Unbiased high precision cell mechanical measurements with microconstrictions

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Abstract

We describe a quantitative, high-precision, high-throughput method for measuring the mechanical properties of cells in suspension with a microfluidic device, and for relating cell mechanical responses to protein expression levels. Using a high-speed (750 fps) CCD camera, we measure the driving pressure $\Delta p$, maximum cell deformation $\varepsilon_{\text{max}}$, and entry time $t_{\text{entry}}$ of cells in an array of microconstrictions. From these measurements, we estimate population averages of elastic modulus $E$ and fluidity $\beta$ (the power-law exponent of the cell deformation in response to a step change in pressure). We find that cell elasticity increases with increasing strain $\varepsilon_{\text{max}}$ according to $E \sim \varepsilon_{\text{max}}$, and with increasing pressure according to $E \sim \Delta p$. Variable cell stress due to driving pressure fluctuations and variable cell strain due to cell size fluctuations therefore cause significant variability between measurements. To reduce measurement variability, we use a histogram matching method that selects and analyzes only those cells from different measurements that have experienced the same pressure and strain. With this method, we investigate the influence of measurement parameters on the resulting cell elastic modulus and fluidity. We find a small but significant softening of cells with increasing time after cell harvesting. Cells harvested from confluent cultures are softer compared to cells harvested from sub-confluent cultures. Moreover, cell elastic modulus increases with decreasing concentration of the adhesion-reducing surfactant pluronic. Lastly, we simultaneously measure cell mechanics and fluorescence signals of cells that overexpress the GFP-tagged nuclear envelope protein lamin A. We find a dose-dependent increase in cell elastic modulus and decrease in cell fluidity with increasing lamin A levels. Together, our findings demonstrate that histogram matching of pressure, strain and protein expression levels greatly reduces the variability between measurements and enables us to reproducibly detect small differences in cell mechanics.
Introduction

Cell mechanical properties have been implicated in a wide range of physiological processes and diseases. For example, cell mechanical properties change systematically in malaria (1, 2), asthma (3), inflammation (4) and cancer (1, 5). The need to measure cell mechanical properties in a high-throughput, quantitative way led to the recent development of various microfluidic techniques, such as the optical stretcher (6), hydrodynamic stretching (7), real-time deformability cytometry (8), microfiltration (9), and microconstriction arrays (10–12). Compared to established low-throughput methods such as atomic force microscopy (13) or optical tweezers (14), these microfluidic systems are easy to implement and, apart from a simple microfluidic chip, do not require expensive specialized equipment other than a microscope, a high-speed camera, and customized image analysis software.

We have recently described a microfluidic method for high-throughput quantitative measurement of cell mechanical properties of suspended cells (15). In this method, we pump cells through an array of 8 parallel microconstrictions and measure the pressure drop $\Delta p$ across the constriction, the cell’s maximum strain $\varepsilon_{\text{max}}$ and the entry time $t_{\text{entry}}$ of the cells into the constriction. The entry time is the time from the moment the cell is lodged in front of a constriction to the moment the cell has been deformed to the width of the constriction. This is also the moment at which the cell exits the constriction, since the length of the constriction channel in our setup is short compared to the cell diameter, and the travel time through the constriction channel is therefore negligible and below the time resolution of our camera (< 1.4 ms).

We have established (15) that pressure, strain and entry time are related through a power-law according to 

$$E \frac{\varepsilon(t)}{\Delta p} = \left( \frac{t}{t_0} \right)^\beta.$$ 

The parameter $E$ describes the elastic modulus of the cells, the power-law exponent $\beta$ describes the cells’ fluidity, and the reference time $t_0$ is set to 1 s (16). Rearranging and considering that $t_{\text{entry}}$ is the time when the cell strain $\varepsilon(t)$ has reached $\varepsilon_{\text{max}}$ gives

$$t_{\text{entry}} = \left( \frac{E \varepsilon_{\text{max}}}{\Delta p} \right)^{1/\beta} \quad \text{(Eq. 1)}.$$ 

A power-law exponent of $\beta = 0$ indicates a purely elastic solid, described by Hooke’s law, in which case there is no time dependency of the creep response, i.e. $\varepsilon(t) = \Delta p/E$. If $\Delta p/E > \varepsilon_{\text{max}}$ the cell entry time is zero, otherwise it is infinite. For $\beta = 1$, Eq. 1 resembles Newton’s law of viscous deformations. Typical values of $\beta$ for cells are in the range of 0.1-0.4, indicating viscoelastic behavior (16, 17). Both $E$ and $\beta$ are strongly influenced by the cytoskeleton (actin, microtubules), but also by cell nuclear properties including chromatin condensation and expression levels of nuclear lamina intermediate filaments.
Moreover, $E$ and $\beta$ in cells after pharmacological treatment are not independent from each other but scale according to predictions from the theory of soft glassy rheology (13, 15–17).

