigma Aldrich), 1% FBS and 1 M NaOH to achieve a final pH of 7.0-7.4. The cell-collagen solution was pipetted into the MVAS-Force and centrifuged to load ~650 cells into each well. The excess collagen was removed and the device was transferred into the incubator for 15 minutes to initiate collagen polymerization. An additional 50,000 cells were added and allowed to adhere to the top of the tissues. Excess cells were washed off. Cell culture media was added and changed every 24 hours.

Force measurements

Microtissue mechanics were deduced from the visible deflection of the forcesensing cantilever while under dynamic loading at 0.25Hz (movie 1). Prior to measurements, microtissues were preconditioned until subsequent loading loops were superimposabale. All measurements were completed at 37°C and 5% CO₂.
To track cantilever deflection, images of both the tops and bottoms of the cantilevers were captured at 15 frames per second for one minute. The bottom positions were measured by finding their centroids on thresholded fluorescent images of the Rho-B dyed cantilevers. The top of each cantilever was tracked using pattern matching with adaptive template learning in Labview on brightfield images. The deflection of the force sensor was calculated from the difference in the top and bottom positions after accounting for the phase lag caused by the camera delay between the top and bottom images. The deflection was then converted into a force measurement using the cantilever spring constant, \( k_{\text{cantilever}} \).

Microtissue strain, \( \varepsilon \), was defined as the percent change in the length between the innermost edges of the tops of the cantilevers (equation 1).

\[
\varepsilon(t) = \frac{\text{length}(t) - \text{length}_0}{\text{length}_0} \times 100
\]  

(2)

The phase lag, \( \delta \), between force and strain was defined as the difference in the phase angles, \( \Phi \), at the oscillatory frequency (equation 2).

\[
\delta = \Phi_{\text{force}} - \Phi_{\text{strain}} \left[ \text{rad} \right]_{0.25\text{Hz}}
\]  

(3)

The storage, \( k' \), microtissue stiffness was defined as the ratio of the magnitudes of the Fourier Transforms of force and strain at the oscillatory frequency multiplied by the cosine of the phase lag between force and strain (equations 3).

\[
k' = \left( \frac{|\text{FFT}[\text{force}(t)]|}{|\text{FFT}[\text{strain}(t)]|} \right)_{0.25\text{Hz}} \cos \delta
\]  

(4)

\( k' \) describes the amount of energy that is elastically stored for a given deformation, and \( \delta \) describes the ratio of energy dissipated to energy stored where in purely elastic samples \( \tan(\delta)=0 \) and in purely viscous samples \( \tan(\delta)=\infty \).

The tension offset or prestress, \( T_0 \), was defined as the magnitude of the Fourier transform of the microtissue force at 0Hz minus the half of the peak-to-peak magnitude of the Fourier transform at 0.25Hz (equation 4).

\[
T_0 = |\text{FFT}[\text{force}(t)]|_{0\text{Hz}} - \frac{1}{2} \left( |\text{FFT}[\text{force}(t)]|_{0.25\text{Hz}} \right)
\]  

(5)

The noise floor for calculating microtissue mechanics is characterized in SI 6.
To assess the response of microtissue mechanics to stretch, measurements were taken at progressively higher strains. After completing measurements at the largest strain, the recovery response was measured by promptly decreasing the strain amplitude. Stiffness recovery was measured by performing Fourier transforms on intervals spanning three loading cycles.

To assess the role of individual cytoskeletal proteins in contributing to the mechanical properties of microtissues and the strain softening behavior, measurements were taken following 20 minute incubations with either 10µM nocodazole (Noco), a microtubule polymerization inhibitor, 5µM blebbistatin (Bleb), a myosin-II inhibitor, or 10 µM cytochalasin D (CytoD), an actin polymerization inhibitor. As mechanical properties can vary between microtissues, each microtissue was compared to its own pre-treatment value where indicated. To prevent crossover in response from multiple drugs, only a single treatment was administrated to a microtissue.

**Quantification of cytoskeletal remodeling and polymerization**

Images were acquired on a TiE A1-R laser scanning confocal microscope (LSCM) (Nikon) with standard LSCM configurations using appropriate laser lines and filter blocks.

To assess actin and microtubule remodeling in living microtissues in response to stretch, cells were loaded with either 0.1µM SiR-actin or SiR-tubulin with 1µM verapamil 6-12 hours before imaging. Z-stacks were taken before and following 4 seconds (one stretch), one minute and five minutes of static resting and then ~9% stretching at 0.25hz. Imaging was completed with a 60x 1.2NA water immersion objective to give a centrally located field of view of 212x106µm (1024x512 pixels). Z-stacks were flattened by integrating slices, divided into sub images with a size of 100x100 pixels with 10-pixel spacing, and compared with cross-correlation. The correlation coefficient is a measure of how closely images matched before and after a given condition, and thus is inversely proportional to the amount of remodeling (i.e. a low correlation coefficient corresponds to a high degree of remodeling).

To assess f-actin expression and microtubule polymerization, microtissues were fixed in situ with 3.5% paraformaldehyde for 15 minutes and permeabilized with 0.5%
Triton-X for 5 minutes. Microtissues were left in blocking buffer (5% FBS in PBS) for 40 minutes. Microtubules were labeled with α-tubulin primary antibody produced in mouse (Sigma, T6074) and a rabbit anti-mouse IgG secondary antibody conjugated to Alexa Fluor 488 (Invitrogen, A11059). The actin cytoskeleton was stained with Alexa Fluor 546 Phalloidin (Fisher, A22283), and the nuclei were stained with DAPI (Fisher, D1306). To quantify f-actin expression per cell, Z-stacks were flattened by integration, averaged and normalized to DAPI fluorescence. Microtubule polymerization was quantified with the same method except images were first thresholded to remove any signal from nonpolymerized tubulin.

