# Lab on a Chip



**View Article Online** 

# PAPER



Cite this: Lab Chip, 2025, 25, 673

# Densimeter-on-chip (DoC): measuring a singlecell mass density by sedimentation in microchannel flows<sup>†</sup>

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Intrinsic biophysical and morphological features are essential for the label-free identification of different cell types. Indeed, apart from object size, density could represent a key parameter for single-cell analysis. However, the measurement of such a parameter is challenging. Therefore, we present a straightforward and versatile microfluidic chip. The densimeter-on-chip (DoC) measures single-cell mass densities thanks to a hydrodynamically induced sedimentation process inside the microchannel. In detail, in-flow buoyant components become more relevant than viscoelastic alignment forces, leading to precise in-flow sedimentation. DoC is based on precise three-dimensional cell alignment, followed by an abrupt change in cross-section to induce calibrated sedimentation. Based on the balance of acting forces and tracking the in-flow cell trajectory, we have developed a self-written mathematical model to precisely measure the single-cell densities of multiple cell types of any shape. Both cell velocity and fall length define the resulting cell density. The working range of object diameters for which density can be estimated is 0.75-22.5 µm. As result, the minimum measured density is 998 kg m<sup>-3</sup> and a sensitivity of 0.001 can be obtained. Great agreement between the computational and the literature findings about red blood cells ( $\sim$ 1159 ± 29.5 kg m<sup>-3</sup>), lymphocytes (~1073  $\pm$  49 kg m<sup>-3</sup>) and neutrophils (~1093  $\pm$  27 kg m<sup>-3</sup>) is obtained without chip modification. Indeed, the computational error between the mean density values is ~1%. Thereby, DoC as an easy-to-use and reproducible solution for label-free single-cell density measurement, provides a universal approach for characterizing a wide range of cell types, independently of their size and shape

Received 31st October 2024, Accepted 20th January 2025

DOI: 10.1039/d4lc00916a

rsc.li/loc

### Introduction

Cell density, regarded as the mass-to-volume ratio, is tightly regulated and varies little within a given cell type. Changes in cell density are determined by the sum of biosynthesis and degradation processes, revealing the status of a cell.<sup>1</sup> Indeed, cells regulate their density during the cell cycle, metabolism, apoptosis, differentiation, chronic diseases or cancer progression, and in response to drugs.<sup>1,2</sup> In fact, in the case of isolation of circulating tumour cells, tracking the physical characteristics of a cell, such as density, cell size, and electrical properties, is necessary to distinguish between healthy and unhealthy cells.<sup>3</sup> Moreover, the way in which cell volumes change after external stimuli, as in the case of induced immune or drug responses, is strictly related to how cell size, structure and density could be rearranged.<sup>4–8</sup> For example, changes in cellular density also occur in lymphocytes during inflammatory responses, when depletion of cytokines inactivates cells and differentiates them into memory cells.<sup>1</sup> Therefore, detailing possible changes in cell density is relevant for both diagnostic and therapeutic applications.

Typically, single-cell density values, comprising nucleus, inner fluids and organelles, range between 1060 and 1120 kg  $m^{-3}$ .<sup>9,10</sup> Of particular interest are changes in cytoplasm density related to alterations in protein concentration that in turn also affect the assembly of protein complexes and cell mechanics.

However, as cells consist of a unique set of morphological and biophysical features, precise characterization of this heterogeneity is still challenging.<sup>11–22</sup> Indeed, density values show very little variation in a given cell type, with a variability 100 times smaller than the variation in cell mass among cells, since there is tight regulation of their density for maintenance of their state.<sup>23</sup> Therefore, having a technique capable of determining, among the same type of cell, single-cell densities at high precision and specificity is crucial.

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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d4lc00916a

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The most widely used method for cell density characterization is density gradient centrifugation, which allows one to obtain a convenient estimation of the average density of cell populations.<sup>24</sup> However, such a technique considers a fixed density threshold (1077 kg m<sup>-3</sup>) mainly to separate lymphoblasts and monocytes from the remaining myeloblasts and red blood cells (RBCs). Moreover, this approach does not provide single-cell results for an inherent cell heterogeneity distinction and there is no correlation between size, density and shape of cells. Therefore, during the last few years, several methods, both biochemical and microfluidic/physical based, have been proposed as valid alternatives to measure the physical properties of cells.<sup>2</sup> Typical examples are fluorescence immunoassays and biochemiluminescence techniques or single-cell mass cytometry,25 magnetic cytometry, levitational image phase-shifting suspended microchannel resonators, interferometry and opto-electrokinetic devices.<sup>26-29</sup> In detail, they can be classified as active or passive methods. Magnetic and electric-based techniques are active methods, which impose external forces to displace cells for separation and characterization.<sup>16,30-32</sup> In contrast, passive applications include hydrophoretic and hydrodynamic filtration and sedimentation.33-36 gravitational Some of them are microfluidic designs, where channels are disposed at different heights, inducing gravity-based cell sedimentation; in others, purely inertia-driven fluid flow conditions allow inflow cell separation based on pinched flow fractionation to profile cells with different sizes and densities.37,38 As an example of these approaches, a microfluidic device where a combined effect of hydraulic jump and sedimentation for size-selective microparticle or cell separation has been proposed. Even if the resulting sorting enrichment based on size was high, the device potential was limited since a complex design and tight control over the fluid-dynamics are required to obtain the desired sorting.39

Nevertheless, other important drawbacks of both active and passive techniques need to be considered. For example, biochemical methods often have problems in achieving noninvasive, high-content and cell-specific measurement. Indeed, label-based analyses could lead to undesired cell reactions due to special preparation procedures. In contrast, physicalbased techniques can measure multiple physical properties simultaneously at the single-cell level. However, the necessity to find a balance between applied flow rates, high throughput, measurement precision and multiparameter measurement, leads to unsatisfactory performance. Moreover, none of the presented techniques allow variable density levels to be recognized and measured among different cell types in the same sample.