Eq. 1 assumes that the elastic and dissipative cell mechanical properties are independent of the applied pressure and the maximum strain. However, previous reports have established that cell mechanical properties can be stress and strain sensitive (18–21). Since the applied pressure drop across the microconstrictions in our device can vary during a measurement due to changes in the occupancy of the channel array, due to accumulation of cell debris in the filter system, and due to user adjustments, and since the maximum cell strain also varies from cell to cell due to variable cell diameters, the measured cell mechanical parameters $E$ and $\beta$ can be subject to a high degree of variability.

In this study, we investigate the influence of stress and strain stiffening and explore how Eq. 1 can be extended to account for these effects. We then describe a method for canceling stress or strain stiffening effects when comparing different cell populations. We achieve this by histogram matching, whereby only those cells from two (or more) measurements are included in the analysis that have experienced the same pressure and the same maximum strain. Moreover, we investigate how cell mechanics is influenced by subtle details of measurement and cell culture conditions, such as cell confluency prior to harvesting, the time since cell harvesting, the choice of the cell suspension medium, or device coating with adhesion-preventing pluronic surfactant. Finally, we explore the effect of protein expression levels in a mixed cell population on the measurement results. Specifically, we transfect cells with a lamin A-green fluorescent protein (GFP) construct and observe them with combined bright-field and fluorescence imaging in our microfluidic device. We then correlate differences in the mechanical properties of individual cells with differences in lamin A-GFP expression levels.

Our results establish that histogram matching of pressure, strain and protein expression levels greatly reduces the variability between measurements and enabled us to reproducibly measure small differences in cell mechanical properties between different groups of cells.
Methods

Cell culture

K562 leukemia cells (No. CCL-243; American Type Culture Collection, Manassas, VA (ATTC)) are cultured at 37°C and 5 % CO₂ in Iscove’s Modified Dulbecco’s Medium (IMDM, Gibco, Cat. No. 12440053) containing 10 % FCS (fetal calf serum, Gibco, Cat. No. 16000036) and 1 % Penicillin-Streptomycin-Glutamine (PSG, Gibco, Cat. No. 10378016). K562 lamin A-overexpressing cells are transfected as described in (15). DLD-1 pMCV colon carcinoma cells are a kind gift of Michael Stürzl (Division of Molecular and Experimental Surgery, University Clinics Erlangen) and are cultured in RPMI Medium (Gibco, Cat. No. 11875093), containing 10 % FCS, 1 % PSG and 1 % G418 (Gibco, Cat. No. 11811098). NIH 3T3 mouse embryonic fibroblast cells (No. CRL-1658; ATCC) are cultured in Dulbecco’s Modified Eagle Medium (DMEM, Gibco, Cat. No. 11885084), containing 10 % FCS and 1 % PSG. Cells are passaged every third day. Actin depolymerization is performed with cytochalasin D (cytoD; Sigma-Aldrich, St. Louis, MO; Cat. No. C8273) at a concentration of 10 µM for a duration of 30 min before measurements. Whole cell fluorescent staining is performed with calcein (Sigma-Aldrich; Cat. No. C0875) at a concentration of 250 nM for a duration of 30 min before measurements.

Design and setup of the device

Each microconstriction device consists of 8 parallel microconstrictions that are surrounded by a pressure-equalizing bypass (Fig. 1a). The pressure across the constriction area is thus clamped by the bypass. The cells enter the device by a punched inlet and first transverse a filter system that holds back debris. After transit through the constriction array, the cells leave the device through a punched outlet. The height and width of the constrications are chosen to be smaller than the cell diameter and are therefore adapted to the mean size of the measured cell population. The height of the whole device, including the constrictions, is on the order of the cell diameter (K562: 13.6 µm – 16.6 µm, DLD-1: 16.6 µm, NIH 3T3: 19 µm). For K562 cells, the width of the constrictions is 5.1 - 6 µm. For DLD-1 cells, a width of 5.1 µm is used. For NIH 3T3 cells, a constriction width of 7.5 µm is used. All device geometries are confirmed with a µ-surf interferometric confocal microscope (NanoFocus AG, Oberhausen, Germany) (SI Fig.1).

