Focusing of mammalian cells under an ultrahigh pH gradient created by unidirectional electropulsoation in a confined microchamber†

Despina Nelie Loufakis,a Zhenning Cao,b Sai Ma,b David Mittelmanc and Chang Lu*a,b

The transport and manipulation of cells in microfluidic structures are often critically required in cellular analysis. Cells typically make consistent movement in a dc electric field in a single direction, due to their electrophoretic mobility or electroosmotic flow or the combination of the two. Here we demonstrate that mammalian cells focus to the middle of a closed microfluidic chamber under the application of unidirectional direct current pulses. With experimental and computational data, we show that under the pulses electrochemical reactions take place in the confined microscale space and create an ultrahigh and nonlinear pH gradient (~2 orders of magnitude higher than the ones in protein isoelectric focusing) at the middle of the chamber. The varying local pH affects the cell surface charge and the electrophoretic mobility, leading to focusing in free solution. Our approach provides a new and simple method for focusing and concentrating mammalian cells at the microscale.

Introduction

Proteins and peptides have both amine and carboxylic acid groups and these groups can either donate or accept protons. Therefore the charge of a protein/peptide molecule varies with the surrounding pH. The pH value at which a protein molecule carries no net surface charge is referred to as the isoelectric point (pI) and is an important physiochemical property. Isoelectric focusing (IEF) has been widely used for protein concentration and separation.1,2 In these experiments, carrier ampholytes (i.e. a mixture of molecules with multiple aliphatic and carboxylate groups) or acrylamido buffers (typically covalently incorporated into the gels) are used to establish a linear pH gradient in gels or capillary tubes. A direct current (dc) field is then applied to focus various molecules to their respective pI values. IEF has also been applied to small bioparticles3 (with diameters <5 μm) such as viruses,4,5 bacteria,6,7 and yeasts.8 In these works, special care was taken to use very low concentrations for these bioparticles to avoid interference due to the interactions between the bioparticles and the ampholyte matrix.8 IEF of mammalian cells (with diameters typically in the range of 10–20 μm) on a simple and practical platform has yet to be demonstrated, due to the potential cell entrapment in matrices.

The surface charge of mammalian cells is typically negative at physiological pH. Cells are covered by a surface coat that is rich in carbohydrate, referred to as glycosmaly. These carbohydrates include oligosaccharide chains covalently bound to membrane proteins (glycoproteins) and lipids (glycolipids).9 The carbohydrate portion of these molecules contains a significant amount of sialic acid residues that carry negative charges at physiological pH.10–12 At the same time, the carbohydrate groups also contain amino sugars which may become positively charged under low pH.10–12 Thus we hypothesized that concentration or focusing of mammalian cells under a very high pH gradient would be potentially feasible. Such method may find applications in microfluidic total analysis systems that require cell manipulations.11–16

Here, by exploiting the interplay among microfluidics, electrochemistry, and electokinetics, we demonstrate IEF of mammalian cells in a simple microfluidic electrochemical chamber containing a low-conductivity buffer. We observed focusing of cells in a closed microfluidic chamber when unidirectional pulses were applied by two surface electrodes. Our experimental and computational results show that an ultrahigh pH gradient (up to 0.14 Unit per μm) was generated by electrolysis of water inside the confined microscale space under the application of the electric pulses. The pH gradient was highly nonlinear and steepest at the center of the chamber. We also studied the correlation between the electrophoretic mobility of cells and the local pH and estimated the pI of Chinese hamster ovary (CHO) cells to be in the range of 4.5–6.5. Our setting offers a simple approach to generate an ultrahigh

†Electronic supplementary information (ESI) available: Figures S1–S5 and videos S1–S2. See DOI: 10.1039/c4sc00319e

Department of Chemical Engineering, Virginia Tech, Blacksburg, Virginia 24061, USA. E-mail: changlu@vt.edu; Fax: +1 540 231 5022; Tel: +1 540 231 8681
School of Biomedical Engineering and Sciences, Virginia Tech-Wake Forest University, Blacksburg, Virginia 24061, USA
 Virginia Bioinformatics Institute and Department of Biological Sciences, Virginia Tech, Blacksburg, Virginia 24061, USA

‡Department of Chemical Engineering, Virginia Tech, Blacksburg, Virginia 24061, USA.
pH gradient at microscale in free solution. Such conditions uniquely permit the focusing of mammalian cells without involving matrices.