Here, we present a densimeter-on-chip (DoC) as a new possibility to in-flow-measure single-cell density with a gravity-based approach. By first inducing a perfect viscoelastic alignment of cells, and then presenting an abrupt change in the channel geometry, in-flow cell sedimentation—where gravity ( $F_{\rm G}$ ) and buoyancy ( $F_{\rm B}$ ) forces matter—is

promoted, since viscoelastic fluid forces dramatically reduce its lift effect. From the known laws regulating the balance of forces for viscoelastic alignment—viscoelastic ( $F_{\rm E}$ ) and drag ( $F_{\rm D}$ ) balance—we provide an analytical set of relations to practically estimate the density, depending on the fall length (L) of cells along the observation channel (Fig. 1).<sup>40</sup> In particular, alignment can be achieved thanks to tight control over the imposed hydrodynamic forces ordering particles/ cells on different equilibrium positions and trajectories, depending on the channel geometry, the fluid properties and the dimensions and shape of the object.<sup>41–44</sup> Indeed, the interplay of size and shape plays a fundamental role in the definition of cell motion for alignment, separation and density measurement purposes.

We achieved precise single-cell analysis from homogenous cell types, by mapping cell diameter *versus* density outcomes, which comprehensively describes the biophysical characteristics of a cell population.

On the other hand, thanks to implementation of the mathematical model, the approach offers the possibility of easily and precisely in-flow-separating differently sized and weighted objects, if diameters and densities are known. With the choice of the best applied pressure conditions, we can predict the length of sedimentation of the object before starting the experiment, therefore allowing sedimentationinduced separation.

In detail, the device consists of one inlet and two outlet capillaries, which are inserted in a sandwich of engraved polymethylmethacrylate (PMMA) slides that contain the observation channel as well as the vertical fluid exit (Fig. 1 and S1<sup>†</sup>). The gravity-based sedimentation can be finely tuned with the fluid flow parameters-such as fluid viscosity and applied pressure in a working range of ~250-2500 mbar -to ensure a precise fall length for different object types. As proof-of-concept, we decided to test both particles and peripheral blood cell sedimentation to predict the fall length and measure the density, respectively. The minimum density that can be measured is 998 kg m<sup>-3</sup>. Thanks to the in-flow observation and the relative computation, we demonstrate the possibility of profiling single-cell densities with great agreement with literature findings. Therefore, DoC is proposed as a valid, highly reproducible and easy-to-use approach to characterize and separate cells, depending on their morpho-physical properties, of prominent interest for diagnostic purposes.

### Materials and methods

#### Densimeter-on-chip (DoC) concept

The working principle of DoC is based on two different fluid conditions, which are consecutively applied to cells of unknown density. First, cells are perfectly aligned to the centre line of a round-shaped capillary, before an abrupt change in cross-section accompanied by a subsequent change in fluid forces results in a triggered dis-alignment in-flow, mainly caused by the mass density signature of a cell. In fact,



**Fig. 1** Design and working principle of DoC. A real image of the device is presented as it is seen down the microscope, as a top view (YX-plane). An initial round capillary is connected to the PMMA-based microfluidic device, and two exits are provided for the lateral (EXIT 1) and straight motion directions (EXIT 2). Cells or particles are first aligned at the centre line of the capillary before an abrupt change in cross-section, which triggers a dis-alignment along the *Z*-direction (in-flow sedimentation effect) due to a change into the force balance. We show a sketch of the side view that would be observed during the experiment. Indeed, depending on the applied pressure drop, cells or particles of different densities and sizes can be aligned and then sediment into a larger section, where viscoelasticity effects are not more relevant while drag dominates. Thanks to the measurement of the length of sedimentation (*L*), diameter of the objects, and fluid density ( $\rho_t$ ), we can calculate the object density ( $\rho$ ), obtaining a mass density signature.

by measuring the fall length of a cell of known cell size and geometrical and fluid properties, we can calculate the density of the investigated cell.

Besides the characterization, we introduced a vertical channel exit in the chip design, which can be used to collect cells of interest, while the remaining sample content can be collected in another exit at the end of the observation channel. The possibility of fine tuning the fall length by changing the flow rate and, as a consequence, the fall velocity allows precise single-cell characterization.

#### Microfluidic chip

The chip for density investigation in-flow consists of a cover and a base part made of PMMA, which are milled to obtain the observation channel for the density measurement (cover) and to hold in place the inlet (cover) and the outlet (cover and base) capillaries. The geometrical parameters of the observation channel are height 500 µm, width 1000 µm and length 8.5 cm (see Fig. S1<sup>†</sup>). The capillaries have a diameter of 75  $\mu m$  for 'INLET' and 'EXIT 2', while 40  $\mu m$  was chosen for lateral 'EXIT 1' (see Fig. S1 and S2<sup>†</sup>). The capillary lengths were optimized for cell alignment in the inlet as well as cell collection at the outlets, resulting in 34 cm for the inlet and 10 cm for both outlet capillaries. Due to the low-cost approach for the device, manufacturing a mismatch of  $\sim 200$ µm between the capillary centre line and the channel bottom had to be solved in designing the device. We decided to introduce into the channel design 5 steps (50 µm in depth) 9 mm before the lateral exit, which successively lowered the channel bottom in the central section of the channel to escort cells or particles of interest into the lateral exit of the device (see Fig. 1 and S2<sup>†</sup>). Furthermore, the lateral capillary enters the observation channel for  ${\sim}50~\mu m$  for smoother extraction of objects.