The microconstriction array is made out of polydimethylsiloxane (PDMS, Sylgard 184, Dow Corning, Midland, MI) and is produced with standard soft lithography as described in (15, 22). After molding and baking the PDMS, the devices are sealed to a glass cover slip through plasma bonding (Zepto, Diener Electronics, Nagold, Germany). Devices are
used after a waiting time of at least 48 h after plasma treatment to ensure stable hydrophobic channel wall properties (12). Before measurements, the device is coated with 1 wt% pluronic (BASF, Ludwigshafen, Germany; Cat. No. P2443, Sigma-Aldrich) in PBS for 30 min to decrease non-specific cell binding to the channel walls (23, 24).

For measurements, we either use naturally suspended cells (K562), or normally adherent cells brought into suspension after trypsinization (DLD-1 and NIH 3T3). 5,000,000 cells are suspended in 1 ml PBS with 1 wt% pluronic. Rigid polyether-ether-ketone tubing (VWR, Radnor, PA) is used to connect the cell reservoirs to the microfluidic device. Cells are then pumped through the channels of the microfluidic device by a hydraulic pump (Bellofram, Newell, WV) using an external pressure range of 1-50 kPa. During experiments, the constriction area of all 8 constrictions (Fig. 1b) is continuously imaged with a 10×, 0.3 NA objective in bright-field mode at a frame rate of 750 fps using a high-speed CCD-camera (G680, Allied Vision Technologies, Stadtroda, Germany) mounted to an inverted microscope (DM-IL, Leica Microsystems, Wetzlar, Germany). Videos are recorded with a custom written C-program for further analysis with MATLAB (The MathWorks, Natick, MA). Typically, cell transits of 5000 cells are recorded during a measurement time of 30 min. All measurements are performed at room temperature (22 °C).

**Fluorescence imaging**

For recording the fluorescence signals from transfected cells, a diode-pumped solid-state laser (wavelength 473 nm; VA-I-N-473, Viaso, Beijing, China) with a maximum power of 100 mW is coupled to the epifluorescence port of the microscope. Through a series of excitation and emission filters (Thorlabs, Germany), light for bright-field imaging (wavelength > 590 nm) is separated from the emission wavelength of GFP (~515-550 nm) (SI Fig. 2). The CCD camera (G680, Allied Vision) in the bright-field channel runs at a frame rate of 750 fps with an exposure time of 100 µs. For every third image, the camera triggers a second, synchronized camera (also G680, Allied Vision) in the GFP-channel at a framerate of 250 fps with an exposure time of 4 ms, which records the GFP-emission light (Fig. 1b). Only fluorescence images of cells that are immobilized at the constriction entrance are evaluated to avoid motion blurring. For computing the fluorescence intensity of a cell, pixel intensities are background-subtracted and summed over a region of interest corresponding to the channel area in front of the constriction. The intensity sum of all pixels is then normalized by the cell area extracted from bright-field images. Thereby, we obtain a cell-size independent average fluorescence intensity of the cell in units of counts per cell pixel.
Calculation of cell mechanical properties

We calculate the average cell mechanical parameters $E$ and $\beta$ from an orthogonal least squares fit of Eq. 1 to the measured entry times $t_{\text{entry}}$ of typically 2000 cells, the maximum cell deformation $\varepsilon_{\text{max}}$ of each cell, and the pressure drop $\Delta p$ that each cell experienced during transit. From the recorded images, the cell size is detected from bright-field images of the undeformed cell before it enters the constriction (Fig. 1b). From the cell size, the maximum deformation $\varepsilon_{\text{max}}$ of the cell is calculated according to $\varepsilon_{\text{max}} = \frac{r_{\text{cell}} - r_{\text{con}}}{r_{\text{cell}}}$, with $r_{\text{cell}}$ being the radius of the undeformed cell, and $r_{\text{con}}$ being half the constriction width.

From the image series of the cell approaching the constriction, the cell speed is measured by tracking the leading cell edge. From the cell speed $v_{\text{cell}}$, the average flow speed $v_{\text{avg}}$ is estimated using an experimentally measured relationship according to $v_{\text{cell}}/v_{\text{avg}} = 2.64 - 1.59 (r_{\text{cell}}/r_{\text{hyd}})$ (SI Fig. 3), with $r_{\text{hyd}}$ being the hydrodynamic radius of the channel, which is calculated from the channel width $w$ and height $h$ according to $r_{\text{hyd}} = \frac{h \cdot w}{h+w}$(15). The flow resistance of each constriction and each segment of the channel network is calculated using Hagen-Poiseuille’s law. The pressure drop across each channel is then solved using Kirchhoff’s circuit laws, whereby we take into account the occupancy of each constrictions. Thus, we continuously update the pressure drop across a constriction during a cell transit, depending on the occupancy of all other constrictions. In addition, every time a new cell enters the constriction region, we measure the local flow speed, and the pressure drop over all constriction regions is updated accordingly (15).

