

1 Intercellular Communication Controls Agonist-Induced Calcium 2 Oscillations Independently of Gap Junctions in Smooth Muscle Cells.

3 Authors

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13 Abstract

14 In this study, we report the existence of a communication system among human smooth
15 muscle cells that uses mechanical forces to frequency modulate long-range calcium waves.
16 An important consequence of this mechanical signaling is that changes in stiffness of the
17 underlying extracellular matrix can interfere with the frequency modulation of Ca²⁺ waves
18 causing smooth muscle cells from healthy human donors to falsely perceive a much higher
19 agonist dose than they actually received. This aberrant sensing of contractile agonist dose
20 on stiffer matrices is completely absent in isolated smooth muscle cells, even though the
21 isolated cells can sense matrix rigidity. We show that the intercellular communication that
22 enables this collective Ca²⁺ response in smooth muscle cells does not involve transport
23 across gap junctions or extracellular diffusion of signaling molecules. Instead, our data
24 support a collective model in which mechanical signaling among smooth muscle cells
25 regulates their response to contractile agonists.

26 **Introduction**

27 Excessive constriction of hollow, tubular transport organs including the airways and the
28 vasculature is a common pathophysiological feature of widespread diseases like asthma and
29 hypertension. The vessel/airway wall undergoes significant pathological changes in the
30 smooth muscle and in the extracellular matrix (ECM) that surrounds and supports the cells
31 with the onset of disease (1, 2). The search for mechanisms that underlie the development
32 of these diseases and the search for novel therapies has largely focused on the smooth
33 muscle cells (SMCs), as they are the primary effectors of constriction (3). Pathological
34 changes in the ECM, on the other hand, have not received much attention (1, 4). More recent
35 studies show that changes in the ECM can impact organ function at the very early stages,
36 and can even precede the thickening of the muscle layer (5). Perhaps, mechanobiological
37 interactions between healthy SMCs and an altered ECM are playing a more critical role in
38 the pathogenesis and progression of diseases like asthma and hypertension than is currently
39 appreciated. In this study, we examined how changes in the matrix stiffness can impact how
40 SMCs sense the dose of an applied agonist.

41 Agonist-induced Ca^{2+} oscillations and long-range Ca^{2+} waves are critical mechanisms that
42 regulate vital parameters such as blood pressure and airway resistance (6, 7). Binding of a
43 muscle agonist, like histamine or acetylcholine, to a surface receptor on the SMC and the
44 subsequent rise in cytosolic Ca^{2+} concentration, is the universal trigger for force generation
45 in the smooth muscle (8). Agonist exposure induces Ca^{2+} oscillations in SMCs that
46 propagate as waves within the smooth muscle layer (9). These agonist-induced Ca^{2+}
47 oscillations serve two critical functions in the smooth muscle: (A) The concentration/dose
48 of agonist detected by the surface receptors is transduced into the frequency of Ca^{2+}
49 oscillations, with a higher concentration of muscle agonist resulting in a higher frequency

50 of Ca^{2+} oscillations, which can then be detected by downstream Ca^{2+} sensors and translated
51 into a dose-dependent increase in smooth muscle contractility (6, 10). (B) Agonist-induced
52 Ca^{2+} oscillations propagate as waves around the circumference of the organ and enable the
53 synchronized contractions of SMCs necessary to constrict the airway/blood vessels (9). At
54 present, little is known about the role of extracellular mechanical factors such as ECM
55 stiffness in regulating agonist-induced Ca^{2+} oscillations and Ca^{2+} waves that can move
56 across an SMC ensemble.

57 In this study, we report a collective phenomenon in clusters of human airway smooth muscle
58 cells, where ECM stiffness alters the intercellular communication between cells in an SMC
59 ensemble causing increased Ca^{2+} oscillation frequencies and synchronized Ca^{2+} oscillations.
60 This altered Ca^{2+} response results in a relative force increase by SMCs on stiff substrates.
61 We examined intercellular transport of Ca^{2+} in SMC cells and we show that contrary to
62 current dogma, the physical mechanism that enables intercellular Ca^{2+} waves does not
63 involve molecular transport across gap junctions or paracrine signaling through
64 extracellular diffusion. Rather, this phenomenon appears to be driven by force-transfer
65 among cells in the cluster. The collective response of SMCs to agonist could be a
66 mechanism by which matrix remodeling can drive disease progression in asthma and
67 hypertension.

68 **Results**

69 **1. Matrix stiffness alters the Ca^{2+} response to agonist in multicellular clusters of SMCs,** 70 **but not in isolated cells.**

71 Modeling the SMC layer in 2D using micropatterning: To study the role of altered matrix
72 stiffness on the frequency of Ca^{2+} oscillations, we used micropatterning to create a 2D
73 approximation of the organization of SMCs seen in lung slices (Fig. 1A, S1A). The substrate

74 we use is NuSil (11), an optically clear, non-porous polydimethylsiloxane substrate whose
75 Young's modulus, E , can be varied in the range 0.3 kPa-70 kPa (11). Based on
76 measurements of ECM stiffness in healthy human airways, we set the ECM stiffness of
77 healthy human airways to $E=0.3$ kPa (12). This matches the ECM stiffness of small airways
78 (inner diameter <3 mm) which are known to collapse in asthma (13). With the onset of
79 airway remodeling, collagen is deposited in the airways and the stiffness of the ECM
80 increases (1). A substrate stiffness of $E=13$ kPa was used to mimic remodeled ECM. These
81 ECM stiffness values are also representative of the two distinct regimes of mechanosensing
82 seen in all adherent cells (14, 15). Using a Ca^{2+} sensitive fluorescent dye (Fluo4-AM), we
83 imaged and quantified the time period of Ca^{2+} oscillations in these SM rings plated on soft
84 and stiff ECM. Images were recorded at a rate of 1 per second for 5 minutes following
85 exposure to 10^{-5} M histamine. On soft ECM ($E=0.3$ kPa), exposure to histamine resulted in
86 Ca^{2+} oscillations with a mean time period of 43.64 ± 1.45 seconds ($N=4$, Fig. 1E). When the
87 substrate stiffness was increased to $E=13$ kPa, the same dose of agonist induced significantly
88 faster oscillations with a mean time period of 22.79 ± 2.42 seconds ($N=4$, t-test, $P < 0.001$,
89 Fig. 1E). Therefore, at the *same* dose of agonist, stiff substrates resulted in a doubling of the
90 cytosolic Ca^{2+} oscillation frequency in healthy smooth muscle cells.

91 Interactions between ECM and isolated cells are insufficient to explain altered Ca^{2+}
92 response: To explain the role of matrix stiffness in regulating the Ca^{2+} response to a low
93 dose of agonist, we first hypothesized that this phenomenon was linked to cell-matrix
94 interactions at the level of the individual cell. With increased matrix stiffness, SMCs
95 develop higher cytoskeletal prestress (15) opening stretch-activated Ca^{2+} channels (16) and
96 potentially increasing the Ca^{2+} flux into the cell. To test this hypothesis, we cultured human
97 airway SMCs at a low density such that individual cells were isolated, spaced at least 100
98 μm from each other (Fig. 1B, S1B). We first measured baseline traction (pre-agonist) stress

99 exerted by these isolated cells on the substrate to confirm that the baseline traction was
100 significantly higher in isolated cells cultured on stiff, with a mean traction stress of
101 15.95 ± 5.02 Pa (N=21), versus isolated cells cultured on soft substrates, with a mean traction
102 stress of 6.98 ± 1.68 Pa (N=16). The corresponding median stresses were 6.89 Pa on soft
103 matrix and 16.00 Pa on stiff substrate (Mann-Whitney Rank Sum test, $P < 0.001$). We then
104 exposed these isolated SMCs to 10^{-5} M histamine and measured the time period of Ca^{2+}
105 oscillations. Contrary to our expectations, ECM stiffness had no impact on the Ca^{2+} response
106 of isolated SMCs to 10^{-5} M histamine (Fig. 1F). Cells cultured on the soft substrate had a
107 mean period of 66.11 ± 10.55 seconds (N=14) and those on the stiff matrix had a mean period
108 of 65.73 ± 11.48 seconds (N=12), which was not statistically significantly different (t-test,
109 $P = 0.930$). Therefore, despite the higher levels of prestress in individual cells, ECM
110 stiffening has no impact on the agonist-induced Ca^{2+} frequency of isolated cells.