**Data analysis and statistics**

All numerical data are presented as mean ± standard error. Statistical tests as described in the results were performed using Originlab 8.5 (Northampton, MA), with p<0.05 considered statistically significant.

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Author Contributions

M.W. performed data acquisition and analysis and wrote the manuscript. P.R. contributed to data acquisition. All authors contributed to the study design and revised the manuscript.

Additional Information

Competing financial interests
The authors declare no competing financial interests.
Fig. 1: The MVAS-Force allows high throughput dynamic mechanical measurements of 3D cell cultures. The MVAS-force was microfabricated from three photolithographic masters (a). It comprises of an array of microtissue wells each bordered by a controllable vacuum chamber (b). A top-down and cross-section view of a microtissue is shown in (c). Microtissues are dense, organized, three-dimensional cell cultures that are freely suspended around the cantilevers. Max projections of confocal stacks, orthogonal views and high magnification images are shown in (b). The actin cytoskeleton is in green and the nuclei are in blue. Both the cytoskeleton and nuclei show a high degree of organization, aligning between the cantilevers. Scale bars in b,c and d, represent 1mm, 100µm and 50µm, respectively.
Fig. 2: 3T3 fibroblast microtissues strain soften. The storage stiffness, $k'$, (a), phase lag between stress and strain, $\delta$, (b) and prestress, $T_0$, (c) all decreased with increasing oscillatory strain amplitude. To see whether this response was reversible, microtissues were oscillated under a large amplitude strain until subsequent loading loops were overlapping (1) and then suddenly switched to small amplitude strain (2) (d). Both the storage stiffness (e) and prestress (f) fully recovered to their initial values over 160 seconds with similar rates.
**Fig. 3: Softening requires an intact actin cytoskeleton.** Images of a microtissue prior to and following CytoD treatment are in panel (a). As shown by the dotted red line and the dotted yellow ellipse that outlines the top of the force-sensing cantilever (left), depolymerization of F-actin with Cytochalasin D (CytoD) visibly moved the cantilever outward, indicating a lower resting tension (a). Importantly, CytoD treatment reduced the stiffness change under large vs. small amplitude stretching (ie. the amount of strain softening) (b). There was also no tension recovery following stretch cessation (c). The scale bar in (a) represents 100µm.
**Fig. 4: Oscillatory stretch increases remodeling of actin filaments in living cells in 3D culture.** The effect of oscillatory stretch on actin remodeling rate was measured across centrally located regions (212x106µm; red rectangle) in living microtissues using live-cell staining and comparing confocal stacks taken immediately before and after various durations of stretching or static culture (a). Representative heat maps of cross-correlation coefficients show that actin remodeling was spatially heterogeneous and increased with large amplitude stretching vs. static conditions (b). The average correlation coefficient was significantly reduced (ie. a greater amount of remodeling had occurred) when stretching vs. static after 1min and 5min (b). The scale bar in (b) represents 50µm. (*P<0.05; N=6 repeated measures t-test)
Fig. 5: F-actin depolymerizes with stretching and repolymerizes upon stretch cessation. Representative images after different durations of stretching show that there were fewer actin filaments with longer stretch duration (a). F-actin expression in average heat maps is similarly reduced with stretch duration (b) (N>14). Moreover f-actin expression recovered to initial values upon stretch cessation (N>11). The average actin expression normalized to the number of cells under various durations of stretching and recovery are shown in (c) and (d), respectively. The scale bars in (a) and (b) represent 50 and 100µm, respectively.
Fig. 6: Microtubules and myosin do not contribute to softening. Microtissues prior to and following nocodazole and blebbistatin treatments are shown in (a). Microtubule depolymerization with nocodazole moved the force-sensing cantilever inward, indicating increased prestress. In contrast, myosin-II inhibition with blebbistatin moved the cantilever outward, indicating decreased prestress. Neither treatment changed the amount of strain softening in terms of percent change (b). Microtubule depolymerization increased the tension recovery while myosin inhibition decreased recovery (c) but neither treatment changed the time constant of the recovery response. The scale bar in (a) represents 100µm.
Fig. 7: Strain softening is a conserved response in microtissue cultures. Microtissues composed of either fibroblasts (3T3), human airway smooth muscle (HASM), or skeletal muscle (C2C12) cells strain softened with similar changes to stiffness (a), phase lag (b), and prestress (c). They also shared similar recovery dynamics upon stretch cessation in terms of the rate of storage stiffness (d) and prestress (e) recovery.
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Cross-Correlation

'How similar?'

(1) Image

(2) Static or Stretch

(3) Image

[Graph showing correlation coefficient over time for static and stretch conditions]
a) 

- **$k'(\% \text{ Decrease})$** vs. **Strain (%)**
- Lines represent:
  - 3T3 (N=22)
  - HASM (N=6)
  - C2C12 (N=12)

b) 

- **$\delta (\text{rad})$**
- Values range from 0 to 0.24

- **$\Delta T_0 (\mu\text{N})$**
- Values range from -2.5 to 0

- **$\text{Time (s)}$**
- Values range from 0 to 150


c) 

- **$k'(\% k'(0))$** vs. **Strain (%)**


d) 

- **$k'(\% k'(0))$** vs. **Time (s)**
- Values range from 0 to 100

- **$\Delta T_0 (\mu\text{N})$** vs. **Time (s)**
- Values range from 0 to 2.0