Sustained Strain Applied at High Rates Drives Dynamic Tensioning in Epithelial Cells

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ABSTRACT

Epithelial cells experience long lasting loads of different magnitudes and rates. How they adapt to these loads strongly impacts tissue health. Yet, much remains unknown about their stress evolution under sustained strain. Here, by subjecting cell pairs to sustained strain, we report a bimodal stress response, where in addition to the typically observed stress relaxation, a subset of cells exhibits a dynamic tensioning process with significant elevation in stress within 100s, resembling active pulling-back in muscle fibers. Strikingly, the fraction of cells exhibiting tensioning increases with increasing strain rate. The tensioning response is accompanied by actin remodeling, and perturbation to actin abrogates it, supporting cell contractility's role in the response. Collectively, our data show that epithelial cells adjust their tensional states over short timescales in a strain-rate dependent manner to adapt to sustained strains, demonstrating that the active pulling-back behavior could be a common protective mechanism against environmental stress.

INTRODUCTION

Epithelial cells are subjected to strains of varied rates and magnitudes¹⁻³. Some deformations are sustained for a period of several minutes rather than acute in the matter of seconds. Cells have effective mechanisms, such as stress relaxation, to dissipate stress and prevent tissue damage^{4,5}. If deformation is applied fast and becomes sustained, stress relaxation may lead to dangerously large deformations, and cells may need to pull back or stiffen actively to prevent these. Such phenomena occur in the muscle where individual myofibers exhibit stiffening behavior by adjusting myosin attachment kinetics under load⁶. However, the generality of such a behavior is unclear and it is unknown if cells in epithelia can respond actively when subjected to high strainrate, sustained deformation. During physiological function, epithelial cells experience strains of widespread rates. Therefore, the timescale of the strain application may be an important factor in determining the response outcome⁷. For instance, force application rate determines the activation of talin-dependent mechanosensing⁷. We have shown that cellular response to ramp strains is rate dependent, with increasing stress levels and higher probability of cell pair fracture at higher strain rates 8 . However, the role of strain rate in determining cellular response to sustained strain is still not fully understood.

Currently, our understanding of how cells and tissues adapt to sustained strain is limited. It is unclear whether the cellular response is cell autonomous or stems from the interactions between neighboring cells in a tissue. Mechanosensation could arise either from intercellular junctions or from focal adhesions that link cells to their substrate. Previous experiments examining the response of suspended monolayers to deformation report stress relaxation mediated by turnover of the actin cytoskeleton but no active response^{5,9}. Yet, other studies point to active responses of cells when they are adherent and subjected to force¹⁰. Thus, focal adhesions may represent a key element in sensing mechanical stress. Here, we examine the respective contributions of intercellular adhesions and focal adhesions by subjecting adherent epithelial cell

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pairs or single cells to a defined strain and monitoring their stress response. This allows us to track stress within the individual cells and investigate the mechanism of adaptation during stress response. Strikingly, we find that epithelial cells indeed exhibit active tensioning in a strain and strain-rate dependent manner in response to a sustained strain. Stress relaxation dominates cellular response at low strain rates during holding, while high strain rates increase the probability of triggering the dynamic tensioning response. This tensioning response depends on actomyosin and is observed in both single cells and cell doublets, suggesting it may be sensed by focal adhesions.

RESULTS

Cells increase their tension in response to a high strain rate deformation

To investigate the change in stress in a cell pair in response to application of strain, we used our Single Cell Adhesion Micro Tensile Tester (SCAµTT) platform (Fig. 1a)⁸, which comprises two islands separated by a 2 μm gap. Cell pairs seeded onto the platform adhere via focal adhesions and form intercellular junctions that span the gap between the two islands. The device allows stretching a cell pair while measuring applied stress and strain. Briefly, a displacement of D imposed on the forcing beam (Island 2) induces a deflection of δ in the sensing beam (Island 1) via forces transmitted across the cell-cell contact (Fig. 1b). Based on beam deflection, the stress and strain in the cell pair can be calculated (see Methods). Importantly, the stiffness of the beams is set such that δ is much smaller than D, and therefore the deformation of the cell pair is assumed to be determined by D alone¹¹. Therefore, by controlling the rate at which D is applied, we can control the rate at which the cell pair is deformed (Fig. 1c).

In this study we investigated the response of A431 epithelial cells to a ramp and hold condition, in which a predetermined strain is applied at a controlled rate and then held constant. Under load, tension is applied to both cell-cell and cell-ECM adhesions as the cytoskeleton is

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deformed, actively responds, and adapts to the loading environment (Fig. 1d). We first subjected cell pairs to a strain of 50% at strain rates varying from 0.5 %s⁻¹ to 50 %s⁻¹. These conditions are within the physiological range (0.5-80 %s⁻¹⁾⁷ and similar to those previously used in monolayer studies^{5,12,13}. Surprisingly, the temporal evolution of stress is quantitatively dependent on the applied strain rate (Fig. 1e, Supplementary Video 1).

