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HIGHLIGHTS
Visco-node-pore sensing (visco-NPS) enables viscoelastic measurements of cells

Visco-NPS can sense changes in cytoskeleton and nucleus structure

SLUG induction in MCF-10A causes partial EMT and increased viscous behavior

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Visco-Node-Pore Sensing: A Microfluidic Rheology Platform to Characterize Viscoelastic Properties of Epithelial Cells

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SUMMARY

Viscoelastic properties of cells provide valuable information regarding biological or clinically relevant cellular characteristics. Here, we introduce a new, electronic-based, microfluidic platform—visco-node-pore sensing (visco-NPS)—which quantifies cellular viscoelastic properties under periodic deformation. We measure the storage ($G'$) and loss ($G''$) moduli (i.e., elasticity and viscosity, respectively) of cells. By applying a wide range of deformation frequencies, our platform quantifies the frequency dependence of viscoelastic properties. $G'$ and $G''$ measurements show that the viscoelastic properties of malignant breast epithelial cells (MCF-7) are distinctly different from those of non-malignant breast epithelial cells (MCF-10A). With its sensitivity, visco-NPS can dissect the individual contributions of different cytoskeletal components to whole-cell mechanical properties. Moreover, visco-NPS can quantify the mechanical transitions of cells as they traverse the cell cycle or are initiated into an epithelial-mesenchymal transition. Visco-NPS identifies viscoelastic characteristics of cell populations, providing a biophysical understanding of cellular behavior and a potential for clinical applications.

INTRODUCTION

Cellular mechanical properties are recognized as important indicators of biologically relevant functions of cells and tissues (Stossel et al., 2001; Janmey and McCulloch, 2007). In particular, viscoelastic properties provide valuable information regarding how cells facilitate their movement and minimize damage from external stimuli by storing and dissipating energy (Lange and Fabry, 2013; Suresh, 2007). Governing the viscoelastic responses of cells are intracellular components such as actin filaments, microtubules, and the nucleus—all of which play important roles in the dynamics of cell proliferation (Provenzano and Keely, 2011), migration (Yamaguchi and Condeelis, 2007), differentiation (Mundel et al., 1997; Woods et al., 2007), and apoptosis (Gourlay and Ayscough, 2005). Mechanical characterization of cells also provides clinical merits in terms of detecting cellular changes that are due to, for instance, cancer malignancy (Kumar and Weaver, 2009; Kim et al., 2018; Prabhune et al., 2012), cell cycle (Otto et al., 2015; Benham-Pyle et al., 2015), stem cell differentiation (Altman et al., 2002; Luu et al., 2009; Saha et al., 2006), or epithelial-mesenchymal transition (EMT) (Cabrera-Benitez et al., 2012; Gjorevski et al., 2012). Thus, analyzing viscoelastic properties of cells provides not only a fundamental understanding of biophysical characteristics but also a tremendous potential for clinical applications.

To quantify the viscoelastic properties of cells, different rheological methods have been developed. To name just a few examples, micropipette aspiration (Sato et al., 1987, 1990), microplate rheometer (Desprat et al., 2006; Thourme et al., 1999), optical stretcher (Guck et al., 2001; Yang et al., 2016), and atomic force microscopy (AFM) (Haase and Pelling, 2015; Alcaraz et al., 2003) have all been used to measure the viscoelastic properties of cells with respect to different loading conditions, such as force magnitude and deformation frequency. These methods, however, have low throughput (only a few cells/hour [Guo et al., 2012; Wang et al., 2015; Cartagena-Rivera et al., 2015]) and consequently are not capable of screening large populations of cells—a critical need for clinical applications. To address this key drawback, a number of higher-throughput microfluidic platforms have been developed, including hydrodynamic stretching cytometry (Gossett et al., 2012; Masaeli et al., 2016), real-time deformability cytometry (RT-DC) (Otto et al., 2015), and mechano node-pore-sensing (mechano-NPS) (Kim et al., 2018). These particular methods, however, focus on mechanically phenotyping cells for screening purposes and do not provide the means to investigate basic cellular mechanics. Recent studies have reported microfluidic platforms that take a rheological approach and drive cells through a confining
channel to measure their cellular viscoelastic properties (Preira et al., 2013; Hu and Lam, 2017). However, these platforms rely on imaging, and consequently, they have a very limited temporal and spatial window for measurement, which in turn, severely restricts sample size and experimental conditions. Key information such as what is the frequency dependence of viscoelastic properties and what is the origin of whole-cell properties cannot be easily obtained. Thus despite the many different platforms for measuring cell mechanical properties that exist today, none thus far are able to provide a comprehensive view of cellular mechanics.

Here, we introduce a novel, all-electronic-based, microfluidic platform to measure the cellular viscoelastic properties. Our platform, visco-node-pore sensing (visco-NPS), employs a rheological approach by utilizing a sinusoidal-shaped contraction channel through which cells undergo an oscillating deformation as a result of the channel’s periodically changing width. By integrating this unique channel with a node-pore sensor (Balakrishnan et al., 2013; Kim et al., 2018), we are able to measure the storage (elasticity) and loss (viscosity) moduli of cells. As we demonstrate, visco-NPS can successfully quantify the differences in the viscoelastic properties between malignant and non-malignant epithelial cells. Moreover, it can ascertain the individual contributions of cytoskeletal components, i.e., actin filaments and microtubules, to the mechanical behavior of cells. Finally, visco-NPS can determine the changes in cell mechanical properties that result from the dynamic transitions that the cytoskeleton and nucleus undergo during the cell cycle or from an initiated EMT. Overall, visco-NPS represents an efficient, simple, and direct means to quantify the mechanical properties of single cells.

RESULTS
Experimental Design
Visco-NPS consists of a microfluidic channel embedded in a polydimethylsiloxane (PDMS) mold that is bonded to a glass substrate with pre-defined platinum (Pt) electrodes and gold (Au) contact pads (Figure 1A). The embedded channel has multiple components: in-plane filters and a sinusoidal contraction channel flanked by a set of pores and nodes (Figure 1B). The in-plane filters have 25-μm-wide gaps and subsequently remove cellular clusters that would otherwise clog the contraction channel. The first node-pore region measures, with sufficient signal-to-noise ratio, the free cell size based on the Coulter principle (Balakrishnan et al., 2013, 2015; Coulter, 1953; Kim et al., 2018) (Table S2). With its periodically changing width ($w_c = w_o + a \cos(\omega t)$, where $w_o$, $a$, and $\omega$ are the initial width, strain amplitude, and deformation frequency, respectively) (Figure S1), the contraction channel measures the viscoelastic properties of transiting cells as they undergo a sinusoidal deformation (Figures 1C and Video S1). For the experiments described here, the contraction channel dimensions were chosen such that MCF-7 and MCF-10A cells were subject to the same strain, $\varepsilon = 0.4 + 0.1 \cos(\omega t)$ (Figure S2 and Table S1). Different periodic contraction channel lengths ($L_p$) were employed to control the deformation frequency applied to each cell type. Pressure-driven flow (13.7–27.6 kPa), via a commercial microfluidic flow controller (Elveflow, France), is used to drive 10–50 cells/min across the channel without coincidence events. By varying the flow rate, a wide range of $\omega$ is achieved even with the same $L_p$. A four-terminal measurement is employed to measure the current across the channel (Balakrishnan et al., 2013, 2015; Saleh and Sohn, 2001; Kim et al., 2018) (Figure S3). After low-pass filtering to remove noise from the measured signal, a derivative cutoff detection method is employed to identify the start and end time points of each sub-pulse ($D_{\text{I}_{np}}$ and $D_{\text{I}_{c}}$ in Figure 1D; Kim et al., 2018).

To validate statistical significance, an unpaired t test, a two-way ANOVA test, or a Pearson correlation coefficient test was performed. Power analysis was employed to ensure that the measured sample size provides adequate power (>0.80) to detect statistical differences compared with their control groups with 95% confidence intervals. All experimental groups with $p < 0.05$ showed sufficient power value with the analyzed sample size ($n = 100$) (Table S3). All data presented in this study were measured using two different microfluidic devices per experimental case to ensure repeatability of results. For example, the storage modulus ($G'$) of MCF-10A cells was measured with two different device replicas. A comparison of the results showed no statistical difference (Figure S4).