Cell entry times are measured from changes in the brightness standard deviation of regions of interest in front of the constrictions (15), as indicated by the red square in Fig. 1b.

Statistical analysis

To calculate an average elastic modulus $E$ and fluidity $\beta$ of a cell population, Eq. 1 is fitted to the measured $\Delta p$, $\varepsilon_{\text{max}}$ and $t_{\text{entry}}$ data from typically 2000 cells using an orthogonal linear least squares fit implemented in MATLAB, which minimizes the perpendicular distances between the data points and the regression line (Fig. 1c). Before fitting, $t_{\text{entry}}$ and the ratio of $\varepsilon_{\text{max}}/\Delta p$ are logarithmically transformed to obtain a linear relationship between $\log(t_{\text{entry}})$ and $\log(\varepsilon_{\text{max}}/\Delta p)$. Standard errors (se) for $E$ and $\beta$ are calculated by bootstrapping, where we repeat the fit 100 times on ensembles of randomly selected cells. This standard error corresponds to one standard deviation between the fitted values. For testing significant differences when comparing pairs of
conditions or cell populations, we compute the area of overlap between the probability density distributions of the fitted parameters. Differences are considered statistically significant for an overlap of less than 5 % (p<0.05).

Results and Discussion

Time dependence of cell deformations

Eq. 1 assumes a power-law creep response of the cells inside the constrictions, \( \varepsilon(t) \sim t^\beta \). To test the validity of this assumption, we observe the cell shape and measure the time-evolution of the long axis \( l \) of NIH 3T3 cells during passage through constrictions (Movie M1, SI Fig. 4). For this measurement, we use a device that only contains a single constriction to avoid short-time pressure fluctuations during the cells’ transit caused by changing clogging configurations in the device. The device also contains an enlarged channel cavity after the constriction to observe cell relaxations. We approximate the strain as \( \varepsilon(t) = (l(t) - l_0)/l_0 \), with \( l_0 \) being the cell diameter in flow direction before the cell has entered the constriction. We confirm that the time evolution of the strain \( \varepsilon(t) \) follows a power-law in time (\( R^2 = 0.98 \)) (Fig. 1d). By contrast, the response of a Kelvin-Voigt model, which is also commonly used to describe visco-elastic cell mechanical properties, fits the creep response with a considerably lower fidelity of \( R^2 = 0.90 \) (25, 26).

Stress and strain stiffening

We fit Eq. 1 to the data (\( t_{\text{entry}}, \varepsilon_{\text{max}}, \Delta p \)) from several thousands of cells and compute a population average for the cell elastic modulus and fluidity. This fit corresponds to fitting a straight line to a scatter plot of \( \log(t_{\text{entry}}) \) versus \( \log(\varepsilon_{\text{max}}/\Delta p) \) (15). Data points from individual cells, however, can deviate substantially from the fit line and form a widely distributed, seemingly randomly scattered cloud (Fig. 2, 3a,d). Yet a surprisingly large part of the data scatter is not random but occurs systematically: When we sort the individual data points into groups that have experienced similar pressure and strain, and then plot the average entry time versus the average ratio of \( \varepsilon_{\text{max}}/\Delta p \) for each group separately, we find that the averaged data also form a cloud but with a highly regular, grid-like structure (black squares, Fig. 2). Lines that connect data points of equal strain are aligned more horizontally and are shifted towards larger entry times as the strain increases, implying strain stiffening of the cells (15). Lines that connect points of equal...
pressure are aligned more vertically but are also shifted towards larger entry times as the pressure increases, implying stress stiffening of the cells (15).

To a good approximation, the cell elastic modulus $E$ in Eq. 1 increases linearly with increasing stress $\Delta p$ and strain $\varepsilon_{\text{max}}$ according $E = E_0 + c_1 \Delta p + c_2 \varepsilon_{\text{max}}$. With the additional fit parameters $c_1$ and $c_2$, we can nearly fully account for the scatter of the binned and averaged data points shown in Fig. 2. The fitted stress-stiffening coefficient $c_1$ is on the order of unity, indicating that for a pressure increase of 1 Pa, the cell elastic modulus also increases by 1 Pa. The fitted strain-stiffening coefficient $c_2$ is around 200 Pa, indicating that for a strain increase of 100 %, the cell elastic modulus increases by 200 Pa.