Depending on the strain rate, characteristics of these stress versus time curves are different. Stress first increased with the ramp deformation and peaked when the target deformation was reached (defining $t = 0$ s). Consistent with the known viscoelastic behavior of cells, peak stress increased from just below 0.5 kPa to 1 kPa with increasing strain rate. Then, regardless of the strain rate, stress briefly relaxed by about 100 Pa following a power law in the first ~10s (Supplementary Fig. 2c-d), possibly due to passive viscoelastic behavior and remodeling of the cytoskeleton, as previously described by others¹⁴. After this short dip in stress, the temporal evolution of stress in the following 2 minutes depended greatly on the applied strain rate, displaying a bimodal behavior. For strain rates \leq 5 %s⁻¹, the stress remained approximately constant at the stress achieved after the relaxation phase, indicative of a solid-like behavior. However, for strain rates $\geq 10\%$ s⁻¹, in some cell pairs, the stress rose linearly (i.e., tensioning) reaching values larger than the stress at $t = 0$ s (Fig. 1e, Supplementary Fig. 1, 2d-f). After this linear phase in the first 30 to 50 seconds, stress started to trend towards a new steady state (Fig. 1e). At the end of the two-minute observation, in cells which were deformed at 10 %s-1, the stress increased by \sim 40% from 0.5 kPa at t = 0 s to 0.7 kPa. For 25 %s⁻¹ and 50 %s⁻¹, the stress approximately doubled. Overall, the rate of stress enhancement increased with strain rate. In conjunction with the stress increase, we also observed a decrease in strain at t = 2 min (as shown in Supplementary Fig. 2b).

These results suggest that cells are actively reacting to the loading conditions and exhibit varied responses dependent on those conditions. When the strain rate is low, the cells relax stress but when the strain rate is high, they appear to exert more tension, perhaps aiming to reduce the deformation that they are subjected to.

Characterizing stress accumulation in cell pairs

Pooling individual curves for each condition revealed two distinct behaviors in cell pairs subjected to high strain rates (10 %s⁻¹, 25 %s⁻¹ and 50 %s⁻¹). In some cell pairs, the stress relaxes and reaches a plateau after a few seconds, whereas, for others, the relaxation phase is followed by a linear increase in stress, i.e., tensioning (Fig. 2a, Supplementary Fig. 1, Supplementary Video 2).

To capture the mechanical characteristics of the stress response, we fitted our experimental stress curves with an empirical fit function for the relaxation modulus of the material, $G(t) = At^{-\alpha} + B + Ct$, based on our characterization of the response to a step strain ε_0 (Fig. 2b, 2c and Supplementary Fig. 2, Methods). This approach assumes that the response is linear with the deformation for the purpose of quantifying the material's response to arbitrary loads. The power law function defines the first relaxation phase, where $A \cdot \varepsilon_0$ sets the amplitude of stress relaxation and α is the power law exponent, $B \cdot \varepsilon_0$ is the residual stress in the relaxed curves and an estimation of the stress in the transition point between the relaxed and tensioned part of the response for tensioned curves, and $C \cdot \varepsilon_0$ captures the slope of the tensioning part of the response (a linear fit for C is shown in Supplementary Fig. 2f). To categorize the two visually distinguishable groups in the stress response, we used the values of B and C which serve as indicators of the second phase of the response and exhibit marked differences between the relaxed and tensioned curves. First, we used an unbiased approach by plotting these two parameters on a semilogarithmic scale and then clustering our experimental data based on the characteristics of the second phase of the stress response (Methods). Remarkably, all the cell pairs subjected to low strain rates (0.5 %s⁻¹, 2.5 %s⁻¹ and 5 %s⁻¹) exhibit a relaxed response (Fig. 2d, hollow circles), whereas the cell pairs that experienced high strain rates (10 %s⁻¹, 25 %s⁻¹ and 50 %s⁻¹) can be categorized into two groups, i.e., relaxed and tensioned, consistent with our qualitative

interpretation (Fig. 2d, and Supplementary Fig. 1) and independent of the optimization and clustering algorithm used (Supplementary Fig. 3).

Based on our classification, the initial stress relaxation captured by the power law was similar for both categories, as the distributions of α and A were not different between tensioning and relaxing cell pairs, except for strain rates of 50 %s⁻¹ (Fig. 2e, f, Supplementary Fig. 4). At low strain rates, cells displayed a power law exponent $\alpha \sim 0.5$, larger and more fluid-like than previously reported for the cortex of single cells¹⁵ or for epithelia devoid of a substrate⁵. This difference may arise because our experiments probe regions of the cell interfaced to the substrate. At 50 %s⁻¹ for cells that displayed tensioning behavior, α was significantly lower, reaching a value of 0.23 ± 0.12 , suggesting that some reinforcement may occur at the highest strain rates to render the cells more solid-like and prevent flow.

When we examined the second part of the response, we found that B and C were significantly larger for tensioning cells than for relaxing cells subjected to the same strain rate (Fig. 2h-i). This indicated that cells with higher B, which represents the elasticity of the system, initiate tensioning in response to high strain rates. We next investigated a potential relationship between B and C. While we could not find any correlation for cells that relaxed (Pearson correlation $r = 0.05$), a positive correlation was detected for cells with a tensioning response (Pearson correlation $r = 0.34$) (Fig. 2d). In addition, when we computed the probability of tensioning for a given strain rate, we found that this increased with strain rate (Fig. 2g and Supplementary Table S1).