Theoretical Model for Visco-NPS
Strain values correspond to the amount of cellular deformation in the contraction channel. By definition, strain ($\varepsilon$) is determined by the free-cell diameter ($D_{\text{cell}}$) and width of the contraction channel ($w_c$). As $w_c$ is periodic along the channel’s longitudinal axis, so, too, is $\varepsilon$. 
3\text{cell} = C_0 \cdot w_0 + \frac{3\pi}{2} \quad \text{(Equation 1a)}

\text{where } w_0, 3\pi, \text{ and } u \text{ denote the strain amplitude, compressive pre-strain, and deformation frequency, respectively. For the experiments and results we present here, we designed and utilized a contraction channel whose width } w_c = 0.4 + 0.1 \cos(ut) \text{ (Table S1).}

\text{To determine the stress value } \sigma \text{ of a cell within our contraction channel, we first analyzed the external forces surrounding a deformed cell. As it transits the contraction channel and deforms, a cell is subject to driving } F_{\text{drive}} \text{ and drag forces (Figure S5). } F_{\text{drive}} \text{ is a result of the pressure distribution around the deformed cell (Equation 2). To calculate this force, we modeled (via COMSOL Multiphysics) the fluid dynamics within our device and used the specific flow rate and channel geometry we employed in our experiments as our parameters. With } F_{\text{drive}} \text{, we then calculated the average pressure difference } \Delta P_{\text{avg}} \text{ across the deformed cell in the channel’s longitudinal axis direction (Equation 2). Frictional forces } F_{\text{fric}} \text{ between the cell surface and channel wall (Figure S5), which are in the opposite direction of the flow, are defined by the frictional coefficient } \mu \text{ and normal force } F_n \text{ (Equation 3).}
As a cell transits the contraction channel with constant velocity (Figure S6), there is a zero net force around the deformed cell (Equation 4). Substituting $F_{\text{drive}}$ and $F_{\text{fric}}$ into Equation 4, we determine $\sigma$ from our fluidic conditions and the compressive deformation of cells (Equation 5).

$$F_{\text{fric}} = \frac{2 \Delta P_{\text{avg}} w_c}{\mu_i \pi D_d}$$  \hspace{1cm} (Equation 3b)

$$F_{\text{drive}} = \int \Delta P dA$$  \hspace{1cm} (Equation 2a)

$$F_{\text{fric}} = \mu_i F_n$$  \hspace{1cm} (Equation 3a)

$$= \frac{D_d^2 \pi \sigma}{4}$$  \hspace{1cm} (Equation 2b)

$$F_{\text{net}} = F_{\text{drive}} - 2 F_{\text{fric}} = 0$$  \hspace{1cm} (Equation 4)

$$\sigma = \sigma_p + G' \omega \cos(\omega t) + G'' \omega \sin(\omega t)$$  \hspace{1cm} (Equation 6)

$$G'(\omega) = \frac{G_0}{\sqrt{1 + (\omega \tau)^2}}$$  \hspace{1cm} (Equation 7a)

$$G''(\omega) = \frac{G_0}{\sqrt{1 + (\omega \tau)^2}} \omega$$  \hspace{1cm} (Equation 7b)

$$G_0 = \frac{G_0}{\sqrt{1 + (\omega \tau)^2}}$$  \hspace{1cm} (Equation 7c)

where $G_0$, $\alpha$, and $\mu$ denote shear modulus at zero frequency, power-law exponent, and viscosity, respectively. $G_0$ and $s_{0B}$ are scaling factors for stiffness and frequency, and $s_{0B} = 1$ Hz in our experiments. All power-law components are numerically calculated by least squares fitting.

**Measuring the Viscoelastic Properties of Epithelial Cells**

We investigated and compared the viscoelastic properties of malignant MCF-7 and non-malignant MCF-10A breast epithelial cells. Both cell types were subject to periodic deformation as they traveled through our platform’s sinusoidal contraction channel. By using contraction channels of different $L_d$ in combination with a range of controlled flow rates, we were able to achieve a wide range of effective deformation frequencies ($\omega$) for both MCF-7 and MCF-10A cell populations we measured. As shown in Figures 2A and 2B, $\omega$ ranged from 20 to 300 Hz for MCF-7 cells and from 20 to 200 Hz for MCF-10A cells. Each $\omega$ range was sufficiently large to quantify the dynamic response of the respective cell types. To determine how cell-channel interactions vary with respect to the cell types utilized in our studies and with respect to frequency, we measured $\mu_i$ (Figure S7). We observed no statistical difference among $\mu_i$ values for MCF-7 and MCF-10A cells throughout the entire frequency regime tested. We subsequently treated our channels with BSA and observed a 1.1% decrease in the mean value of $\mu_i$ for both MCF-7 and MCF-10A cells when compared with the untreated control case (Figure S7). Based on these results, we concluded that utilizing bare, untreated PDMS devices is appropriate under our given experimental conditions and that cell-surface interactions, although present, ultimately do not contribute significantly to the quantitative analysis of the cellular viscoelastic properties that our platform performs.

Figure 2C shows the storage ($G'$) and loss ($G''$) moduli (i.e., elasticity and viscosity, respectively) of MCF-7 and MCF-10A cells. Throughout the entire range of the effective frequency we tested, $G'$ for MCF-10A cells is greater than that for MCF-7 cells, indicating that these cells are generally stiffer. To investigate the frequency-dependent viscoelastic response of cells, we employed the power-law structural damping model (solid and dashed lines in Figure 2C). Table 1 provides the measured power-law components—$G_0$ (scaling factor), $\alpha$ (power-law exponent), and $\mu$ (Newtonian viscosity)—that we derived from measuring both cell types. As indicated in Table 1, $\alpha$ is much larger for MCF-10A than MCF-7 cells (0.69 versus 0.44), reflecting
the fact that their $G'$ increased more rapidly with deformation frequency. Physically, MCF-10A cells have a far stiffer mechanical response to increasing $u$ when compared with MCF-7 cells. With regard to viscous behavior, both cell types have exceedingly small $G''$ values when compared with $G'$ throughout the entire frequency regime tested, with MCF-10A cells having a $G''$ value only slightly greater than that of MCF-7 cells (Figure 2C and inset). The very small $G''$ values we obtain indicate that both malignant and non-malignant breast epithelial cell lines have "solid-like behavior," i.e., they have low viscosity. This is remarkably different from that observed at the tissue level, where malignant breast tissue has been shown to exhibit a more viscous response to oscillating stimuli when compared with normal tissue (Sinkus et al., 2005).