#### Rheological fluid properties and fluid-flow losses

The viscoelastic measurement fluid consists of 0.3, 0.4 and 0.5 wt% of polyethylene oxide (PEO, 4 MDa, Sigma Aldrich) in phosphate-buffered saline. We investigated the fluids with a stress-controlled rheometer to obtain their rheological properties (see Fig. S4b<sup>†</sup>). The zero-shear viscosity ( $\eta_0$ ) was measured with 0.00683, 0.01002, 0.01998 Pa s for PEO 03, 04 and 05, respectively. The reported flow curves (ESI,† Fig. S4b) show that there is well-defined Newtonian behaviour up to a shear rate  $\gamma$  of ~500 s<sup>-1</sup> for PEO 04, the one we chose for the experiments. In our device, even inside the capillary, the highest estimated  $\gamma$  is ~200 s<sup>-1</sup>. Therefore, non-Newtonian effects on viscosity are absent.45,46 The constant viscosity was helpful to ensure that the influence of temperature was also negligible. Indeed, there is a critical concentration (~1.6 wt%) above which both characteristic non-Newtonian behaviour and the direct influence of temperature on viscosity values are observed.45,46

Next, we investigated the hydraulic resistance of the microfluidic system, where  $R_1$  and  $R_2$  (inlet capillary and channel before EXIT 1) as well as  $R_4$  and  $R_5$  (channels after EXIT1 and EXIT 2 capillaries) are considered in series, while  $R_3$  (capillary of EXIT 1) and  $R_{4-5}$  (all resistance after EXIT 1) are calculated in parallel (see Fig. S3†). Note that  $R_3$  shows a ~12-fold increase in hydraulic resistance compared to  $R_5$  when considering PEO 04 fluid. The interplay of hydraulic resistance is scaled according to the chosen PEO dilution. We optimized the capillary geometries according to the length and diameter of human bloodstream cells.

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### Calibration beads

Polystyrene latex beads (PSL) of different sizes (4.16  $\pm$  0.062  $\mu m$ , 6.08  $\pm$  0.082  $\mu m$  and 8.13  $\pm$  0.09  $\mu m$  from FLUKA, as well as 10.70  $\pm$  0.25  $\mu m$  and 15.66  $\pm$  1.43  $\mu m$  from PolyScience) were used for calibration of the microfluidic device (Fig. 2b). Note that PSL are known for their perfectly spherical shape and monodispersed size distribution. Beads below 10  $\mu m$  have a density of 1050 kg  $m^{-3}$ , while bigger beads were provided with a density of 1055 kg  $m^{-3}$ .

### Cell sample preparation

Cells were recovered from a healthy donor after obtaining informed consent in accordance with relevant guidelines and regulations. At the time of blood donation, the donor signed an informed consent (model no. 5526 of Azienda Ospedaliera Universitaria "FEDERICO II", Naples, Italy), which specifies that waste parts of the blood, not useful for the medicaltherapeutic purposes, can be used for scientific research purposes. All experiments were performed and analysed anonymously. The collection was performed to divide peripheral blood mononuclear cells and red blood cells (RBCs) from the whole sample. To do this, we diluted the blood with phosphate-buffered saline (PBS, SIAL Srl) in a 1:1 proportion. To this solution, we added a Ficoll-Paque (Fisher Scientific) with  $\rho = 1077$  kg m<sup>-3</sup>, still in a 1:1 proportion. Thanks to a density gradient, Ficoll remains on the bottom of the blood-PBS solution. Then, we centrifuged the sample at  $200\vec{g}$  for 30 minutes, in no-brake mode. After centrifugation, we removed the plasma and platelets, which are useless for our experimental purposes. The peripheral blood mononuclear cell (PBMC) ring was picked up and placed inside another tube rinsed with Roswell Park Memorial Institute (RPMI) 1640 medium (SIAL Srl). Note that  $\sim$ 15% of the collected cells are monocytes, and although most of them adhere during the measurement procedure to the surface of the sample container, we will consider only the lymphocytes (LYM) for the microfluidic experiments. For the neutrophil (NEU) class, cells separated below the gradient medium were collected and washed with RBC lysis solution three times for 5 minutes. We centrifuged the sample three times at  $200\vec{g}$  for 10 minutes, then removed the supernatants and rinsed the sample with pure RPMI. Such cell populations will also contain basophils and eosinophils, although they represent only ~3% of the collected population. In fact, we consider only NEU for our experiments. For the RBCs, a remaining part of fresh blood was used. We centrifuged it at  $200\vec{g}$  for 10 minutes, in no-brake mode. Again, removing



**Fig. 2** DoC calibration. (a) We investigated the theoretical fall length (*L*) of PSL 6 (star) and 10 (circle) beads for PEO 03 (low  $\eta$ ), 04 (medium  $\eta$ ) and 05 (high  $\eta$ ) for fluid  $\Delta$ Ps from 400 up to 2000 mbar. (b) Particle velocity was investigated theoretically (line) and experimentally (circles) for two different  $\Delta$ Ps in measurement position X1, which is placed 2 mm after the inlet capillary directly in the measurement channel. *e%* is 7% (*n* = 14) and 8% (*n* = 26) for 400 mbar and 700 mbar, respectively. (c) The vertical position of PSL 10 beads was investigated over distance for different  $\Delta$ Ps and compared with fluid simulations. Mean *e%* is 24% (*n* = 20) and 21% (*n* = 36) for 400 mbar and 700 mbar, respectively. (d) A schematic representation of PSL 6 versus PSL 10 beads in DoC for PEO 04 and 700 mbar of applied pressure.

plasma and platelet parts, we collected only RBCs. In-flow measurements were performed with cell concentrations of  $\sim 10^6$  cells per mL to avoid cell interaction or aggregation.

#### Object tracking in-flow

We recorded objects in the XY-plane with an inverted brightfield microscope (IX81, Olympus) using 10× magnification and a CMOS camera (ORCA flash 4.0, Hamamatsu Photonics K.K.) with 6.5 µm pixel size. Next, we tracked cells or particles with a self-written MATLAB R2023a (MathWorks Corp.) routine, which first filtered the input frames with a set of Gaussian filters. Afterwards, we applied a threshold-based bandpass filter to obtain the object centroids for each recorded frame. Subsequently, the routine combines the tracked coordinates into single tracking information. In the case of missing centroid data in one or a maximum of 3 consecutive frames, the routine can reconstruct the missing information to guarantee the best tracking performance. However, such a tracking approach will work only in a certain range of Z-positions ( $\sim 100 \mu m$ ), which is limited by the diffraction-limited focus in the recorded bright-field images. In other words, we changed the focal plane for each observed video, limiting the observation to a maximum range of  $\sim 50$  $\mu$ m above and  $\sim$ 50  $\mu$ m below the optimal object focus. Note that the object diffraction is size dependent. Finally, the object velocity was calculated for trajectories above a minimum threshold of 5  $\mu$ m s<sup>-1</sup>, which was considered to be a quiescent flow condition.