111 SMCs sense matrix as a collective and alter their Ca^{2+} response to agonist: To probe this
112 phenomenon further, starting with the isolated SMCs (Fig. 1B), we increased the seeding
113 density (Fig. 1C, S1C) until we had a confluent cluster of SMCs (Fig. 1D, S1D). At each
114 seeding density, we measured the time period of Ca^{2+} oscillations for SMCs adhering to soft
115 ($E = 0.3$ kPa) and stiff ($E = 13$ kPa) substrates in response to 10^{-5} M histamine (Fig. 1F-1H).
116 Sparsely seeded cells (Fig. 1G) responded similarly to isolated cells (Fig. 1F), exhibiting no
117 statistically significant difference between the agonist-induced Ca^{2+} oscillations on soft and
118 stiff matrix (t-test, $P = 0.481$). The mean time period of oscillations of sparse SMCs on the
119 soft substrate was 51.74 ± 8.11 seconds (N=4) and 55.63 ± 6.48 seconds (N=4) on the stiff
120 substrate (Fig. 1G). Confluent cells behaved like those patterned in a ring, with cells plated
121 on stiffer substrate exhibiting a significantly higher frequency of Ca^{2+} oscillations in
122 response to 10^{-5} M histamine. Confluent SMCs on soft matrix had a mean time period of
123 58.36 ± 5.89 seconds (N=5), and the same healthy, confluent cells on stiff matrix had a mean

124 period of 35.03 ± 1.03 seconds ($N=7$, Fig. 1H). The corresponding median time periods were
125 58.87 seconds on soft matrix and 35.11 seconds on stiff matrix (Mann-Whitney Rank Sum
126 Test, $P=0.003$). These results suggest the existence of a collective phenomenon in smooth
127 muscle cells, where clusters of confluent SMCs alter their agonist-induced Ca^{2+} oscillations
128 in response to changes in matrix stiffness while isolated cells do not. In the supplementary
129 information (Fig. S2), we include additional measurements made on two intermediate
130 stiffnesses, $E=0.6$ kPa, $E=3$ kPa, and glass ($E \rightarrow \infty$). These experiments confirm that Ca^{2+}
131 oscillations resulting from exposure to 10^{-5} M histamine are unaffected by ECM stiffness in
132 isolated SMCs, whereas in confluent cells, the same dose of agonist can evoke a
133 significantly different Ca^{2+} response in SMCs depending on the stiffness of the underlying
134 matrix. This finding is extremely significant in all smooth muscle pathologies because all
135 downstream Ca^{2+} dependent molecular processes rely on Ca^{2+} oscillations to perceive the
136 external concentration of contractile agonist detected by the G-protein coupled receptors on
137 the cell surface. Here we show that the combination of confluence and ECM stiffness can
138 alter how cells perceive contractile agonist.

139 **2. Matrix stiffening synchronizes Ca^{2+} oscillations within a multicellular SMC cluster.**

140 We next explored the nature of intercellular communication underlying the collective
141 agonist-induced Ca^{2+} response in SMCs. To do this, we first analyzed the time series of
142 histamine-induced Ca^{2+} oscillations for signs of interactions among the different SMCs
143 within a confluent cluster. After correcting for drift due to photobleaching of the
144 fluorophore, we calculated the cross-correlation coefficient ($\rho_{i,j} \in [-1,1]$) of the Ca^{2+}
145 oscillations occurring in the i^{th} cell and the j^{th} cell in the cluster for all the cells in a cluster.
146 The measured values of $\rho_{i,j}$ in a typical SMC cluster is depicted in Fig. 2A as a representative
147 24×24 matrix, with the extreme values $\rho_{i,j} \rightarrow 1$ (pink) indicating that the Ca^{2+} levels in the

148 i^{th} and j^{th} cell rise and fall perfectly in sync with each other (perfectly correlated) (Fig. 2B),
149 $\rho_{i,j} \rightarrow -1$ (green) indicating that when the Ca^{2+} levels in i^{th} cell rises, the Ca^{2+} levels in the j^{th}
150 cell falls and vice-versa (anti-correlated), $\rho_{i,j} \rightarrow 0$ (white) indicating no correlation in the
151 Ca^{2+} oscillations occurring in i^{th} and j^{th} cells (Fig. 2B). To avoid the effects of histamine
152 diffusion, the first 60 seconds immediately after application of histamine was not considered
153 in the correlation calculation. Histograms of $\rho_{i,j}$ measured in isolated SMCs and confluent
154 clusters of SMCs cultured on soft and stiff matrices are shown in Figs. 2C and 2D,
155 respectively. Isolated SMCs (Fig. 2C), did not exhibit correlated Ca^{2+} oscillations, with a
156 mean ρ of zero, regardless of whether they were cultured on soft (0.01 ± 0.06 , $N=5$) or stiff
157 (0.03 ± 0.06 , $N=4$) substrates. However, for cells in a confluent cluster, matrix stiffening
158 caused a statistically significant shift in ρ towards positive correlation (Fig. 2D). The mean
159 ρ for confluent clusters on soft matrices was 0.03 ± 0.04 , $N=4$, versus on stiff matrices, ρ
160 increased to 0.26 ± 0.10 , $N=5$. A two-way ANOVA test with confluence and ECM stiffness
161 as independent factors showed a significant interaction between confluence and ECM
162 stiffness ($P=0.007$). Post-hoc pairwise comparisons using the Holm-Sidak test showed a
163 significant difference in the pairwise correlations in confluent clusters due to ECM stiffness
164 ($P<0.001$). In the isolated cells, there was no statistical difference in the pairwise
165 correlations due to ECM stiffness ($P=0.733$). We also did not observe any systematic
166 relationship between the pairwise correlation in Ca^{2+} oscillations and the distance between
167 the SMCs.

168 Next, we investigated the time it takes after the addition of histamine for the Ca^{2+}
169 oscillations to synchronize. In order to do this, we repeated the previous calculation of cross-
170 correlations, but instead of using the entire time series, we used a time window of 120
171 seconds starting at $t=60$ seconds after addition of histamine and repeated the $\rho_{i,j}$ calculation
172 for the Ca^{2+} time series within this 120 seconds window for all cells in the cluster. The time

173 window was then shifted in increments of 15 seconds over the rest of the 5-minute time
174 frame over which we measured Ca^{2+} oscillations. Shown in Fig. 2E and Fig. 2F are the
175 average through time, $\rho_{i,j}(\tau)$ of cross-correlation coefficients between all cells either isolated
176 or confluent, soft and stiff matrix, respectively. A two-way ANOVA with time after addition
177 of histamine and ECM stiffness as the two independent factors shows no difference due to
178 stiffness ($P=0.303$) or time ($P=0.274$) in isolated cells. However, in confluent cells on stiff
179 ECM, the correlation coefficients gradually increase over time, for about 1 minute, which
180 matches the time course for force generation in airway SMCs (17). In conjunction with our
181 findings from the previous section, these results demonstrate that not only does the
182 combination of a stiff matrix and confluent clusters of cells lead to faster Ca^{2+} oscillations,
183 but it also causes histamine-induced Ca^{2+} oscillations to synchronize across the cells in the
184 cluster.

185 **3. Gap junctions do not play a role in regulating the collective Ca^{2+} response of SMC** 186 **clusters.**

187 The most well studied long-range communication mechanism in multicellular systems is
188 the intercellular waves of Ca^{2+} that are capable of propagating over distances much longer
189 than a cell length through a regenerative Ca^{2+} induced Ca^{2+} release mechanism (18). This
190 mode of communication is mediated through two critical pathways: (i) gap junction
191 channels, which connect the cytoplasm of neighboring cells and allow for the transport of
192 signaling molecules from one cell to its neighbor and (ii) extracellular diffusion of a
193 signaling molecule like ATP, which can diffuse and bind to purinergic receptors on
194 neighboring cells, causing Ca^{2+} release in these cells (19).

195 Transport through gap junctions is unaffected by ECM stiffness: We first fluorescently
196 labeled gap junctions by staining for connexin-43 (Cx43). The expression of Cx43 (red) for

197 confluent clusters of SMCs on soft and stiff substrates are shown in Figs. 3A and 3B,
198 respectively. The actin filaments (green) and the nucleus (blue) are also labeled. From these
199 images, we quantified the number of gap junctions per cell for confluent cells plated on soft
200 and stiff substrates. There was no significant difference between Cx43 expression on soft
201 and stiff substrates from N=5 independent samples per stiffness, each containing⁽¹⁾
202 approximately 35 cell measurements (t-test, P=0.977) (Fig. 3C). Next, we tested whether
203 ECM stiffness induced a change in the efficiency of transport through gap junctions. To this
204 end, we employed a commonly used technique to quantitatively assess the efficiency of
205 transport through gap junctions called gap-FRAP (20). Briefly, a confluent layer of SMCs
206 was incubated with membrane-permeable calcein-AM. Upon entering the cell, the acetyl
207 methyl ester bond was hydrolyzed by intracellular esterases, trapping the hydrophilic
208 calcein molecule within the cell (Fig. 3D, column 1). The calcein in one cell selected at
209 random was then bleached using a high-intensity laser (Fig. 3D, column 2). The bleached
210 calcein molecules and the unbleached calcein molecules in neighboring cells diffuse
211 through gap junctions leading to a recovery in fluorescence (Fig. 3D, column 3). The
212 kinetics and extent of recovery in fluorescence in the bleached cell reflects the efficiency of
213 transport across gap junctions via diffusion. A typical recovery curve is shown in Fig. 3E,
214 where F_0, F_b, F_r indicate the fluorescence intensity in the target cell at baseline, after
215 bleaching and after recovery. To quantify the extent of recovery, we calculated the mobile
216 fraction, Γ , given by

$$\Gamma = \frac{F_r - F_b}{F_0 - F_b} \times 100\%$$

218 for confluent SMCs cultured on soft and stiff matrices (N=16, N=18 independent trials,
219 respectively). We found no difference in the mobile fraction due to ECM stiffness. SMCs
220 on soft matrix had a mean mobile fraction, Γ of $42.19 \pm 10.50\%$, and those on stiff matrix

221 had a Γ of $40.05 \pm 9.18\%$ (t-test, $P=0.532$) (Fig. 3F). We also did not observe any statistically
222 significant difference in the rate of recovery in fluorescence k_r , which was calculated by
223 fitting an exponential function $F(t) = F_b + F_r(1 - e^{-k_r t})$ to the recovery curve. On soft ECM,
224 k_r was 0.01 ± 0.02 (N=16) and on stiff ECM, k_r was 0.01 ± 0.05 (N=18). There was no
225 statistical difference in the rate of recovery due to ECM stiffness (t-test, $P=0.55$).

