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# Single-cell impedance spectroscopy of nucleated cells<sup>†</sup>

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Single-cell microfluidic impedance spectroscopy is widely used to characterise single cells, but the intrinsic electrical properties are rarely determined owing to the limited number of data points across a wide frequency bandwidth. To address this shortcoming, we have developed a system with an extended frequency range (to 550 MHz) that measures the impedance spectrum of single nucleated cells at high throughput. The system was evaluated using HL60 cells treated with glutaraldehyde or cytochalasin D, and THP-1 cells differentiated into macrophages. The impedance data was fitted to the double-shell model to obtain cell membrane capacitance and cytoplasm conductivity. It is shown that reducing the conductivity of the suspension media significantly enhances the dielectric relaxations of the cell membrane, allowing small differences between control and chemically modified cells to be discriminated.

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## 1. Introduction

The electrical properties of single cells have been widely used as potential label-free markers of cell phenotype, and microfluidic impedance cytometry (MIC) can rapidly characterise the electrical properties of individual cells, for a recent review see.<sup>1</sup> MIC has been used to explore a range of cell properties including viability,<sup>2,3</sup> identification of leukocyte types in whole blood,<sup>4,5</sup> activation state of macrophages,<sup>6,7</sup> parasite invasion of red blood cells,<sup>8,9</sup> identification of cancer cells,<sup>10,11</sup> and antimicrobial susceptibility.<sup>12,13</sup>

In microfluidic impedance cytometry, cells are usually measured at two or at most three discrete frequencies, and categorisation is typically made according to the value of the impedance signals, for example, amplitude, phase, and/ or electrical opacity (the ratio of high to low frequency impedance). However, this limited range of measurements is insufficient to determine the intrinsic electrical properties of each single cell (*e.g.* membrane capacitance), which requires fitting the frequency-dependent impedance data to a set of Debye relaxations,<sup>14</sup> using, for example, a multi-

shell model. Too few frequency points and/or a limited frequency range leads to poor fits that return cell electrical properties with a high degree of uncertainty, limiting the applicability of the method to distinguish small changes between different cells. To address this, the properties of single cells need to be measured across a wide range of frequencies. Early work by Sun et al.15 used maximum length sequence (MLS) analysis to measure the properties of polystyrene beads with 512 evenly distributed frequencies between approximately 1 kHz and 500 kHz, but this method had a limited frequency bandwidth. Some groups have attempted to extrapolate particle dielectric parameters from a limited range of frequencies. For example, Chai et al.<sup>16</sup> used giant liposomes of known radius, membrane permittivity and cytoplasm conductivity and fitted a single dispersion to four discrete frequencies up to 1.2 MHz. The dielectric properties of the liposomes were extracted using the singleshell model.<sup>17</sup> Zhao et al.<sup>18,19</sup> used a constriction channel to measure the capacitance of single cells at two low frequencies, up to 250 kHz. Using equivalent electrical circuits and without considering the nucleus in the model, they claimed to extract the cell membrane capacitance. Haandbæk et al.20 characterised different yeast cell phenotypes at frequencies up to 20 MHz, fitting data to a shell model including the cell wall.

recently described а single-cell dielectric We spectroscopic method capable of characterising the properties of thousands of single cells in the frequency range from 200 kHz to 50 MHz with 8 simultaneous frequencies.21 The system was evaluated using model particles that have no nucleus (red blood cells (RBCs) and ghosts), where the 50 MHz upper

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frequency limit was sufficient to fit the data to the singleshell model. However, this frequency range is insufficient to fully characterise the electrical properties of nucleated mammalian cells suspended in physiological saline, where the frequency needs to extend to >500 MHz. In this paper, we describe an improved system capable of highspeed multi-frequency characterisation of nucleated cells. In order to extract the dielectric properties of each individual cell, single-cell spectra were fitted to the double-shell model by post-processing the data, although real-time machine learning methods could also be used.<sup>22</sup> Throughput is around 250 cells per second and the system performance was demonstrated by measuring the intrinsic dielectric properties of different cells.

Classically, the dielectric properties of nucleated (mammalian) cells are usually measured in low-conductivity media. This is done for several reasons: to minimise issues with electrode polarisation which is much worse for a high conductivity media (thin double layer), and also to minimise Joule heating in the medium, particularly for single-cell electrorotation studies where the electric field can be very high. Furthermore, because the cell membrane characteristic relaxation frequencies decrease in a lowconductivity suspending medium, it is easier to generate the maximum frequency of the measurement system e.g. 10 MHz. To avoid osmotic stress, cells are resuspended in a single-cell non-metabolising sugar solution. However, impedance cytometry is generally performed using physiological saline to maximise the current through the micro-channel and the signal to noise (SNR). To avoid issues with electrode polarisation reference beads can be used to calibrate the transfer function of the system.<sup>20,21</sup> However, as shown here, there is a trade-off between optimising the suspending medium conductivity for SNR (high conductivity) and the ability to resolve discrete Debye relaxations (lower conductivity).