#### Velocity field computation

Since we are interested in reconstructing the in-flow motion of particles and cells inside the microchannel, we wrote the Poiseuille velocity profile (according to the hypothesis of the Oldroyd-B fluid model, ESI†). For rectangular cross-sections with respect to a fixed Cartesian system (see Fig. 1), the profile is:<sup>47</sup>

$$U_X(y,z) = U_{\text{avg}} \left[ 1 - \left(\frac{z}{H}\right)^n \right] \left[ 1 - \left(\frac{y}{W}\right)^m \right]$$
(1)

where  $U_{avg}$  is the average velocity of the fluid in the two directions of variation, *y* and *z*. The characterizing parameters are obtained from:

$$n = 2 \text{ if } AR \ge 3$$
  
 $n = 2 + 0.3 \left( AR^{-1} - \left( \frac{1}{3} \right) \right) \text{ if } AR \le 3$   
 $m = 1.7 + 0.5 (AR^{-1})^{-1.4}$ 

by varying AR, where AR is the aspect ratio of the channel (ESI<sup>†</sup>). From that, we wrote the existing relation between the maximum velocity  $(U_{\text{max}})$  and the  $U_{\text{avg}}$ .<sup>47</sup>

$$U_{\max} = \left(\frac{m+1}{m}\right) \left(\frac{n+1}{n}\right) U_{\text{avg}}.$$
 (2)

From a comparison of experimental and computational data, we calculated the error percentage (e%) for each case as:

$$e\% = \frac{\text{Exp}_{\text{Data}} - \text{Comp}_{\text{Data}}}{\text{Exp}_{\text{Data}}} \cdot 100.$$

### Results and discussion

#### Governing forces for DoC

Before performing the density-based cell characterization, the capability of the DoC was investigated with calibration beads. We used density standard beads (PSL) similar in size and density to peripheral blood cells. Therefore, a viscoelastic pressure-driven flow with an applied inlet pressure ( $\Delta P$ ) is reported by modelling the fluid according to Oldroyd-B (ESI†). In such fluid-flow conditions, suspended particles/ cells experience a force  $F_{\rm E}$  that emerges from an imbalance of the positive  $N_1$  (ESI†), which moves them toward their equilibrium position at the channel centre line. In our case, the alignment condition has already been reached and kept stable inside the inlet capillary (Fig. 1 and 2d). When particles move from the capillary inside the bigger rectangular cross-section, a decrease in  $F_{\rm E}$  contribution occurs, which triggers an in-flow sedimentation effect.

In detail, we describe such phenomena by defining different sedimentation falling lengths (L) for different-sized microparticles. Assuming the height (H, Fig. 1) of the rectangle is smaller than the width, we can neglect the influence of side walls and focus on forces applied along the vertical direction (*z*-axis). At the beginning of the observation channel (the centre line of the rectangular section), the balance of forces can be defined as follows:

$$F_{\rm Ez} + F_{\rm g} - F_{\rm b} - F_{\rm Dz} = 0 \tag{3}$$

where  $F_{Dz}$  and  $F_{Ez}$  are the drag and elastic components along z, respectively. The difference  $F_g - F_b$  is the buoyant force. By assuming the particle is already at the centre line of the microchannel with a velocity comparable to the maximum of the fluid itself, we can write the expression for  $F_{Ez}$ :<sup>48</sup>

$$F_{\rm Ez} = Ca^3 \frac{1}{\zeta} \nabla N_1 = Ca^3 2\eta \lambda \nabla (\dot{\gamma}^2) \frac{1}{\zeta} = C128 U_{\rm max}{}^2 \eta \lambda \beta^3 \frac{1}{\zeta} \qquad (4)$$

where  $C = 16\pi$  and  $\zeta$  is the non-dimensional channel gap, which varies according to the confinement ratio ( $\beta$ ) of the particle we are considering.<sup>48</sup>  $U_{\text{max}}$  is the maximum fluid velocity from eqn (2). The profile is the typical Poiseuille flow, with parabolic trend (Fig. 2b). This velocity dramatically reduces from the inlet capillary ( $U_{\text{max}} \sim 4.2 \text{ mm s}^{-1}$ ) to the bigger observation channel ( $U_{\text{max}} \sim 0.046 \text{ mm s}^{-1}$ ), leading to a progressive loss of the alignment condition in the *z*-direction. Indeed, buoyant forces become competitive with respect to the other acting forces and the particles start falling along the flow direction.

Therefore, we want to compute the limit velocity of the process to estimate the horizontal length at which particles reach the channel bottom (L). To do this, the following

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assumptions need to be made: the settling is not affected by the presence of other particles in the fluid; the fluid can be considered a continuous medium. There is no slip interaction between the particles and the fluid molecules. The settling velocity becomes greater as both particle size and density are increased; the walls of the containing duct do not exert an appreciable retarding effect: this means that the drag coefficient is not enhanced.<sup>49</sup>

When the Reynolds number (Re, ESI<sup>†</sup>) is much less than 1, such a coefficient can be expressed as:

$$C_{\rm D} = \frac{24}{\rm Re} \left( 1 - \frac{9}{32} \left( \frac{1}{\zeta} \right) \right) \tag{5}$$

 $C_{\rm D}$  is the drag coefficient that we defined, for the first time, as a function of  $\zeta$  to establish a direct dependency on the particle diameter. This is reasonable since it has already been demonstrated that sedimentation processes in viscoelastic fluids are related to both object size and the relative position on the side and bottom walls of the channel where it moves.<sup>50–52</sup>