226 Blocking gap junctions does not affect agonist-induced Ca^{2+} oscillations: To further explore
227 the role of gap junctions in the collective response of the SMCs to agonist, we used $30 \mu\text{M}$
228 of β -glycyrrhetic acid (βGA) to block transport across gap junctions. βGA causes
229 dephosphorylation of connexins and disassembly of gap junctions (21). We first used gap-
230 FRAP to verify that a $30 \mu\text{M}$ dose of βGA was sufficient to completely block transport
231 across gap junctions. The mobile fraction, Γ , measured in a confluent layer of SMCs after
232 application of $30 \mu\text{M}$ βGA dropped significantly from a mean of $40.05 \pm 9.18\%$ to
233 $6.84 \pm 2.89\%$ (N=18, with corresponding medians 40.41% and 7.02% , respectively, Mann-
234 Whitney Rank Sum Test, $P < 0.001$) (Fig. 3F). This recovery is similar to the 7.89% recovery
235 we measured in an isolated cell. Similar minimal recovery has been noted in the literature
236 (22), indicating that transport across gap junctions was blocked. After 30 minutes of
237 incubation with $30 \mu\text{M}$ dose of βGA , we measured histamine-induced Ca^{2+} oscillations in
238 confluent clusters of SMCs. Much to our surprise, we found that blocking gap junctional
239 transport had no impact on the histamine-induced Ca^{2+} oscillation periods in multicellular
240 clusters of SMCs, regardless of the stiffness of the ECM (N=6 soft, N=4 stiff) (Fig. 3G). A
241 two-way ANOVA with treatment (βGA) and ECM stiffness as the two independent factors
242 showed no difference in the time period of histamine-induced Ca^{2+} oscillations due to
243 treatment ($P=0.784$) for a given stiffness. Post-hoc pairwise comparisons with the Holm-
244 Sidak method showed the Ca^{2+} oscillations remained significantly faster on the stiff matrix
245 ($P < 0.001$) even after blocking gap junctions.

4. Mechanical force transfer among cells regulates the collective Ca²⁺ response of SMC clusters.

Ca²⁺ wave propagation follows the contractile axis of the SMCs: Next, we considered extracellular diffusion of signaling molecules, such as ATP, as a possible mode of intercellular communication that enables the collective Ca²⁺ response of SMC clusters on stiff matrices (18). In order to do this, we measured the direction in which the Ca²⁺ wave propagates from one cell to the next in SMC clusters cultured on stiff substrates. If we select a cell in an SMC cluster, a wave that passes through it will appear as a localized increase in Ca²⁺ in the cell which is followed by a localized increase in Ca²⁺ in one of its neighbors. We reasoned that if the intercellular Ca²⁺ transport was being enabled by extracellular diffusion of signaling molecules, then the resulting Ca²⁺ wave should have an equal chance of moving in all directions (isotropic). We split the direction of propagation of the Ca²⁺ wave from an SMC into two directions: a direction parallel to the contractile axis of the SMC, and a direction perpendicular to the contractile axis of the SMC. For the cell labeled 1 shown in Fig. 4A (insets), cells 2 and 4 were considered parallel to cell 1's contractile axis and cells 3 and 5 were considered perpendicular to cell 1's contractile axis. We calculated the conditional probability for a localized increase in Ca²⁺ in one cell to be followed by a localized increase in a parallel neighbor versus its perpendicular neighbor. This conditional probability quantifies the isotropy in Ca²⁺ wave propagation with respect to the contractile axis of the SMC, with equal probability (0.5) in parallel and perpendicular direction indicating isotropy in Ca²⁺ wave propagation. Instead, we found that there was an 80.5±7.53% chance for the Ca²⁺ wave to follow the contractile axis of the SMC (Fig. 4B, N=10, t-test, P<0.001). The high probability of the Ca²⁺ wave to follow the contractile axis of the SMCs rules out extracellular diffusion as the dominant mechanism for intercellular communication in our experiments.

271 Mechanical force transfer between SMCs regulates their Ca^{2+} response: Thus far, we have
272 ruled out transport through gap junctions and extracellular diffusion, the two widely
273 accepted mechanisms responsible for intercellular communication in confluent cell clusters
274 using Ca^{2+} waves (18). The propensity of Ca^{2+} waves to follow the direction of the
275 contractile axis of the SMC suggested force transfer between neighboring SMCs as a
276 potential mechanism that regulates the collective Ca^{2+} response of SMC clusters. To
277 quantify SMC forces in our confluent clusters, we used Fourier transform traction force
278 microscopy (23) to measure the traction stress generated by cells exposed to 10^{-5} M
279 histamine. Traction stress is the force exerted by the cell normalized by the cross-sectional
280 area over which the force acts. Before agonist stimulation, mean traction stresses for isolated
281 cells were 6.98 ± 1.68 Pa on soft (N=16) and 15.95 ± 5.02 Pa on stiff (N=21) substrates, and
282 13.56 ± 1.56 Pa on soft (N=12) and 14.26 ± 1.89 Pa on stiff (N=21) for confluent cells. We
283 found that histamine had generated a normalized traction (post-histamine traction stress/pre-
284 histamine traction stress) of 1.22 ± 0.11 in confluent cells on soft substrates (N=12). On
285 stiffer matrix, a confluent cluster of SMCs from the same healthy donor and passage
286 generated normalized force of 2.24 ± 0.26 (N=21) (Fig. 4C). There was a statistically
287 significant difference ($P < 0.001$, t-test) between the force generated on soft and stiff
288 substrates in response to the same dose of agonist. To test the possibility that higher force
289 transfer among cells on stiff substrates was responsible for this collective phenomenon, we
290 experimentally measured the effect of reducing muscle force on Ca^{2+} oscillations with two
291 independent inhibitors of force generation: (1) Rho-associated kinase (ROCK) inhibitor, Y-
292 27632, which reduces SMC force by inhibiting myosin light chain phosphorylation and by
293 removing the inhibitory effect of ROCK on the activity of myosin light chain phosphatase
294 and (2) myosin light chain kinase (MLCK) inhibitor, ML-7, which reduces force by
295 inhibiting myosin light chain phosphorylation. Confluent SMC clusters on $E=13$ kPa matrix

296 were pre-treated with increasing doses of either Y-27632 for 1 hour prior to histamine
297 exposure or increasing doses of ML-7 for 5 minutes prior to histamine exposure (Fig. 4D).
298 We found that increasing the concentration of Y-27632 from 10 μ M (N=3) to 100 μ M (N=4)
299 abolished the stiffness-induced reduction in time period of Ca^{2+} oscillations in a dose-
300 dependent manner (One-Way ANOVA with treatment as the independent variable,
301 $P<0.001$). Similarly, increasing the dose of ML-7 from 0.01 μ M to 0.05 μ M also abolished
302 the stiffness-induced reduction in time period of Ca^{2+} oscillations in a dose-dependent
303 manner (One-Way ANOVA, $P<0.001$). With both these inhibitors, the higher concentration
304 resulted in histamine-induced Ca^{2+} oscillation periods on stiff matrices that were not
305 statistically different from the histamine-induced Ca^{2+} oscillation periods on soft matrices.
306 These results clearly demonstrate that the exaggerated histamine-induced Ca^{2+} response due
307 to matrix stiffening can be *systematically*, and *completely* abrogated by reducing mechanical
308 forces in the multicellular ensemble.

309 **5. Confining the SMCs to a line reduces the variability in Ca^{2+} oscillation periods:**

310 Comparing the alignment of SMCs in Fig. 1B to Fig. 1D and Fig. 4A, we observed that with
311 the onset of confluence, SMCs naturally tend to organize themselves into spatial clusters of
312 aligned cells. To test the effect of SMC alignment on the higher frequency of Ca^{2+}
313 oscillation on stiff substrates, we compared the time period of Ca^{2+} oscillations in confluent
314 SMC clusters to the time period of Ca^{2+} oscillations in SMC clusters micropatterned in a
315 line which was 1000 μ m long and 15 μ m wide (~1 cell wide and ~10 cell lengths long) (Fig.
316 5A, S1E). The idea here was to eliminate the possibility of intercellular communication
317 occurring perpendicular to the contractile axis in confluent clusters; through mechanisms
318 that are less likely to be influenced by force. We found that while the mean time period of
319 oscillations was nearly identical in both confluent SMC clusters and lines of SMCs, the

320 probability of cells with a higher time period of Ca^{2+} oscillations decreases when SMCs are
321 aligned (Fig. 5B). An F-test shows a significant decrease in the variability of the time period
322 distribution when the cells are aligned in a line (N=8 soft, N=6 stiff, $P < 0.001$). This result
323 is consistent with the idea that higher Ca^{2+} frequencies are being driven by force transfer
324 along the contractile axis.