Tensioning is suppressed by large deformations

We next investigated the dependency of the tensioning behavior on the applied strain magnitude. The strain magnitude was increased to 125%, a strain level that epithelial cells such as keratinocytes can tolerate without losing integrity or viability¹⁶, and strain rates of 0.5 %s⁻¹, 25 %s⁻¹

 $¹$, and 50 %s⁻¹ were used. Interestingly, for this large strain, the average stress curve shows</sup> predominantly a relaxation behavior across these strain rates (Fig. 3a). We theorized that high strain applied in one step could result in cytoskeleton damage and thus prevent active tensioning responses, and therefore compared the behavior of cell pairs stretched at 50 %s-1 subjected to strains of 25%, 50%, and 125% (Fig. 3b). Qualitatively, tensioning appears to dominate at 50% strain, but for both 25% and 125% strain, cell pairs appear to predominantly exhibit a relaxing behavior. To understand this phenomenon further, we plotted B vs C and clustered our data to determine the probability of tensioning and relaxing for each strain (Fig. 3c). This reveals that 77% of cell pairs stretched to 50% display tensioning compared to just 25% and 20% for strains of 25% and 125%, respectively, suggesting the existence of a functional strain range of the response (Supplementary Fig. 6e-g, and Supplementary Table S1). The probability of tensioning for all tested combinations of strain and strain rate are shown in Fig. 3d to visualize the combined influence of these parameters on tensioning (Supplementary Table S1, see Supplementary Fig. 5-7 for summary of all data). The plot shows that tensioning is most likely to occur at an intermediate strain magnitude of 50% at strain rates above 25 %s-1. Tensioning is unlikely to occur at low strain rates regardless of the applied strain, as well as at the low and high ends of the strain magnitude.

To further investigate the hypothesis that high strains applied at high rate may suppress tensioning behavior due to actin disruption, we conducted a stretch-hold-stretch experiment. In these, a total strain of 100% was applied to the cell pairs at a rate of 25 $\%$ s⁻¹, but strain was subdivided into two steps of 50% with a hold time of 5 minutes between the steps (Fig. 3e). We reasoned that such a pause may provide cells with sufficient time to remodel their cytoskeleton and prevent damage because the duration of the pause is commensurate with the characteristic turnover time of actomyosin^{5,17}. Consistent with our initial experiments (Fig. 1), tensioning behavior is seen in some cells in the first step of the experiment. In the second step of the experiment, cell pairs that exhibited tensioning in the first step continued with this behavior, while those that initially relaxed continued to relax. In contrast, when the 100% strain is applied in a single step, all cell pairs exhibit relaxing behavior (Fig. 3e, f and Supplementary Table S1). These results support the idea that the tensioning behavior can be observed for larger strains when cell pairs are allowed to respond and adapt, and that application of large, sudden strains may lead to damage that prevents adaptation.

Tensioning depends on actomyosin contractility, not on cell-cell junctions

Next, we investigated the molecular mechanisms underlying sensing of strain and transduction of the tensioning response. In epithelial cells, mechanosensitive processes have previously been reported to take place at focal adhesions^{18,19} and at intercellular junctions²⁰. To elucidate the role of intercellular junctions in sensing the applied strain, we modified our platform such that a single cell could bridge the gap between islands. Strikingly, single cells displayed tensioning responses similar to cell pairs (Supplementary Fig. 8), indicating that intercellular adhesions are not necessary to sense strain and trigger tensioning.

As actomyosin plays a key role in generating the substrate stress in single cells and cell pairs²¹⁻²³, we investigated its role in controlling the tensioning behavior. We first examined how Factin organization was influenced by the applied strain rate by tracking its localization and intensity with a live stain for 10 minutes after stretching the cells to 50% strain at a strain rate which does not lead to tensioning $(0.5\%s^{-1})$ and at one that does $(50\%s^{-1})$ (Fig. 4a). Further, as a control in which active responses are abrogated and to account for changes in fluorescence intensity due to imaging-induced photobleaching, pairs of cells fixed with paraformaldehyde were also stretched at both rates (Supplementary Video 3). In response to stretch, F-actin intensity in live cell pairs increased in the confinement region relative to the fixed control for both strain rates, although the increase was larger at 50 %s⁻¹ (Fig. 4b). Near the cell-cell junction, there was no difference between the live cells stretched at either rate although the intensity did increase relative

to the fixed controls (Fig. 4c). These results support the previous findings that the cell-cell junction might not be required for triggering the tensioning behavior and support the idea that reorganization of the actin cytoskeleton occurs concomitantly with tensioning.

One reason that cells may no longer be able to tension above 50% strain could be that the actin filaments rupture in response to stretch and prevent myosin mini filaments from applying tension. Therefore, we theorized that by stabilizing the actin cytoskeleton, tensioning could occur even at larger strains. To test this hypothesis, we repeated our experiments with cells treated with jasplakinolide (Jas), which stabilizes actin filaments. When we applied a strain of 125% at 25 %s- 1 , Jas had a minimal influence on the magnitude of stress in cell pairs compared to controls (Fig. 4d). However, a tensioning behavior was qualitatively apparent in Jas treated cells. After analyzing individual curves, we found that the average C with Jas treatment is significantly higher than in the control group (Fig. 4e, Fig. 4e inset, and Supplementary Fig. 9b). Additionally, with Jas treatment, the probability of tensioning more than doubled, increasing from 20% to 46% (Fig. 4f), suggesting that, with a stable F-actin cytoskeleton, tensioning persists at large strain. These results point to the actin cytoskeleton as an important component in controlling the tensioning behavior of cell pairs.

We next asked if myosin contractility is required for tensioning. To test this idea, we examined the effect of drugs that destabilize the F-actin scaffold and inhibit myosin directly or indirectly. Tests were conducted at a strain of 50% at a rate of 25 %s⁻¹, a condition shown to induce tensioning behavior in a large portion of cell pairs (Fig. 2d, Fig. 3d). Latrunculin B (LatB, which inhibits actin polymerization), Y27632 (which inhibits myosin contractility by inhibiting Rhokinase), and blebbistatin (Bleb, which directly inhibits myosin contractility), all reduced the average magnitude of stress experienced by the cell pairs compared to controls, with the average value of B for all treatment conditions being significantly reduced (Fig. 4g, Supplementary Fig. 9e). When we categorized individual curves into tensioning and relaxing behaviors (Fig. 4h), we

found that all three treatments resulted in a dramatic reduction in tensioning probability. Both LatB and Y27632 reduced the probability to 0%, and Bleb reduced the probability to 25% (Fig. 4i). Myosin contractility and an intact F-actin network therefore are important components in the tensioning process.