Due to sedimentation, particles assume a velocity different from that of the fluid, and the drag force can be determined from such a velocity difference. By solving the drag force expression (ESI<sup>†</sup>) as a function of the Re number, we get:

$$F_{\rm D} = 6\pi U_{\rm S} \eta a \left( 1 - \frac{9}{32} \left( \frac{1}{\zeta} \right) \right)^{-1}$$

where  $U_{\rm S}$  is the velocity at which falling occurs. When the stationary condition is reached (ESI<sup>†</sup>), the average  $U_{\rm Sz}$  expression becomes:

$$U_{\rm Sz} = \frac{F_{\rm Ez} + F_{\rm g} - F_{\rm b}}{6\pi\eta a} \left(1 - \frac{9}{32} \left(\frac{1}{\zeta}\right)\right) \tag{6}$$

As long as  $F_{Ez}$  is not more relevant for alignment ( $F_{Ez} < 10^{-14}$  N),  $U_{Sz}$  depends on the competition between drag and buoyant forces. Indeed, a smaller and lighter particle will fall more slowly to the bottom of the microchannel than a larger and heavier particle. To compute the sedimentation length *L*, we estimate the time needed from particles to move according to the fluid along *x* direction and the time to sediment along *z*.<sup>53,54</sup> Indeed, from  $U_S$ , we can compute the time  $t_z$  required for their motion along the *z*-direction:

$$t_z = \frac{H/2}{U_{\rm S}}$$

and the time  $t_{\rm L}$  required to move along the *x*-direction:

$$t_{\rm L} = \frac{L}{U_{\rm max}}$$

From *H*/2 to the bottom,  $t_z = t_L$  and:

$$L = \frac{U_{\text{max}}}{U_{\text{Sz}}} \frac{H}{2} \tag{7}$$

defined as the fall length.

#### **DoC** calibration

Thanks to the presented computation, we calculate first the *L* of PSL 6 and 10 beads for changing viscoelastic fluids as well as applied  $\Delta P$  (see Fig. 2a). As expected, PSL 6 show a significantly longer *L*, which scales with applied  $\eta_0$ . In addition,  $\eta_0$  influences the scaling of *L* with applied  $\Delta P$ . Note that we limited our investigations to fluid conditions, which prevent fluid turbulence during the alignment section as well as the transection phase from the inlet in the observation channel. For the present microfluidic design, fluid viscosity is significant for the alignment capability in the inlet capillary, while *L* remains less affected (see Fig. S5†). Next, we investigated PSL 10 and 6 shortly after the channel entrance (*X*1 position, see Fig. 2d) for 400 and 700 mbar of applied  $\Delta P$ .

As shown in Fig. 2b, good agreement between calculated and measured particle velocities was observed. Note that particles are perfectly aligned at the channel centre line  $(Y_{X1})$ , where the fluid velocity is the maximum of the parabolic profile. The accord between analytical and experimental results for  $U_{X1y}$  indicates that the change in cross-section between the capillary and observation section does not affect the alignment conditions. A knowledge of the starting position as well as the velocity is needed for a precise mass density signature calculation of single cells.

To better characterize the sedimentation process, we investigated PSL 10 beads for different values of  $\Delta P$  at consecutive distances from the channel inlet (see Fig. 2c). To follow the particle trajectory, the focus was altered by manually changing the distance between the microscope objective and the channel.

In this way, we were able to compare the settling dynamics modelled by the equations with the experimental observations. We defined the settling z-displacement of the beads as a 'reversed-migration' of the object, since the alignment is lost as the viscoelastic lifting force capability. Such a z-position depends on the channel dimension and on the difference between the local velocity of the object compared to that of the fluid.<sup>51,52</sup> Tracking different x locations, we estimated the variation in the object position along the channel height, since it moves at a velocity different from that of the fluid. It will migrate in reverse towards the wall, where it reaches zerovelocity and complete sedimentation. Such a dis-alignment will be faster or not, depending on the applied pressure (Fig. 2c). Higher pressure means a more sustained alignment of beads (4 cm is the length of sedimentation for the application of 700 mbar; Fig. 2c).

Therefore, from eqn (1) and the experimental bead velocity, we defined the *z*-coordinate at each *x*-position as follows:

$$z = \sqrt[n]{\left(1 - \left[\frac{U_{Xy}}{U_{\max}}\left(\left(\frac{m+1}{m}\right)\left(\frac{n+1}{n}\right)\right)\right]^{-1}\left(1 - \left(\frac{y}{W/2}\right)\right)^{-1}\right)\left(\frac{H}{2}\right)^{n}}$$
(8)

here reported as a generic formulation for any values of m and n.  $U_{Xy}$  is the local velocity of particles, which changes

along *x* as the particles fall, until the PSL are out-of-focus (the lowest detectable velocity is 5.134 µm s<sup>-1</sup>). This velocity can be regarded as a vectorial component of  $U_{Sz}$  computed for the estimation of *L* (eqn (7)). Therefore, eqn (8) solves the *z*-position of a particle which moves at a velocity  $U_{Xy}$ , which is different from the fluid  $U_{max}$ , as expected from the migration process due to the sedimentation conditions. Note that the observed *L* is in agreement with the value computed from eqn (7), with e% < 10 (Table S1†), further confirming the validity of the model prediction.

Good agreement between the computed and the experimental value of z is shown in Fig. 2c. However, as the presence of the change in cross-section (EXIT 1) becomes relevant, the difference between measurement and computation is greater far away from the inlet, since analysis of the particle motion is more challenging. Indeed, there is a match between simulated and observed particle positions up to 1.5 cm from the channel inlet.

Moreover, for higher concentrations of viscoelastic medium, we observed lower sedimentation particle velocities (see Fig. S5 and S6†). From the *z*-position and *L* computation, we show for the first time the possibility of measuring the unknown density of objects of different shapes, from their in-flow motion. To the best of our knowledge, both drag and density of objects have been studied only for streaming flows over spheres, freely falling into the fluid along a vertical direction.<sup>55</sup> Here, we propose the possibility of computing the density of moving objects inside Poiseuille fluid-flow conditions.

