Cells Under Stress: An Inertial-Shear Microfluidic Determination of Cell Behavior

Fern J. Armistead,1 Julia Gala De Pablo,1 Hermes Gadêlha,2 Sally A. Peyman,1 and Stephen D. Evans1,*

1Molecular and Nanoscale Physics Group, Department of Physics and Astronomy, University of Leeds, Leeds, United Kingdom and 2Department of Mathematics, University of York, York, United Kingdom

ABSTRACT The deformability of a cell is the direct result of a complex interplay between the different constituent elements at the subcellular level, coupling a wide range of mechanical responses at different length scales. Changes to the structure of these components can also alter cell phenotype, which points to the critical importance of cell mechanoresponse for diagnostic applications. The response to mechanical stress depends strongly on the forces experienced by the cell. Here, we use cell deformability in both shear-dominant and inertia-dominant microfluidic flow regimes to probe different aspects of the cell structure. In the inertial regime, we follow cellular response from (visco-)elastic through plastic deformation to cell structural failure and show a significant drop in cell viability for shear stresses >11.8 kN/m². Comparatively, a shear-dominant regime requires lower applied stresses to achieve higher cell strains. From this regime, deformation traces as a function of time contain a rich source of information including maximal strain, elastic modulus, and cell relaxation times and thus provide a number of markers for distinguishing cell types and potential disease progression. These results emphasize the benefit of multiple parameter determination for improving detection and will ultimately lead to improved accuracy for diagnosis. We present results for leukemia cells (HL60) as a model circulatory cell as well as for a colorectal cancer cell line, SW480, derived from primary adenocarcinoma (Dukes stage B). SW480 were also treated with the actin-disrupting drug latrunculin A to test the sensitivity of flow regimes to the cytoskeleton. We show that the shear regime is more sensitive to cytoskeletal changes and that large strains in the inertial regime cannot resolve changes to the actin cytoskeleton.

INTRODUCTION

Cell deformability is linked to the structure and mechanical properties of its biological constituents, which includes the cytoskeleton, nucleus, and cytoplasm. Disease-induced changes to the cytoskeleton can alter many cellular processes, and cell mechanoresponse is a key biophysical indicator of these changes (1), with distinct mechanical responses being correlated to many diseases (2–5). In recent years, several techniques have been developed to measure cell deformability; these include atomic force microscopy (AFM) (6), optical stretching (7), magnetic twisting cytometry (8,9), micropipette aspiration (3,10), and microfluidics.

Different methods measure localized or whole-cell deformation as well as deforming over different timescales, resulting in widely varying mechanical properties being reported (11,12). Several of these techniques also have limited throughput because the preselection of each single cell is required (6,7,9,10). Mechanical properties can differ on a cell-by-cell basis depending on a cell’s physiological state and its stage in the cell-division cycle (13). To determine the heterogeneity of a sample and accurately compare the deformability of different cell types, a high-throughput approach is therefore needed for collection of statistically relevant deformation events. This problem motivated the development of several microfluidics-based methods, which display high-throughput potential (N > 1000) and require a small sample volume (13–17).

The mechanical response of cells is affected by the magnitude of the force, the timescale over which the force is applied, and the method of investigation. Previous studies have shown that different microfluidic flow regimes can alter the mechanical response of cells. Gossett et al. (14,17,18) developed the technique deformability cytometry (DC) (19,20) in which cells are hydrodynamically stretched at the stagnation point (SP) of an extensional flow device at rates up to 2000 cells/s. A compressional force (FC) due to fluid inertia and a shear force (FS) due to fluid viscosity act on the cells. The FC contribution was estimated to be ~1000 times greater than FS, resulting in an inertia-dominant flow regime and high Reynold’s number (Re ∼ 1) (14). Their
study showed increased deformability in lymphocyte activation and stem cell pluripotency; these states are characterized by loose, open chromatin structures (18). However, treatment with several cytoskeletal-altering drugs showed negligible changes to cell deformability (14,21). Comparatively, Guillou et al. (22) also used an extensional flow device for single cell deformation but in a regime dominated by shear forces \((Re \ll 1)\). They utilized a high-shear, low-velocity, lower-strain regime compared to DC. Here, cells were treated with the actin-disrupting drug cytochalasin D, and they saw an increase in deformability.

Otto et al. (13,23–26) developed real-time DC (RT-DC), which passes cells through a channel slightly larger than the cell, where the strong velocity gradient in the channel causes deformation. RT-DC is also dominated by shear forces \((Re \ll 1)\). RT-DC is able to detect deformability changes in cells treated with various cytoskeleton-altering drugs but was not sensitive to changes to the nuclear structure.

It is clear that the sensitivity of deformation cytometry techniques is highly dependent on the flow regime as well as the device geometry, strain, and strain rates applied to the cells. Previous works remain in either a purely shear-dominant or inertia-dominant regime for all studies, and cell deformation is often probed over a small range of flow rates (22,27–30). Here, we deform from low to high strains in both flow regimes, using a single device geometry (Fig. 1a), bridging a critical gap between distant mechanoresponses of the cell.

Microfluidic deformation assays were performed to phenotype two different cell lines. HL60 is a circulating leukemia cell line expected to exhibit a more deformable response compared to SW480 cells, which originate from a solid colorectal cancer tumor. SW480 cells were also treated with an actin-cytoskeleton-disrupting drug, latrunculin A (LatA; Cayman Chemical, Ann Arbor, MI), to determine the sensitivity of the different flow regimes to changes in the actin cytoskeleton. By studying both regimes, we show that specific flow conditions probe different aspects of the cell structure, demonstrating that a shear-dominant and low-strain regime is most sensitive to cytoskeletal changes. Additionally, we found that in the inertial regime, we can achieve a high-strain response resulting in cytoskeletal fluidization and ultimately to failure in the structural integrity of the cell. However, changes caused by LatA could not be resolved in this regime. Viability studies show that cells can remain viable post deformation below the “failure point,” meaning the cells could be mechanically phenotyped and continue to be studied.