325 The effect of localized ECM stiffening can be sensed by SMCs over long distances: ECM

326 remodeling in airways and blood vessels often occur as spatially localized processes. How
327 does this pathological change in the ECM spread? Current theories require cells to migrate
328 into the region of stiffer ECM for them to sense the altered matrix and respond by excessive
329 secretion of matrix proteins thereby creating a positive feedback loop that leads to more
330 ECM remodeling (24). However, given the collective nature of ECM stiffness sensing in
331 SMC clusters, it may be possible for SMCs located far away from the site of ECM
332 remodeling to detect this localized change, even though these SMCs are not physically in
333 contact with the stiff ECM. To test this hypothesis and to quantify the distance over which
334 a localized increase in ECM stiffness would be felt by an ensemble of SMCs, we created a
335 dual-stiffness substrate (Fig. 5C), where the region marked in green has Young's modulus
336 of 13 kPa and the region in black has Young's modulus of 0.3 kPa. We then patterned SMCs
337 in a line starting from the stiff region and extending into the soft region. Cells on soft and
338 stiff ECM were simultaneously exposed to 10^{-5} M histamine, and we measured the time
339 period of Ca^{2+} oscillations in SMCs on the soft ECM for 5 minutes. The change in Ca^{2+}
340 oscillation time period was not sudden as one moved from the stiff to the soft ECM (Fig.
341 5D). Rather, the mean time period increased at a slow rate of 1.83 s/100 μm . We grouped
342 the cells by distance from the edge into 400 μm bins and statistically tested the difference
343 in the time period of Ca^{2+} oscillations between each bin and cells in physical contact with
344 the stiff substrate. We found that there was no statistically significant difference between

345 histamine-induced Ca^{2+} oscillations for cells on stiff substrates and cells up to 800 μm
346 (approximately 8 cell lengths) away from the edge (N=4, Mann-Whitney Rank Sum Test,
347 P=0.140). Cells beyond 800 μm had significantly different time periods (N=4, t-test,
348 P<0.001). This result demonstrates that spatially localized alterations in the ECM can be
349 detected by SMCs far from the site of ECM remodeling, suggesting that ECM pathology
350 can spread through the organ much faster than currently believed.

351 Discussion

352 Increased stiffness of the extracellular matrix (ECM) that surrounds and supports cells in
353 tissue is associated with a number of disease conditions ranging from cancer and fibrosis
354 (25) to cardiovascular (26), lung (1), eye (27), and age-related diseases (28). Traditionally,
355 pathological alterations in the ECM were thought to be the consequence of disease
356 progression. However, it is now becoming increasingly apparent in many diseases that
357 matrix stiffening precedes disease development and could, therefore, contribute to disease
358 progression (5, 29). Recognizing the importance of ECM remodeling, clinical trials were
359 undertaken to restore the healthy, homeostatic state of the ECM (30). These early efforts
360 were unsuccessful (31) and attention has now turned to understanding and targeting the
361 mechanisms by which cells perceive and respond to changes in the ECM (29).

362 Here, we demonstrate a collective phenomenon in smooth muscle cells (SMC) in which
363 matrix stiffness alters the intercellular communication between cells in an ensemble
364 resulting in elevated contractility at low doses of agonist. We show that this collective
365 mechanosensing phenomenon is enabled by crosstalk among cells using Ca^{2+} waves (18). A
366 common mechanism of communication involves the molecular transport of Ca^{2+} and
367 inositol trisphosphate (IP_3) across gap junctions. However, our measurements showed no
368 tendency for molecular transport through gap junctions to differ depending on ECM

369 stiffness. We blocked transport through gap junctions using 18 β -glycyrrhetic acid (β GA)
370 (32) and confirmed that the dose we used was sufficient to completely disrupt molecular
371 transport across gap junctions. Contrary to our expectations based on current models (18),
372 blocking transport through gap junctions had no impact on the collective agonist response
373 of SMCs to ECM stiffening. There was no change in the Ca^{2+} oscillation for SMCs on soft
374 and stiff substrates after β GA treatment. Following a recent finding that mechanical forces
375 can synchronize contraction of cardiac myocytes in the developing heart independent of gap
376 junctions (32), we tested whether ECM stiffening led to a switch in the mode of
377 communication between cells from molecular signaling through gap junctions to
378 mechanical force based signaling.

379 Ca^{2+} waves can travel distances far greater than a cell length because Ca^{2+} waves regenerate
380 in each cell by “calcium-induced calcium release” from the endoplasmic reticulum (ER). In
381 the most widely accepted theory of long-range Ca^{2+} transport, Ca^{2+} release from the ER is
382 enabled by a messenger molecule, IP_3 , which diffuses faster than Ca^{2+} and activates the
383 receptors on the ER, so when the Ca^{2+} wave arrives, it can release more Ca^{2+} . This theory
384 had its basis in measurements of IP_3 diffusivity in the *Xenopus* extract model of the
385 cytoplasm (33), which put the diffusivity of IP_3 at $400 \mu\text{m}^2/\text{s}$ and that of Ca^{2+} at $40 \mu\text{m}^2/\text{s}$.
386 However, more recent measurements made in human cells show that IP_3 diffuses at a slower
387 rate than Ca^{2+} (34). Our finding that molecular diffusion through gap junctions is not
388 necessary to sustain Ca^{2+} waves in SMC clusters is consistent with the challenge to the
389 dogma of Ca^{2+} wave propagation through gap junctions.

390 In contrast to Ca^{2+} wave propagation in other cell types, measurements in confluent SMC
391 clusters (Fig. 4A,B) show that Ca^{2+} waves follow the direction of the contractile axis of the
392 SMCs. Further, reducing the SMC force in our experiments with two independently-acting

393 muscle relaxants slowed the Ca^{2+} oscillations and intercellular Ca^{2+} waves (Fig. 4D),
394 suggesting that mechanical force transfer from one cell to its neighbor enables the
395 intercellular Ca^{2+} transport. Can force transfer between cells also serve as a mechanism that
396 amplifies and modulates the frequency of oscillations as the Ca^{2+} wave moves from cell-to-
397 cell? The mechanisms that underlie the findings of the present study can be understood in
398 light of previous work from Felix et al (35) and Tanaka et al (36) who show that forces
399 applied to a cell membrane cause PIP_2 to be hydrolyzed resulting in an increase in cytosolic
400 IP_3 concentration. This would mean that for every SMC cell within a confluent cluster, there
401 are two ways by which IP_3 can be released into the cytosol: (i) from agonist binding to
402 GPCR receptors on the cell surface. (ii) from IP_3 generated by a contracting SMC pulling
403 on its neighbor causing IP_3 release in the neighboring cell (35, 36). This *additional* method
404 of IP_3 release only exists for cells in a confluent cluster. Further, we have previously shown
405 that force transfer between SMCs increases 8-fold with matrix stiffening (12). Such a force-
406 based IP_3 release mechanism can explain how SMC clusters alter their agonist-induced Ca^{2+}
407 oscillations in a matrix stiffness-dependent manner while isolated cells, which lack the
408 second source of IP_3 , do not.

409 Dysfunction in the smooth muscle has long been thought to be responsible for the
410 exaggerated narrowing of transport organs like as asthma, hypertension and Crohn's
411 disease. Asthma is an example of a disease where remodeling of the ECM is well
412 characterized (1), but its effects are not considered in therapy or drug development.
413 Asthmatics can be free of inflammation, and have spirometry and respiratory mechanics
414 within the range of healthy individuals, up until they are exposed to a smooth muscle agonist
415 at which point airways in an asthmatic will hyper-constrict (37). This exaggerated response
416 of the airway is currently thought to result from the sensitization of force-generating
417 pathways in the SMC due to prolonged exposure to inflammatory agents. Here, we

418 demonstrate that the fault may instead lie in changes in ECM stiffness that can regulate how
419 smooth muscle cells perceive agonist dose. To assess the overall magnitude and potential
420 physiological impact of the observed effect, we measured the Ca^{2+} response to different
421 doses of histamine at two different stiffnesses (Fig. S3). The data shows that if the ECM
422 stiffness of small airways (inner diameter < 3 mm) in a healthy human lung with $E=0.3$ kPa
423 (*12*) were to remodel and become stiff ($E=13$ kPa), healthy airway SMCs would now
424 responds to a dose of 10^{-5} M histamine as though they received 10^{-3} M histamine (two log
425 dose higher concentration). We also demonstrate that the corresponding agonist-induced
426 force is also substantially higher (Fig. 4C).

427 There are several confounding factors to consider as one extrapolates these findings at the
428 length scale of cells to the length scale of the airway/lung. These confounding factors
429 include, but are not limited to (i) connectivity among smooth muscle cells within the airway:
430 to constrict an airway, SM cells need to connect with each other to form a percolating force
431 transfer path that winds around the circumference of the airway, and generates a force
432 sufficient to narrow the airway lumen. These physical connections between cells can occur
433 through direct cell-cell (cadherin-based contacts) or cell-ECM (focal adhesion) connections,
434 which are in turn dependent on mechanical forces (*12*). (ii) Transmural pressure: the
435 mechanical load presented by the ECM which will vary depending on the size of the airway
436 and the transmural pressure. (iii) Additional confounding factors: surface tension, airway
437 tone, parenchymal tethering forces, etc. will also play a role in dictating the degree to which
438 airway lumen narrows in response to smooth muscle activation by inhaled agonist.
439 Experimentally validating these results in lung tissue will require the development of new
440 technology to alter ECM stiffness of airways. With careful calibration, it might be possible
441 to adapt methods from tissue engineering like UV-induced crosslinking of collagen to study
442 the effects of ECM stiffening *in situ* in precision-cut lung slices.

443 At first glance, it appears contradictory that stretch decreases airway smooth muscle force
444 (38). Yet, stretch has also been shown to trigger a rise in cytosolic Ca^{2+} (39). The key to this
445 apparent paradox may be found in the single stretch experiments of Krishnan et al (40) as
446 well as computational models of the effect of stretch on muscle force (15). These studies
447 show that the decline in force occurs almost instantly upon stretching and is related to
448 rupture events in the cytoskeleton. The recovery, however, is slow and occurs at a timescale
449 of the order of 100s in human smooth muscle. When a cell is being subjected to cyclic
450 stretch (such as the length oscillations), the cell is attempting to recover its original prestress
451 which it lost immediately after the onset of stretch. To regain prestress, myosin activity is
452 required, and this necessitates higher Ca^{2+} levels.