In detail, according to the hypothesis of an absence of viscoelastic contribution (eqn (3) and (4)) to the fluid flow, we determined the density ( $\rho$ ) of spherical objects from the expression of velocity migration due to sedimentation. As input parameters for the measure, we used the *z* obtained from eqn (8) compared to the initial position (*H*/2) of the objects at *X*1, *U*<sub>avg</sub> of the fluid (eqn (1)) and the experimental  $U_{X3y}$  as an estimate of the lag-motion of the particle at position *X*3, immediately before EXIT 1. Then,  $\rho$  was computed as:

$$\rho = \left(18\eta \frac{U_{\text{avg}} - U_{X3y}}{L_{\text{exp}}} \left(\frac{H}{2} - z\right) D^{-2}\right) + \rho_{\text{f}}$$
(9)

where *D* is the object diameter and  $\rho_f$  is the fluid density. A percentage error of less than 5% reveals a good match between the data sheet (from the bead manufacturer) values of PSL density values, for different-sized diameters, and the computed values (Table S4†).

To further establish the precision of the measurement, we defined the sensitivity of the system, which is related to any possible variation in the output density result due to a change into the object velocity. In detail, sensitivity can be estimated as the ratio between the variation in the output signal (density) and the change in the input velocity, which is the experimental parameter most involved in density measurement.<sup>56</sup> The sensitivity is found to be equal to  $\sim 0.001$ . The lower the sensitivity, the better is the performance of the approach. Moreover, we found that,

regardless of the imposed fluid-dynamic conditions, the limit of measurement of the density is dictated by the density of the fluid used, due to the desired buoyant effect. In our case, the limit is 998 kg m<sup>-3</sup>. Conversely, an upper limit to the measurement is essentially related to the maximum object diameter that we can align with a viscoelastic contribution. This means that the upper limit is related to the range of blockage ratios (see ESI†) useful for alignment purposes, in the range  $0.01 < \beta < 0.3$ , equivalent to a diameter of 0.75–22.5 µm.<sup>57</sup>

#### Cell sample characterization

Mass density signatures of human cells at single-cell level are difficult to obtain. Gao et al. recently demonstrated the first method based on magnetic levitation. Nevertheless, a more precise measurement approach is needed, which is also capable of separating cells of interest. Therefore, we investigated different peripheral blood stream cells (Fig. 3: LYM, RBC and NEU) and observed their fall length and velocity in the DoC microchannel. From particle observations, we concluded that different concentrations of viscoelastic medium would result in different z-positions and consequently also different cell velocities at fixed channel positions (X-position) but would not significantly change the cell-specific fall length. Therefore, we used a viscoelastic medium concentration (PEO 04), which ensured perfect cell alignment for the whole size range of peripheral blood cells. Moreover, we applied a  $\Delta P$  of 700 mbar, as the best compromise between cell alignment and measurement time.

To obtain unknown cell density values, a previous knowledge of the investigated cell size is needed. Therefore, we investigated the major axis of different cell classes by bright-field observation (see Fig. S4a<sup>†</sup>). Major cell diameters of 8.70 µm (LYM), 10.98 µm (NEU) and 8.23 µm (RBC) were calculated via ImageJ observations.58 We first investigated single-cell velocities  $(U_{Xy})$  in different observation channel positions while applying a constant pressure of 700 mbar (see Fig. 3). Directly after the inlet capillary, all cell types show similar cell velocities, in good agreement with particle investigations (see Fig. 2b). From X1 to X2, all 3 cell types significantly reduce their velocity. From measurement positions X2 to X6, RBC shows a reduction in cell velocity of  $\sim 10 \ \mu m \ s^{-1}$ , which indicates a long fall length, in good agreement with our simulated length. The observed cell velocity for LYM reduces significantly until the lateral exit position (X3). Some of the cells arrived at position X5, but no cells were observed in X6. Due to sample heterogeneity in the investigated lymphocyte class (T-LYM, B-LYM, natural killer cells and small fractions of monocytes) as well as the intercell type heterogeneity, no precise fall length could be established. In fact, by reducing the applied  $\Delta P$ , we can tune DoC to separate all cells from the LYM class or only a part (see Fig. 5b). In the case of the NEU cell class, the major cell fraction (4.4-fold) exits laterally for an applied  $\Delta P$  of 700 mbar.



**Fig. 3** Cell investigations with DoC. We investigated cell velocity at different positions of the measurement channel, where X1 to X6 correspond to 2, 40, 52, 57, 67 and 75 mm, respectively, from the inlet capillary. Histograms are normalized over sample number: 302, 417 and 165 for RBCs, LYM and NEU, respectively. In more detail, for RBC X1 = 6, X2 = 14, X3 = 61, X4 = 12, X5 = 31, and X6 = 178; for LYM X1 = 55, X2 = 167, X3 = 144, X4 = 47, and X5 = 4; for NEU X1 = 23, X2 = 51, X3 = 74, and X4 = 17. The microfluidic illustration indicates a measurement condition of 1000 mbar at PEO 04.

Besides the cell velocity, we were also interested in the change in the *Y*-position of cells, which is a good indicator of whether a cell will exit laterally or not. We plotted cells for each measurement position in X in one single scatter plot (see Fig. 4) per cell class. A clear transition in cell velocity reduction

to higher *Y*-position values was observed, which clearly shows the movement of cells in the direction of EXIT 1. Such phenomena were not observed for cell velocities greater 40  $\mu$ m s<sup>-1</sup>. In fact, a cell velocity of less than 40  $\mu$ m s<sup>-1</sup> at *X*3 will guarantee a higher possibility of lateral collection of cells. LYM



**Fig. 4** Cell velocity *versus Y*-position. The tracking of cells with PEO 04 at 700 mbar in the observation channel at different *X*-positions indicate the in-flow dynamics of cells. The number of analysed cells: 302, 417 and 165 for RBCs, LYM and NEU, respectively. In more detail, for RBC X1 = 6, X2 = 14, X3 = 61, X4 = 12, X5 = 31, and X6 = 178, for LYM X1 = 55, X2 = 167, X3 = 144, X4 = 47, and X5 = 4, for NEU X1 = 23, X2 = 51, X3 = 74, and X4 = 17.