We also considered which deformation parameters have potential as biophysical markers of the cell’s mechanical phenotype. By tracking the deformation and relaxation of the cells, multiple characteristic parameters were extracted, including strain \(\varepsilon\), cell velocity profiles, and deformation and relaxation times. The Kelvin-Voigt model was also used to extract an elastic modulus, \(E\), for each cell type, giving us an intrinsic mechanical parameter comparable to previous works using AFM (23,31). Results verified that HL60 are significantly softer than SW480 and that treatment with LatA also reduced the stiffness of SW480. Interestingly, the determination of the different cell types based on relaxation time had the lowest associated error compared with the strain and elastic modulus. These results show the potential of relaxation time as a biophysical marker for mechanical phenotyping and that multiparameter analysis is vital for furthering understanding of cell mechanics.

**MATERIALS AND METHODS**

**Microfluidic devices**

Microfluidic devices were fabricated in polydimethylsiloxane (PDMS) using a silicon master as a mold. A silicon wafer (3 inches) was cleaned using
piranha wet etch (using H$_2$SO$_4$ and H$_2$O$_2$) and then rinsed with deionized water. The 25-μm photore sist layer of SU-8 2025 (Microchem, Warwickshire, UK) was applied to the wafer. Direct-write laser lithography was used to etch the channel designs into the SU-8 using a laser of wavelength 375 nm (MicroWriter ML; Durham Magneto-Optics, Durham, UK).

A 1:10 ratio of PDMS base and a cross-linking agent (Sylgard 184) were poured onto the master creating a negative replica of the SU-8 structures in PDMS, which was cured in an oven at 75°C for ~1 h, becoming a hydrophobic elastomer. The PDMS layer was then peeled away from the master, and the fluid inlet and outlet access holes are punched using a biopsy puncher. The PDMS was sealed to a glass slide using oxygen plasma treat ment. The channel dimensions at the cross-flow junction had a width of 35 μm and a height of 25 μm.

Characterizing flow regime

Cells were deformed at the SP of the extensional flow, defining the deformability using the deformation index $Dl = H/W$, where $H$ is the height of the cell and $W$ is the width of the cell (Fig. 1 b). The forces acting on a cell can be estimated from the shear and compressive components $F_s$ and $F_c$. The compressive force $F_c$ was determined from Eq. 1, where $\rho$ is the density of the suspension media, $U$ is the fluid velocity, $A_p$ is the cross-sectional area of the cell. The drag coefficient, $C_D$, is highly dependent on the Reynolds number, $Re$. The calculation of $C_D$ is detailed in Supporting Materials and Methods (32,33). The shear force $F_s$ was determined from Eq. 2, where $\mu$ is the viscosity of the suspension media, $r$ is the cell radius, and $\gamma$ is the shear strain rate (14,22). Fig. S1 shows how flow rate and viscosity of the suspension medium can be adjusted to achieve a shear-dominant or inertia-dominant regime. For a solution with a viscosity of 1 centipoise (cP), $Re > 40$ for flow rates $\geq 11$ μL/min. Given that inertial effects start for $Re$ above 20–40, we use $Re = 40$ as the boundary for the inertial regime (17,34,35). For $\mu = 33$ cP, the Reynolds number is low ($Re < 6$) and for the entire range of flow rates used in this body of work, which we define as the shear regime.

Fig. S2 further describes the dependence of $F_s$ and $F_c$ as a function of flow rate and the Reynolds number for $\mu = 1$ cP and $\mu = 33$ cP, where the total force $F_T$ is the sum of the two force components, $F_T = F_s + F_c$. $F_c$ increases with density, whereas $F_s$ increases with viscosity. Adding methylcellulose to the suspension buffers led to only a small increase in density but a significant increase in viscosity, resulting in $F_s$ being dominated by $F_s$. However, $F_c$ increases as $U^2$ compared to $U$ for $F_s$. Thus, $F_T$ is dominated by $F_c$ at low viscosity and high flow rates.

$$F_c = \frac{1}{2} \rho U^3 C_D A_p$$

$$F_s = \gamma \mu (4\pi r^3) = 2\pi U \mu r.$$  

Experimental procedure

The microfluidic device was mounted above an inverted brightfield microscope (Eclipse Ti-U, Nikon, Tokyo, Japan) with a 10× objective to use for cell deformation events with an additional 1.5× magnification for flow rates ($Q < 100 \mu$L/min). A high-speed camera (Photron, Tokyo, Japan) at a frame rate of 7500–260,000 fps and exposure time of 0.37–6.67 μs was used to capture cell deformation events. An external light source was mounted over the setup to capture images at higher frame rates and reduce exposure times to prevent motion blurring.

Automated image analysis was performed offline using ImageJ and MATLAB, with the position and shape of each cell event tracked and parameters such as initial size, velocity, circularity, and maximal deformation index ($DI = H/W$) extracted. This precision tracking used a mathematical image processing algorithm adapted from flagellar image tracking (36). Cells that did not travel centrally down the inlet channel and did not deform at the SP were excluded to calculate the average DI of a sample, ensuring all included events experience the same stress during deformation. Methods for calculating DI are described in the Supporting Materials and Methods, including Fig. S3.

Calculation of cell elastic modulus

A Kelvin-Voigt model was fitted to the time-dependent deformation of cells to determine the elastic modulus. This model comprises an elastic element (linear spring) and a viscous element (dashpot) arranged in parallel.