Tensioning is compatible with an active rheological model for actomyosin

We next sought to provide a rheological framework that encompasses all our observations. Based on our experiments, we posited the existence of a cellular detection mechanism responsible for triggering the tensioning of the cells that is sensitive to strain rate. Following detection, the signal is transduced into a response that increases tension at a constant rate through a mechanism that depends on actomyosin activities (Fig. 4). We therefore adapted a model of the active rheology of actomyosin (ARA) by Etienne et al. 24 that accounts for both the contractility of myosin and turnover of the F-actin network to capture the dynamics of stress generation (Fig. 5a). Since we proved already that the parameters A and α controlling the short-term power-law relaxation do not correlate with the probability of tensioning, we focused our attention on the longer-term response, in relation to B and C. We assume that when deformation is applied, the detection mechanism elevates the steady state tension in cells to a new value, σ_A . The ARA model uses this stress generation element in parallel to a dashpot, η_A , and in series with an elastic component, E_1 . It provides us a simple way of testing this hypothesis and interpreting the rate of tensioning over time: upon a step change in contractile activity σ_A , the system's stress is expected to evolve towards a new plateau value at a rate controlled by both the compliance of the system and the turn-over of actomyosin (encompassed in the η_A term of the model). We could therefore interpret the growing tension as a transient but slow evolution of the system towards a new state of higher contractile tension. In the tensioning cases, where the stress can visibly curve towards a plateau, all parameters can be calculated; otherwise, in relaxed curves with no plateau where η_A does not converge, the mean of η_A from the tensioning cases is used to calculate σ_A and E_1 .

When we fitted our experimental data for 50% strain with the ARA model, we were able to accurately describe the prolonged response of the cell pairs in ramp and hold experiments and found good agreement between our model predictions and the data (Fig. 5b). We could also show that the ARA model parameters could be mapped to our B and C values in the transient linear phase ($B=E_{eq}$ and $C=\left(\frac{1}{\varepsilon_0}\right)\cdot\frac{\sigma_A-E_{eq}\cdot\varepsilon_0}{\tau}$, where $E_{eq}=\frac{E_1\cdot E_b}{E_1+E_b}$ represents the spring equivalent of the cell pair and the sensing island and $\tau=\frac{\eta_A}{E_{eq}}$ is the characteristic time determining the transition between the elastic-like to fluid-like response, see Methods, and Supplementary Fig. 10). When we examined the model parameters obtained by fitting individual curves, we found that the tension σ_A is significantly larger when the cell-pair is tensioning than when it is relaxing, whereas no obvious trend was apparent for η_A (Fig. 5c, d). Interestingly, the characteristic timescale of the contractile force generation $(\frac{\eta}{\epsilon})$ $\frac{dI}{dA}$ is on the order of 100 seconds, consistent with the reported actomyosin turnover timescale^{5,17}. This analysis shows that the steady tensioning process may result from a comparatively rapid but sustained change in the active stress state of the cell.

DISCUSSION

Cells exhibit different behaviors in response to mechanical stimuli, including stress relaxation and reinforcement^{5,25-28}. Stress relaxation is commonly observed in epithelial tissue and individual cells⁵. Another outcome resulting from mechanical stimulation is reinforcement, which can manifest as an increase in stiffness or larger force generation^{26,29,30}. Our findings on epithelial cells reveal the existence of a new and different behavior in single cell pairs in response to strain. Specifically, we identified an active tensioning response, where stress dynamically increases in response to ramp-and-hold strains at high rates resulting in a higher stress than that arising passively from deformation. This response resembles the behavior of muscle fibers which stiffen in response to pulling at different speeds 31 and it may serve as a protective mechanism

physiologically to prevent tissue damage or fracture, by making further deformations difficult or even decreasing deformation (as shown in Supplementary Fig. 2b).

Our study shares similarities with the study by Khalilgharibi et al⁵, where they reported stress relaxation in MDCK epithelial monolayers and single cells with similar relaxation patterns as those we observed. Strikingly, in addition to stress relaxation, we observed that a significant proportion of cells increase the tension they generate over time. In contrast to the suspended anchorless monolayers which only exhibit relaxation, cell pairs and single cells in our study are fully anchored to a substrate via cell-ECM adhesions. These adhesions are known to participate in mechanosensing $32-34$, and our study points towards this adhesion site as the point of detection for cells to initiate their tensioning response. In addition to mechanosensing, a fully anchored cell can use its focal adhesion sites to physically balance the actomyosin generated tension in the actin network $35,36$. Importantly, the tension response is sensitive to strain and strain rate. Our observations reveal clear functional limits to the tensioning response, as too high deformations or deformation rates lead to a muted or absent tensioning response. Whether this is due to limitations in sensing or in transduction remains unclear. Force transmission to cell-ECM adhesions could be disrupted due to rupture at cell-cell junctions under high deformations. Preliminary results show E-cadherin reorganization at the junction in response to the elevated stress at high rates in the functional range and point towards its role in maintaining the mechanical link to facilitate sensing at the cell-ECM adhesion (Supplementary Video 4). In all cases, tensioning arose within ~10s of application of deformation. Such a time-scale points to mechanisms involving signalling in the form of post translational modifications, ion channels, or emergent mechanisms such as the catch bond properties of myosin³⁷.