Contribution of Cytoskeletal Components to Cellular Properties

The cytoskeleton is an important component governing cellular mechanical behavior (Fletcher and Mullins, 2010). To investigate the individual contributions of cytoskeletal actin filaments and microtubules to the mechanical properties of cells, we analyzed the viscoelastic response of MCF-7 and MCF-10A cells when they were subject to different pharmacological treatments. Specifically, we stabilized actin filaments with jasplakinolide (Jas, Figures 3A and 3D) and de-activated them with latrunculin B (LatB, Figures 3A and 3D). In parallel, we stabilized and destabilized microtubules with paclitaxel (TAX) and nocodazole (Noc) treatment, respectively. We performed all measurements at $u > 100$ Hz to provide a sufficient window to detect different viscoelastic responses between the two cell types. As shown in Figures 3B and 3E, stabilized actin filaments led to an increased $G'$ for both cell types versus the respective controls, indicating that the treated cells had become stiffer. In contrast, de-activated actin filaments led to a decreased $G'$ and the treated cells had become softer. With regard to specific cell type, Jas and LatB treatment had the strongest effect on $G'$ at different frequencies, $u_{Jas} = 275$ Hz for MCF-7 cells and $u_{LatB} = 100$ Hz for MCF-10A cells, indicating that actin filaments have different frequency ranges in which they are actively engaged. The difference in actin filament behavior between MCF-7 and MCF-10A cells may be a result of their structural and functional differences, which in turn, are correlated with malignancy and metastatic potential (Lindberg et al., 2008; Proietti et al., 2018; Geltmeier et al., 2015). In terms of cell viscosity, both Jas- and LatB-treated MCF-7 and MCF-10A cells have extremely small values of $G''$ when compared with $G'$ (Figures 3C and 3F). Interestingly, both Jas and LatB treatment enhanced the viscous behavior of MCF-10A cells (noted as larger
untreated cells, indicating that the cells had softened. Similar to our result on cellular elasticity, LatB-compared with actin filaments in each cell-cycle phase via LatB treatment. In general, we measured lower To analyze the contribution of the nucleus to the viscoelastic properties measured, we de-polymerized spindle to separate the sister chromatids. Further reverses upon transitioning from G2 to M as the nuclear envelope breaks down, enabling the mitotic structuring of the nuclear envelope, all in preparation for division. Furthermore, this mechanical stability cells experience the maximum accumulation of cytoplasmic material, assembly of microtubules, and nucleus in this phase are mechanically the most stable. This is consistent with the fact that, in the G2-phase, response in the G2-phase, suggesting that the combined contribution of cytoskeletal filaments and the elasticity of MCF-7 cells is highly dependent on the cell-cycle phase. Cells have the stiffest mechanical

|                  | G₀ [Pa]            | α             | µ [Pa s]        |
|------------------|--------------------|---------------|----------------|
| MCF-7            | 396.42 ± 12.438    | 0.44 ± 0.006  | 2.17 ± 0.045   |
| MCF-10A          | 235.02 ± 5.789     | 0.69 ± 0.005  | 2.15 ± 0.074   |

Table 1. Parameters of the Power-Law Structural Damping Model with Corresponding Standard Error
G₀, α, and µ denote the shear modulus at zero frequency, power-law coefficient, and viscosity, respectively.

Gᵦ' values in our data when compared with the control untreated cells, Figure 3F), but further study is necessary to understand the biological basis of this phenomena.

Figures 4A and 4D show the viscoelastic properties of MCF-7 and MCF-10A cells after microtubule stabilization and de-activation. Stabilization resulted in an increased G' for MCF-7 cells, whereas disruption of microtubule formation led to a reduced G' for both cell types throughout all applied frequencies (Figures 4B and 4D). Both pharmacological microtubule treatments on MCF-7 cells showed the most significant effects at ω̅avg = 125 Hz (Figure 4B). Noc treatment, especially, revealed a bipolar effect on elasticity (G' ) measurements. At ω̅avg = 125 Hz, cells softened, which was reflected in the significantly smaller G' values when compared with untreated cells. At ω̅avg = 200 Hz, the difference in G' is relatively smaller than other frequencies. At ω̅avg = 275 Hz, G' of cells significantly decreased again by Noc treatment. For MCF-10A cells, disrupting microtubules via Noc treatment showed stronger effects on cellular elasticity when compared with stabilizing them with TAX treatment throughout the frequency range we tested (Figure 4D). Significantly, at ω̅avg = 180 Hz, the overall values of G' for MCF-10A cells decreased by TAX treatment, in contrast to other frequency conditions. This suggests that the dynamics of cytoskeletal filament formation induce different cellular mechanical behavior of cells and depend on loading conditions, i.e., frequency (Gardel et al., 2008; Brangwynne et al., 2006; Fletcher and Mullins, 2010). Similar to our results on the viscosity of cells after Jas and LatB treatments, both TAX- and Noc-treated cells showed a much smaller G' when compared with G' (Figures 4C and 4F), indicating that the cells consistently have a solid-like behavior. Finally, whereas TAX-treated MCF-7 cells have an increased G' in the high-frequency range, TAX-treated MCF-10A cells have a decreased G' with respect to increasing deformation frequency (Figure 4F). Cellular Mechanical Transition through Cell-Cycle Phases
Cells experience dramatic changes to their cytoskeleton and nucleus during the cell cycle (Baluska and Barlow, 1993; Rusan et al., 2001), and correspondingly these changes should induce different cellular mechanical behaviors. We thus analyzed the viscoelastic properties of MCF-7 cells that were synchronized in their cell cycle via a double thymidine block and release protocol (see Transparent Methods) and evaluated the effects of actin filament de-activation during each cell-cycle phase. Figures 5A and S8 show the morphological changes that MCF-7 cells undergo as they progress through the different cell-cycle phases, from G1- to S- to G2- to M-phase. As expected, the relative size of the nuclei increases during S-phase and stained nuclei show the classical spike shape of mitosis in M-phase. By immunostaining nuclear DNA, we validated cell synchronization and confirmed that DNA content increased as the cells traversed their cell cycle (Figure 5A). We screened synchronized MCF-7 cells with visco-NPS at ω = 200 Hz. Our results show that the G' of MCF-7 cells clearly increases as they go from G1- to S- to G2-phase. As the cells progress from G2- to M- back to G1-phase, there is a significant decrease in G' (Figure 5B). Our results show that the elasticity of MCF-7 cells is highly dependent on the cell-cycle phase. Cells have the stiffest mechanical response in the G2-phase, suggesting that the combined contribution of cytoskeletal filaments and the nucleus in this phase are mechanically the most stable. This is consistent with the fact that, in the G2-phase, cells experience the maximum accumulation of cytoplasmic material, assembly of microtubules, and structuring of the nuclear envelope, all in preparation for division. Furthermore, this mechanical stability reverses upon transitioning from G2 to M as the nuclear envelope breaks down, enabling the mitotic spindle to separate the sister chromatids.

To analyze the contribution of the nucleus to the viscoelastic properties measured, we de-polymerized actin filaments in each cell-cycle phase via LatB treatment. In general, we measured lower G' values versus untreated cells, indicating that the cells had softened. Similar to our result on cellular elasticity, LatB-treated cells also showed the largest G' value in G2 phase (Figure 5B). We evaluated the statistical difference among all experimental conditions based on correlation coefficient (rho) and p values (Table 2). G1 + LatB versus G2 + LatB, G2 + LatB versus M, S + LatB versus M + LatB, G2 versus M + LatB, and G2 + LatB
versus M + LatB showed significant statistical difference. From this, we conclude that the cell nucleus in G1-, G2-, and M-phases greatly contributes to cellular elasticity. To quantify the statistical effect of the two variables, i.e., cell-cycle phase and LatB treatment, we performed a two-way ANOVA test and showed that both cell-cycle phase ($p = 6.781 \times 10^{-8}$) and LatB treatment ($p = 6.806 \times 10^{-37}$) had a statistically significant effect on the $G'$ of MCF-7 cells. In addition, the combined effect of these two variables also had a significant effect on the $G'$ of MCF-7 cells ($p = 6.781 \times 10^{-8}$). Cellular viscosity ($G''$) showed a trend similar to elasticity ($G'$) regarding cell-cycle phase and LatB treatment. MCF-7 cells in G2-phase exhibited the largest $G''$ value in both untreated and LatB-treated groups (Figure 3C). As well, the perturbation of actin filaments by LatB

Figure 3. Contribution of Actin Filaments to Cellular Viscoelastic Properties

(A) Fluorescence images of MCF-7 cells after jasplakinolide (Jas) and latrunculin B (LatB) treatment. For comparison, fluorescence images of untreated cells (Ctrl) are also included. 4',6-diamidino-2-phenylindole (DAPI, blue) and rhodamine phalloidin (red) were used to stain the cell nucleus and actin filaments, respectively. Scale bar, 10 μm.

(B) Storage modulus ($G'$) of treated MCF-7 cells per the different ranges of effective deformation frequency ($\omega$) ($n = 100$ for all cases). Jas treatment stabilizes actin filaments, and LatB treatment de-activates them. Untreated cells were used as a control (Ctrl) case. Statistical differences were determined by an unpaired t test. Within each box, the central line is the median, the red cross is an outlier, and the edges of the box correspond to 25% and 75% of the population.