There is a considerable covariance between the fit parameters $c_1$ and $c_2$, however, and the grid-like systematic scatter of the binned data points can be accounted for by considering only stress-stiffening according to $E = E_0 + c_1 \Delta p$ (black lines in Fig. 2), or only strain stiffening according to $E = E_0 + c_2 \varepsilon_{\text{max}}$ (not shown). We therefore opted to fit Eq. 1 to the data with only 2 constant fit parameters, $E$ and $\beta$. These parameters must be interpreted as an effective cell elastic modulus and fluidity that are valid only in the range of pressure and strain values of that particular measurement. To prevent stress or strain stiffening from introducing bias, e.g. when comparing cell populations with different cell sizes or when comparing two measurements obtained with different constriction sizes, it is crucial to control for the strain and stress that the cells have experienced. This can be achieved with a histogram matching method.

**Histogram matching**

This method selects only those cells from two or more different cell populations for comparison that have experienced both the same range of pressure drop $\Delta p$ and the same maximum strain $\varepsilon_{\text{max}}$. In brief, we compute 2-dimensional histograms of distributions for $\varepsilon_{\text{max}}$ with a bin width of 0.05, and for $\Delta p$ with a bin width of 50 Pa, similar to one grid tile in Fig. 2. For every $\varepsilon_{\text{max}}/\Delta p$ bin, we then randomly exclude cells from the population with the larger number of cells in that particular bin until the number of cells in each histogram bin match. This ensures that also the histograms of the ratios of $\varepsilon_{\text{max}}/\Delta p$ for each of the cell populations match approximately. The random cell exclusion process is repeated for every bootstrap repetition (i.e. 100 times) to ensure that no data point within the overlap region of the histogram is thrown away.

We demonstrate the validity and importance of the histogram matching method by comparing two measurements conducted with two constrictions of differing sizes (6.0 x 13.6 µm vs. 5.1 x 16.6 µm) but on the same cell type (K562 leukemia cells). From the
scatter plots of $\log(t_{\text{entry}})$ vs. $\log(\varepsilon_{\text{max}}/\Delta p)$ data (Fig. 3a) and from the histograms of measured $\varepsilon_{\text{max}}$ and $\Delta p$ (Fig. 3b left and right), systematic differences between the two measurements are noticeable: cells measured with 5.1 µm constrictions experience a higher maximum deformation than cells measured with 6.0 µm constrictions (Fig. 3b left). Cells measured with the smaller constrictions also experience a higher pressure drop $\Delta p$, as the experimenter tends to increase the pressure across the narrower device to compensate for the greatly increased entry time (Fig. 3b right). The combined effect of an increased strain and pressure cause the elastic modulus to increase due to strain and stress stiffening, whereas the power-law exponent remains similar (Fig. 3c left and right). After $\Delta p$ and $\varepsilon_{\text{max}}$ histogram matching (exemplary bootstrap fit shown in Fig. 3d, e), the cell elastic moduli for the two measurement conditions match nearly perfectly (Fig. 3f). The remaining mismatch of the elastic modulus of 3% (7 Pa) is within the systematic margin of error caused by measurement inaccuracies of the channel geometries (± 0.25 µm) between the different devices.

**Influence of culture conditions and measurement parameters**

We next test to what degree cell mechanical properties are further influenced by measurement parameters such as cell confluency prior to harvesting, time between cell trypsinization and measurements, choice of cell suspension medium, or device preparation with pluronic. All measurements for a given parameter or measurement condition are conducted on the same day with DLD-1 colon carcinoma cells harvested from identically prepared flasks. Between 2-3 independent measurements (i.e. with cells from 2-3 different flasks), each with at least 5000 cells, are performed for each condition. The measurement data for each condition are pooled and histogram matched for comparison. Note, however, that absolute cell elastic modulus values between different types of parameters cannot be compared, as slightly different deformation and pressure histograms have been chosen for different parameter types to allow for an optimal histogram overlap within the parameter group.

*Reproducibility:* To investigate the measurement reproducibility of our system, we conduct four independent measurements of DLD-1 cells on one day under identical conditions. We then randomly pool two of the measurements into one group (control 1) and compare the resulting mechanical properties with the pooled data from the remaining two measurements (control 2). We find only negligible differences between the elastic modulus (SI Fig. 5a, Fig. 4a) or the power-law exponent (SI Fig. 5b) between the two groups, demonstrating the reproducibility of the method.

*Confluency:* Cells are split 24 h prior to measurements and are seeded at different densities such that they reach different confluencies (20 %, 60 %, and 100 %) before
harvesting. We find a significant decrease in cell elastic modulus between 20 % versus 60 % confluency, and between 60 % and 100 % confluency (SI Fig. 5c, Fig. 4b). The power-law exponent $\beta$ also shows a slight decrease from 20 % to 100 % confluency (SI Fig. 5d). Based on this result, we recommend that cell confluency should be tightly controlled in all experiments.