#### 2. Theory

#### 2.1. Shell model and mixture theory

The multi-shell model developed by Irimajiri *et al.*<sup>23</sup> has been widely used to determine the intrinsic dielectric properties of cells from experimental impedance data, both for the case of cell suspensions<sup>24</sup> and for single-cell electrorotation measurements.<sup>25</sup> In this model the complex permittivity  $\varepsilon_{mix}^*$  of a cell in suspension is related to the cell ( $\varepsilon_{cell}^*$ ) and suspending media ( $\varepsilon_{med}^*$ ) by Maxwell mixture's theory (MMT):

$$\varepsilon_{\text{mix}}^{*} = \varepsilon_{\text{med}}^{*} \frac{1 + 2\varphi_{\text{cell}} f_{\text{CM(cell)}}^{*}}{1 - \varphi_{\text{cell}} f_{\text{CM(cell)}}^{*}} = \varepsilon_{\text{med}}^{*} \frac{1 + 2\varphi_{\text{cell}} \left(\frac{\varepsilon_{\text{cell}}^{*} - \varepsilon_{\text{med}}^{*}}{\varepsilon_{\text{cell}}^{*} + 2\varepsilon_{\text{med}}^{*}}\right)}{1 - \varphi_{\text{cell}} \left(\frac{\varepsilon_{\text{cell}}^{*} - \varepsilon_{\text{med}}^{*}}{\varepsilon_{\text{cell}}^{*} + 2\varepsilon_{\text{med}}^{*}}\right)} \quad (1)$$

where a general complex permittivity is defined as  $\varepsilon_i^* = \varepsilon_i \varepsilon_0 - j \frac{\sigma_i}{\omega}$ ,  $\varphi_{cell}$  is the volume fraction of the cell in the measurement volume, and the Clausius–Mossotti factor of a cell is defined as:

$$f_{\rm CM(cell)}^* = \frac{\varepsilon_{\rm cell}^* - \varepsilon_{\rm med}^*}{\varepsilon_{\rm cell}^* + 2\varepsilon_{\rm med}^*}$$
(2)

The complex permittivity of the cell is modelled as a series of concentric shells, as illustrated in Fig. 1(a), which shows the double-shell model as applied to a nucleated cell. This model relates the dielectric parameters of each individual shell to the complex dielectric properties of the cell according to:<sup>14,26</sup>

$$\varepsilon_{\text{cell}}^{*} = \varepsilon_{1}^{*} \frac{2(1-\gamma_{a}) + (1+2\gamma_{a})E_{a}}{(2+\gamma_{a}) + (1-\gamma_{a})E_{a}}$$
(3)

where the subscript "1" indicates cell membrane. The factor  $y_a$  is given by  $y_a = (R_a/r_{cell})^3$ , with  $r_{cell}$  the cell radius,  $d_1$  the thickness of the cell membrane and  $R_a = r_{cell} - d_1$ . The parameter  $E_a$  is a factor that depends on layer dimensions and dielectric parameters, see ESI† for full derivation of the double-shell model. Note that the cell membrane capacitance  $C_{mem}$ , cytoplasm conductivity  $\sigma_{cyt}$ , and cell diameter d are determined from  $C_1 = \varepsilon_1/d_1$ ,  $\sigma_2$ , and  $2r_{cell}$  in the model shown in Fig. 1(a). Substituting eqn (3) into eqn (2) gives the Clausius–Mossotti factor for the cell  $\left(f_{CM(cell)}^*\right)$  as a function of the suspending media, cell dielectric properties and dimensions.

Fig. 1(b) and (c) shows example plots of  $f^*_{CM(cell)}$  calculated using the double-shell model for a "typical" nucleated cell (for details, see figure legend) suspended in electrolytes of different conductivities (corresponding to four-fold dilutions of physiological saline). The figure demonstrates how the spectrum changes with suspending medium conductivity. The real part of the polarizability is always negative except for the lowest conductivity, and the magnitude of the frequency dispersion (peak in the imaginary part) is the greatest at the lowest conductivity.

Fig. 1(d)–(g) demonstrate how variations in cell dielectric properties manifest themselves in different frequency windows for the highest and lowest suspending medium conductivities. The low-frequency limit of the Clausius–Mossotti factor (~100 kHz) is dominated by the cell volume (assuming the particle to be a perfect insulator) and to a lesser degree by the membrane capacitance. For a fixed specific membrane capacitance, the low frequency relaxation is particle size dependent, moving to higher frequencies for smaller cells. Higher membrane capacitance shifts this relaxation peak to a lower frequency whilst the cytoplasm conductivity impacts the spectra at high frequencies >10 MHz.