(C) Loss modulus ($G''$) of Jas- and LatB-treated MCF-7 cells compared with untreated (Ctrl) cells ($n = 100$ for each case). Error bars indicate standard error.

(D) Fluorescence images of MCF-10A cells after Jas and LatB treatment. For comparison, fluorescence images of untreated cells (Ctrl) are also included. DAPI (blue) and rhodamine phalloidin (red) were used to stain the cell nucleus and actin filaments, respectively. Scale bar, 10 μm.

(E) Storage modulus ($G'$) of treated MCF-10A cells per the different ranges of effective deformation frequency ($\omega$) ($n = 100$ for all cases). Statistical differences were determined by an unpaired t test. Within each box, the central line is the median, the red cross is an outlier, and the edges of the box correspond to 25% and 75% of the population.

(F) Loss modulus ($G''$) of Jas- and LatB-treated MCF-7 cells compared with untreated (Ctrl) cells ($n = 100$ for each case). Error bars indicate standard error. For all graphs $^{*}p < 0.05$, $^{**}p < 0.01$, $^{***}p < 0.001$, and $^{****}p < 0.0001$. See also Table S3.
treatment resulted in a decrease in $G''$ compared with untreated cells during each cell-cycle phase. Strikingly, LatB-treated MCF-7 cells showed a more dramatic decrease of $G''$ in S-phase (Figure 5C inset). In S-phase, cellular DNA is actively being unwound by helicases and replicated and might therefore be less mechanically stable than DNA packaged around histones and in chromosomes. In general, visco-NPS provides sufficient sensitivity to measure the contributions of actin filaments and nuclear structure to the cellular viscoelastic properties throughout the different cell-cycle phases.

Figure 4. Contribution of Microtubules to Cellular Viscoelastic Properties
(A) Fluorescence images of MCF-7 cells after paclitaxel (TAX) and nocodazole (Noc) treatment. For comparison, fluorescence images of untreated cells are shown as the control case (Ctrl). DAPI (blue) and green fluorescence protein (GFP, green) stain cell nucleus and tubulin, respectively. Scale bar, 10 μm.
(B) Storage modulus ($G'$) of treated MCF-7 cells with respect to pharmacological treatments per the different range of effective deformation frequency ($\omega$) (n = 100 for all cases). TAX treatment stabilizes microtubules, whereas Noc treatment de-activates them. Untreated cells were used as a control (Ctrl) case. Statistical differences were determined by an unpaired t test. Within each box, the central line is the median, the red cross is an outlier, and the edges of the box correspond to 25% and 75% of the population.
(C) Loss modulus ($G''$) of TAX- and Noc-treated MCF-7 cells compared with untreated (Ctrl) cells (n = 100 for each case). Error bars indicate standard error.
(D) Fluorescence images of MCF-10A cells after TAX and Noc treatment. For comparison, fluorescence images of untreated cells are shown as the control case (Ctrl). DAPI (blue) and green fluorescence protein (GFP, green) stain cell nucleus and tubulin, respectively. Scale bar, 10 μm.
(E) Storage modulus ($G'$) of treated MCF-10A cells with respect to pharmacological treatments per the different range of effective deformation frequency ($\omega$) (n = 100 for all cases). Statistical differences were determined by an unpaired t test. Within each box, the central line is the median, the red cross is an outlier, and the edges of the box correspond to 25% and 75% of the population.
(F) Loss modulus ($G''$) of TAX- and Noc-treated MCF-10A cells compared with untreated (Ctrl) cells (n = 100 for each case). Error bars indicate standard error. For all graphs, *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, and ****p ≤ 0.0001. See also Table S3.
SLUG-Initiated Epithelial-Mesenchymal Transition Induces Cellular Mechanical Changes

To highlight further the power and sensitivity of our platform in detecting dynamic state changes in cells, we initiated EMT re-programming in MCF-10A cells and studied the effects of this stimulation on cellular viscoelastic properties. EMT is a cellular program that assumes diverse roles in biology, from embryogenesis to wound healing to tumor progression, in which epithelial characteristics of tight cell-cell junctions and polarity are attenuated in favor of a migratory and invasive, mesenchymal-like phenotype (Kalluri and Weinberg, 2009; Yang and Weinberg, 2008). SLUG, a member of the Snail family transcription factors, has been identified as one of many dynamic regulators of EMT, driving a key hallmark of this transition—namely, E-cadherin downregulation by binding to the E-boxes of the E-cadherin promoter and repressing transcription (Shih and Yang, 2011; Yang and Weinberg, 2008). Here, we activated the EMT cascade by transducing MCF-10A cells with a retroviral vector expressing SLUG. As EMT has been described as a dynamic spectrum with transitional states in which epithelial cells adopt any permutation of mesenchymal characteristics, we first characterized the phenotypic changes caused by the induction of SLUG, comparing against untreated MCF-10A cells (Aiello et al., 2017; Nieto et al., 2016; Brabletz et al., 2018; Micalizzi et al., 2010). One of the first changes that we observed was in terms of morphology (Figure S9). Normal MCF-10As adopted classical epithelial morphology with “cobblestone” appearance and apical-basal polarity, whereas our virus-infected cells exhibited a greater degree of cell spreading, increased cell lamellipodia and filopodia, and disassembly of intercellular junctions, all of which are consistent with EMT (Lamouille et al., 2014). We then proceeded to compare the expression patterns of a panel of common EMT markers: SLUG, E-cadherin, vimentin, and N-cadherin (Zeisberg and Neilson, 2009). As shown in Figure 6A, the key difference that we observed was in terms of morphology (Figure S9). Normal MCF-10As adopted classical epithelial morphology with “cobblestone” appearance and apical-basal polarity, whereas our virus-infected cells exhibited a greater degree of cell spreading, increased cell lamellipodia and filopodia, and disassembly of intercellular junctions, all of which are consistent with EMT (Lamouille et al., 2014). We then proceeded to compare the expression patterns of a panel of common EMT markers: SLUG, E-cadherin, vimentin, and N-cadherin (Zeisberg and Neilson, 2009). As shown in Figure 6A, the key difference that we observed was in terms of morphology (Figure S9). Normal MCF-10As adopted classical epithelial morphology with “cobblestone” appearance and apical-basal polarity, whereas our virus-infected cells exhibited a greater degree of cell spreading, increased cell lamellipodia and filopodia, and disassembly of intercellular junctions, all of which are consistent with EMT (Lamouille et al., 2014). We then proceeded to compare the expression patterns of a panel of common EMT markers: SLUG, E-cadherin, vimentin, and N-cadherin (Zeisberg and Neilson, 2009). As shown in Figure 6A, the key difference that we observed was in terms of morphology (Figure S9). Normal MCF-10As adopted classical epithelial morphology with “cobblestone” appearance and apical-basal polarity, whereas our virus-infected cells exhibited a greater degree of cell spreading, increased cell lamellipodia and filopodia, and disassembly of intercellular junctions, all of which are consistent with EMT (Lamouille et al., 2014). We then proceeded to compare the expression patterns of a panel of common EMT markers: SLUG, E-cadherin, vimentin, and N-cadherin (Zeisberg and Neilson, 2009). As shown in Figure 6A, the key difference that we observed was in terms of morphology (Figure S9). Normal MCF-10As adopted classical epithelial morphology with “cobblestone” appearance and apical-basal polarity, whereas our virus-infected cells exhibited a greater degree of cell spreading, increased cell lamellipodia and filopodia, and disassembly of intercellular junctions, all of which are consistent with EMT (Lamouille et al., 2014).
a mesenchymal state (Zeisberg and Neilson, 2009). However, SLUG transduction did not result in E-cadherin to N-cadherin switching (Figure 6A(iv) and S10). Our SLUG-transduced MCF-10As, therefore, appeared to have acquired an intermediate EMT phenotype, which is consistent with the work of others within the field that overexpressed SLUG and also reported decreased E-cadherin and increased vimentin expression (Bolós et al., 2003; Come et al., 2004; Lee et al., 2008; Vuoriluoto et al., 2011). Having validated a transitioned state, we then investigated the mechanical cell properties of untreated versus EMT-initiated cell states using our device. We found that SLUG MCF-10A cells exhibited significantly larger Gʺ (i.e., they were more viscous) than untreated MCF-10A cells, despite both cells possessing a similar level of Gʹ (Figures 6C and 6D). Although further in-depth study is required to dissect fully the underlying biological contributions, we, nonetheless, have observed morphological, phenotypic, and viscoelastic changes of cells as they undergo an induced EMT via SLUG (Voulgari and Pintzas, 2009; Zhang et al., 2014).