**Fig. 5** Cell dynamics at EXIT 1. (a) The colour code indicates the *X*-position of cells during their travel in the *X*3 section of the observation channel. (b) Top-view of EXIT 1 with highlighted cell tracking outcome. Fraction of cells exiting lateral (coloured) *versus* cells continuing (black) in the inflow direction are shown. The number of analysed cells for 400 mbar: 14 and 53 for RBCs and LYM, respectively. For 700 mbar: 61, 144 and 74 for RBCs LYM and NEU, respectively.

measured at  $Y_{X3}$  below 500 µm are likely to be RBCs, showing significantly higher cell velocities compared to previous *X*-positions. Note that cell types were not separated with a cellspecific selection approach. A certain percentage of different cell types could be present in the measurement sample. Moreover, the self-written MATLAB routine considered all passing objects, without using any object exclusion related to the observed object size or shape. This guarantees a clear sight of possible outliers, which is fundamental for a proof-ofconcept study. However, cells which reduced their velocity below 15 µm s<sup>-1</sup> are considered as having fallen (touching the bottom of the observation channel). Note that cells tend to roll on the bottom of the microfluidic channel, so a minimum cell velocity is always present.

We investigated the cell dynamics at EXIT 1 for 400 as well as 700 mbar of  $\Delta P$  (see Fig. 5a). Interestingly, LYM accelerate in the last part of journey before exiting the observation channel (see Fig. S8<sup>†</sup>). This phenomenon of lateral capillary aspiration must be considered for velocity-based cell trajectory analysis. In general, we can confirm that for higher  $Y_{X3}$  values, a lower  $U_{X3y}$  can be observed. In the case of NEU a  $\Delta P$  of 400 mbar is insufficient to reach the lateral exit position (computed L = 3 cm), while tuning of the applied  $\Delta P$ can be used to find the optimal cell class separation. For instance, in the case of LYM cells, 550 mbar showed the threshold value for EXIT 1 separation of ~40% of cells (see Fig. S9<sup>†</sup>). Lower  $\Delta P$  (e.g. 400 mbar) results in perfect separation efficiency. In the case of NEU, a value of 630 mbar was experimentally investigated, while for RBCs a significant reduction in the applied  $\Delta P$  was needed (see Fig. S11<sup>†</sup>). The balance between applied  $\Delta P$  and effective cell velocity during their journey in the observation channel is limited by a minimum  $\Delta P$  of *circa* 250 mbar for the PEO 04 case (see Fig. S10<sup>†</sup>). To reduce the applied  $\Delta P$ , the fluid viscosity can be

reduced, which in the case of PEO 03 resulted in an applied  $\Delta P$  of 150 mbar (data not shown).

We also analyzed the fractions of separated and not separated cells under two different  $\Delta P$  conditions (400 and 700 mbar). NEU show a separation of 90% for 700 mbar, while LYM show their highest separation efficiency for 400 mbar, at 100% (see Fig. 5b). RBCs were found never to exit from EXIT 1. Note that cell tracking information was not filtered according to different cell classes during the measurement process. A more precise cell class selection could improve the fractioning of cells at EXIT 1. Interestingly, cells which pass EXIT 1, but significantly reduce their velocity are also likely to increase their  $Y_{X3}$  position. An investigation of the final  $Y_{X3}$  coordinate *versus* resulting  $U_{Xy}$  cell velocity could be used as an additional density-based indication (see Fig. S11†).

Therefore, from the combination of the applied fluid-flow conditions and the relative motion of cells, a single-cell characterization with diameter and density measurements was performed.

First, to predict where cells sediment, we computed the relative *L* for the different cell types, using density values obtained from the literature. In more detail, we used density values of 1.072 kg m<sup>-3</sup> (LYM), 1.086 kg m<sup>-3</sup> (NEU) and 1.099 kg m<sup>-3</sup> (RBC), which resulted in fall lengths (see Fig. S12†) of 7.2 (LYM), 4.8 (NEU), and 9.5 cm (RBC). To guarantee the highest possible exit of cells at EXIT1 and/or at EXIT2, we used  $\Delta P$  of 400 mbar, 550 mbar and 700 mbar for RBC, LYM and NEU, respectively. Secondly, we compared the obtained results with the real fall lengths measured at the same pressures used for computation. With e% < 20%, good agreement between the experimental and estimated results was found (see Fig. S13†). Then, we decided to perform single-cell measurement of cell densities, starting from an

evaluation of cell diameters and velocities. The fall length used is the one measured for an entire cell population.

We found good agreement with literature values for both D and  $\rho$  (from eqn (9); see also Tables S2 and S3†). A mean value of 1093 kg m<sup>-3</sup> for NEU was found. In the case of LYM (the most heterogenic cell class), a density between 1.005 and 1.145 kg m<sup>-3</sup> with a mean value of 1063 kg m<sup>-3</sup> was calculated (Fig. 6).

Despite the higher density, RBCs fall at a longer distance (EXIT 2) than other cells. This is due to the cell-specific shape being oblate. In detail, these cells have a third axis smaller than the other two axes that are equal to each other (Fig. 1). In the case of an oblate shape, the drag force is less than would be exerted on a sphere equal to the equatorial radius of the spheroid.<sup>55,59</sup> Therefore, a scaling factor affecting the way in which drag effect changes on an oblate shape was necessary to compute  $\rho$  for RBCs. Happel and Brenner presented the correction factor in a hypothesis for a flow streaming past an oblate spheroid. The expression is:<sup>55</sup>

$$K = \frac{1}{\frac{3}{4}\sqrt{\left(\lambda^2 + 1\right)\left[\lambda - \left(\lambda^2 - 1\right)\cot^{-1}(\lambda)\right]}}$$

where  $\lambda = \left[ \left( \frac{D/2}{b} \right)^2 - 1 \right]^{-1/2}$ , for which *D* and *b* are the diameter

and the third semi-axis of the RBC, respectively. Then, a scaling of the drag coefficient (eqn (5)) as  $C_{\text{DCorr}} = KC_{\text{D}}$  can be written. The final expression of  $\rho$  for an oblate can be reported as:

$$\rho_{\rm Obl} = \left(18\eta \frac{U_{Xy}}{L_{\rm exp}} \left(\frac{H}{2}\right) \frac{1}{K} D^{-2}\right) + \rho_{\rm f} \tag{10}$$