Comparing the spectra for these two conductivities shows that the relative changes due to variations in cell dielectric parameters are more pronounced in the lower conductivity. In terms of impedance measurements, the absolute value of the current between the two electrodes is lower (by approximately a factor of 5), but the relative change in impedance is greater at 0.32 S m<sup>-1</sup> than in physiological saline (1.6 S m<sup>-1</sup>). The single-shell model (Fig. S1†), shows

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Fig. 1 Double-shell model and calculated Clausius–Mossotti factor  $(f_{CM(cell)}^*)$  (a). Diagram showing the multi-shell model used to characterise a nucleated cell, subscript 1: cell membrane, 2: cytoplasm, 3: nuclear envelope, 4: nucleoplasm, med: media – see ESI† for further details. Note that the cell membrane capacitance  $C_{mem}$ , cytoplasm conductivity  $\sigma_{cyt}$  and cell diameter *d* correspond to  $C_1$  ( $\varepsilon_1/d_1$ ),  $\sigma_2$ , and  $2r_{cell}$ . (b) and (c). Plots of the real and imaginary parts of the Clausius–Mossotti factor for a "typical" cell as a function of frequency for different suspending media conductivities, modelled using the double-shell model. Cell parameters were: diameter = 12  $\mu$ m; membrane capacitance = 12.4 mF m<sup>-2</sup>; cytoplasm conductivity = 0.3 S m<sup>-1</sup>; for other parameters, see ESI† Table S1. (d)–(g) Illustration showing how changes in cell dielectric parameters influence the real and imaginary parts of  $f_{CM(cell)}^*$ . Diameter range (orange region), 8–24  $\mu$ m, membrane capacitance (red region), 5.3–19.5 mF m<sup>-2</sup>, and cytoplasm conductivity (blue region), 0.1–0.5 S m<sup>-1</sup>. Plots are calculated for 0.32 S m<sup>-1</sup> and 1.6 S m<sup>-1</sup> conductivity electrolyte. The black curves are for "typical" values of cell dielectric parameters, as in (b) and (c).

similar trends in terms of conductivity but also illustrates that this is inadequate for modelling a nucleated mammalian cell and would lead to incorrect values of cell membrane and cytoplasm dielectric properties. Fig. 1 also demonstrates that in order to measure and characterise nucleated cells, it is imperative that the measurement frequency extends to hundreds of MHz to allow accurate data fitting with the double-shell model.

#### 2.2. Experimental principle

The microfluidic impedance cytometer along with the workflow is shown in Fig. 2. The measurement chip is made

from glass and consists of a microchannel 30  $\mu$ m in height and 40  $\mu$ m in width, with several micrometer-sized electrodes. An AC voltage is applied to one set of electrodes while the opposite sensing electrodes measure the differential current (ESI† Fig. S2). As a cell transits through the channel, a bipolar Gaussian differential signal is generated, with an amplitude related to the cell impedance at that applied frequency.

The current in the empty channel should ideally be invariant with frequency, but the actual system consists of many circuit elements, which means that the impedance of the empty cell is non-linear. To overcome these nonlinearities, polystyrene beads of known size and dielectric



Fig. 2 System overview and measurement principle. Cells suspended in saline flow along the microchannel where microelectrodes measure the differential current at different frequencies ranging from 250 kHz to 550 MHz. After gating and calibration, the frequency spectrum was fitted to the double-shell model to give cell diameter and cell dielectric parameters (membrane capacitance:  $C_{mem}$ , cytoplasm conductivity:  $\sigma_{cyt}$ ).

properties are used as internal calibration particles.<sup>20,21</sup> Because the complex permittivity of the bead is known and fixed, the complex permittivity of the cell can be determined from the ratio of particle volumes.<sup>21</sup> For spherical cells dispersed in a suspending media at a low volume fraction,  $(\varphi_{cell} < 0.01)$ , the differential complex current of the mixture is given by:<sup>21</sup>

$$\Delta i_{\rm mix}^* = \varepsilon_{\rm med}^* \varphi_{\rm cell} f_{\rm CM(cell)}^* \tag{4}$$

and the ratio of the complex current between a cell (unknown) and a bead (known) is:

$$\frac{\Delta t_{\text{cell}}^*}{\Delta t_{\text{bead}}^*} = \frac{\varphi_{\text{cell}} f_{\text{CM}(\text{cell})}^*}{\varphi_{\text{bead}} f_{\text{CM}(\text{bead})}^*} \tag{5}$$

Because  $\varphi_{\text{bead}}/\varphi_{\text{cell}} = r_{\text{bead}}^3/r_{\text{cell}}^3$ , the Clausius–Mossotti factor can be expressed as

$$f_{\rm CM(cell)}^* = \frac{r_{\rm bead}^3}{r_{\rm cell}^3} f_{\rm CM(bead)}^* \times \frac{\Delta i_{\rm cell}^*}{\Delta i_{\rm bead}^*}$$
(6)