**DISCUSSION**

Visco-NPS enables the analysis of populations of single cells for their viscoelastic properties. By combining a sinusoidal contraction channel with node-pore sensing (Balakrishnan et al., 2013) or mechano-NPS (Kim et al., 2018), we can quantify the mechanical response of cells as they periodically deform. Based on just two mechanical properties, storage (elasticity) and loss (viscosity) moduli, malignant and non-malignant breast epithelial cells show distinctly different viscoelastic behavior and frequency-dependent responses to dynamic loading conditions. Yet, for all frequencies measured, each cell type’s viscosity was smaller than its elasticity. These results are in direct contrast to prior AFM studies measuring the same cell types, which report that viscosity has a greater value than elasticity above a threshold frequency (Charras and Horton, 2002; Alcaraz et al., 2003; Haase and Pelling, 2015; Rother et al., 2014). Likewise, our results that centered on the effects of EMT on cell viscoelastic behavior differ from those studies in which EMT had been induced in cells by transforming growth factor-β1 and subsequently measured via AFM (Schneider et al., 2013; Efremov et al., 2017). In these studies, EMT-induced cells were shown to exhibit more solid-like behavior and greater energy storage, which is in contrast with our findings that such cells exhibit more viscous behavior. The discrepancy between these results more than likely rests upon the fundamental differences between the two characterization modalities—AFM and visco-NPS—with regard to how they probe cells. Visco-NPS measures whole-cell mechanical properties, whereas AFM targets localized, cellular sub-regions. Equally important, visco-NPS measures cells in suspension and, arguably, in a more unbiased state, as AFM requires cells to be adhered to a substrate. In the latter case, substrate stiffness, topography, and porosity are all key material parameters that are capable of re-modeling the numerous components of
the cytoskeleton and, thus, modulating cell mechanics. This is especially important to consider with regard to our EMT experiments in which we observed a dramatic upregulation of vimentin expression, which is well known to affect the mechanical properties of epithelial cells (Guo et al., 2013; Hu et al., 2018). Mendez et al. report that the biomechanical contributions of vimentin intermediate filaments depend greatly on the stiffness of the substrate upon which cells are immobilized for AFM (Mendez et al., 2014). Thus, in general, compared with visco-NPS, AFM requires an extra step of de-coupling substrate properties from observed cell mechanical phenomena.

As we have shown, visco-NPS can determine the individual contributions of cytoskeletal components, such as actin filaments and microtubules, to cellular viscoelastic properties. Furthermore, through pharmacological treatments, in which we stabilized or de-activated cytoskeletal filaments or microtubules, visco-NPS can quantitatively measure the subsequent effects on the elasticity and viscosity of the treated cells. Finally, the sensitivity of visco-NPS enables measurement of the changes in mechanical properties of cells as they traverse their cell cycle or are initiated into EMT. In general, visco-NPS provides a comprehensive understanding of the dynamic mechanical behavior of cells.

Attractive features of visco-NPS are that it is label free, does not require a high-speed camera or advanced fluidics, and has high-throughput capability. Its simplicity allows it to be combined potentially with other cell analyses, including cell-surface marker screening (Balakrishnan et al., 2015) and single-cell whole-transcriptome analysis (Shapiro et al., 2013; White et al., 2011; Streets et al., 2014). This, in particular, would provide opportunities to correlate mechanical properties with biochemical properties. The design flexibility of visco-NPS allows us to utilize a variety of experimental conditions, models, and mechanical tests for further studies. By varying the width and period of the sinusoidal contraction channel, one can apply different levels of strain to, and can access different frequency regimes in which to measure, cells. Moreover, contraction channels with different periodic lengths placed in series would allow one to measure a single
cell at multiple deformation frequencies. In so doing, one can determine the frequency dependence of the cell’s viscoelastic properties, which ultimately would provide a more in-depth understanding of the cellular mechanical properties at the single-cell level.

Although we have focused on the power-law structural damping model, other mechanical models (Lim et al., 2006) are applicable to this platform. For example, cortical shell-liquid core (Yeung and Evans, 1989), linear viscoelastic solid (Karcher et al., 2003; Schmid-Schönbein et al., 1981), and biphasic models (Leterrier, 2001; Haider, 2004) could all be applied to different cell types and their corresponding biophysical function. By modifying the microfluidic channel design, one can perform different types of mechanical tests. For example, one could perform a mechanical creep test by applying a constant compressive stress to the deformed cells through pneumatic side channels positioned parallel to the sides of the sinusoidal contraction channel. Such a test could provide an understanding of the biophysical relationship between mechanical properties and cell recovery characteristics (Bonakdar et al., 2016). Even in its present form, visco-NPS, can successfully measure the viscoelastic properties of cells under various biological conditions, as we have demonstrated here.

Cellular mechanics is an emerging area of research, especially within the context of cancer (Kumar and Weaver, 2009; Goldmann et al., 2013; Peyton et al., 2007). Quantifying the viscoelastic properties of cell populations, visco-NPS introduces new possibilities in basic biology and clinical applications, such as cancer diagnosis, cytoskeleton-targeted drug screening, studies for cell cycle as a therapeutic target of cancer, and EMT-induced cellular mechanical transition. By combining the clear relationship between malignancy and viscoelastic properties of cells with numerous features of our platform (e.g., label-free analysis and high-throughput capability), visco-NPS could potentially be used as a cancer screening and diagnostic tool, especially for those cancer cells, e.g., triple-negative breast cancer, that are otherwise difficult to identify with traditional methods (Dent et al., 2007; Schneider et al., 2008; Hudis and Gianni, 2011). As it can quantify the contribution of cytoskeletal filaments to whole-cell mechanical properties, visco-NPS could be employed as a screening method for the cytoskeleton-targeted drugs that are often used in cancer therapies (Martin et al., 2014; Checchi et al., 2003; Tommasi et al., 2007). Recently, a number of studies have shown that cell-cycle pathways, such as regulation of CDK (cyclin-dependent kinases), ATM (ataxia telangiectasia mutated), and ATR (AMT- and Rad3-related), lead to aberrant cell proliferation and are critical to carcinogenesis (Hartwell and Kastan, 1994; Kastan and Bartek, 2004; Abraham, 2001; Smith et al., 2010). By analyzing the cellular mechanical transitions during the cell cycle and quantifying the mechanical properties of actin filaments as they are perturbed during each phase, visco-NPS could yield a new approach to the development of cancer therapies that target these, and other, cell-cycle pathways.

Visco-NPS is an electronic-based, high-throughput, single-cell analysis method that can quantify viscoelastic properties of cell populations with respect to a wide range of deformation frequencies. As demonstrated, measurement of a cell’s storage and loss moduli in combination with the power-law structural damping model provides a quantitative mechanical metric for comparing different cell types, evaluating cytoskeletal contributions to cellular properties, and analyzing cellular mechanical transitions through the different phases of the cell cycle or other significant morphological and phenotypic changes, such as EMT. Visco-NPS thus has great potential to be utilized as an efficient measurement tool for those basic research and clinical applications that require characterization of cellular mechanics.