453 ECM remodeling in airways and blood vessels often occur as spatially localized processes.
454 We also show that a spatially localized change in the ECM might be sufficient for
455 exaggerated force generation in SMCs far away from the site of ECM remodeling.
456 Consequently, the effects of ECM remodeling may manifest much earlier in disease
457 progression than currently believed. Our findings are relevant to discovering new pathways
458 by which ECM remodeling can be the primary driver of disease and for ongoing efforts to
459 develop new drugs that target mechanosensory pathways (29). For airway smooth muscle,
460 and asthma, in particular, these results show that the current treatment regimen may need to
461 shift to new paradigms that target matrix remodeling and mechanosensory pathways for a
462 more lasting cure.

463 **Materials and Methods**

464 **Fabrication of optically clear substrates of tunable stiffness:** NuSil is an optically clear,
465 biologically inert PDMS substrate with Young's Modulus tunable in the range from 0.3 kPa
466 to 70 kPa (11). Equal parts of NuSil gel-8100 parts A and B (NuSil, Carpinteria, CA, USA)

467 were mixed with various amounts of the crosslinking compound of Sylgard 184 (Dow
468 Corning, Midland, MI, USA) to adjust substrate stiffness. A crosslinker volume 0.36% of
469 the combined parts A and B volume was added for substrates with Young's modulus $E = 13$
470 kPa, and no crosslinker was added to the 1:1 A:B mixture for substrates with Young's
471 modulus $E = 0.3$ kPa. After mixing, the substrate was spin coated onto 30 mm diameter, #1.5
472 glass coverslips for 50 seconds to produce a 100 μm -thick layer. They settled on a level
473 surface at room temperature for 1 hour before curing at 60° C overnight. These cured
474 substrates on coverslips were secured in sterile 40 mm Biotech dishes (Biological Optical
475 Technologies, Butler, PA, USA) to be used for cell culture.

476 **Matrix protein coating:** In order to coat the entire silicone substrate with protein, a volume
477 of 0.1% gelatin solution was added and incubated at room temperature in the biosafety
478 cabinet for 1 hour. To create protein patterns, we utilized the Alvéole's PRIMO optical
479 module and the Leonardo software, a UV-based, contactless photopatterning system
480 (Alvéole, Paris, France). The substrate surface was first coated with 500 $\mu\text{g}/\text{mL}$ PLL (Sigma
481 Aldrich, St. Louis, MO, USA) for 1 hour at room temperature. The substrate was washed
482 with PBS and 10 mM HEPES buffer adjusted to pH 8.0, and then incubated with 50 mg/mL
483 mPEG-SVA (Laysan Bio, Inc., Arab, AL, USA) at room temperature for 1 hour, and washed
484 with PBS once more. The PRIMO system was calibrated using fluorescent highlighter on
485 an identical substrate. The PBS was replaced by 14.5 mg/mL PLPP (Alvéole, Paris, France),
486 and then the desired pattern, previously created with graphic software, was illuminated with
487 UV light focused on the substrate surface for 30 seconds. Patterned surfaces were washed
488 again with PBS and then incubated with 0.1% gelatin for 1 hour at room temperature. The
489 substrate was washed and maintained hydrated in PBS at 4° C overnight.

490 **Human airway smooth muscle cell culture:** Primary human airway smooth muscle cells
491 (SMCs) were acquired through the Gift of Hope foundation (via Dr. Julian Solway, M.D.,

492 University of Chicago) and through ATCC (<https://www.atcc.org>). Both these sources are
493 public and pertinent medical information about the donor was relayed to us, but all donor
494 identifiers are removed. The donor remains anonymous and cannot be identified directly or
495 through identifiers linked to the subjects, meeting NIH guidelines. In this study, we used 4
496 healthy human donors with no history of asthma: 2 male, ages 59 and 34, and 2 female, ages
497 18 and 82. This study was carried out in accordance with the guidelines and regulations
498 approved by the Institutional Biosafety Committee at Northeastern University. Cells were
499 grown under standard culture conditions of 37° C and 5% CO₂ and utilized by P6 for all
500 experiments. Culture medium: Cells were cultured in 10% fetal bovine serum, DMEM/F12
501 (Fisher Scientific), 1x penicillin/streptomycin (Fisher Scientific), 1x MEM non-essential
502 amino acid solution (Sigma Aldrich), and 25 µg/L Amphotericin B (Sigma Aldrich). Prior
503 to any measurements, the growth medium was switched to serum-free media for at least 24
504 hours. The serum-free medium was comprised of Ham's F-12 media (Sigma Aldrich), 1x
505 penicillin/streptomycin, 50 µg/L Amphotericin B, 1x glutamine (Fisher Scientific), 1.7 mM
506 CaCl₂ 2H₂O, 1x Insulin-Transferrin-Selenium Growth Supplement (Corning Life Sciences;
507 Tewksbury, MA), and 12 mM NaOH. Both patterned and non-patterned gelatin-coated
508 substrates were UV sterilized for 1 hour, then incubated at 37° C for 1 hour before seeding
509 human airway SMCs, passage 3-6. For patterned substrates, cells were seeded in Biotech
510 dishes at 10⁴ cells per cm² and incubated for 10 minutes in 10% serum media to allow cells
511 to adhere to patterns. Next, the dishes were washed with PBS to remove excess cells, and
512 then filled with 10% serum media and incubated for 6 to 24 hours. For non-patterned
513 substrates, cells were seeded at the desired density and then incubated in 10% serum media
514 for 6 to 24 hours. After this time, media was replaced with serum-free media and incubated
515 for at least 24 hours prior to measurements. For isolated cells, the seeding density was 10²

516 cells per cm^2 . For sparse cells, the seeding density was 10^3 cells per cm^2 . For confluent cells,
517 the seeding density was 10^4 cells per cm^2 .

518 **Fluorescent imaging of Ca^{2+} :** Serum-starved airway SMCs were loaded with a fluorescent
519 cytosolic Ca^{2+} indicator to record changes in $[\text{Ca}^{2+}]$. FLIPR Ca^{2+} 6 (Molecular Devices, San
520 Jose, CA, USA) was used for all Ca^{2+} measurements except Fig. 1A, S1A, where we used
521 Fluo4-AM (Sigma Aldrich, St. Louis, MO, USA). The commonly used Fluo4-AM is prone
522 to photobleaching, and the measured Ca^{2+} traces must be bleach-corrected prior to
523 measurements of the time period. FLIPR Ca^{2+} 6, on the other hand, did not photobleach
524 even after 30 minutes of continuous imaging at 1 Hz. Fluo4-AM was prepared according to
525 the manufacturer's standards. Cells were loaded with $0.2 \mu\text{M}$ Fluo4-AM solution, diluted
526 in HBSS, and incubated at room temperature for 1 hour. Next, the cells were washed with
527 HBSS and incubated in the dark in HBSS for an additional 30 minutes. The cells were
528 washed once more before imaging. FLIPR Ca^{2+} 6 was also prepared according to
529 manufacturer's standards. Cells were incubated with a 1:1 solution of FLIPR Ca^{2+} 6 and
530 serum-free media at 37°C and 5% CO_2 for 2 hours before imaging. Both Ca^{2+} indicators
531 use acetoxymethyl esters to pass through the cell membrane, which are then hydrolyzed by
532 cytosolic esterases, trapping the fluorescent dye inside the cell. Cells were imaged with a
533 Leica DMI8 inverted microscope, a Leica DFC6000 camera (Leica, Wetzlar, Germany), and
534 a Lumencor Sola SEII LED light source (Lumencor, Beaverton, OR, USA). A FITC filter
535 cube (excitation: 480/40 nm, emission: 527/50 nm) was used to image the fluorescent dye.
536 Fluorescent intensity increases with increasing cytosolic $[\text{Ca}^{2+}]$. 16-bit images were
537 recorded at 1 Hz for 1 minute before agonist addition, and for at least 5 minutes after 10^{-5}
538 M histamine exposure. In order to analyze data, each image sequence was loaded in Fiji
539 ImageJ, and regions of interest (ROIs) were hand-selected in the cytoplasm of each cell to
540 obtain mean grayscale intensities over the area of the ROI for each frame in time. A custom

541 MATLAB (MathWorks, Natick, MA, USA) code was written to process the data and
542 measure mean Ca^{2+} oscillation periods. This code measured mean Ca^{2+} oscillation periods
543 by finding peaks in the time series data above a certain prominence and taking the mean of
544 the time between all sequential peaks (Fig. S4).

545 **Cell traction force measurements:** The base NuSil substrates were coated with a layer of
546 fluorescent beads as fiducial markers for traction force microscopy. A 5% solution of 0.2
547 μm diameter red fluorescent carboxylate-modified microspheres (FluoSpheres, Invitrogen,
548 Carlsbad, CA, USA) in PBS was vortexed for 10 seconds. 2 mL of solution was added to
549 each substrate in a Bioptech dish and left at room temperature for 1 hour to allow the beads
550 to adhere. The bead solution was poured off, substrates were washed 3x with PBS, and then
551 PBS was poured off. NuSil solution was prepared as described before, with the appropriate
552 amount of crosslinker to match the stiffness of the base substrate. NuSil was spin-coated
553 onto the newly bead-coated base substrate at 2500 RPM to create a 1 μm -thick layer and
554 seal the beads. These substrates rested on a flat surface for 1 hour before curing overnight
555 at 60° C. Substrates were protein-coated and seeded with cells as before. After a 24 hour
556 incubation in serum-free media, the SMC tractions were recorded by imaging the
557 fluorescent beads with a 20x/0.55 dry objective and the Leica DMi8 microscope in an
558 environmental chamber maintained humidified at 37° C. Images were taken at baseline,
559 after a 15 minute incubation with 10^{-5} M histamine, and after cells were removed using RLT
560 Lysis Buffer (Qiagen, Hilden, Germany). Using these images, cellular forces were
561 calculated with a custom MATLAB (MathWorks, Natick, MA) software program using
562 Fourier Traction Force Microscopy (23).