Equation 4 shows the variation of strain rate $t(t)$ as a function of applied stress $\sigma(t)$, where $E$ is the elastic modulus associated with the linear spring and $\eta$ is the viscosity associated with the dashpot. In the cross-flow, the stress increases from zero to a maximum of $\sigma_0$ as the cell enters the extensional flow junction and reaches the SP. Fig. S4 shows the velocity profile calculated along the central axis within the cross-flow section of the device. This suggests that $\sigma(t)$ varies approximately as a sine function, for a period $T$. Equation 5 shows $\sigma(t)$, where $\omega = 2\pi/T$, and is used to solve Eq. 4. The analytical solution is shown by Eq. 6, from which the elastic modulus can be directly extracted from the cell deformation dynamics, discussed below.

$$\dot{\epsilon}(t) = \frac{1}{\eta} (\sigma(t) - E\epsilon(t))$$

$$\sigma(t) = \sigma_0 (1 + \sin(\omega t))$$

$$\epsilon(t) = \frac{\sigma_0}{(\eta^2 \omega^2 + E^2)} \left( (\eta^2 \omega^2 - E\eta \omega + E^2)e^{-\frac{E \omega t}{\eta}} - E \eta \omega \cos(\omega t) + \omega^2 \eta^2 + E^2 \sin(\omega t) + E^2 \right).$$  

Cell culture

The HL60 cell line was purchased as a frozen stock (ECACC 98070106) and cultured in Roswell Park Memorial Institute growth media supplemented with 10% fetal bovine serum (Sigma, Welwyn Garden City, UK), 2 mM GlutaMax (Thermo Fisher Scientific, Waltham, MA), and penicillin (10 units/mL) and streptomycin (100 μg/mL) (Sigma). HL60 cells are a nonadherent cell line. Centrifuging at 100 × g for 4 min was sufficient to visibly pellet the cells, which were then gently resuspended in the desired suspension medium. Cells were either suspended in Roswell Park Memorial Institute media or resuspended in phosphate-buffered saline (PBS) with 0.24, 0.35, or 0.50% (w/v) methylcellulose (Sigma) to increase viscosity. The viscosity of the cell suspension mediums was measured using a Rheometrics SR-500 Dynamic Stress Rheometer in the parallel plate configuration with a diameter of 25 mm.

The SW480 cell line was provided by St James’s University Hospital and cultured in Dulbecco’s Modified Eagle Medium (DMEMF-12; Gibco, Rockville, Maryland) supplemented with 10% fetal bovine serum, 2 mM GlutaMax, and penicillin (100 units/mL) and streptomycin (100 μg/mL). Passage numbers were below 50 for all experiments. SW480 is an adherent cell line and was detached by incubating in TrypLE (Thermo Fisher Scientific) for 5 min. The cells were then centrifugated at 100 × g for 4 min and then resuspended in PBS with 0.5% (w/v) methylcellulose (Sigma).

Drug treatment

SW480 cells were detached by incubation with TrypLE for 5 min, then resuspended in DMEM with varying concentrations of LatA for 2 h. Concentrations of 0.01, 0.1, and 1 μM were compared to a control. Confocal
fluorescence of live cells was used to visualize the effect of LatA on the actin structure of SW480 (Fig. S5). The images show that the actin cortex underwent disruption with increased concentration of LatA because of inhibition of actin polymerization (37).

For subsequent cell deformation experiments, a concentration of 1 μM of LatA was used. Detached cells were incubated with LatA for 2 h before performing microfluidic deformation experiments. Cells were deformed while suspended in DMEM or by resuspension in 0.5% PBS methyl cellulose buffer while maintaining a constant concentration of the drug throughout the measurement period.

RESULTS

Cell deformability in shear and inertial regimes

High-speed imaging was used to capture the maximal deformation of HL60 cells at or near to the SP for a range of flow rates, Q. This was repeated using cell suspension media of increasing viscosity, ranging from 1 to 33 cP, where the 1 cP data set represents an inertia-dominant flow regime, and the 11–33 cP data sets represent an increasingly shear-dominant regime.

Fig. 2 shows deformation in the inertia-dominant regime. For $Q \leq 400 \ \mu L/min$, the DI tended toward a plateau, $DI_{max} = 1.70 \pm 0.13$. For $Q > 400 \ \mu L/min$, the DI further increased nonlinearly until ~600 $\mu L/min$, beyond which cells rupture and visibly break apart in the cross-flow junction, and deformation could not be measured. The critical deformation $DI_{crit}$ before cell rupture was found to be $DI_{crit} = 2.84 \pm 0.27$. We can define the stress corresponding to 400 $\mu L/min$ as the yield stress of the cell and 600 $\mu L/min$ as the failure point of the cell. Videos S1, S2, and S3 show example cell deformations occurring at flow rates corresponding below the yield stress at the yield stress and at the failure point. The deformation regime for $Q > 400 \ \mu L/min$ is associated with the breakdown of the cells internal structure (i.e., actin filament breakup) (38).

Trypan blue staining was used to measure the viability of HL60 cells deformed at various flow rates. The results (Fig. S6) show that cell viability was within the error of the undeformed control for $Q \leq 600 \ \mu L/min$. However, for $Q > 600 \ \mu L/min$, the viability dropped to <50%. This substantial drop in cell viability occurs at the failure point where cell rupture occurred on-chip. Fig. S7 a shows images of samples collected after deformation at 600 and 800 $\mu L/min$; for 800 $\mu L/min$, there was a reduction the number of viable cells and an associated increase in the amount of debris compared to the control and the 600 $\mu L/min$ samples. Further, in this regime, the cell did not recover their shape, which was analyzed using the deviation from circularity $D_C$, defined as $D_C = (1 - c) = 1 - 2\sqrt{\pi A/\ell}$, where $\ell$ is the cell perimeter and $c$ is the circularity; a perfect circle would have $D_C = 0$. Fig. S7 b shows low $D_C$ for the control cells (undeformed) and cells deformed at 600 $\mu L/min$. For the 800 $\mu L/min$ values, there is a general increase in the scatter of $D_C$ values resulting from cell debris caused by cell destruction at large flow rates. These results suggest high deformation can be achieved for 400 $\mu L/min < Q < 600 \ \mu L/min$ without adversely affecting cell viability.