Limitation of Study
There are some possible drawbacks of the present study. First, we focused on cells from breast tissue to analyze cellular viscoelastic properties. Studying cells from other tissues could provide more broad information regarding the relationship between different cancers and cellular mechanical properties. Second, all cells tested in this study were epithelial cell lines (MCF-7 and MCF-10A) rather than patient-derived cells. Third, visco-NPS in its current platform design measures a single cell at only one specific deformation frequency. Thus the frequency dependence of cellular viscoelastic properties can only be assessed for populations of cells versus individual cells. Fourth, measuring the whole-cell properties via visco-NPS resulted in significantly smaller values of cellular viscosity (i.e., loss modulus, \(G''\)) when compared with elasticity (i.e., storage modulus, \(G'\)) for most of experimental cases, which could limit the resolution of measurement.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2019.02.021.

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AUTHOR CONTRIBUTIONS

J.K. conceived of the device and method and performed the visco-NPS measurements; O.J.S. performed the viral packaging and viral transduction of SLUG; B.L. and O.J.S. contributed to the cytimmunoochemistry characterization of SLUG-infected MCF-10A cells; Y.K. assisted with the fabrication of the devices and performed tissue culture; J.K., B.L., and L.L.S. analyzed the results; J.K., B.L., and L.L.S. devised experiments; J.K. and B.L. performed FACS; and J.K., B.L., O.J.S., and L.L.S. wrote the manuscript.

DECLARATION OF INTERESTS

L.L.S. and J.K. are inventors of a pending patent (U.S. Patent: 62/571,255 PCT/US17/5642), which covers the design of the visco-NPS platform introduced in this article.

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Supplemental Information

Visco-Node-Pore Sensing: A Microfluidic Rheology Platform to Characterize Viscoelastic Properties of Epithelial Cells

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Supplementary Figures

Supplementary Figure 1, related to Figure 1. Fluid flow in visco-NPS. A, Schematic of the microfluidic channel with a sinusoidal contraction channel (Green Dashed box). In the inset, the red dashed line corresponds to the microfluidic channel’s longitudinal axis. \( w_c, L_p, w_0, \alpha \) and \( \omega \) are the contraction channel’s width, periodic length, initial width, strain amplitude, and deformation frequency, respectively. B, Fluid-velocity profile along the longitudinal axis of a channel described by \( w_c=8.71+1.5\cos(\omega t) \), \( L_p=1000\mu m \), and an inlet pressure of 10 Pa. Comsol Multiphysics was employed to calculate the velocity magnitude. The sinusoidal shape of the velocity magnitude confirms that the contraction channel does indeed have a periodically changing channel width, even though it is not visible with naked eye due to the gradual change of \( w_c \).

Supplementary Figure 2, related to Figure 1. Distribution of free cell diameter. A, Box plot of free cell diameter for MCF7 and MCF10A cells, measured from phase images. Within each box, the central line is the median, the red cross is an outlier, and the edges of the box correspond to 25% and 75% of the population (MCF7; \( n=447 \) MCF10A; \( n=459 \)). B, Size distribution of MCF-7 cells measured by visco-NPS
Supplementary Figure 3, related to Figure 1. Experimental set up for visco-NPS. A commercial microfluidic pump applies a non-pulsatile pressure to drive a suspension of cells through the channel. A schematic drawing on the right side shows the electrical circuit model for the DC voltage, four-point measurement performed (Kim et al., 2018, Saleh, 2003). The yellow dashed box shows the equivalent circuit elements of the fluid flowing through the microfluidic channel and the cell that transits the channel. \( R_c \) and \( C_c \) are the resistance and capacitance of the microfluidic channel, respectively. \( R_H \) and \( R_L \) correspond to the resistance of the fluid in the inlet and outlet reservoir, respectively. \( V_H - V_L \) is the voltage difference across the channel. In this circuit, the electric current flows from \( I_H \), to \( I_L \), to \( I_{out} \). \( R_f \) and \( C_f \) are additional elements that remove oscillations in the circuit.

Supplementary Figure 4, related to Figure 2. Storage modulus (\( G' \)) of MCF-10A cells from different replicas of the visco-NPS device at a deformation frequency of 140 Hz. \( G' \) of MCF-10A cells as measured by the two different device replicas show no statistical difference through an un-paired t-test (MCF-10A(1): \( n=92 \), MCF10A(2): \( n=76 \), and \( p=0.956 \)). Within each box, the central line is the median and the edges correspond to 25% and 75% of the \( G' \) distribution.
Supplementary Figure 5, related to Figure 2. Schematic drawing of the forces around the deformed cell while in the contraction channel. $F_{\text{drive}}$, $F_n$, $F_{\text{flic}}$, and $D_d$ correspond to the driving force, normal force, frictional force, and the diameter of the deformed cell while in the contraction channel, respectively. Under a cell’s constant velocity, the driving force generated by the pressure difference around the cell is counter-balanced by the frictional forces between cell surface and the channel wall.

Supplementary Figure 6, related to Figure 2. Stress-strain relationship of a MCF7 cell measured by visco-NPS. A, Applied strain and B, stress of an MCF-7 cell transiting the contraction channel with $L_p=500\ \mu\text{m}$ and $V_{\text{pore}}=65\ \text{mm/s}$. During the cell’s transit through the contraction channel, $\Delta T_{n}=1,2,\ldots,5$ represent the period of stress it experiences ($\Delta T_{n} = 5.5 \pm 0.17\ [\text{ms}]$). The constant period of stress indicates that the cell transits the contraction channel with constant velocity.
Supplementary Figure 7, related to Figure 2. Interaction between cell surface and channel wall. A, Measured frictional coefficient ($\mu_f$) of MCF-7 (blue) and MCF-10A (red) cells as they transit through bare PDMS devices ($n=100$ for all cases). No statistical difference was found in $\mu_f$ for both cell types throughout the entire $\omega$ regime measured. B, Schematic drawing of the experimental conditions (bare PDMS vs. BSA-treated PDMS) used to measure the effects of cell-surface interaction on $\mu_f$. BSA treatment minimizes cellular adhesion to the channel wall. C, $\mu_f$ of MCF7 and MCF10A cells as they transit through bare PDMS and BSA-treated channels ($n=100$ for each case). For all graphs, error bar represents standard deviation. **** indicates $p \leq 0.0001$.

Supplementary Figure 8, related to Figure 5. Representative fluorescence images of MCF-7 cells in each cell-cycle phase. 4',6-diamidino-2-phenylindole (DAPI, blue) and rhodamine phalloidin (red) stain the cell nucleus and actin filaments, respectively. Scale bar = 5 $\mu$m
Supplementary Figure 9, related to Figure 6. Representative phase contrast images of SLUG induction of MCF-10A. Untreated MCF-10A cells at low confluence (top-left, 10X; top-middle, 20X) and high confluence (top-right, 20X) form regularly-structured monolayers with rectangular morphology. SLUG-induced MCF-10A cells are spindle-shaped at low confluence (bottom-left, 10X; bottom-middle, 20X) and high confluence (bottom-right, 20X), with notable protrusions at low confluence (bottom-middle, 20X). Scale bar = 25 μm

Supplementary Figure 10, related to Figure 6. Flow cytometric analysis of SLUG induction of MCF-10A. SLUG induction in MCF-10A cells results in decreased E-cadherin expression but not an increase in N-cadherin expression.
**Supplementary Table 1**, related to Figure 1. Microfluidic channel dimensions and the applied strain to cells. \( L_p, w_{\text{pore}}, \) and \( w_c \), correspond to the periodic length of the contraction channel, pore width, and the contraction channel width, respectively. \( D_{\text{cell}} \) is the free cell diameter (mean±standard deviation) and \( \varepsilon \), the applied strain to the cells.

| Cell type | \( L_p \) [\( \mu \text{m} \)] | \( w_{\text{pore}} \) [\( \mu \text{m} \)] | \( w_c \) [\( \mu \text{m} \)] | \( D_{\text{cell}} \) [\( \mu \text{m} \)] | \( \varepsilon \) |
|-----------|-------------------------------|---------------------------------|-----------------|------------------|--------|
| MCF-7     | 1000                          | 22                              | 10.5±1.5cos(\( \omega t \)) | 17.0±2.63       | 0.4±0.1cos(\( \omega t \)) |
| MCF-10A   | 1000                          | 18                              | 8.75±1.5cos(\( \omega t \)) | 15.2±1.48       |        |