563 **Fluorescent labeling of connexin-43/actin/nuclei:** Cells were fluorescently labeled for
564 connexin-43 (Cx43) and filamentous actin (F-actin). Cells were fixed in 4%
565 paraformaldehyde in PBS at room temperature for 10 minutes. Then, cells were

566 permeabilized with 100% ethanol for 10 minutes at 20° C. Following permeabilization, cells
567 were blocked using 1x PBS containing 0.1% Tween-20, 1% bovine serum albumin (BSA),
568 and 22.52 mg/ml glycine for 30 minutes at room temperature. Next, cells were stained for
569 Cx43 (ab11370; Abcam, Cambridge, UK) at a dilution of 1:200 in 1x PBS containing 1%
570 BSA for 1 hour at 37° C. Secondary antibody labeling and phalloidin staining were done
571 simultaneously at dilutions of 1:200 and 1:40, respectively, using Alexa Fluor 594
572 (ab150080; Abcam, Cambridge, UK) and Alexa Fluor 488 Phalloidin (A12379; Invitrogen,
573 Carlsbad, CA, USA). Lastly, cells were labeled with NucBlue (Fisher Scientific, Waltham,
574 MA, USA) to label cell nuclei. Images were acquired using a 63x/1.4 oil-immersion
575 objective (Leica, Wetzlar, Germany).

576 **The gap-FRAP assay:** Gap Fluorescent Recovery After Photobleaching (FRAP) is an
577 experimental technique that has been established as an effective method of observing gap-
578 junctional communication of small fluorescent molecules between adjacent cells (22).
579 SMCs were cultured on NuSil gels of E=0.3 kPa and E=13 kPa until confluence was
580 achieved, and then serum-starved for at least 24 hours prior to the experiment. Cells were
581 loaded with 1 μ M calcein-AM solution diluted in warmed 1x PBS solution and incubated at
582 37° C and 5% CO₂ for 15 minutes. Calcein-AM is a cell-permeable dye that is hydrolyzed
583 into fluorescent calcein by cytoplasmic esterases upon entry through cell membrane and has
584 been shown to permeate through gap junctions due to its low molecular size (622 Da) (20).
585 After dye incubation, samples were washed with warm 1x PBS solution and returned to
586 serum-free medium for experiments. FRAP was performed using a ZEISS confocal laser
587 scanning microscope system equipped with a 20x/0.8 objective and a 488 nm Argon laser.
588 Fluorescence data was captured using ZEN 2012 SP5 imaging software (ZEISS,
589 Oberkochen, Germany). Samples were placed in an incubation chamber maintaining 37° C
590 during experiments to preserve cell viability during imaging. Prior to photobleaching, a

591 manual ROI was drawn around the border of a target cell visibly connected to adjacent cells.
592 Whole cells were selected for photobleaching to ensure that fluorescence recovery could
593 only be attributed to the diffusion of calcein from adjacent cells. Laser power was adjusted
594 to 1% and images were acquired every 5 seconds for 50 seconds using scan speed 12 (pixel
595 dwell = 0.42 μ s) to provide baseline fluorescence measurements. After baseline scans were
596 acquired, the laser power was adjusted to 100% and cells were bleached to at least 20% of
597 their initial fluorescence using scan speed 2 (pixel dwell = 40 μ s). Following the bleaching
598 step, fluorescence recovery images were collected every 5 seconds for approximately 5 mins
599 at 1% laser power. The images from gap-FRAP experiments were analyzed using Fiji
600 ImageJ software. First, baseline fluorescence intensities from the target cell were averaged
601 over the first 10 frames captured to establish a reference value for fluorescence recovery.
602 Next, fluorescence intensities were measured during the recovery period and divided by the
603 average baseline intensity to normalize the data. To account for any photobleaching
604 occurring during the recovery period, fluorescence intensities were collected from a region
605 at the edge of the field of view. These values were used to adjust the intensity of the target
606 cell over time to account for fluorescence degradation due to repeated scanning of the
607 microscopic field, since the edge region was not affected directly during the bleaching step.
608 The normalized fluorescence values of the bleached cell during recovery were plotted as a
609 function of time. To compare fluorescence recovery across multiple sample groups, the
610 mobile fraction of fluorescent molecules was calculated. Mobile fraction (Γ) measures the
611 fraction of fluorescent molecules that contribute to recovery of fluorescence in bleached
612 cells and is calculated with Eq. 1, where F_0 , F_b , F_r indicate the fluorescence intensity in the
613 cell being bleached at baseline, after bleaching and after recovery, respectively (Fig. 3E).

614 **Gap junction blocker experiments:** Gap junctions between confluent SMCs were blocked
615 with 18 β -glycyrrhetic acid (β GA) (Sigma Aldrich, St. Louis, MO, USA). Confluent SMCs

616 on soft ($E=0.3$ kPa) and stiff ($E=13$ kPa) substrates were incubated with $30\ \mu\text{M}$ βGA at 37°
617 C and 5% CO_2 for 30 minutes. Although βGA is a common gap junction blocker, it has been
618 reported to affect cell viability at higher concentrations (21). We used gap-FRAP to find the
619 lowest possible dose of βGA that still blocked gap junctions between confluent cells, which
620 we used here in our experiments. This treatment was used in conjunction with the Ca^{2+}
621 imaging protocol to investigate the role of gap junctional diffusion in agonist-induced Ca^{2+}
622 oscillations.

623 **Force-inhibitor experiments:** Smooth muscle cell contractile force was reduced with two
624 separate inhibitors of force: Y-27362 (Sigma Aldrich, St. Louis, MO, USA), a specific
625 inhibitor of ROCK, and ML-7 (Sigma Aldrich, St. Louis, MO, USA), a selective inhibitor
626 of MLCK. Confluent SMCs on soft (0.3 kPa) and stiff (13 kPa) substrates were pre-
627 incubated with $10\ \mu\text{M}$ or $100\ \mu\text{M}$ of Y-27632 for 1 hour, or $0.01\ \mu\text{M}$ or $0.05\ \mu\text{M}$ of ML-7
628 for 5 minutes, in tandem with the 2 hour FLIPR Ca^{2+} 6 indicator incubation, before
629 experiments. Ca^{2+} imaging and 10^{-5} M histamine addition was added as previously
630 described.

631 **Data presentation and statistical testing:** Throughout the manuscript, we represent data
632 as the average of N independent trials. The error bars indicate standard deviation over
633 independent trials. The individual data points are shown in all bar plots. Each data point is
634 in turn the average of measurements made over multiple cells within one trial. Sigmatat
635 (Systat Software, San Jose, CA) was used to perform statistical tests. Two-Way ANOVAs
636 followed by posthoc pairwise comparisons were used to test for significant differences in
637 datasets which were influenced by two independent factors. One-Way ANOVAs followed
638 by posthoc pairwise comparisons were used to test for significant differences in datasets of
639 three or more groups which were influenced by one independent factor. Pairwise
640 comparisons used the t-test when the data was normally distributed. Otherwise, the Mann-

641 Whitney Rank Sum Test was used to compare the median values. The specific tests used,
642 the number of samples and the P-value are described along with the corresponding results.
643 A P-value of 0.05 was used as the threshold for a statistically significant difference between
644 data sets.

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768

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773 junction staining. SS, RR, JB, EJC & HP contributed to writing the manuscript and analysis of the

774 data. HP is the corresponding author who conceived and directed this project.

775

776 **Competing interests:** The authors declare that they have no competing interests.

777

778 **Data and materials availability:** Data supporting the findings of this study are available within

779 the manuscript. All other relevant data are available from authors upon reasonable request.

780

781 **Figures and Tables**

782 **Fig. 1. Effect of matrix stiffness on agonist-induced Ca^{2+} oscillations in SMCs.** (A) To

783 study the role of altered matrix stiffness on the time period of agonist-induced

784 Ca^{2+} oscillations, we micropatterned a 2D approximation of the in-situ organization

785 of SMCs. Cells were also cultured on non-patterned surfaces in three different

786 densities: (B) isolated, (C) sparse, and (D) confluent. The colors in Figs 1A-D

787 correspond to cytosolic $[\text{Ca}^{2+}]$ concentrations as indicated by the color bar in Fig.

788 1A. Scale bars=200 μm . (E) Increasing matrix stiffness from 0.3 kPa to 13 kPa

789 caused a significant decrease in Ca^{2+} oscillation period in SMCs rings (t-test,

790 $P < 0.001$, $N = 4$ each). (F, G) The periods of SMC Ca^{2+} oscillations were not affected

791 by matrix stiffness in both isolated (soft N=14, stiff N=12) and sparse (N=4 each)
792 conditions (t-test, P=0.93, P=0.481, respectively). **(H)** Confluent cells behaved like
793 those patterned in a ring, with cells plated on stiff matrix exhibiting significantly
794 faster Ca²⁺ oscillations in response to 10⁻⁵ M histamine compared to those on a soft
795 matrix (soft N=5, stiff N=7, Mann-Whitney Rank Sum Test, P=0.003). These results
796 demonstrate that matrix stiffness can modulate the agonist-induced Ca²⁺ response of
797 confluent SMCs, but not that of isolated cells.