Fig. 3 a shows DI as a function of $Q$ for solutions of increasing viscosity. For each viscosity, the DI was found to increase asymptotically toward a maximal deformation value $DI_{max}$, which was determined by fitting an exponential function (see Supporting Materials and Methods). Fig. 3 b shows images of cells deformed at $DI_{max}$ under each flow condition. Each image is accompanied by a superimposed color contour plot, which shows how the deformation varies

![Figure 2](image1.png)

FIGURE 2 Deformation index, DI ± standard error of HL60 cells versus flow rate, at $\mu = 1$ cP. DI ± standard error was averaged from multiple cell events combined from $N = 3$ repeats; each data point includes $30 > n > 500$ cell events. For $Q < 400 \ \mu L/min$, deformation can be fitted by an exponential, which tends toward a maximal deformation of $DI_{max}$. To see this figure in color, go online.

![Figure 3](image2.png)

FIGURE 3 (a) Deformation index versus $Q$ for HL60 cells in four different media with viscosity changing between 1 and 33 cP. DI ± standard error was averaged from multiple cell events combined from $N = 3$ repeats; each data point includes $30 > n > 500$ cell events. The data is fitted with an exponential. (b) Images of a cell deformation for each flow condition where $DI = DI_{max}$ are shown. They are accompanied by superimposed color contour plots that show how the deformation changes as a function of time. To see this figure in color, go online.
as a function of time, going from blue, where the cell approaches the cross-flow junction, to red, where the cell is deformed at the SP. At higher viscosities, the back pressure in the channels increased, reducing the upper limit of $Q$ achievable without device failure. Despite the limitations in terms of maximal attainable flow rates for the higher viscosity solutions, it is evident that significantly larger deformations can be achieved compared to the inertia-dominant regime. For example, $DI_{\text{max}} = 2.35$ for flow rates less than 100 $\mu$L/min at 33 cP.

Fig. S8 a shows the $DI$ data sets plotted as a function of force, $F_T$, rather than $Q$. In general, this shows that for the same applied force, the cells were more deformable in a shear-dominant regime. However, this is only true below the previously determined yield stress in the inertia-dominant regime; above this, the $DI$ in the inertia-dominant regime exceeds the shear-dominant regime. Additionally, the limiting deformation $DI_{\text{max}}$ varied linearly with viscosity (Fig. S8 b).

In addition to measuring $DI$, we also plotted strain (defined as $\epsilon = \frac{H - H_0}{H}$) versus time to determine the deformation and relaxation times as well as allowing the application of simple models to extract parameters such as elastic modulus $E$.

Fig. 4 a shows the average strain of 50 HL60 cells as a function of time, deformed at 5 $\mu$L/min in a shear-dominant regime ($\mu = 33.4 \pm 0.3$ cP) (Video S6). A shear-dominant and low-velocity flow condition was chosen to reduce the frame rate (7000 fps) required for tracking and maximize the field of view available. The sign of the strain value describes the cell direction, which changes as the cell moves from the inlet to the outlet, and the magnitude describes the amount of strain. As cells traversed from the inlet to the SP, the strain increased; this was fitted with an exponential with an associated deformation time $\tau_d$.

Further, it continued to increase as the cell moved from the SP toward the outlet, reaching a maximal strain of $\epsilon_{\text{max}} = 0.18 \pm 0.04$ (Fig. 4 b). The strain then decreased exponentially as the cells traveled toward the outlet and was fitted to an exponential associated with a relaxation time $\tau_r$. Additionally, the initial strain (before entering the SP) $\epsilon_0$ was found and compared to the final strain value $\epsilon_{\infty}$, which was found by extrapolation of the exponential fit of the relaxation.

Fig. S9 shows the cell strain and velocity as a function of time. The velocity profile can be roughly approximated to a single period of a sine wave, shown in red in Fig. S9 a. The minimum in the velocity profile occurs when the cell is closest to the SP. The sine-oscillating Kelvin-Voigt model (Eq. 5) can then be used to fit the strain trace, shown in red in Fig. S9 b.

By equating to the sine-oscillating Kelvin-Voigt model (described in Materials and Methods) the elastic modulus of HL60 was found to be $E = (0.30 \pm 0.03)$ kPa. To demonstrate the potential of this additional parameterization of cell deformation for classifying cell types and understanding disease states, we compared HL60 cells from the circulating leukemia cell line to SW480 cells originating from a primary adenocarcinoma, Dukes stage B.

Fig. 5 a shows the $DI$ of the cells for a range flow rates up to $\sim 100$ $\mu$L/min in a shear-dominant regime ($\mu = 33$ cP; Video S4). Firstly, we note that the HL60 cells have significantly a higher $DI$ for all flow rates compared to the SW480. Secondly, we observed that treatment of the SW480 cells with the known actin disruptor LatA led to an increase in $DI$ at low flow rates (Fig. S5 shows fluorescence images of LatA-induced disruption of the actin cytoskeleton) (Video S5). In contrast, the $DI$ in the inertial regime (Fig. 5 b) increases approximately linearly for $Q < 400$ $\mu$L/min, with the HL60 cells being slightly more deformable than the SW480 cells. However, on treatment with LatA, the SW480 cells become softer and are similar to the HL60 cells. At $\sim 400$ $\mu$L/min, the point at which the actin scaffold undergoes significant disruption, there is a change in slope for the HL60 and untreated SW480 cells, whereas the $DI$ for the LatA-treated cells continues undeviated. For $Q > 400$ $\mu$L/min the LatA-treated cells and untreated SW480 have comparable values of $DI$.