We speculate that deformation applied at different rates could induce sensing at cell-ECM adhesions via a mechanism similar to substrate stiffness detection. This may be facilitated by force-induced unfolding of proteins such as talin, followed by vinculin recruitment^{7,38-40}. Under

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actomyosin generated tension, protein unfolding and subsequent mechanosensing is induced only for stiff ECM, which allows sufficient force build-up⁴⁰. Under applied strain, tension is applied to this same adhesion complex through stretch of the basal side of the cell and activates potentially the same detection mechanism. This activation occurs during the ramp strain application and influences cell behavior during holding. More importantly, studies have shown that detection molecules are not only sensitive to the magnitude of the force, but also the rate at which it is applied and this can be described by the interplay between the rheology of the cells and the molecular clutch model⁷. Our observation of higher tensioning probability at high strain rates also agrees with this model, in which high rates (50 %s⁻¹ compares to 40 %s⁻¹ in Andreu et al.⁷) allow faster force build-up in sensing molecules, and thus higher probability of triggering a response. The exact conditions at the cell-ECM and cell-cell adhesion sites, such as force-sensing protein distribution and degree of clustering, will influence how stress is distributed across these proteins and can influence the likelihood of triggering the range of responses.

The higher propensity of stress relaxation at large strain may be a sign of actin network disruption or damage due to the strain application, resulting in cells failing to perform force generation in response to high strain rates even when stimuli is detected^{7,22,27}. Consistent with this idea, when we applied large strain in two steps separated by a pause rather than a single step, the tensioning response was preserved, suggesting that the pause allows the cell to adapt, perhaps through remodeling of the cytoskeleton. Damage to the actomyosin cytoskeleton could arise because of loss of connectivity within the actin network $41,42$, similar to the loss of contact between myosin thick filaments and F-actin thin filaments when sarcomeres are overextended43,44. Alternatively, damage could arise from direct breakages in actin filaments due to the large and rapid application of strain 7,15,45, . In support of this, stabilization of F-actin with Jas enabled tensioning responses to persist at high strain. However, further work will be necessary to determine the mechanism underlying loss of tensioning.

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METHODS

TPP fabrication. Fabrication of stretchers was carried out as described in ref.⁸. Briefly, standard tessellation language (STL) files were generated from the geometry of the stretcher made with COMSOL 4.2 software and were imported into the Describe software (Nanoscribe, GmbH) to compile job files for the Photonic Professional (GT) tool (Nanoscribe, GmbH). The stretchers were fabricated from IP-S photoresist (Nanoscribe) with a 25x immersion microscope objective. Glass coverslips with diameters between 11 and 25 mm and thickness of ~160 μm were coated in indium tin oxide (ITO) to allow autofocusing. To promote adhesion of the stretchers to the glass slide, a ~2 μm thick layer of porous silicon oxide (PSO) was deposited onto the ITO-coated slides using an in-house developed protocol. Arrays of stretchers varying from 5x5 to 6x6 were fabricated on each slide.

Cell culture. A431 cells were cultured in growth medium composed of Dulbecco's modified Eagle's medium (ThermoFisher) supplemented with 10% FBS and 1% P/S in an incubator at 37°C with 5% CO₂.

Stretcher preparation and cell deposition. The glass slide on which the stretchers were fabricated was adhered to a glass bottom petri dish using an optically clear UV curable adhesive (Norland Products). To prepare stretchers for cell deposition, a 100 μL drop of Geltrex (Thermofisher) was placed on the stretchers and incubated for 1 hour to promote adhesion of cells to the stretchers. Finally, the remaining Geltrex solution was rinsed off and the petri dish was filled with $CO₂$ independent media (Thermofisher) supplemented with 10% FBS, 1% P/S, and 1% GlutaMAX (Thermofisher). Cells were deposited on the stretching platforms using the protocol described in ref.⁸. Briefly, an Eppendorf single-cell isolation setup was used to pick up and place cells within the confinement on the platform. The setup consisted of a microcapillary (Piezo Drill Tip ICSI, Eppendorf) connected to a pressure controller (CellTram 4r Air/Oil, Eppendorf), which was mounted on a three-axis micromanipulator (Transferman 4R, Eppendorf). After a drop of cells was placed within the petri dish containing the stretchers, cells were picked up and placed on the stretchers one at a time. Cell deposition was carried out on a Nikon Eclipse Ti-S microscope fitted with a custom heating chamber maintained at 37°C. After cells were deposited the petri dish was kept on the microscope overnight to allow cells to adhere to the stretchers and form cell-cell adhesions with adjacent cells on the same stretchers.

Mechanical stretching test. Cell pairs were subjected to mechanical stretch using an AFM as described previously⁸. Briefly, a through hole was drilled into the end of an AFM cantilever tip (NanoAndMore) with a focused ion beam to engage with the pillar on the forcing island. With the hole engaged with the pillar, the displacement and displacement rate were selected in the AFM software and stretch was initiated. The tests were visualized with a Zeiss Axio Observer 7 microscope and recorded using a screen-recording software (Camtasia). After stretching experiments were complete, the petri dish was incubated in trypsin for 1 hour to detach cells from the stretchers. The stretchers were rinsed with PBS to remove cells, and finally submerged in 70% ethanol for 1 minute to sterilize the stretchers in preparation for the next experiment. Stretchers could be reused up to 10 times before they began detaching from the substrate, at which point new stretchers were fabricated.