Results and discussion

The microfluidic device used in our study is shown in Fig. 1A and ESI Fig. S1.† Cells (typically 10–150) were loaded into a microfluidic chamber of 1.5 mm × 400 μm × 13 μm. These cells were sealed inside the chamber by closing the pneumatic valves at the two ends. Two long planar gold electrodes (120 μm wide, and 1 mm apart) were inside the chamber and 30 ms-long dc pulses were applied via these electrodes every 10 s for 50 times with the pulse intensity in the range of 400–1800 V cm⁻¹. CHO cells were initially tested. Interestingly, under unidirectional pulses, the cells did not move in a consistent direction. The cells eventually focused into a narrow region around the center of the chamber (Fig. 1B and ESI Video S1†). Such movement could be tracked by both phase contrast imaging and fluorescence imaging (after labeling the cells with Hoechst 33342 which penetrates the cell membrane and intercalates within their DNA). In general, the focusing of the cells became more rapid when higher field intensity pulse was used (in the range of 400–1800 V cm⁻¹). The cells finished focusing after 50 pulses under 1000 V cm⁻¹ while the same process essentially was completed after 10 pulses of 1800 V cm⁻¹. Fig. 1C reveals that in general, high pulse intensity also led to improved focusing. For example, a 1000 V cm⁻¹ or higher field intensity was required for complete focusing of cells within a ~30 μm-wide region.

Next, we analyze the mechanistic details involved in such isoelectric focusing of mammalian cells. The use of a closed microscale chamber in our design amplified the impact of electrochemical lysis of water. Hydroxides were generated at the cathode (the negative terminal) and protons were produced at the anode (the positive terminal) under sufficiently high voltage. Thus during the course of electropulsation, the local pH at the cathode became basic while the pH close to the anode turned acidic. The generation of a pH gradient under this mechanism was a significant process given the tiny volume of the microfluidic chamber. We conducted COMSOL modeling to examine the electrochemical process. A couple of simplifications needed to be made in order for the modeling to converge. First, we neglected the electrophoresis of ions. We will discuss the implications of this simplification below. Second, we considered only two important reactions (neutralization between H⁺ and OH⁻ and conversion between the anions in the

![Fig. 1](image-url)
buffer \( \text{H}_2\text{PO}_4^- \) and \( \text{HPO}_4^{2-} \)). The other reactions associated with phosphoric acid and phosphate ion are only important in extreme pH (<3 or >12, respectively), thus have smaller impact on the buffering.