798 **Fig. 2. Effect of matrix stiffness and confluence on the correlated nature of Ca²⁺**

799 **oscillations.** **(A)** The cross-correlation coefficients ($\rho_{i,j}$) range in values from 1
800 (positively correlated, pink) to 0 (uncorrelated, white) to -1 (negatively correlated,
801 green), as shown in a representative 24x24 cell matrix. **(B)** Examples of Ca²⁺
802 oscillations measured in two cells with high pairwise-correlation ($\rho=0.78$), and low
803 pairwise correlation ($\rho=0.02$). **(C)** In isolated cells, the Ca²⁺ oscillations were
804 generally uncorrelated with a probability density function centered around zero.
805 ECM stiffness did not affect pairwise correlations in isolated SMCs (N=5 soft, N=4
806 stiff, P=0.733, Two-Way ANOVA). **(D)** In confluent cells, there is a statistically
807 significant increase in ρ , indicating that the Ca²⁺ oscillations were more
808 synchronized on stiffer ECM (N=4 soft, N=5 stiff, Two-Way ANOVA, P=0.007).
809 **(E, F)** To evaluate the time it takes after agonist addition for Ca²⁺ oscillations to
810 become synchronized, we calculated $\rho_{i,j}(\tau)$ within a 120s moving window. **(E)** $\rho_{i,j}(\tau)$
811 in isolated cells does not change over time (N=5 soft, N=4 stiff, P=0.274) or with
812 stiffness (P=0.303) (Two-Way ANOVA). However, **(F)** in confluent cells on stiff
813 matrix, $\rho_{i,j}(\tau)$ shows a significant increase after 30 seconds (N=4 soft, N=5 stiff)
814 (Two-Way ANOVA, P<0.001). Error bars indicate standard deviation.

815 **Fig. 3. Effect of matrix stiffness on intercellular communication via gap junctions.**

816 Confluent SMCs on soft (A) and stiff (B) matrix were stained for the gap junction
817 protein Cx43, actin, and the nucleus. (C) Gap junctions were counted for each SMC
818 in individual frames on soft (N=5) and stiff (N=5) matrix, represented by the mean
819 and standard deviation of these measurements, and we found no statistical difference
820 due to matrix stiffening (t-test, P=0.977). (D) Diffusion through gap junctions was
821 quantified by bleaching a small fluorescent molecule within a cell and measuring
822 signal recovery due to intercellular diffusion, a technique called gap-FRAP. (E) A
823 representative recovery curve for one cell shows a recovery of ~40% its initial
824 fluorescence. These values were used to calculate mobile fraction, a measure of
825 diffusion efficiency through gap junctions. (F) Matrix stiffness had no effect on
826 mobile fraction (t-test, soft N=16, stiff N=18, P=0.532). The gap junction blocker
827 β GA significantly reduced the recovery (D) and mobile fraction (F) (N=18, Mann-
828 Whitney Rank Sum Test). (G) Despite blocking gap junctions with β GA, there was
829 no effect on Ca^{2+} oscillation periods in confluent cells on either soft (N=6) or stiff
830 (N=4) matrix (Two-Way ANOVA, P=0.784 treatment within stiffness). Scale bars=
831 30 μm .

832 **Fig. 4. Role of mechanical force in Ca^{2+} wave propagation through multicellular**

833 **ensembles of SMCs.** (A) SMCs in confluent layers form organized clusters of cells,
834 with certain cells aligned end-to-end along their contractile axis (parallel), and others
835 branching off at an angle (perpendicular). Scale bar=250 μm . Insets (a), (b) show
836 cells 2 and 4 parallel to the contractile axis of cell 1, whereas cells 3 and 5 are
837 perpendicular. Inset scale bars=50 μm . The conditional probability for a localized
838 increase in Ca^{2+} in cell 1 to be followed by an increase in a parallel or perpendicular
839 neighbor is plotted in (B) with mean and standard deviation. Ca^{2+} waves were

840 statistically more likely to propagate along the contractile axis (t-test, N=10). **(C)**
841 Histamine caused a significantly greater increase in traction stress in confluent cells
842 on stiff matrix (N=21) rather than on soft (N=12, t-test). **(D)** The faster agonist-
843 induced Ca^{2+} oscillations on stiff ECM were systematically abrogated in a dose-
844 dependent manner by preincubating the SMCs with the ROCK inhibitor Y-27632
845 for 1 hour prior to histamine exposure (N=3, N=4, $P < 0.001$, One-Way ANOVA).
846 Identical results can be obtained with MLCK inhibition by preincubating with
847 increasing doses of ML-7 for 5 minutes (N=3 each, $P < 0.001$, One-Way ANOVA).

848 **Fig. 5. Effect of confining SMCs in a line using micropatterning.** **(A)** When SMCs were
849 patterned in lines, **(B)** the probability of finding SMCs with high time periods
850 decreased and the variance of the time periods from aligned cells was significantly
851 smaller than confluent cells (F-test, $P < 0.001$). Matrix stiffness still affected the
852 oscillation period (soft N=8, stiff N=6, Mann-Whitney Rank Sum Test). To probe
853 the limits of the SMC cluster's collective matrix sensing abilities, we simulated
854 localized ECM stiffening by **(C)** patterning SMCs in lines spanning a dual-stiffness
855 PDMS substrate (E=13 kPa highlighted in green, left, and E=0.3 kPa, right). **(D)** The
856 Ca^{2+} oscillation time period of cells along lines is plotted as a function of the distance
857 from the stiff matrix. Binning cells in 400 μm intervals, only those over 800 μm
858 from stiff matrix had different Ca^{2+} oscillation time periods from cells directly in
859 contact (t-test, N=4). Scale bars=250 μm .

1 **Supplementary Materials**

2 **Materials and Methods**

3 **Fabrication of NuSil substrates at additional stiffnesses:** Equal parts of NuSil gel-8100 parts A
4 and B (NuSil, Carpinteria, CA, USA) were mixed with various amounts of the crosslinking
5 compound of Sylgard 184 (Dow Corning, Midland, MI, USA) to adjust substrate stiffness.
6 Crosslinker volumes of 0.07% and 0.15% of the combined volumes of parts A and B were added
7 to the 1:1 A:B mixture to create substrates with Young's modulus $E = 0.6$ kPa and 3 kPa,
8 respectively. The spinning and curing protocol to finish the substrates can be found in the main
9 manuscript.

10 **Histamine dose-response experiment:** Confluent SMCs cultured on soft (0.3 kPa) and stiff (13
11 kPa) substrates were incubated with FLIPR Ca^{2+} 6 indicator and imaged following the protocol in
12 the main manuscript. For each sample, histamine concentration was gradually increased and
13 followed by 5 minutes of imaging. Stiff samples were exposed to concentrations increasing from
14 10^{-6} M, to 10^{-5} M, to 10^{-4} M, and soft samples were exposed to concentrations of 10^{-5} M, 10^{-4} M,
15 and 10^{-3} M histamine.

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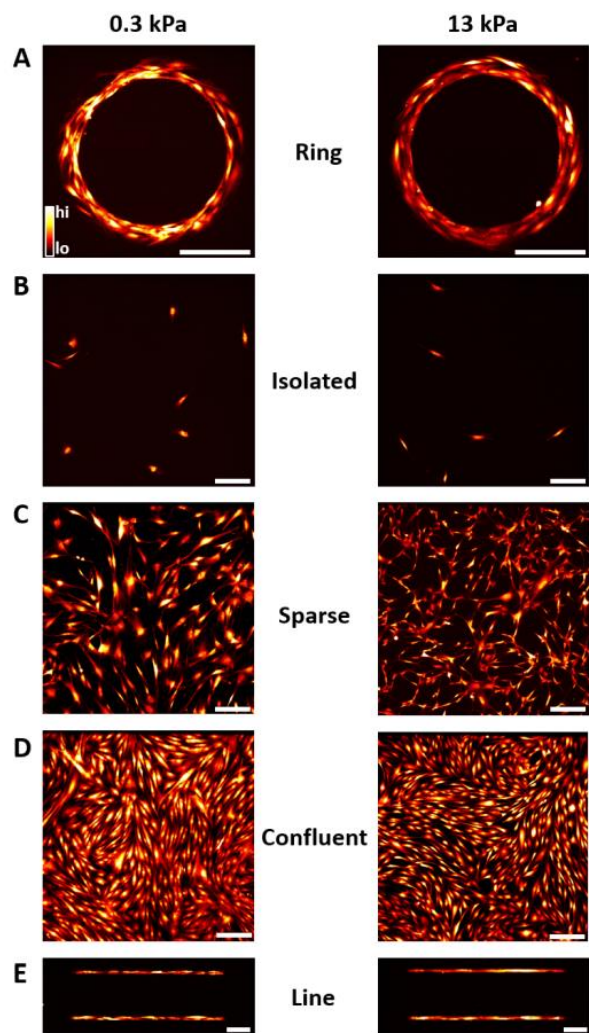
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22 **Figures**

23 **Fig. S1.**



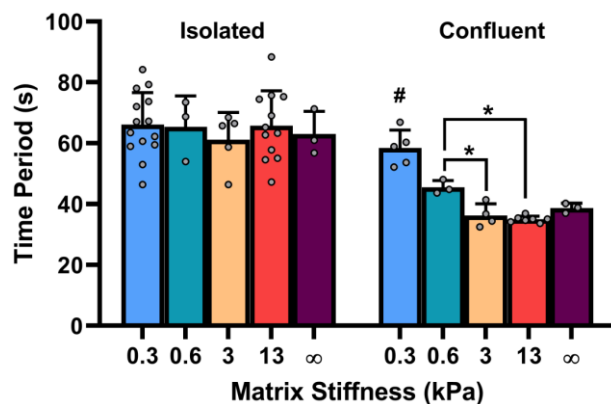
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25 **Fig. S1. Morphology of SMCs on soft and stiff substrates.** Fluorescent images of cytosolic Ca²⁺
26 in SMCs show similar appearance and density of cells in the different culture conditions used in
27 this paper: (A) rings, (B) isolated, (C) sparse, (D) confluent, and (E) lines. The cytosolic [Ca²⁺] is
28 pseudo-colored following the color bar, as in Fig. 1. Scale bar = 200 μ m.