Eqn (2) and (6) are two different expressions of the same parameter,  $f^*_{\rm CM(cell)}$ . The equations show that the dielectric spectrum of the cell can be determined from the experimentally measured differential currents ( $\Delta i^*_{\rm cell}$  and  $\Delta i^*_{\rm bead}$ ) provided the frequency-dependent dielectric properties of the beads ( $r_{\rm bead}$  and  $f^*_{\rm CM(bead)}$ ) are known according to:

$$f_{\rm CM(cell)}^* \frac{r_{\rm bead}^3}{r_{\rm cell}^3} = f_{\rm CM(bead)}^* \times \frac{\Delta i_{\rm cell}^*}{\Delta i_{\rm bead}^*}$$
(7)

The impedance signal is the differential current for the cell/bead multiplied by the Clausius–Mossotti factor for the bead (right-hand side of eqn (7)). Given that the radius and dielectric properties of the bead are known then from eqn (7), the radius and unknown dielectric parameters of the cell  $f_{CM(cell)}^*$  can be determined by fitting the data to the shell model. In this case, the experimental data was fitted using the least-squares minimisation algorithm "pattern-search" (MATLAB). The fit determines cell size and the volume ratio of nucleus to cell, along with dielectric parameters ( $\varepsilon_1$  to  $\varepsilon_4$ , and  $\sigma_2$  to  $\sigma_4$ ). For the fitting, these parameters were allocated starting values and boundaries as in ESI† Table S2. The cell membrane conductivity,  $\sigma_1$  was fixed to  $10^{-10}$  S m<sup>-1</sup> (assuming a viable cell).

#### 3. Methods

#### 3.1. Experiment setup

The impedance chips (Fig. 2) were fabricated using standard photolithography and thermo-compression bonding as outlined elsewhere.<sup>27</sup> Electrodes (Pt) were patterned onto 700  $\mu$ m glass wafers followed by deposition and patterning of a thick resist to create channels 40  $\mu$ m wide and 30  $\mu$ m high. A pair of wafers were bonded together and the final

microfluidic chips were clamped into a custom PEEK holder to provide electrical and fluidic connections. An overview of the chip fabrication protocol is shown in Fig. S3.<sup>†</sup>

Cells were suspended at a concentration of  $5 \times 10^5$  ml<sup>-1</sup> in saline of different conductivities and mixed with polystyrene reference beads (latex carboxylate-modified polystyrene beads, Sigma Aldrich) at a concentration of around  $2.5 \times 10^5$ ml<sup>-1</sup>. The suspension was pumped through the microfluidic chip at a flow rate of 30 µl min<sup>-1</sup>. A lock-in amplifier (UHFLI, Zurich Instruments) generated an applied voltage  $V_{AC}$  in the frequency range from 250 kHz to 550 MHz. The current signals from the chips were amplified with custom electronics and demodulated with the lock-in. For single-cell spectroscopy, eight frequencies were applied simultaneously to the impedance electrodes. Data was processed using custom MATLAB scripts to extract the signal amplitude at different frequencies for each particle.

Two different conductivity saline solutions were used (a) normal physiological saline at 1.6 S m<sup>-1</sup> (0.9% w/v) with an osmolarity of 280 to 300 mOsm and (b) 5× diluted saline (0.32 S m<sup>-1</sup>) with p-mannitol (in water) to adjust the osmolarity to the range of 280 to 300 mOsm. Prior to use, the solutions were filtered through a 0.22  $\mu$ m filter.

#### 3.2. Cells

Two different cells were used in the experiments, HL60 and THP-1. The cells were grown in the same culture media, consisting of RPMI 1640 + Glutamax (Gibco) media, 10% fetal bovine serum (FBS, Gibco), and 1% penicillin-streptomycin (Sigma Aldrich) in a humidified incubator at 37 °C and 5% CO<sub>2</sub>. Cells were harvested for treatment and measurement at a concentration of  $5 \times 10^5$  cells per ml.

HL60 cells were treated with glutaraldehyde (GA, 11428743, Thermo Scientific Chemicals) as follows. GA-DPBS solution at a concentration of 1% v/v was added to a 1 ml suspension of cells in DPBS (11590476, Gibco). The sample was kept at room temperature for 30 minutes and then resuspended in saline (conductivity 0.32 S m<sup>-1</sup> or 1.6 S m<sup>-1</sup>). Cytochalasin D (CytoD, C2618, Sigma Aldrich) was used to disrupt HL60 cell cytoskeletons. Stock solutions of CytoD were dissolved in DMSO (A13280.36, Thermo Scientific Chemicals) and added to 1 ml cell samples in 0.32 S  $m^{-1}$  or 1.6 S  $m^{-1}$  media to adjust the concentration to 10  $\mu M.$  After incubating the cells at 37 °C incubator for 10 minutes, the sample was kept at room temperature for around 1 minute so that the conductivity would return to the correct value. The untreated group was added to the same amount of DMSO and also kept at 37 °C for 10 minutes, following the same protocol as the treated group.