Drug treatment. For conducting cell stretching experiments with drug treatments, cells were treated with the drug immediately before stretching them. For latrunculin B (Abcam) treatment, LatB was diluted in DMSO and then diluted in DMEM to a final concentration of 200 nM and exposed to the cells for 30 minutes. For blebbistatin (Sigma Aldrich) treatment, stock bleb was diluted in DMSO and then diluted in DMEM to a final concentration of 3.4 μM and exposed to cells for 2 hours. For Y27632 (Abcam) treatment, stock Y27632 was diluted in DMSO and then diluted in DMEM to a final concentration of 30 μM and exposed to cells for 10 minutes. For Jasplakinolide (Thermofisher) treatment, stock Jasplakinolide was diluted in DMSO and then diluted in DMEM to a final concentration of 0.1 μM and exposed cells for 10 minutes. For all drug treatments the media containing the drug was replaced with fresh media before beginning the stretching experiment.

Displacement tracking and stress-time curve calculation. The displacement of each island and the calculation of stress were carried out as described previously 8 . Briefly, the location of each island was determined using a modified version of MATLAB DIC. A rectangular region was defined along the confinement bowtie on each island and the location of each pixel within this region was tracked on every tenth frame to reduce computation time. The displacement of each pixel relative to the first analyzed frame was determined and then the average of the displacement of each pixel was calculated for each island to yield the displacement of each island. The MATLAB code is available at github.com/YangLabUNL.

To calculate stress, first the force within the cell pair system was determined based on the displacement of the sensing island and Hooke's Law, $F = k\delta$, where k is the previously calculated stiffness of the bottom island, 0.11 N/m. Engineering stress was calculated based on the calculated force and the assumed cross-sectional area of the cell-cell contact, which was determined to be 120 μm² with little variation due to the geometry of the confinement. Finally, the time for each data point was determined based on the framerate of the original video and the percentage of analyzed frames to give the stress-time curve.

To calculate strain, the deformation of the cell pair or single cell was divided by its initial length, following the equation $\varepsilon = D/L_0$, with the deflection of the sensing island δ assumed to be negligible compared to the forcing island displacement. For a cell pair, the initial length was 20 μ m, considering the nucleus-to-nucleus spacing determined previously⁸, and for a single cell was 17 μm, or half the length of the confinement along the stretching direction.

Tracking of F-actin intensity. After cell deposition, cells were treated with SPY650-FastAct (Cytoskeleton) at a 1X concentration for 2 hours prior to imaging. For control experiments, cells were fixed with 4% paraformaldehyde for 10 minutes prior to imaging. Imaging was carried out on a Ziess LSM 800 confocal microscope, and images of the cell pairs were taken once a minute. Intensity change was quantified by determining the average intensity within the given region of interest (either within the confinement or between the islands) at each timepoint and calculating the slope of the line of best fit. Therefore, positive values represent an increase in intensity and negative values represent a decrease in intensity.

Data analysis and modeling. The stretching process applied to our cell pairs involved two phases. Initially, a stretch was applied at a constant rate to reach the desired strain magnitude ε_0 . followed by keeping the strain constant for a period of two minutes. To account for the impact of the rate of strain application during the initial phase, we utilized a convolution integral to accurately model the temporal evolution of stress $\sigma(t)$ in our experiments:

$$
\sigma(t) = \int_0^t G(t - t') \frac{d\varepsilon(t')}{dt'} dt'
$$
\n(1)

In our model, we define the relaxation modulus $G(t) = \frac{\sigma(t)}{\varepsilon_0}$ as $G(t) = At^{-\alpha} + B + Ct$. This equation incorporates several terms: a power law term $At^{-\alpha}$ to capture the relaxation phase occurring within the first few seconds, $B \cdot \varepsilon_0$ representing the residual stress in the relaxed curves or an estimate of the point at which stress starts to increase in the tensioned curves, and finally,

the linear function C_t to account for the stress increase in the tensioned curves. We employed the RHEOS software implemented in the Julia programming language to fit our experimental data⁴⁶.

To initiate the fitting procedure, we assign initial values to each parameter of the $G(t)$ function. The initial value of B is defined as the lowest stress level occurring between t_{hold} , which represents the second time point after maintaining the applied stretch constant, and the final point at the end of the two-minute hold (Supplementary Fig. 2a). The initial value for α is calculated as the slope of a line fitted on the ($\sigma(t_{hold} < t < t_{omin}) - B$) on a log-log scale (Supplementary Fig. 2c-d). Here, $t_{\epsilon min}$ refers to the time at which the initial value for B was calculated. Then, the curves are cut at $\sigma_{cut} = 1.4 \sigma_{min}$, and the slope of the line fitted on the $\sigma(t_{\text{cmin}} < t < t_{\text{ccut}})$ is defined as the initial value for C (Supplementary Fig. 2f). Finally, the initial value for A is obtained by fitting the curve with the function $\sigma(t) = A' t^{-\alpha} + B' + C' t$, where the values of α, B' , and C' are the initial values calculated in the previous steps. To ensure the consistency of the outcomes for the fittings, we employed three optimization methods, namely COBYLA, BOBYQA, and SBPLX. These techniques yielded comparable errors in most cases (Supplementary Fig. 2g-h). The fitting parameters presented in this study are derived from the fitting process conducted with the BOBYQA optimizer.

Given the observation of two visually distinguishable behaviors in the experimental data presented in Fig. 2a and Supplementary Fig.1, we aimed to identify a mathematical approach to categorize the data. To accomplish this, we initially plotted the values of B vs C on a semilogarithmic scale (Fig. 2d), which highlighted the existence of two distinct groups. Subsequently, we employed the NbClust package to determine the optimal number of clusters for our dataset, and the analysis confirmed that two clusters were the most appropriate choice. In the next step, to eliminate bias in the results of our clustering, we applied two clustering algorithms, namely Kmeans and K-medoids, to classify the dataset into two groups. The outcomes from both clustering algorithms exhibited similarity, effectively categorizing the experimental results into two groups: relaxed and tensioned (Supplementary Fig. 3).