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30

31 **Fig. S2.**



32

33 **Fig. S2. Effect of incremental matrix stiffening on agonist-induced Ca^{2+} oscillations in SMCs.**

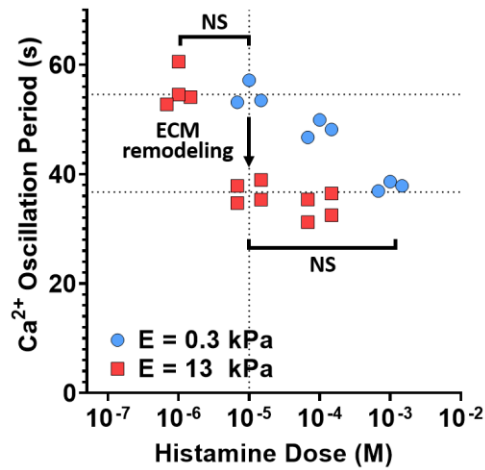
34 PDMS substrate stiffness was tuned between our limits for soft (0.3 kPa) and stiff (13 kPa) matrix
35 to create two intermediate stiffness matrices at 0.6 kPa and 3 kPa. Glass coverslips were used to
36 represent matrix of infinite stiffness. In confluent cells, ECM stiffness had a significant impact on
37 agonist-induced Ca^{2+} oscillation time periods (One-Way ANOVA, $P < 0.001$). Pairwise
38 comparisons with a Holm-Sidak test are indicated by symbols. The time period at 0.3 kPa was
39 significantly different from all other substrate stiffnesses (indicated by #, $P < 0.001$). The time
40 period at 0.6 kPa was also significantly different from that at 3 kPa and 13 kPa ($P = 0.003$, $P < 0.001$,
41 respectively). There was no statistical difference between time periods at 3 kPa, 13 kPa and glass
42 (3 kPa vs. 13 kPa $P = 0.566$, 3 kPa vs. glass $P = 0.379$, 13 kPa vs. glass $P = 0.017$, respectively). Matrix
43 stiffness had no effect on the agonist-induced Ca^{2+} response in isolated cells (One-Way ANOVA,
44 $P = 0.907$).

45

46

47

48 **Fig. S3.**



49

50 **Fig. S3. Effect of histamine dose on Ca²⁺ oscillation time period for cells on soft and stiff**

51 **ECM.** Confluent SMCs on soft and stiff matrix were exposed to multiple doses of histamine.

52 Increasing concentration of histamine from 10⁻⁵ M to 10⁻³ M systematically decreased the mean

53 Ca²⁺ oscillation time periods of cells on soft matrix (One-Way ANOVA, P<0.005). On stiff matrix,

54 increasing histamine dose from 10⁻⁶ M to 10⁻⁵ M caused a dramatic decrease in time period, but

55 increasing the dose further had no effect (One-Way ANOVA, P<0.001, P=0.174, respectively).

56 For SMCs on soft (E=0.3 kPa) substrate, the histamine dose needs to be increased from 10⁻⁵ M to

57 10⁻³ M (2 log scales higher) to match the Ca²⁺ response of SMCs on the stiff substrate.

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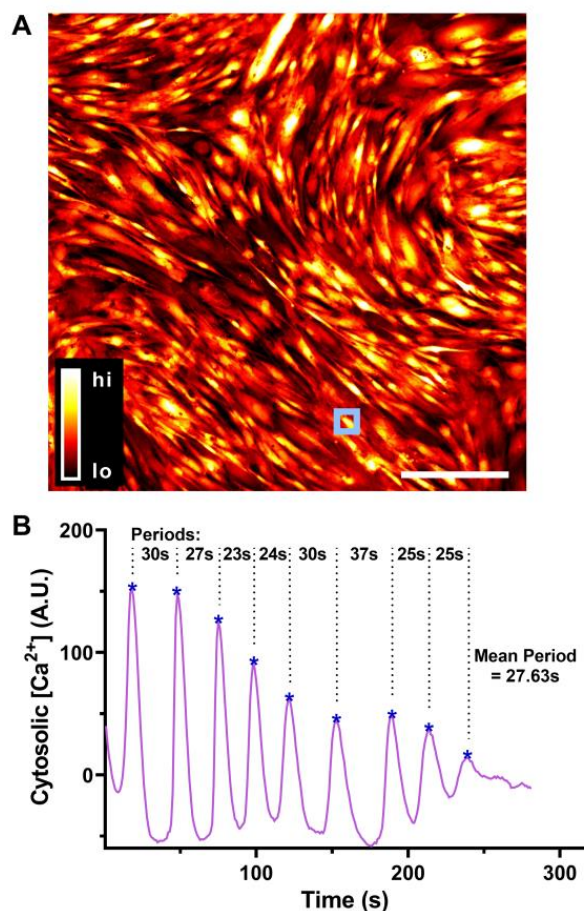
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64 **Fig. S4.**



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66 **Fig. S4. Method of calculating Ca²⁺ oscillation periods.** (A) Images for each experiment were
67 loaded into Image J, and 5x5 pixel regions of interest (ROIs) were hand-selected in the cytosol of
68 the smooth muscle cell. The mean gray intensity of each ROI for each frame was imported into
69 MATLAB and plotted to show changes over time for each cell. The Ca²⁺ indicator used in our
70 experiments indicates cytosolic [Ca²⁺], therefore changes in grayscale intensity correspond with
71 changes in cytosolic [Ca²⁺]. (B) The portion of data after the initial spike in Ca²⁺ due to histamine
72 addition was selected for further processing. First, the code subtracted the mean [Ca²⁺] value of
73 each time series from the time series. Next, local peaks above a threshold value were identified
74 (indicated by *), and the time between sequential peaks were recorded as oscillation periods. Time

75 periods are measured for a minimum of 40 cells/trial in confluent cells and for every SMC in
76 isolated cells. The average time period of all measured cells is reported for each independent trial.

77 **Movies S1 to S4 Captions**

78 **Movie S1. Cytosolic Ca²⁺ oscillations in confluent smooth muscle cells on stiff**
79 **matrix.** Confluent SMCs on stiff matrix (E=13 kPa) loaded with a fluorescent Ca²⁺ indicator were
80 imaged before and during exposure to a 10⁻⁵ M dose of the contractile agonist histamine, added at
81 the time indicated. This movie shows the first 2.5 minutes of the experiment. The movie is
82 compressed to 8 bit and 0.25 frames/second to match journal data guidelines. The full 5 minutes
83 of Ca²⁺ oscillations recorded at 1 frame/second, 16-bit, 2048 x 2048 images used to make
84 measurements is available from the authors on reasonable request. Scale bar = 200 μm.

85 **Movie S2. Cytosolic Ca²⁺ oscillations in confluent smooth muscle cells on soft**
86 **matrix.** Confluent SMCs on soft matrix (E=0.3 kPa) loaded with a fluorescent Ca²⁺ indicator were
87 imaged before and during exposure to a 10⁻⁵ M dose of the contractile agonist histamine, added at
88 the time indicated. This movie shows the first 2.5 minutes of the experiment. The movie is
89 compressed to 8 bit and 0.25 frames/second to match journal data guidelines. The full 5 minutes
90 of Ca²⁺ oscillations recorded at 1 frame/second, 16-bit, 2048 x 2048 images used to make
91 measurements is available from the authors on reasonable request. Scale bar = 200 μm.

92 **Movie S3. Cytosolic Ca²⁺ oscillations in isolated smooth muscle cells on stiff matrix.** Isolated
93 SMCs on stiff matrix (E=13 kPa) loaded with a fluorescent Ca²⁺ indicator were imaged before and
94 during exposure to a 10⁻⁵ M dose of the contractile agonist histamine, added at the time indicated.
95 This movie shows the first 2.5 minutes of the experiment. The movie is compressed to 8 bit and
96 0.25 frames/second to match journal data guidelines. The full 5 minutes of Ca²⁺ oscillations
97 recorded at 1 frame/second, 16-bit, 2048 x 2048 images used to make measurements is available

98 from the authors on reasonable request. Scale bar = 200 μm .

99 **Movie S4. Cytosolic Ca^{2+} oscillations in isolated smooth muscle cells on soft matrix.** Isolated

100 SMCs on soft matrix ($E=0.3$ kPa) loaded with a fluorescent Ca^{2+} indicator were imaged before and

101 during exposure to a 10^{-5} M dose of the contractile agonist histamine, added at the time indicated.

102 This movie shows the first 2.5 minutes of the experiment. The movie is compressed to 8 bit and

103 0.25 frames/second to match journal data guidelines. The full 5 minutes of Ca^{2+} oscillations

104 recorded at 1 frame/second, 16-bit, 2048 x 2048 images used to make measurements is available

105 from the authors on reasonable request. Scale bar = 200 μm .