THP-1 differentiation was achieved by adding phorbol 12-myristate 13-acetate (PMA, P1585, Sigma Aldrich) into the cell suspension (in culture media) followed by incubation at 37 °C, 5% CO<sub>2</sub> for 48 h. THP-1 cells were harvested at a concentration of  $5 \times 10^5$  ml<sup>-1</sup> and resuspended in fresh culture media. Stock PMA dissolved in DMSO was added to



Fig. 3 Plots of the normalised Clausius-Mossotti factor (real and imaginary) for HL60 cells suspended in low and high-conductivity saline. The spectra (blue: real part, red: imaginary part) for untreated (solid lines with dots) and fixed cells (dashed lines with circles) were measured in two different saline conductivities (a) 0.32 S  $m^{-1}$  with 7  $\mu$ m diameter beads and (b) 1.6 S m<sup>-1</sup> with 5  $\mu$ m diameter beads. (c) Normalised impedance spectra (blue: real part, red: imaginary part) of untreated HL60 cells (solid lines with dots) and CytoD-treated (dashed lines with circles) cells in 0.32 S  $m^{-1}$  media with 7  $\mu$ m beads. This is a representative plot for three repeats (n = 3). In (a), (b) and (c), dots and circles are individual data points. Solid and dashed lines fit to the double-shell model. (d) Mean and S.D of  $C_{\rm mem}$  for untreated and CytoD treated cells for three independent repeats (n = 3), demonstrating a significant change (\*\*\*,  $p \leq 0.001$ , Students' t-test) in membrane capacitance. The suspending media relative permittivity was set to 80, cell membrane conductivity was fixed to  $10^{-10}$  S m<sup>-1</sup> membrane thickness to 5 nm and nuclear envelope thickness to 20 nm. Bead permittivity =  $2.5\varepsilon_0$ , conductivity = 0.57 mS m<sup>-1</sup>.

adjust the concentration to 200 ng ml<sup>-1</sup>. The untreated THP-1 group was added to the same amount of DMSO as a control. The two different cell suspensions were both moved to 6-well flasks and placed in an incubator. After 48 h, the differentiated macrophages (M0) were bathed in a thin layer of TrypLE Express (12605010, Gibco) and placed in an incubator for 10 minutes, then resuspended in culture media without PMA at  $5 \times 10^5$  ml<sup>-1</sup> for further measurement. A control group was also resuspended in

culture media at the same concentration. Samples were resuspended in 0.32 S  $m^{-1}$  conductivity saline.

## 4. Results

# 4.1. Mean dielectric properties and influence of suspending media conductivity

Fig. 3(a) and (b) show example impedance-frequency spectra for a population of untreated HL60 cells (solid dots) and with glutaraldehyde (unfilled cells fixed circles). Glutaraldehvde treatment of cells cross-links both membrane lipids and proteins,28 making the cells nonviable. The data was collected using two frequencies, a fixed reference frequency of 18 MHz and a second frequency that was swept between 250 kHz and 550 MHz at 18 discrete frequencies. Beads and cells were gated using the 18 MHz phase scatter data to maximise the signal to noise ratio (SNR), defined as the mean square ratio of noise to particle sub-populations (Fig. S4<sup>†</sup>). The spectrum is, therefore, the mean value of impedance at each frequency for the population of cells. The data is the normalised Clausius-Mossotti factor determined using eqn (7) scaled by the ratio of the volume of cells to beads. The numbers of cells at each frequency point are summarised in Table S3.† Data was collected using independent batches of cells on different days. The solid curves (untreated cells) and dashed curves (fixed cells) are best fits to the double-shell model for the real (blue) and imaginary (red) parts of the spectrum. The cell diameter (d), membrane capacitance  $(C_{\text{mem}})$  and cytoplasm conductivity  $(\sigma_{\text{cvt}})$  were extracted from the fit as described in section 2.2. Table 1 summarises the data set for untreated and fixed cells suspended in saline at 0.32 S m<sup>-1</sup> ( $R^2 > 0.98$ ), and 1.6 S m<sup>-1</sup> ( $R^2 > 0.90$ ). For physiological saline, the fixed and unfixed cell diameters were similar. The two key dielectric parameters ( $C_{mem}$  and  $\sigma_{\rm cvt}$ ) were also close ( $C_{\rm mem}$ : 11.07 mF m<sup>-2</sup> vs. 11.52 mF m<sup>-2</sup>,  $\sigma_{\text{cvt}}$ : 0.31 S m<sup>-1</sup> vs. 0.33 S m<sup>-1</sup>). In the lower-conductivity media, there were slight differences between untreated and fixed cell diameter;  $C_{
m mem}$  and  $\sigma_{
m cyt}$  decreased after glutaraldehyde exposure ( $C_{mem}$ : 9.14 mF m<sup>-2</sup> reducing to 6.32 mF m<sup>-2</sup>;  $\sigma_{cvt}$ : 0.33 S m<sup>-1</sup> reducing to 0.07 S m<sup>-1</sup>). The mean  $\pm$  SD of  $\sigma_{cvt}$  calculated from three repeat experiments decreased from 0.29  $\pm$  0.04 S m<sup>-1</sup> to 0.08  $\pm$  0.02 S m<sup>-1</sup> after glutaraldehyde exposure (Table S4<sup>†</sup>). This is consistent with previous studies showing that  $C_{\rm mem}$  and  $\sigma_{\rm cyt}$  of RBCs (no nucleus) decreased after fixing.29-31 A plot demonstrating that the single-shell model does not fit the data is presented