Statistical analysis. The excluded outliers were defined as the values outside of the range $[Q_1 1.5(Q_3 - Q_1)$, $Q_3 + 1.5(Q_3 - Q_1)$], where Q_1 and Q_3 are the 25th and 75th percentile, respectively. The normality of each dataset was tested using Shapiro-Wilk tests, and equality of variances was tested using F-tests. When the data was drawn from a normally distributed population, the selection of the analysis method depended on the equality of variances among different data groups. When examining two groups of data, we conducted an independent sample t-test in cases of equal variances, and for cases of unequal variances, we employed Modified t-tests with Welch correction. For groups consisting of three or more, One-way ANOVA was utilized when variances were equal, while Welch's ANOVA was applied in instances of unequal variances. Subsequently, to identify statistically significant differences among group means, we conducted Tukey's post hoc tests for groups with equal variances and Games Howell's post hoc tests for groups with unequal variances.

If the data did not exhibit a normal distribution, we employed the Mann-Whitney test for the analysis of two data groups and the Kruskal-Wallis nonparametric ANOVA test with Dunn's post hoc test for three or more data groups. We conducted the statistical analysis using the OriginLab software. Datasets with $p < 0.05$ were considered to have significantly different means and are denoted by a single asterisk (*). Data sets with $p < 0.01$, $p < 0.001$, and $p < 0.0001$ were highly significantly different and were denoted by double $(**)$, triple $(***)$ and quadruple $(****)$ asterisks, respectively. The edges of the box plots represent the 25th and 75th percentiles of the data, the line in the middle marks the median, and whiskers extend to include maximum and minimum values in the dataset.

Active phenomenological model. Here, we employed an active phenomenological model (Fig 6a), which has been previously utilized by Etienne et al.²⁴ to explain how single cells respond to changes in the stiffness of their environment. Our goal was to apply this model to characterize the strain rate dependent response of cell pairs to a sustained strain. This model comprises two main components: a spring represented by E_1 , signifying the elastic modulus of the cell pair, and an active portion that includes an active element denoted as σ_A , representing the contractile stress generated by actomyosin activity, in parallel with a dashpot labeled as η_A , which accounts for the dissipation of force generated by myosin motor proteins through a mechanism resembling viscosity. Using this model, we were able to accurately describe the prolonged response of cell pairs in ramp and hold experiments.

The sensing island of the microstructure is modeled as a spring E_b in series with the model of the cell pair. These two springs (E_1 and E_b) are combined into a single equivalent spring of $E_{eq} = \frac{E_1 \cdot E_b}{E_1 + E_b}$. Stress and strain in the active section and the equivalent spring are presented below:

$$
\sigma = E_{eq} \varepsilon_s = \sigma_A + \eta_A \cdot \dot{\varepsilon}_a \tag{2}
$$

$$
\varepsilon = \varepsilon_s + \varepsilon_a \tag{3}
$$

where ε_s and ε_a are displacements in the equivalent spring and the active section, respectively. Constitutive equation of the model is therefore the following equation:

$$
\frac{\dot{\sigma}}{E_b} + \frac{\sigma}{\eta_A} = \dot{\varepsilon} + \frac{\sigma_A}{\eta_A} \tag{4}
$$

And the response of the model to a step strain ε_0 is:

$$
\sigma(t) = (E_{eq} \cdot \varepsilon_0 - \sigma_A)e^{-\frac{t}{\tau}} + \sigma_A
$$
\n(5)

with the characteristic time scale $\tau=\frac{\eta_{A}}{E_{eq}}.$ The relaxation modulus of the system is therefore:

$$
G(t) = \frac{1}{\varepsilon_0} \cdot \left[(E_{eq} \cdot \varepsilon_0 - \sigma_A) e^{-\frac{t}{\tau}} + \sigma_A \right]
$$
(6)

This relaxation modulus was then used in Equation (1) to fit the average relaxed and tensioned curves of the experimental data collected at 50% strain using the RHEOS software (Fig. 6b).

We have also shown that when $t \ll \tau$, stress in the system can be defined as:

$$
\sigma = E_{eq} \cdot \varepsilon_0 + \frac{\sigma_A - E_{eq} \cdot \varepsilon_0}{\tau} \cdot t \tag{7}
$$

which is equivalent to the empirical model we presented before, $\sigma(t) \ = \ B + C t,$ with $B = E_{eq}$ and $C = (\frac{1}{\varepsilon_0}) \cdot \frac{\sigma_A - E_{eq} \cdot \varepsilon_0}{\tau}$.

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COMPETING INTERESTS

None to declare.

AUTHOR CONTRIBUTIONS

A.M.E., J.R., and B.T.S. performed the tensile tests on single cell pairs. A.M.E., J.R. and A.O.M. analyzed the tensile tests and produced the stress-time and strain-time curves. B.T.S. and A.K. developed the phenomenological data analysis method and performed data analysis. G.M. conducted the AFM characterization of the TPP structures. J.R. performed imaging and image analysis. N.V.L. fabricated the microstructures used for the tensile test. R.Y. conceived the project and designed the experiments. C.H., A.K. and G.C. provided conceptual advice. B.T.S, J.R., A.K. and R.Y. wrote the manuscript. All authors discussed the results and the manuscript.