In the shear-dominant regime, the increase in $DI$ of SW480 treated with LatA compared to control cells was more prominent at low flow rates ($Q < 40 \mu$L/min), with the behavior at higher flow rates asymptotically tending toward untreated behavior. Comparatively, the increase in $DI$ of HL60 compared to SW480 increases with flow rate and is more prominent at high flow rates ($Q > 40 \mu$L/min). In the inertia-dominant regime, there is a small increase in $DI$ of SW480 treated with LatA compared to control cells for $Q < 400 \mu$L/min. The treated and untreated cells are indistinguishable above this flow rate. Contrasting this, HL60 and SW480 have comparable $DI$ for $Q < 400 \mu$L/min; above
this flow rate, the DI of HL60 is significantly higher. Fig. S10 uses the DI ratio to show these behaviors explicitly.

Deformation traces for SW480 control and SW480 treated with LatA are shown in Fig. 6, where cells were deformed in a shear-dominant regime \((\mu = 33.450.3 \text{ cP})\) at 5 \(\mu\)L/min. Additionally, the velocity and strain profiles are shown in Fig. S11 and were used to fit the Kelvin-Voigt model. The deformation traces show distinct differences, namely, the derived values in the deformation and relaxation times; strain and elastic modulus \(E\) are shown in Table 1.

**DISCUSSION**

In the inertia-dominant regime, we identified the point of inflection in the DI versus \(Q\) data (Fig. 2) \((Q > \sim 400 \mu\)L/min) with the yield stress for HL60 cells (equating to a force of \(\sim 0.58 \mu\)N). Below the yield stress, the cells undergo modest changes in deformation, whereas, above the yield stress, the microstructure associated with the actin cytoskeleton breaks down leading to increased deformation (18). Cells that were deformed in this regime are able to recover their original shape and remained viable post-deformation (Fig. S6). However, if cells were deformed beyond the failure point (Fig. 2) \((Q > 600 \mu\)L/min or \(DI > 2.84)\), then they suffered a significant drop in viability and often did not recover to their original shape post-deformation (Fig. S7). The failure point represents a limit below which live cells can be mechanically phenotyped and collected for further studies, such as chemical phenotyping via Raman spectroscopy (39). For flow rates between 50 and 400 \(\mu\)L/min, only modest changes in DI were observed for HL60 cells \((1.37–1.70)\) and similarly for SW480 \((1.23–1.47)\). This regime provides only weak sensitivity to changes in the actin cytoskeleton, as demonstrated by the treatment of

![FIGURE 5](image-url) (a) The DI as a function of flow rate, \(Q\), of HL60 cells, SW480 cells, and SW480 cells treated with 1 \(\mu\)M of LatA. The flow regime was shear dominant, and the viscosity of the cell suspension buffer was 33 cP. (b) DI versus flow rate \(Q\) for HL60 cells, SW480 cells, and SW480 cells treated with 1 \(\mu\)M of LatA. The flow regime was inertia dominant, the viscosity of the cell suspension buffer was \(\sim 1 \text{ cP}\). To see this figure in color, go online.

![FIGURE 6](image-url) Strain \(\varepsilon\) was tracked for SW480 \((N = 56)\) and SW480 treated with LatA \((N = 30)\) as a function of time, with the standard error shown. \(Q\) was fixed at 5 \(\mu\)L/min, and the suspension medium viscosity was 33 cP. The dashed lines represent the extrapolated final strain \(\varepsilon_\text{N}\) for both samples. To see this figure in color, go online.

**TABLE 1** Multiple Characteristic Parameters Extracted from the Deformation Traces of HL60, SW480, and SW480 LatA Cells, Deforming at the Stagnation Point of an Extensional Flow at 5 \(\mu\)L/min in the Shear Regime

|                | HL60 \((N = 50)\) | SW480 \((N = 56)\) | SW480 LatA \((N = 30)\) |
|----------------|-------------------|-------------------|---------------------|
| \(A\) (\(\mu\)m) | 12.3 ± 0.3        | 15.1 ± 0.2        | 15.4 ± 0.1          |
| \(\varepsilon_\text{max}\) | 0.18 ± 0.01      | 0.08 ± 0.01       | 0.11 ± 0.01         |
| \(\tau_\text{f}(\text{ms})\) | 3.52 ± 0.14     | 1.36 ± 0.06       | 0.67 ± 0.09         |
| \(\tau_\text{d}(\text{ms})\) | 1.04 ± 0.05      | 1.19 ± 0.20       | 0.78 ± 0.24         |
| \(E(\text{Pa})\) | 301 ± 29         | 542 ± 66          | 419 ± 54            |
| \(\varepsilon_0\) | –0.012 ± 0.004   | –0.012 ± 0.006    | –0.007 ± 0.014      |
| \(\varepsilon_\text{N}\) | +0.03 ± 0.009    | +0.010 ± 0.003    | +0.059 ± 0.001      |

Where \(A\) is the cell diameter, \(\varepsilon_\text{max}\) is the maximal strain, \(\tau_\text{f}\) is the relaxation time, \(\tau_\text{d}\) is the deformation time, \(E\) is the elastic modulus, \(\varepsilon_0\) is the magnitude of the initial strain, and \(\varepsilon_\text{N}\) is the magnitude of the final strain.
the SW480 cells with LatA, after which their DI values increased slightly to be similar to those of the intrinsically softer HL60 cells (Fig. 5 b). The LatA-treated cells show no obvious change in behavior at \( Q \leq \sim 400 \mu L/min \), whereas the HL60 and untreated SW480 both show increase in gradient of DI with \( Q \). This observation supports the suggestion that DI in the inertial regime is relatively insensitive to the actin scaffold post-degradation of the actin scaffold.