**Time between cell trypsinization and measurements:** Typically, measurements start around 15-20 min after harvesting the cells by trypsinization. This time span includes 4-5 min trypsinization, a 4 min centrifugation, 5 min transport to the microscope and setup, and a 5 min flushing time in which the cells travel through the tubes to the microconstrictions. Measurements typically last for another 30 min. Here, we find that an increase in the waiting time from 20 min to 60 min between harvesting and start of the measurement (cells are stored in a falcon-tube in the incubator) results in a significant decrease of the cell elastic modulus and a slight but significant increase of the power-law exponent (SI Fig. 5e,f, Fig. 4c). There is no further decrease in the cell elastic modulus for a waiting time of 100 min. Based on this result, we recommend that measurements should be terminated one hour after trypsinization at the latest.

**Cell suspension medium:** After trypsinization and centrifugation, cells are usually resuspended in PBS (pH=7.25) and kept therein for the time of the measurement. Resuspending cells in HEPES-buffer (50 mM HEPES, 10 mM NaCl, 10 mM glucose, pH=7.0) instead of PBS significantly increases the cell elastic modulus and decreases the power-law exponent (SI Fig. 5g,h, Fig. 4d). Resuspending cells in DMEM (1 g glucose, pH=8.13) significantly increases the cell elastic modulus, but not the power-law exponent, compared to PBS. It is unclear whether differences in the pH or in the composition of the cell suspension medium have caused these changes. Phenol-red containing DMEM is a reasonable choice for a cell suspension medium as it is also the medium in which the cells are cultured. However, PBS improves the image quality and reduces cell clumping compared to DMEM, which justifies its use in our study.

**Device coating with pluronic:** To prevent the adhesion of cells onto the walls of the microfluidic device, we flush the devices with a 1 wt% pluronic surfactant solution in PBS for 30 min prior to measurements. Here, we compare non-coated devices with devices coated for 20 min or 40 min with 1 wt% pluronic. We find a significant increase in cell elastic modulus for the devices coated for 20 min and for 40 min (SI Fig. 5i, Fig. 4e) compared to the non-coated devices. We also observe that a large portion of cells become stuck in the inlet filter and in the constrictions of the uncoated microfluidic devices. Thus, non-specific adhesion and increased friction between the cells and the walls of the uncoated microfluidic device may have caused a preferential retention of stiffer cells. The power-law exponent $\beta$ does also show a systematic increase between the three pluronic coating times (SI Fig. 5j).
We also investigate the influence of pluronic coating concentration on cell mechanical properties. We compare the usual coating concentration of 1 wt% with a 0.25 wt% and a 4 wt% concentration of pluronic in PBS, each for a 30 min coating time prior to measurements. We find a significantly higher cell elastic modulus for the 0.25 wt% coating concentration compared to 1 wt% and 4 wt% coating concentration (SI Fig. 5k, Fig. 4f), but no changes in the number of cells that become stuck in the inlet filter. Nonetheless, we expect that friction and non-specific adhesion between the cells and the walls of the device including the inlet filter increases at low pluronic concentrations, and thus we speculate that this may have caused a small but detectable mechanical activation and stiffening of the cells (27). The power-law exponent $\beta$ shows small and non-systematic but statistically significant changes for the 1 wt% and 4 wt% compared to the 0.25 wt% coating concentration (SI Fig. 5l). Thus, pluronic concentration and coating time both need to be tightly controlled for all measurements.

In summary, although the cell mechanical responses to altered culture and measurement conditions are in most cases relatively small and physiologically irrelevant when considered in isolation, the combined effect of poorly controlled conditions can lead to statistically significant bias.

**Influence of protein expression levels**

We have demonstrated above that histogram matching effectively compensates for systematic changes in cell mechanical properties that are caused by pressure and strain effects. In the following, we extend the histogram matching approach to evaluate systematic changes in cell mechanical properties that are caused by different protein expression levels. This is relevant foremost in the case of transiently transfected cells where only a part of the cell population, typically between 10 – 90 %, is likely to be transfected (28), and where also the amount of expressed proteins varies greatly between transfected cells. We have extended our setup with a second high-speed CCD camera that acquires fluorescence images. This camera is synchronized with the CCD camera for the bright-field images, and thus we can measure for each cell not only the entry into the constriction but also the expression level of a fluorescently tagged protein. This allows us to exclude non-transfected cells from the analysis, or to correlate the GFP-signal intensity with altered mechanical properties to measure a dose-response curve.