Table 1 Parameter set from fits for untreated and fixed HL60 cells as in Fig. 3(a) and (b). For confidence intervals (CI) see Table S6 in ESI†

		Diameter ( <i>d</i> , μm)	Membrane capacitance $(C_{\text{mem}}, \text{mF m}^{-2})$	Cytoplasm conductivity ( $\sigma_{ m cyt}$ , S m <sup>-1</sup> )
1.6 S m <sup>-1</sup>	Untreated	11.6	11.07	0.31
	Fixed	11.7	11.52	0.33
$0.32 \text{ Sm}^{-1}$	Untreated	12.0	9.14	0.33
	Fixed	12.6	6.32	0.07
	CytoD	11.7	7.11	0.33

in ESI<sup>†</sup> Fig. S5 and Table S5. It shows that the double-shell model is required to fit the data for nucleated cells.

HL60 cells were also treated with cytochalasin D (CytoD), a compound that disrupts the cytoskeleton. Cells were measured in 0.32 S m<sup>-1</sup> saline and an example data set (n = 3) is shown in Fig. 3(c). The electrical opacity  $(|Z_{3MHz}|/|Z_{150kHz}|)$  of treated cells was also measured separately (n = 3), see data in Fig. S6.<sup>+</sup> The treated groups (unfilled circles) have a small but significant difference in the spectra compared with the control cells (solid dots). The optimal frequency for discrimination between the treated and untreated groups was found to occur at a frequency of around 6 MHz (for methodology, see ESI<sup>†</sup>). The best fits using the double-shell model (solid and dotted lines) are summarised in Table 1.  $C_{\text{mem}}$  is reduced after treatment, but  $\sigma_{\text{cvt}}$  remains unchanged. The results summarised in Fig. 3(d) demonstrate a statistically significant difference (data from three independent biological repeats). The membrane capacitance  $C_{\text{mem}}$  = 9.27 ± 0.30 mF m  $^{-2}$  for the untreated group and 7.01  $\pm$  0.19 mF m  $^{-2}$  for the CytoD group. These findings contrast with previously reported impedance measurements of CytoD-exposed cells suspended in physiological saline where no significant change was observed.<sup>32</sup> However, Jaffe and Voldman<sup>33</sup> used a dielectrophoresis spring system to measure a few hundred cells treated with CytoD and demonstrated a small but significant difference in cell polarizability. Liu et al.<sup>34</sup> used a constriction channel to measure cell electrical properties and concluded that treated cells had a higher membrane capacitance. However, this method applies significant stress on a cell, changing its shape and potentially influencing the membrane capacitance.<sup>35</sup> Importantly, the data in Fig. 3(c) demonstrates that measuring the impedance of cells suspended in a lower conductivity electrolyte increases the sensitivity of the system allowing small changes in cell electrical properties to be determined.

#### 4.2. Single-cell spectroscopy

Single-cell spectra were measured by applying eight simultaneous frequencies spaced logarithmically apart (251 kHz, 761 kHz, 2.31 MHz, 6.99 MHz, 21.2 MHz, 64.1 MHz, 194.3 MHz, 575 MHz). A suspension of HL60 cells consisting of a 50/ 50 mixture of normal and GA fixed, along with reference beads was measured in 0.32 S m<sup>-1</sup> or 1.6 S m<sup>-1</sup> saline. Cells were obtained from the same batch for each conductivity. Each single-cell spectrum was normalised against the mean bead impedance and then fitted to the double-shell model as described in section 2.2. Fig. 4(a) and (b) show examples of single-cell spectra for cells in the two different conductivities. Similar to the population mean data (Fig. 3(a) and (b)), the change in the normalised  $f^*_{\rm CM(cell)}$  for the untreated and fixed cells in 0.32 S m<sup>-1</sup> saline is very clear. However, this difference is barely visible for cells suspended in physiological saline. The fitting results from Fig. 4(a) and (b) are summarised in Table 2. The spectrum for 1000 individual cells from one sample of each type was fitted to the double-shell model and the results are shown on the scatter plots in Fig. 4(c) and (d). For the lower of the two suspending conductivities (Fig. 4(c)), the cell



**Fig. 4** Single-cell impedance spectra for HL60 cells in low and high-conductivity saline. (a) and (b) are examples of normalised  $f^*_{CM(cell)}$  (blue: real part, red: imaginary part) for untreated (solid line with dots) and fixed cells (dashed line with circles) measured in saline of conductivity 0.32 S m<sup>-1</sup> and 1.6 S m<sup>-1</sup>, respectively. (c)-i and (d)-i are scatter plots (*d vs. C*<sub>mem</sub>) for 1000 individual untreated (blue) and fixed (red) HL60 cells. (c)-ii and (d)-i il Histogram of  $\sigma_{cyt}$  in two different conductivity media. Mean ± SD are summarised in Table 3.