For \( Q \leq 400 \mu L/min \), the HL60 cells show an increase in the slope of DI with flow rate compared to SW480 cells; this change is also characterized by the DI ratio (Fig. S10 b). This could be indicative of the smaller nuclear size in these cells (Figs. S12 and S13). The nuclear ratio \((A_{nucleus}/A_{cell})\) of HL60 is 0.55 \pm 0.02 and of SW480 is 0.72 \pm 0.01. The nucleus is known to be significantly stiffer than the surrounding cytoplasm, which is pervaded by the cytoskeleton (40). For example, Caille et al. (41) showed a 10-fold increase in the nucleus elastic moduli of endothelial cells compared to their cytoplasm.

Fig. 3 a showed that for HL60 cells, the DI tended toward a maximal value, DI_{max}, with flow rate. The value of DI_{max} increased linearly with solution viscosity, indicating that at higher viscosities, the same deformation is achieved at lower flow rates (Fig. S7 b). It also indicates that inertial and shear forces act very differently on a cell and that for the same magnitude of force, a significantly larger deformation is observed in the shear-dominant regime (\( \mu = 33 \) cP) (Fig. S8 a). Further, the cell shape at maximal deformation changes from elliptical at low viscosity to tear-shapes at higher viscosities, with the cell perimeter changing from convex to concave (Fig. 3 b). Deformation in a shear-dominant regime occurs at lower Re, and the tear-shape is indicative of the shear force being dominant, causing the pointed ends of the cell.

In the shear-dominant regime as well as in measuring the DI, we captured “deformation traces,” which show the variation of cell shape on approach to the SP and relaxation after passing through it (Fig. 4). By plotting these as strain versus time, we obtained a number of parameters related to the cellular state. In particular, the deformation and relaxation times, maximal strain, and through a fit of the Kelvin-Voigt model the elastic modulus. For the HL60 cell line, the elastic modulus, \( E \), was found to be 0.30 \pm 0.03 kPa, which falls between those previously reported for HL60 cells obtained from AFM (0.17 \pm 0.03 kPa) by Rosenbluth et al. (31) and RT-DC (1.48 \pm 0.03 kPa) by Mietke et al. (23). Although we would expect our value to be closer to that determined by RT-DC because of the similar timescales of measurement (\( \sim 1 \) ms), we note that RT-DC probes much lower strains and hence potentially different aspects of the cells.

Trikritis et al. (42) used AFM to determine the elastic modulus of SW480 to be 1.39 kPa. Comparatively, Palmieri et al. (43) noted that SW480 cells have two appearances in culture, an epithelial-type morphology and a rounded morphology. Using AFM, they found the elastic modulus of SW480 epithelial-type morphology to be 1.06 kPa and SW480 rounded-type morphology to be 0.58 kPa. The elastic modulus determined here for SW480 rounded-type-morphology cells was within error (i.e., the same as that found by Palmieri et al. (43)). After treatment with LatA, the elastic modulus was reduced to (420 \pm 54) Pa.

Table 1 provides a summary of the multiple parameters extracted from the deformation traces of the different cell types. The values for \( E \) and \( \varepsilon_{\text{max}} \) confirm that HL60 are significantly softer than SW480 and that treating SW480 with LatA reduced their stiffness. The deformation times \( \tau_d \) for the three cell types were all within error. However, the relaxation time \( \tau_r \) can be used to distinguish them from each other. HL60 had the largest relaxation time of \( \sim 3.5 \) ms (which is \( \sim 2.5 \) times longer than for SW480), an expected result considering these are the softest cells. However, the LatA-treated SW480 recover at a faster rate than untreated SW480 despite being more deformable.

The initial strain \( \varepsilon_0 \) was \( \sim 0 \) for each cell type. The extrapolated final strain \( \varepsilon_{\infty} \) was within error of \( \varepsilon_0 \) for the HL60 and SW480 cell lines \( (\varepsilon_0 = \varepsilon_{\infty}) \), whereas the LatA-treated cells showed a larger \( \varepsilon_{\infty} \) compared to untreated cells \( (\varepsilon_0 < \varepsilon_{\infty}) \). Actin disruption using LatA led to a shorter relaxation time being measured, tending to a nonzero strain. This possibly indicates that actin disruption leads to an additional, slower relaxation process occurring over a longer timescale and not fully recovered in our experiments.

**CONCLUSIONS**

Deformation of HL60 cells as a function of flow rate in the inertia-dominant and shear-dominant regimes show cell response is dependent upon the nature of the applied force (shear, compressive) and not simply the amplitude of the force. Cells appear stiffer in an inertial regime (low viscosity, high flow rate) compared to a shear regime (high viscosity, low flow rate). This behavior indicates that different deformation regimes are likely to be sensitive to different subcellular components. To explore this, we compared two different cell types. The HL60 cells are circulating leukemia cells that exist in the circulatory system, navigating the vasculature and capillary beds as isolated cells. In contrast, the SW480 cells are derived from a primary colorectal adenocarcinoma, a solid tumor. The inertial regime showed several distinct behaviors: at low flow rates, DI increased almost linearly with flow rate until the yield point was reached (\( Q = \sim 400 \mu L/min \)) at which point the actin scaffold undergoes significant disruption and possible fluidization (38). After this point, 400 \( \mu L/min < Q < 600 \mu L/min \) the HL60 become significantly more deformable, whereas the change for the SW480 cells is less significant. For these flow rates, the higher stiffness of the SW480 cells is attributed to the larger cell nucleus (and nuclear ratio), being more resistant to deformation. Flow rates above 600 \( \mu L/min \) led to irreversible cell damage with \( \sim 50% \) reduction in cell viability and poor shape recovery post deformation.
In the shear regime, larger $D$Is were attained with significantly lower flow rates; here, the fluid flow profile probes “stretching deformations” of the cell membrane and the cytoskeleton as opposed to “compressive” behavior of the cytosol and nucleus in the inertial regime. To probe this further, SW480 cells were treated with the actin-disrupting agent LatA. In the shear regime at low flow rates, the LatA-treated SW480 cells were significantly more deformable than the untreated cells; however, as the flow rate was increased, the difference was reduced (see Figs. 5a and S10a), with the SW480 cells approaching that of the actin disrupted cells, thus indicating that the cytoskeleton is probed at the lower flow rates.