FIGURES

Figure 1. Mechanical response of epithelial cell pairs is strain rate dependent in ramp and hold experiments. a. SEM image of the single cell adhesion micro tensile tester platform used to stretch cell pairs. **b.** Schematic diagram presenting the method used to apply strain to the cell pair. Cells (blue) were deposited on each part of the bowtie and had time to make mature junctions. Then, they were stretched at different rates, and using the deflection in island 1 (δ) and island 2 (D), the stress and strain in the cell pair was calculated. **c.** Time course of the applied 50% strain with different rates. **d.** Confocal images of the cells prior and post stretch application (red: F-actin; green: structure autofluorescence). The cells are stretched at 25 %s⁻¹. **e.** Temporal evolution of stress (average of all curves for each condition \pm s.e.) for different rates. To make the comparison easier to visualize, we aligned the initiation of the holding phase to $t = 0$ s.

Figure 2. At 50% strain, the mechanical response of the cell pair to high strain-rate stretch is bimodal: relaxed and tensioned. a. Temporal evolution of the stress in the cell pair following a 50% strain applied at 10 %s-1 (N = 30). **b.** The example experimental relaxed and tensioned curves (gray) are fitted with the empirical model (red). **c.** Average stress-time curves obtained in ramp-hold experiments for various levels of strain rate (gray) are fitted with the empirical model (red). To make the comparison easier to visualize, we aligned the initiation of the holding phase to t = 0 s. **d.** B vs C values for all strain rates applied to reach 50% strain are plotted and clustered on a semi logarithmic scale. The results of clustering are represented by solid triangles and hollow circles to represent tensioned and relaxed curves, respectively. **e-f, h-i.** Box plots representing parameters $A \cdot \varepsilon_0$ (amplitude of relaxation), α (power law exponent), $B \cdot \varepsilon_0$ (residual stress), and $C \cdot \varepsilon_0$ (captures the slope of tensioning) for the cell pairs experiencing various levels of strain rate. **g.** Probability of tensioning for each strain rate, which was calculated by dividing the number of tensioned curves by the number of all curves for each condition.

Figure 3. The effect of the strain magnitude on the mechanical response of cell pairs subjected to high strain rates. a. Temporal evolution of stress (average of all curves for each condition ± s.e.) over time for cell pairs stretched at different rates to reach 125% strain. To make the comparison easier to visualize, we aligned the initiation of the holding phase to t = 0 s. **b.** Temporal evolution of stress for cell pairs stretched at a strain rate of 50 %s-1 to reach various strain magnitudes. **c.** B vs C values calculated from fitting the temporal evolution of stress over time in response to different strain magnitudes applied at 50 %s⁻¹ are plotted and clustered on a semi logarithmic scale. Solid triangles and hollow circles represent tensioned and relaxed responses, respectively. **d.** The probability of tensioning versus the strain magnitude and strain rate. The stars indicate the probability of tensioning calculated from the experimental results. **e.** Temporal evolution of the stress in the cell pair in response to stretch-hold-stretch (N = 14), and a single stretch to 100% strain (N = 12). In the stretch-hold-stretch experiments, the cell pairs were stretched to 50% at 25 %s⁻¹ strain rate. After 5 minutes of hold, the cell pairs were stretched at 25 %s-1 to reach a total of 100% strain. **f.** Probability of tensioning for stretch-hold-stretch experiments and a single stretch to 100% strain. The inset shows the applied strain over time for a stretch-holdstretch experiment and a single stretch to 100% strain.

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Figure 4. Actomyosin regulates the response of cell pairs in the ramp-and-hold experiments. a. Time series fluorescent images of F-actin in cells stretched to 50% strain at 50 %s⁻¹ and 0.5 %s⁻¹. Scale bar: 10 μm. **b, c.** Relative change in F-actin intensity within the confinement region (**b**) and near the cell-cell junction (**c**) while the strain was held constant. Scale bar: 10 μm. **d.** The evolution of stress over time in control (N = 10, orange) and Jasplakinolide (Jas) treated (N = 13, brown) cell pairs stretched at 50 %s⁻¹ to 125% strain (average of all curves for each condition ± s.e.). **e.** The average of the B vs C values calculated from fitting individual curves for the Jas treated cell pairs and the control condition on a semi logarithmic scale. Inset shows the clustered B vs C

values where solid triangles and hollow circles represent tensioned and relaxed responses, respectively. **f.** The probability of tensioning for the control and Jas treated cell pairs in response to 125 % strain applied at 50 %s-1. **g.** The evolution of stress over time in control (N = 40, green), Bleb (N = 16, gray), Y-27632 (N = 7, purple), and Lat B (N = 16, pink) treated cell pairs stretched at 25 %s⁻¹ to 50% (average of all curves for each condition \pm s.e.). **h.** The corresponding B vs C values calculated and clustered for the data presented in (**g**). **i.** The probability of tensioning for the cell pairs treated with actomyosin modulators and the control condition in response to 50% strain applied at 25 %s-1.

Figure 5. An active phenomenological model characterizes the bimodal stress response of the cell pair under different strain rates. a. A schematic illustration of the active phenomenological model which includes a spring labeled E_1 symbolizing the elastic modulus of the cell pair. The model also incorporates an active branch, consisting of an active element σ_A to signify the contractile stress generation in the actomyosin network, along with a dashpot labeled η_A , to represent the dissipation of the force generated by the myosin motor proteins through a viscous-like mechanism. **b.** The average relaxed and tensioned stress-time curves of the cell pairs stretched to 50% strain at 25 %s⁻¹ are fitted using Eq. (6) according to the active model. **c.** The values of η_A are calculated for tensioned curves at high rates (i.e., 10 %s⁻¹, 25 %s⁻¹, and 50 %s⁻¹). These values are then averaged and kept as a constant for the relaxed curves. d . The values of σ_A obtained from fitting the individual curves for different rates with Eq. (6).