For our imposed fluid-flow conditions, RBCs still move along the center line of the channel from the alignment reached inside the Inlet capillary. Therefore, the velocity  $(U_{X4,5y})$ measured at X4 and X5 (Fig. 3), where motion analysis was performed, is still equal to  $U_{\text{max}}$  and no lag motion is observed. Indeed, H/2 also does not scale with respect to a



**Fig. 6** Cell density computation from experimental in-flow motion. The number of analysed cells: 22, 78, 15 for RBCs, LYM and NEU, respectively. Mean  $\pm$  std. dev.:  $D = 8.25 \pm 0.97 \ \mu\text{m}, \ \rho = 1159 \pm 29.5 \ \text{kg} \ \text{m}^{-3}$  for RBCs;  $D = 7.25 \pm 0.73 \ \mu\text{m}, \ \rho = 1073 \pm 49 \ \text{kg} \ \text{m}^{-3}$  for LYM;  $D = 10.66 \pm 1.15 \ \mu\text{m}, \ \rho = 1093 \pm 27 \ \text{kg} \ \text{m}^{-3}$  for NEU.

different *z*-position. As a result, the single-cell mass density signature was obtained for RBCs (Fig. 6).

A scaling of  $\rho$ -values with respect to D and  $U_{X4,5y}$  is presented. As expected, from the oblate shape, despite the higher density, RBCs do not fall and reach the end of the microfluidic device. Moreover, even if RBCs are similar in size range to LYM, eqn (10) well demonstrates that sedimentation behavior is influenced not only by the size but also by the shape of the object. Indeed,  $U_{X4,5y}$  is also the highest for the RBC class (Fig. 6). NEU and LYM have lower densities than RBCs and their falling motion is dictated only by a difference in size and intrinsic  $\rho$ , which are higher for NEU than for LYM.

### Conclusions

The presented densimeter-on-chip (DoC) is a label-free and easy-to-use tool for single-cell mass density measurement. During the last few years, knowledge of cell density (lower cell-to-cell variation among the same types compared to other biophysical features) has been regarded as a key factor for single-cell classification, due to the great relevance of density in cell state control and regulation. However, having an approach capable of recognizing and analysing different density levels of cells belonging to the same cell type is still challenging.

Here, thanks to a triggered imbalance of hydrodynamic components—as viscoelastic, buoyant and drag forces—DoC proposes to map both single-cell diameters and densities, thanks to a non-invasive, label-free in-flow sedimentation process.

In detail, we induced the traceable gravity-based sedimentation of cells into different device exits. Additionally, the chip is designed to be versatile, with the possibility of choosing the fluid flow characteristics, such as viscosity, relaxation time and applied in-flow velocity, to achieve precise control over the sedimentation process. Moreover, since the chip is conceived to be a passive type of technique, no other operational parameters, such as external magnetic, electric or acoustic field intensities, need to be controlled. Depending on dimension and shape, the approach allows us to measure and distinguish different density levels for multiple cell types. The working range of the device is related to the applied inlet pressure and object diameters. The former varies from ~250 mbar up to ~2500 mbar, with the lowest detectable value of 5.134  $\mu$ m s<sup>-1</sup>. The latter has a wide range of possible dimensions, from 0.75 to 22.5 µm, according to the needs of the microchannel blockage ratio. Instead, the value of 998 kg m<sup>-3</sup> is the lowest value of measurable density. In detail, multiple types of object, such as PSL microspheres or peripheral blood stream cells-namely RBCs, LYM and NEU-are forced to pass through an inlet round capillary, where the alignment along the centre line is guaranteed by the action of a lifting viscoelastic fluid force-greater than the buoyant components-counterbalanced by the drag force. Next, cells

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pass from the capillary to a larger rectangular cross-section, which triggers a dramatic reduction in viscoelastic forces, which on the other hand leads to an enhancement in gravitydriven motion for a fall of cells along the vertical direction of the channel. According to the hypothesis of spherical shape, cells sediment according to their size and density. On the other hand, despite their intrinsically higher density, RBCs stably follow the centre line of the channel, thanks to a spheroidal oblate shape for which the viscoelastic force still applies a lifting effect, which results in an extended fall length. Thanks to a new formulation of governing forces, we implemented a set of equations that describe fluid flow conditions and the imbalance in forces, by setting only cell dimension and shape. In such a way, we were able to extrapolate the single-cell mass density of PSL, RBCs, LYM and NEU with a great agreement with the data in the literature. Moreover, the results from the approach are highly precise and reproducible with an intrinsically low sensitivity in density measurement of 0.001. However, in a future perspective, we would further enhance the precision and reproducibility of the system by improving the hardware for complete motion reconstruction with object acquisition in both Y and Z directions to allow an accurate reconstruction of both velocity and sedimentation profiles.

Moreover, the creation of a platform where both density and deformability measurements can be performed would allow a comprehensive characterization of cell properties, providing a complete phenotyping.

Indeed, with the capability of efficient and simple operating conditions, we believe that the proposed DoC approach provides a valid alternative for next-generation microfluidic density based single-cell characterization and manipulation platforms.

## Data availability

The data supporting this article have been included as part of the ESI.<sup>†</sup> For any additional requests or queries regarding the data, please contact David Dannhauser at david.dannhauser@unina.it.

## Author contributions

D. D.: conceptualization, methodology, validation, writing original draft, formal analysis. M. I. M.: conceptualization, mathematical model, validation, writing original draft, formal analysis. P. A. N.: supervision. F. C.: conceptualization, writing-review, & editing, supervision.

# Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors thanks Claudia DeClemente for valuable discussion.

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