As might be expected, the disruption of the actin scaffold using LatA in the inertial regime led to a softening of the cells such that for $Q < 400 \, \mu L/min$, the SW480-LatA cells were indistinguishable in their deformability from the HL60 cells. However, at $Q = 400 \, \mu L/min$ where the actin is disrupted, the HL60 cells become much softer, giving a change in gradient, with a $\Delta D/\Delta Q$ of $\sim 5.9 \times 10^{-3}$ compared with $\sim 1.1 \times 10^{-3}$ for the lower flow rates. The gradient for the untreated SW480 cells also changes abruptly at this point although with a lower slope. In contrast, the SW480-LatA shows no change in gradient.

In the shear-dominant (cytoskeletal sensitive) regime, we also measured deformation traces and determined multiple characteristic parameters, including maximal strain $\varepsilon_{\max}$, initial strain $\varepsilon_0$, final strain $\varepsilon_\infty$, elastic modulus $E$, and relaxation time $\tau_r$. Interestingly, the elastic modulus values of each cell line were of the same order of magnitude as previous AFM measurements despite the different modes and timescales of operation. The fast relaxation of the LatA-treated cells to a nonzero extrapolated final strain suggests that actin disruption causes additional relaxation processes on timescales not recovered in our experiments. Our results show that the multiple parameters have the accuracy to distinguish different cell types and that there is merit in measuring in the shear as well as the inertial regimes to characterize cell response to applied force. The microfluidic approach offers a high-throughput technique for the cell mechanophenotyping as well as increases the range of deformability ($1.3 < D/I < 2.8$) and strain rates ($10^5 - 10^7 \, Hz$) that can be achieved. The data used in the figures of this article will be available at https://doi.org/10.5518/397.

SUPPORTING MATERIAL

Supporting Materials and Methods, thirteen figures, and six videos are available at http://www.biophysj.org/biophys/journal/supplemental/S0006-3495(19)30104-3.

AUTHOR CONTRIBUTIONS

F.J.A. performed research, analyzed data, and wrote the manuscript. J.G.D.P. helped with culturing of cells and provided code for data analysis. H.G. also provided analytical tools. S.A.P. and S.D.E. helped to design the experimental plan, analyse data, and write the manuscript.

ACKNOWLEDGMENTS

We would like to acknowledge Matthew Hughes for making the viscosity measurements on bulk solutions. We also acknowledge Catriona Marshal (St James’s University Hospital) for providing the cell lines and short tandem repeat validation.

F.J.A. thanks the University of Leeds for financial support. S.D.E. acknowledges the following funders: Medical Research Council (MR/M009084/1) and Engineering and Physical Sciences Research Council (EP/P023266/1). S.D.E. is supported by the National Institute for Health Research infrastructure at Leeds. The views expressed are those of the author(s) and not necessarily those of the National Health Service, the National Institute for Health Research, or the Department of Health.

REFERENCES

1. Zheng, Y., J. Nguyen, ..., Y. Sun. 2013. Recent advances in microfluidic techniques for single-cell biophysical characterization. Lab Chip. 13:2464–2483.
2. Baskurt, O. K., D. Gelmont, and H. J. Meiselman. 1998. Red blood cell deformability in sepsis. Am. J. Respir. Crit. Care Med. 157:421–427.
3. Guo, Q., S. J. Reiling, ..., H. Ma. 2012. Microfluidic biomechanical assay for red blood cells parasitized by Plasmodium falciparum. Lab Chip. 12:1143–1150.
4. McMillan, D. E., N. G. Utterback, and J. L. Puma. 1978. Reduced erythrocyte deformability in diabetes. Diabetes. 27:895–901.
5. Stuart, J., and G. B. Nash. 1990. Red cell deformability and haematological disorders. Blood Rev. 4:141–147.
6. Cross, S. E., Y. S. Jin, ..., J. K. Gimzewski. 2007. Nanomechanical analysis of cells from cancer patients. Nat. Nanotechnol. 2:780–783.
7. Guck, J., R. Ananthakrishnan, ..., J. Käs. 2001. The optical stretcher: a novel laser tool to micromanipulate cells. Biophys. J. 81:767–784.
8. Puig-de-morales-marinkovic, M., K. T. Turner, ..., S. Suresh. 2007. Viscoelasticity of the human red blood cell. Am. J. Physiol. Physiol. 293:C597–C605.
9. Bausch, A. R., F. Ziemann, ..., E. Saxmann. 1998. Local measurements of viscoelastic parameters of adherent cell surfaces by magnetic bead microrheometry. Biophys. J. 75:2038–2049.
10. Hochmuth, R. M. 2000. Micropipette aspiration of living cells. J. Biomech. 33:15–22.
11. Mceendrarry, E., and A. R. Harris. 2014. Cell mechanics: principles, practices, and prospects. Wiley Interdiscip. Rev. Syst. Biol. Med. 6:371–388.
12. Suresh, S. 2007. Biomechanics and biophysics of cancer cells. Acta Biomater. 3:413–438.
13. Otto, O., P. Rosendahl, ..., J. Guck. 2015. Real-time deformability cytometry: on-the-fly cell mechanical phenotyping. Nat. Methods. 12:199–202, 4, 202.
14. Gossett, D. R., H. T. Tse, ..., D. Di Carlo. 2012. Hydrodynamic stretching of single cells for large population mechanical phenotyping. Proc. Natl. Acad. Sci. USA. 109:7630–7635.
15. Forsyth, A. M., J. Wan, ..., H. A. Stone. 2010. The dynamic behavior of chemically “stiffened” red blood cells in microchannel flows. Microvasc. Res. 80:37–43.
16. Faustino, V., D. Pinho, ..., R. Lima. 2014. Extensional flow-based microfluidic device: deformability assessment of red blood cells in contact with tumor cells. Biochip J. 8:42–47.
17. Dudani, J. S., D. R. Gossett, ..., D. Di Carlo. 2013. Pinched-flow hydrodynamic stretching of single-cells. Lab Chip. 13:3728–3734.
18. Tse, H. T., D. R. Gossett, ..., D. Di Carlo. 2013. Quantitative diagnosis of malignant pleural effusions by single-cell mechanophenotyping. Sci. Transl. Med. 5:212ra163.