Table 2 Fitting results for untreated and fixed single-cells in 0.32 S m<sup>-1</sup> and 1.6 S m<sup>-1</sup> saline (Fig. 4(a) and (b)). For confidence intervals (CI), see Table S7 in ESI†

		Diameter ( $d$ , $\mu$ m)	Membrane capacitance ( $C_{ m mem}$ , mF m $^{-2}$ )	Cytoplasm conductivity ( $\sigma_{cyt}$ , S m <sup>-1</sup> )
0.32 S m <sup>-1</sup>	Untreated	12.5	10.31	0.24
	Fixed	11.3	6.26	0.07
1.6 S m <sup>-1</sup>	Untreated	11.4	10.23	0.36
	Fixed	11.6	9.26	0.43

Table 3 Single-cell fitting results (mean ± SD) for untreated and fixed cells in two different conductivity saline (Fig. 4(c) and (d)). CV: coefficient of variation

		Diameter ( <i>d</i> , µm)	Membrane capacitance ( $C_{\text{mem}}$ , mF m <sup>-2</sup> )	Cytoplasm conductivity ( $\sigma_{cyt}$ , S m <sup>-1</sup> )
0.32 S m <sup>-1</sup>	Untreated	$11.8 \pm 0.7$ , CV = 0.06	$10.47 \pm 1.26$ , CV = 0.12	$0.22 \pm 0.10$ , CV = 0.45
	Fixed	$12.6 \pm 0.8$ , CV = 0.06	$6.02 \pm 0.84$ , CV = $0.14$	$0.07 \pm 0.03$ , CV = $0.43$
$1.6 \text{ S m}^{-1}$	Untreated	$11.9 \pm 0.6$ , CV = 0.05	$10.64 \pm 3.68$ , CV = $0.35$	$0.37 \pm 0.08$ , CV = $0.22$
	Fixed	$12.0 \pm 0.5$ , CV = 0.04	$9.15 \pm 3.20$ , CV = $0.35$	$0.38 \pm 0.09$ , CV = $0.24$

distribution has a low coefficient of variance, with clear separation in size and membrane capacitance. The fixed cells had lower  $C_{\text{mem}}$  and  $\sigma_{\text{cvt}}$  (mean ± SD) compared with the untreated cells; see Table 3. However, the values of  $C_{\text{mem}}$  and  $\sigma_{\rm cvt}$  determined from single-cell data were slightly different from those obtained by the population mean method (section 4.1). For example, the mean  $\pm$  SD of  $C_{\text{mem}}$  determined from three repeats using the population mean method is  $9.28 \pm 0.30$ mF m<sup>-2</sup> (Table S4<sup>†</sup>), while the mean value calculated from 1000 individual cells using the single cell method is  $10.47 \pm 1.26$  mF  $m^{-2}$  (Table 3). This difference may be because the limited set of frequencies (8) in the single cell data set reduces the accuracy of the fits, particularly at the low frequency end of the spectrum where the size of the cell is determined. These results contrast with the higher conductivity saline results where the subpopulations overlap, making discrimination impossible.

#### 4.3. Differentiated macrophages

Macrophages play an essential role in the immune system as they recognise, engulf and digest pathogens. Monocytes migrate to the site of inflammation, where they differentiate into macrophages,<sup>36</sup> and identifying differentiated macrophages from monocytes is important for the study of immune processes and diseases. In this work, a monocytic leukaemia cell line (THP-1) was differentiated into a macrophage (M0) by exposure to phorbol 12-myristate 13- acetate (PMA).<sup>37,38</sup>

Protocols for differentiating THP-1 cells vary widely.<sup>38–41</sup> In this work, cells were exposed to PMA (200 ng ml<sup>-1</sup>) for 48 h followed by washing and resuspension, as in Fig. 5(a). Fig. S7(a) and (b)† show images of THP-1 and M0 cells in culture media. THP-1 cells float and have regular round shapes, while M0 cells attach to the flask bottom and have a spreading morphology. To measure cell size, THP-1 and M0 cells were dyed with Trypan blue and imaged with a microscope (Fig. 5(b) and (c)). The average diameter of THP-1 cells (30 cells) was 14.3  $\pm$  1.4 µm and M0 cells was 18.0  $\pm$  2.9 µm (30 cells); M0 cells become enlarged after differentiation. Fig. 5(d) shows a scatter plot of diameter (*d*) vs.  $C_{\rm mem}$  for the THP-1 and M0 population (see Fig. S8<sup>+</sup> for other data sets). The differentiated M0 cells have a broader distribution in size and membrane capacitance. Fig. 5(e) shows a histogram of  $\sigma_{\rm cyt}$ , showing that the two groups have a similar range but differ in their mean value (mean  $\pm$  SD: THP-1: 0.15  $\pm$  0.04 S m<sup>-1</sup>; M0: 0.14  $\pm$  0.04 S m<sup>-1</sup>). Fig. 5(f) shows mean  $\pm$  SD for three repeats with 1000 cells in each group.