As a proof of principle, we measure a mixed population of K562 leukemia cells. Half of the cells are treated for 30 min with the actin-depolymerizing drug cytochalasin D (10 µM) together with the green fluorescent dye calcein (250 nM). These labeled cells are then mixed with unlabeled, untreated cells and are measured in our microconstriction
setup. For the analysis, we split the data into a fluorescent group (> 8 count/cell pixel, median = 28 counts/cell pixel) and a non-fluorescent group (≤ 8 count/cell pixel, median = 0.3 counts/cell pixel) and perform Δp and εmax histogram matching between the two groups (Fig. 5a,b). As expected, we find a significantly (p<0.05) decreased cell elastic modulus (Fig. 5c) and an increased power-law exponent (Fig. 5d) in the green fluorescent, cytochalasin D-treated group, which is in accordance with the literature (8, 21).

We next measure the mechanical properties of K562 cells transiently transfected with a lamin A-GFP protein. Lamin A-overexpression has been shown to increase cell elastic modulus (15, 29), which protects the cell nucleus from damage during compression at high deformations (30–32). After transfection, individual cells show the typical wide distribution of GFP and thereby lamin A-expression levels (Fig. 5e). To measure a dose-response curve, we split the data into three subgroups corresponding to “low”, “medium” and “high” levels of GFP (bins as indicated in Fig. 5e). The median fluorescence cell pixel intensities for the three groups are 1.2, 11.2 and 25.5 counts/pixel, respectively. We determine E and β separately for the three groups with Δp and εmax histogram matching (Fig. 5f). With increasing GFP levels, we find a significant (p<0.05) increase of the cell elastic modulus (Fig. 5g) and a decrease of the power-law exponent (Fig. 5h). These data are consistent with a dose-dependent nuclear stiffening for increasing expression levels of lamin A.

**Conclusion**

In this report, we analyze the influence of stress and strain stiffening on cell mechanical properties measured with a microfluidic microconstriction device. Stress and strain stiffening is a major confounding factor when comparing cell populations that have different mean cell sizes, when comparing measurements from devices with constrictions of different sizes, or when comparing measurements conducted with different mean working pressures. To correct for stress and strain stiffening effects, we introduce the method of stress and strain histogram matching and demonstrate that it greatly reduces the variability between different measurements and ensures high reproducibility. Moreover, we demonstrate that also seemingly minor cell culture details or measurement parameters can confound the resulting data and therefore need to be tightly controlled, such as cell confluency prior to harvesting, time between harvesting and measurements, choice of cell suspension medium, as well as details of the device preparation with pluronic. Lastly, we introduce a new fluorescence extension to our setup that allows us to measure transiently and incompletely transfected cell populations by excluding untransfected cells from the analysis, or by correlating protein
expression levels with cell mechanical properties. We show that we can thereby measure dose-response-relationships between the expression of proteins and mechanical cell properties.

Author Contributions

JRL, BF, WS and GW designed the setup and experiments; JRL, SR and GW developed the data acquisition and analysis software; JRL and MS performed the measurements; JRL, CM and BF analyzed the data; TK generated transfected lamin A-GFP cell line; JRL and BF wrote the paper.

The authors declare no conflict of interest.

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Supporting Citations

Figure Captions

**Fig. 1** (a) Schematic of the microfluidic device with inlet (top), debris filter, constriction area surrounded by a bypass, and outlet (bottom). (b) Bright-field (top) and fluorescence (bottom) image of the constriction region with K562 leukemia cells stained with calcein. Images are contrast enhanced. Scale bar is 20 µm. The standard deviation (SD) of the brightness within each ROI (indicated by a red square in front of a constriction) is used to calculate cell entry time. (c) \( t_{\text{entry}} \) vs. \( \frac{\varepsilon_{\text{max}}}{\Delta p} \) of 19,991 K562 leukemia cells. Colors indicate the bivariate kernel density estimate of the data points. The black line is the orthogonal least squares fit of Eq. 1 to the data. (d) Strain evolution \( \varepsilon(t) \) of NIH 3T3 cells (black circles, mean ± se of 59 cells) during entry into a constriction can be fitted by a power-law (orange line). Time \( t \) and strain \( \varepsilon \) are normalized by \( t_{\text{entry}} \) and \( \varepsilon_{\text{max}} \) before averaging. \( t = 0 \) corresponds to the time point when the cell first encounters the constriction, and \( t = 1 \) corresponds to the time point when the cell has reached its maximum strain \( \varepsilon = \varepsilon_{\text{max}} \) and leaves the constriction. Inset: Normalized strain vs. normalized time of individual cells.