19. Che, J., Y. Yu, ..., D. Di Carlo. 2017. Biophysical isolation and identification of circulating tumor cells. Lab Chip. 17:1452–1461.

20. Lin, J., D. Kim, ..., D. Di Carlo. 2017. High-throughput physical phenotyping of cell differentiation. Microsystems and Nanoeng. 3:17013.

21. Masaïli, M., D. Gupta, ..., D. Di Carlo. 2016. Multiparameter mechanical and morphometric screening of cells. Sci. Rep. 6:37863.

22. Guillou, L., J. B. Dahl, ..., S. J. Muller. 2016. Measuring cell viscoelastic properties using a microfluidic extensional flow device. Biophys. J. 111:2039–2050.

23. Mietke, A., O. Otto, ..., E. Fischer-Friedrich. 2015. Extracting cell stiffness from real-time deformability cytometry: theory and experiment. Biophys. J. 109:2023–2036.

24. Gollier, S., P. Rosendahl, ..., O. Otto. 2017. High-throughput cell mechanical phenotyping for label-free titration assays of cytoskeletal modifications. Cytoskeleton (Hoboken). 74:283–296.

25. Xavier, M., P. Rosendahl, ..., O. Otto. 2016. Mechanical phenotyping of primary human skeletal stem cells in heterogeneous populations by real-time deformability cytometry. Integr. Biol (Camb). 16:616–623.

26. Chan, C. J., A. E. Elkenyong, ..., O. Otto. 2015. Myosin II activity softens cells in suspension. Biophys. J. 108:1856–1869.

27. Bae, Y. B., H. K. Jang, ..., J. M. Kim. 2016. Microfluidic assessment of mechanical cell damage by extensional stress. Lab Chip. 16:96–103.

28. Cha, S., T. Shin, ..., J. M. Kim. 2012. Cell stretching measurement utilizing viscoelastic particle focusing. Anal. Chem. 84:10471–10477.

29. Reymond, L., E. D. Este, ..., K. Johnsson. 2014. Fluorogenic probes for live-cell imaging of the cytoskeleton. Nat. Methods. 11:731–733.

30. Deng, Y., S. P. Davis, ..., A. J. Chung. 2017. Inertial microfluidic cell stretcher (iMCS): fully automated, high-throughput, and near real-time cell mechanotyping. Small. 13:1700705.

31. Rosenbluth, M. J., W. A. Lam, and D. A. Fletcher. 2006. Force microscopy of nonadherent cells: a comparison of leukemia cell deformability. Biophys. J. 90:2994–3003.

32. Brown, P. P., and D. F. Lawler. 2003. Sphere drag and settling velocity revisited. J. Environ. Eng. 129:222–231.

33. Di Carlo, D., D. Irimia, ..., M. Toner. 2007. Continuous inertial focusing, ordering, and separation of particles in microchannels. Proc. Natl. Acad. Sci. USA. 104:18892–18897.

34. Haward, S. J., T. J. Ober, ..., G. H. McKinley. 2012. Extensional rheology and elastic instabilities of a wormlike micellar solution in a microfluidic cross-slot device. Soft Matter. 8:536–555.

35. Kim, J. M. 2015. Kinematic analyses of a cross-slot microchannel applicable to cell deformability measurement under inertial or viscoelastic flow. Korean J. Chem. Eng. 32:2406–2411.

36. Smith, D. J., E. A. Gaffney, ..., J. C. Kirkman-Brown. 2009. Bend propagation in the flagella of migrating human sperm, and Its modulation by viscosity. Cell Motil. Cytoskeleton. 66:220–236.

37. Brenner, S. L., and D. Korn. 1987. Inhibition of actin polymerization by latrunculin A. FEBS Lett. 2:316–318.

38. Janney, P. A., U. Euteneuer, ..., M. Schliwa. 1991. Viscoelastic properties of vimentin compared with other filamentous biopolymer networks. J. Cell Biol. 113:155–160.

39. De Pablo, J. G., M. Lones, ..., S. D. Evans. 2018. Biochemical fingerprint of colorectal cancer cell lines using label-free live single-cell Raman spectroscopy 49:1323–1332.

40. Lammerding, J. 2015. Mechanics of the nucleus. Compr. Physiol. 1:783–807.

41. Caille, N., O. Thoumine, ..., J. Meister. 2002. Contribution of the nucleus to the mechanical properties of endothelial cells. J. Biomech. 35:177–187.

42. Trikritis, D., S. Richmond, ..., A. Downes. 2015. Label-free identification and characterization of living human primary and secondary tumor cells. The Analyst. 140:5162–5168.

43. Palmieri, V., D. Lucchetti, ..., M. De Spirito. 2015. Mechanical and structural comparison between primary tumor and lymph node metastasis cells in colorectal cancer. Soft Matter. 11:5719–5726.