There is a significant difference in cell diameter (\*\*,  $p \leq$ 0.01), accompanied by a change in cytoplasm conductivity (\*,  $p \leq 0.05$ ), but the mean change in membrane capacitance is not significant. The variability in this parameter may reflect differences in cell states since the differentiation ability of THP-1 cells is influenced by passage number. These results differ from the work of Soe et al.42 who used dielectrophoresis (DEP) to characterise macrophage phenotypes and reported that the membrane capacitance of THP-1 monocytes and M0 cells were different (11.1 and 12.8 mF  $m^{-2}$ ), but these values were determined from a single DEP cross-over frequency measurement in very low conductivity media.

### 5. Conclusion

This work shows that the complete dielectric properties of nucleated cells suspended in a high conductivity electrolyte can be measured in a system with an extended frequency range of up to 550 MHz. Data for nucleated cells cannot be accurately fitted to a single-shell model; only to the double-shell model. HL60 cells modified with GA or CytoD were measured both in low (0.32 S m<sup>-1</sup>) and high-conductivity (1.6 S m<sup>-1</sup>) saline, demonstrating that the lower conductivity media significantly improves the resolution of the system, making it possible to discriminate untreated from treated cells. Experimentally this is consistent with the work of Ostermann *et al.*<sup>43</sup> who showed that a low conductivity measurement buffer (1:4 PBS: 0.28 M sucrose) enhanced



**Fig. 5** Single-cell analysis of THP-1 cells differentiated into macrophages (M0) following 48 h exposure to PMA in saline (0.32 S m<sup>-1</sup>). (a) Experimental workflow for THP-1 cells differentiation. (b) and (c) are images of cells dyed with Trypan blue after harvesting from the plates and ready to be measured. The average diameter (mean  $\pm$  SD) of 30 THP-1 cells is 14.3  $\pm$  1.4 µm, and 18.0  $\pm$  2.9 µm for 30 M0 cells. (d) Shows a scatter plots (*d vs. C*<sub>mem</sub>) for 1000 individual THP-1 cells (blue) and M0 cells (red). (e) Show a histogram of  $\sigma_{cyt}$  for two different cell subpopulations. (f) Plots of the mean  $\pm$  SD of diameter, membrane capacitance and cytoplasm conductivity for each set of 1000 cells (three-repeat experiments). Blue dots are THP-1 cells and red are M0 cells. Statistical analysis of *d*, *C*<sub>mem</sub> and  $\sigma_{cyt}$  from Student's *t*-test (\*,  $p \le 0.05$ ; \*\*,  $p \le 0.01$ ; ns, not significant).

discrimination of viable and necrotic U937 cells. Fitting to the double-shell model at 0.32 S  $m^{-1}$  showed that the membrane capacitance of the cross-linked cells was 30% less than untreated cells in saline. The reduction in cytoplasm conductivity was greater, demonstrating a decrease in cytoplasmic ion concentration in the non-viable cells. Cells treated with CytoD had a significantly reduced membrane capacitance but unchanged cytoplasm conductivity. These differences could not be observed in cells suspended in physiological saline.

Single-cell measurements of nucleated cells were also made by applying 8 simultaneous frequencies. In 0.32 S m<sup>-1</sup> saline, untreated and fixed cells could be easily discriminated through changes in membrane capacitance and cytoplasm conductivity, consistent with the data for the mean population. Analysis of the electrical properties of THP-1 cells

and their differentiation into macrophages demonstrated moderate changes in size and cytoplasm conductivity, with a higher variance in the membrane capacitance.

In conclusion, we have developed a new high-frequency single-cell impedance cytometer capable of analysing many hundreds of single nucleated cells in a short time frame. Resuspension of cells in an electrolyte with conductivity 5× lower than physiological saline improves discrimination between different cell types and significantly increases the accuracy of the double shell fits. Analysis of many individual nucleated cells by impedance spectroscopy can be used to extract dielectric data using the double-shell model. The work highlights the shortcoming of the singleshell model for a nucleated cell and demonstrates the requirement for enhanced frequency bandwidth in order to provide accurate fits.

(a)

# Data availability

Data for this article can be accessed at [https://doi.org/ 10.5258/SOTON/D3368].

# Conflicts of interest

There are no conflicts of interest.

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