**Supplementary Table 2**, related to Figure 1. Quantification of the channel’s effective diameter. Polystyrene microspheres (Polysciences, #64155) were used to measure the effective diameter of the node-pore channel region \( (D_{\text{eff,np}}) \) \( (n=20\) for all cases). Within the table, \( d, \Delta I/I, w_{\text{pore}}, L, \) and \( L_p \), correspond to the average diameter + standard deviation of the spheres, width of pore, total channel length, and periodic length of the contraction channel, respectively.

| \( d \) [\( \mu \text{m} \)] | \( \Delta I/I \) | \( w_{\text{pore}} \) [\( \mu \text{m} \)] | \( L \) [\( \mu \text{m} \)] | \( L_p \) [\( \mu \text{m} \)] | \( D_{\text{eff,np}} \) [\( \mu \text{m} \)] |
|-----------------------------|----------------|-------------------------------|-----------------|------------------|------------------|
| 14.73±1.36                 | 5.39e-4        | 22                            | 6500            | 1000             | 31.30            |
|                            | 9.09e-4        |                               | 4000            | 500              | 27.70            |
|                            | 8.17e-4        | 18                            | 6500            | 1000             | 26.10            |
|                            | 14.54e-4       |                               | 4000            | 500              | 25.45            |

**Supplementary Table 3**, related to Figure 3 and 4. Power of experimental groups with \( p<0.05 \). The 2-sample and 2-sided power analysis with 95% of confidence interval was employed to ensure experimental groups have adequate power compared to their own control groups. With the analyzed sample size \( (n=100\) for all), all experimental cases show adequate power (\( >0.08 \)). \( \omega_{\text{avg}} \) corresponds to the averaged deformation frequency.

| Cell type | MCF-7_Jas | MCF-7_Jas | MCF-7_Jas | MCF-7_LatB | MCF-10A_Jas |
|-----------|-----------|-----------|-----------|------------|-------------|
| \( \omega_{\text{avg}} \) [Hz] | 125       | 200       | 275       | 275        | 100         |
| Power     | 0.987     | 1.000     | 0.999     | 1.000      | 1.000       |

| Cell type | MCF-10A_Jas | MCF-10A_LatB | MCF-10A_LatB | MCF-7_TAX | MCF-7_TAX |
|-----------|-------------|--------------|--------------|-----------|-----------|
| \( \omega_{\text{avg}} \) [Hz] | 140         | 100          | 140          | 125       | 200       |
| Power     | 0.995       | 0.978        | 0.976        | 1.000     | 0.999     |

| Cell type | MCF-7_Noc | MCF-7_Noc | MCF-7_Noc | MCF-10A_TAX | MCF-10A_Noc |
|-----------|-----------|-----------|-----------|-------------|-------------|
| \( \omega_{\text{avg}} \) [Hz] | 125       | 200       | 275       | 180        | 100         |
| Power     | 1.000     | 0.966     | 1.000     | 0.990      | 1.000       |

| Cell type | MCF-10A_Noc | MCF-10A_Noc |
|-----------|-------------|-------------|
| \( \omega_{\text{avg}} \) [Hz] | 140       | 180         |
| Power     | 1.000     | 1.000       |
Transparent Methods

Device fabrication

Standard soft-lithography is employed to fabricate the PDMS mold for the visco-NPS microfluidic device. Briefly, SU-8 3025 resist (MicroChem, USA) is employed to fabricate negative-relief masters onto polished silicon wafers. A prepared mixture of PDMS (1:9 weight ratio of elastomer to curing agent, Dow Corning, USA) is poured onto the masters and subsequently cured at 85°C for 2 hours. A slab of PDMS with the embedded microfluidic channel is cut and excised from the master, and inlet and outlet holes are created using a 1-mm biopsy punch (Integra LifeSciences, USA). To make the Pt electrodes and Au contract pads, standard photolithography with positive photoresist (S1813, MicroChem, USA) is used to pattern the glass substrates. Electron-gun evaporation is employed to deposit a 100/250/250 Å of Ti (Titanium)/Pt/Au film onto the patterned substrate. A gold wet etch (Gold Etchant TFA, Transene Company, USA) exposes the Pt electrodes. To create a permanent bond between the two, the PDMS mold and the glass substrate with pre-defined electrodes are both exposed to an oxygen plasma (470 mTorr, 80W, 1min). Once the PDMS mold and glass substrate are aligned, mated, and sealed, the completed device is baked at 85°C on a hotplate for 2 hours. For the specific surface-treatment experiments described, 2% (w/v in PBS) of bovine serum albumin (BSA) was injected into sealed devices. After incubation at 37°C for 2 hours, the channels were flushed with 1X PBS, and cells were immediately injected into the devices for measurement.

Electronic-based measurement of visco-NPS

We quantified multiple biophysical properties of cells, including free-cell diameter and deformation, by employing node-pore sensing (NPS) (Balakrishnan et al., 2013, Balakrishnan et al., 2015, Kim et al., 2018), a method based on the Coulter principle (Saleh and Sohn, 2001, Coulter, 1953, DeBlois and Bean, 1970, Carbonaro et al., 2008). As shown in Figure 1D, a unique current pulse is measured when a cell transits the channel. This unique pulse reflects the channel geometry and is the result of the cell partially blocking the flow of the electric current in the channel. Because the current density changes from node to pore to channel, the amount of current that the cell blocks in these different regions correspondingly changes, leading to the current pulse with the overall periodic structure shown in Figure 1D. When the cell specifically transits the sinusoidal contraction channel and is periodically deformed, the resulting sub-pulse also oscillates.

To determine the size a cell, the following relationship is employed,

\[
\frac{\Delta I}{I} = \frac{d^3}{D_{\text{eff}}^2 L} \left[ \frac{1}{1 - 0.8 \left( \frac{d}{D_{\text{eff}}} \right)^3} \right]
\]

where \(\Delta I/I\), \(d\), \(D_{\text{eff}}\), \(L\) correspond to current drop, baseline current, cell diameter, effective channel diameter, and overall channel length, respectively (Saleh and Sohn, 2001, DeBlois and Bean, 1970). \(D_{\text{eff}}\) is numerically determined by measuring the \(\Delta I/I\) caused by polystyrene microspheres of known size, \(d\), traversing the channel (Supplementary Table 2). The effective diameter of the first node-pore region, \(D_{\text{eff, np}}\), can thus be determined and used with \(\Delta I_{\text{np}}/I\) in Equation 1 to quantify the free-cell dimeter (\(D_{\text{cell}}\)). The effective contraction channel diameter (\(D_{\text{eff, c}}\)) is determined by a ratio of the hydraulic diameter of the node-pore and the contraction channel (\(D_{\text{eff, c}} = D_{\text{eff, np}} \sqrt{w_{\text{np}}/w_{\text{pore}}}\)). To quantify cellular deformation in the contraction channel, we assume that the cell undergoes an isometric deformation in a plane parallel to the channel’s side walls and is consequently disk shape (Supplementary Figure 4). We determine the deformed cell diameter (\(D_{\text{d}}\)) from \(\Delta I/I \sim V_{\text{deform}}/V_{\text{contraction}}\), where \(V_{\text{deform}}\) and \(V_{\text{contraction}}\) are the volume of the deformed cell and contraction channel, respectively (Kim et al., 2018).