**Fig. 2** Stress and strain stiffening: Scatter plot of \( t_{\text{entry}} \) vs. \( \frac{\varepsilon_{\text{max}}}{\Delta p} \) for 19,991 K562 cells as in Fig. 1c. Black squares (connected by red lines) show median values of \( t_{\text{entry}} \) vs. \( \frac{\varepsilon_{\text{max}}}{\Delta p} \) for different pressure and strain bins with at least 250 cells in each bin, with a strain bin width of 0.03, a pressure bin width of 50 Pa, and median pressure and strain values as indicated by the numbers next to the grid lines. Black grid shows the fit of Eq. 1 to the binned data, with stress stiffening according to \( E = E_0 + c_1 \Delta p \). Fit parameters are \( E_0 = 58 \) Pa, \( c_1 = 1.11 \) Pa\(^{-1}\), \( \beta = 0.07 \).

**Fig. 3** Histogram matching: (a) Scatter plot \( t_{\text{entry}} \) vs. \( \frac{\varepsilon_{\text{max}}}{\Delta p} \) for two K562 leukemia cell populations measured with wide (6 µm) and narrower (5.1 µm) constrictions. Black lines show the fit of Eq. 1 to the data. (b) Strain histograms (left) and stress histograms (right) are shifted to higher values for cells in narrow constrictions (blue) compared to cells in wider constrictions (yellow). Areas where histograms overlap are marked in green. (c) Without histogram matching, cells measured in the narrower constrictions (blue) appear significantly (p<0.05) stiffer (left) but have a similar fluidity (right) compared to cells measured in the wider constrictions (yellow). (d) Exemplary scatter plots and fit lines of the same measurements as (a) after \( \Delta p \) and \( \varepsilon_{\text{max}} \) histogram matching. (e) Exemplary strain histograms (left) and pressure histograms (right) after histogram matching show perfect overlap for both cell populations. (f) After histogram matching, cells measured in the narrow (blue) and wider (yellow) constrictions both have similar elastic modulus (left) and fluidity (right) values.

**Fig. 4** Influence of culture and measurement conditions on cell mechanical properties of DLD-1 colon carcinoma cells: (a) Elastic modulus of cells from four independent measurements, arbitrarily split into two groups (control 1 and control 2). (b) Cell elastic modulus measured for cell populations with different culture confluency (20 %, 60 %, 100 %) prior to harvesting. (c) Cell elastic modulus in response to different detachment times before measurement (20 min, 60 min, 100 min). (d) Cell elastic modulus measured for three different measurement media: PBS, HEPES-buffer and DMEM cell culture medium. (e) Cell elastic modulus measured after coating the devices with 1 wt% pluronic for 0 min, 20 min and 40 min. (f) Cell elastic modulus measured with devices coated for 30 min with pluronic
concentrations of 0.25 wt%, 1 wt%, and 4 wt%. n > 1200 for each condition. $\varepsilon_{\text{max}}$ and $\Delta p$ histogram matching was performed separately for each panel (a-e). Numbers show mean ± se. Significant (p<0.05) differences to first parameter values (green bars) are indicated by asterisks.

Fig. 5 (a-d) Measurement of a mixed K562 leukemia cell population, containing 50 % untreated cells and 50 % cells treated with cytoD (10 µM, 30 min) and stained with calcein (250 nM, 30 min). (a) Histogram of average fluorescence intensities per cell pixel. Dashed green line indicates the fluorescence threshold for separating cells into a "zero" and a "high" fluorescence intensity group. (b) Exemplary histogram-matched scatter plots of $t_{\text{entry}}$ vs. $\varepsilon_{\text{max}}/\Delta p$ for cells with "zero" fluorescence (non-treated) and "high" fluorescence (cytoD and calcein-treated) after histogram matching. Black lines indicate the fit of Eq. 1 to the data. (c) Cell elastic modulus and (d) cell fluidity for the two groups. (e-h) Measurement of lamin A-GFP transfected K562 leukemia cells. (e) Histogram of average fluorescence intensities per cell pixel. Dashed green lines indicate the fluorescence threshold for separating cells into a "low", "medium", and "high" fluorescence intensity group. (f) Exemplary histogram-matched scatter plots of $t_{\text{entry}}$ vs. $\varepsilon_{\text{max}}/\Delta p$ for cells with "low", "medium" and "high" expression levels of lamin A-GFP. Black lines indicate the fit of Eq. 1 to the data. (g) Cell elastic modulus and (h) cell fluidity for the three groups. Significant (p<0.05) differences compared to the cell group with "zero" (a-d) and "low" (e-h) intensity are indicated by asterisks.

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