Cell culture

MCF-7 cells (ATCC® HTB-22™) were cultured in DMEM (Fisher Scientific, BW12719F), supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine, and 1% Pen-Strep. MCF-10A cells (ATCC® CRL-
10317™) were cultured in MEBM medium (ATCC® PCS-600-030™), supplemented with 0.1% insulin, 0.1% human Epidermal Growth Factor, 0.4% hydrocortisone, and 10% cholera toxin. All cell cultures were maintained at 37°C in 5% CO₂ and routinely passaged, per published protocols (Freshney, 2005), once they reached 80% confluence. Cells were dissociated with 0.25% trypsin/EDTA for either 3 min (MCF-7 cells) or 5 min (MCF-10A cells) at 37°C (Sugarman et al., 1985, Giard et al., 1973, Reddel et al., 1989), washed with respective growth media, centrifuged at 0.2 RCF for 4 min, and re-suspended at a concentration of ~500,000 cells/mL in PBS. To ensure cell viability, cells were injected into the prepared devices for measurement immediately following re-suspension.

Pharmacological treatment for cytoskeletal components

Jasplakinilide (Jas, Abcam, USA) and Latrunculin B (LatB, Enzo Life Science, USA) were used to stabilize and disrupt actin polymerization, respectively. Paclitaxel (TAX, Abcam, USA) and Nocodazole (Noc, Thermo-Fisher Scientific, USA) were used to stabilize and perturb microtubule formation, respectively. All drugs were dissolved in dimethyl sulfoxide (DMSO) and then added to the respective growth medium for each cell type to achieve a final concentration of 200nM Jas, 5 µg/mL LatB, 2 µM TAX, and 20 µM Noc.

Prior to measurement, MCF-7 and MCF-10A cells were incubated with Jas, LatB, TAX, or Noc solution for 6, 2, 18, or 10 hours at 37°C, respectively. The concentration and incubation time of the pharmacological treatments were chosen to have an adequate effect on the cells and are based on previously published work (Kubitschke et al., 2017, Kim et al., 2018). Cells were detached from their culture flask with 0.25% trypsin/EDTA, rinsed once with growth medium, centrifuged at 0.2 RCF for 4 min, and re-suspended in PBS at a concentration of ~500,000 cell/mL. To confirm that actin polymerization was successfully stabilized or disrupted by the treatment, the cells were fixed with 4% (w/v) of paraformaldehyde in PBS for 15 min. They were then permeabilized with 0.1% Triton-X 100 (Sigma-Aldrich, USA) in PBS for 5 min. To visualize the effects of Jas and LatB treatments, cell nuclei and actin filaments were counter-stained with 4',6-diamidino-2-phenylindole (DAPI, Sigma-Aldrich, USA) and rhodamine phalloidin (Thermo-Fisher Scientific, USA), respectively, using the protocols provided by the manufacturer. Stabilization and de-activation of microtubules were confirmed by staining with CellLight® Tubulin-GFP (Life Technology, USA) for live cells. For DAPI nuclear staining, cells were fixed and permeabilized using the same protocol described above. All stained cells were imaged with a Zeiss LSM780 confocal microscope.

Cell-cycle synchronization

A standard double-thymidine block and release protocol was used to synchronize MCF-7 cells to the border of Gap 1 (G1)/Synthesis (S) phase (Bostock et al., 1971). Briefly, cells were first treated with 2.5 mM of thymidine (Sigma-Aldrich, USA) for 20 hours, released into growth medium for 9 hours, and then treated once again with thymidine for an additional 16 hrs. Once synchronized, cells were captured in S- or G1-phase by incubating in fresh medium for 30 min or 12 hrs, respectively. To capture cells in Gap 2 (G2) phase, synchronized cells were incubated with 9 µM of CDK1 inhibitor RO3306 (VWR, USA) for 5 hours. Cells in Mitosis (M) phase were captured by incubating synchronized cells with 10 µM of dimethylenastron (VWR, USA) for 10 hours. All incubations were performed at 37°C with 5% of CO₂. To validate cell-cycle synchronization, cells in each phase were fixed and stained with DAPI (Sigma-Aldrich, USA) and rhodamine phalloidin (Thermo-Fisher Scientific, USA). The DNA content in the nucleus was quantified using DAPI fluorescence intensity. All synchronized cells were analyzed with visco-NPS using the methods described above.

Viral Packaging and Transduction of pPGS-hSLUG.fl.flag

HEK 293Ts were cultured at 37°C and 5% CO₂ in DMEM (Corning Cellgro), supplemented with 10% fetal bovine serum (FBS, Life Technologies) and 1% penicillin/streptomycin (P/S, Life Technologies). pPGS-hSLUG.fl.flag (#25696, Addgene) plasmid DNA was transfected into 293Ts along with retroviral helper plasmids using polyethylenimine (PEI) at a 4:1 ratio (PEI (ug): DNA (ug)). Media from virus-producing 293Ts was collected days 2 and 3 post-transfection, pooled, and filtered through a 0.45-µm syringe filter. To separate virus from 293T contamination, virus was centrifuged in a SW28 swinging bucket rotor in a
Beckman Dickinson ultracentrifuge in the presence of a 20% sucrose layer (2 hr, 24,000 RPM, 4°C). The resulting pelleted virus was re-suspended in sterile PBS and stored at -80°C until ready to use. MCF10As were infected with SLUG-expressing retrovirus and selected with 400 μg/mL G418 (InvivoGen) for 7 days. Following selection, cells were expanded, aliquoted, and frozen down.

Immunostaining

Both MCF10As and SLUG-expressing MCF10As were fixed with 4% (w/v) paraformaldehyde for 10 minutes at room temperature and subsequently washed 3x with 1X PBS. Cells were blocked and permeabilized for 1 hour at room temperature with 5% donkey serum and 0.1% Triton X-100 prepared in PBS (PBS-DT). The following primary antibodies were then diluted in PBS-DT and incubated with cells overnight on a shaker at 4°C: 1:100 rabbit polyclonal IgG for SLUG (ab27568, Abcam), 1:500 rabbit monoclonal IgG for E-cadherin (ab40772, Abcam), 1:250 mouse monoclonal IgG for vimentin (ab8978, Abcam), 1:100 mouse monoclonal IgG for N-cadherin (33-3900, Thermo Fisher Scientific). The next day, primary antibody solution was removed, cells were washed 3X with PBS, and the following secondary antibodies were prepared in PBS-DT and allowed to incubate for 1 hour at room temperature on a shaker: 1:250 AlexaFluor488 donkey anti-rabbit IgG (H+L) (A-21206, Thermo Fisher Scientific) and 1:250 AlexaFluor 488 donkey anti-mouse IgG (H+L) (A-21202, Thermo Fisher Scientific). DAPI was added during the second of four 5-minute washes with PBS. Samples were stored in PBS before and during imaging.

Flow Cytometric Analysis

Cells were disassociated according to previously described methods and resuspended in 1X PBS at a concentration 1,000,000 cells/mL. Unstained MCF-10A and SLUG-induced MCF-10A cells were used as negative controls and stored in 1X PBS supplemented with 1% FBS on ice during immunostaining. Cell samples for immunostaining were then pelleted via centrifugation at 1000 RCF for 4 min. Samples were blocked with 100 uL of 5% normal goat serum (ThermoFisher Scientific, USA). The two primary antibodies – rabbit monoclonal anti-E-cadherin antibody (ab40772, Abcam) and mouse monoclonal anti-N-cadherin antibody ((#33-3900, Thermo Fisher Scientific) – were added to cell suspensions at a dilution of 1:200. Blocking and primary antibody staining took place simultaneously by incubation for 30 minutes at room temperature in the dark. Samples were then washed 3 times with 1X PBS and resuspended in 100 uL of 1X PBS. For secondary staining – using AlexaFluor 488-conjugated goat anti-rabbit secondary antibody (A-11034, ThermoFisher Scientific) and AlexaFluor 594-conjugated goat anti-mouse secondary antibody (A-11032, ThermoFisher Scientific) – antibodies were added at a dilution of 1:200 and incubated for 35 minutes at room temperature in the dark. Samples were washed 3 more times before flow cytometric analysis. 50,000 events were collected per sample using a BD LSR Fortessa X20, with whole cells gated according to forward scatter and side scatter (approx. 30,000 cells per sample). Fluorescence channel gates were made according to the fluorescence intensity from unstained samples.
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