We modeled three buffer concentrations under the application of 50 pulses (each with a 30 ms duration and 1000 V cm\(^{-1}\) intensity): (1) \( [\text{HPO}_4^{2-}] = 0, [\text{H}_2\text{PO}_4^-] = 0 \) (Fig. 2A); (2) \( [\text{HPO}_4^{2-}] = 4.8 \text{ mM}, [\text{H}_2\text{PO}_4^-] = 1.2 \text{ mM} \) (Fig. 2B); (3) \( [\text{HPO}_4^{2-}] = 9.6 \text{ mM}, [\text{H}_2\text{PO}_4^-] = 2.4 \text{ mM} \) (Fig. 2C). As presented in the inset graphs of Fig. 2A-C, after the first pulse (with the onset at \( t = 0 \) and a duration of 30 ms) at \( t = 0.04 \text{ s} \), hydroxides and protons are generated at the locations of the electrodes inside the chamber. These ions then diffuse during the interval between the first and second pulse \((9.97 \text{ s})\) into the rest of the chamber volume, forming a sharp pH gradient close to the center of the chamber. Such pH gradient continues to increase with the application of first several additional pulses. In all three cases, the pH profile reaches its steady state after roughly the first 10 cycles (with one cycle referring to one 30 ms pulse followed by 9.97 s interval). After reaching the steady state, the pH profile still experiences the dynamics caused by the production of protons and hydroxides by additional pulses. However, the pH profile at the end of each additional cycle is identical. With increasing buffer concentrations, the pH profile in the chamber shows more resistance to pulse-induced pH change. Furthermore, the buffer in general has more buffering power in the acidic region (the right side of the chamber) than in basic region (the left side of the chamber) due to the higher starting concentration for \( \text{HPO}_4^{2-} \) (4 times higher than that of \( \text{H}_2\text{PO}_4^- \)).

Electrophoresis of ions affects the pH profile. The electrophoresis of protons and hydroxides only has minor impact on the location of the neutral pH interface in the middle of the chamber. On the other hand, the electrophoresis of \( \text{HPO}_4^{2-} \) and \( \text{H}_2\text{PO}_4^- \) may have more significant influence on the pH profile. The majority of these anions is swept to the right side of the chamber after the first 4 cycles by electrophoresis and stay there for the rest of the process, as shown in ESI Fig. S2.† Thus, after the first 4 cycles, the buffering effect is primarily present on the right side of the chamber (with roughly doubled buffer concentration) while the left side of the chamber has few buffering ions.

To summarize, the actual pH profiles in the chamber over the course of the 50 cycles are best described by mixing the results under various buffer concentrations shown in Fig. 2. For a buffer with the concentration of \( [\text{HPO}_4^{2-}] = 4.8 \text{ mM} \) and \( [\text{H}_2\text{PO}_4^-] = 1.2 \text{ mM} \) (which was used in most of our experiments), the pH profiles are similar to the ones in Fig. 2B for the first 4 cycles. Then for cycles 5 to 50, the left side of the pH profiles will be similar to those in Fig. 2A and the right side will be similar to those of Fig. 2C. The pH gradient forms in the chamber in spite of the presence of the buffer. The buffer provides some resistance to the pH change within the initial 10 cycles, with such buffering effect more pronounced on the right side than on the left side. The location of cell focusing observed experimentally (at \( x \approx 750 \mu \text{m} \), as shown in Fig. 1B) matches the location of the steep pH gradient suggested by the modeling when the buffer is considered (Fig. 2B and C).

---

**Fig. 2** The formation of pH gradient due to water electrolysis. The pH profiles along the length of the microfluidic chamber (x axis) at multiple time points are computed by COMSOL Multiphysics for three buffer concentrations (A) no buffer, (B) 4.8 mM Na\(_2\)HPO\(_4\), 1.2 mM KH\(_2\)PO\(_4\), and (C) 9.6 mM Na\(_2\)HPO\(_4\), 2.4 mM KH\(_2\)PO\(_4\). The cathode is located at \( x = 130–250 \mu \text{m} \), the anode is located at \( x = 1250–1370 \mu \text{m} \), and both electrodes are 120 \( \mu \text{m} \) wide. The pulses are applied once every 10 s and last for the first 30 ms of each 10 s cycle (e.g. in the first cycle, the pulse lasts during \( t = 0–0.03 \text{ s} \)). The insets show the pH profile change during the first cycle (\( t = 0–10 \text{ s} \)).
Based on the data generated by the modeling, we estimate that a pH gradient of 0.07 Unit µm⁻¹ was formed after the first pulse and it reached and stabilized at ~0.14 Unit µm⁻¹ after the 10th pulse (Fig. 2B) at a location around the center of the chamber. Such a pH gradient was roughly 2 orders of magnitude steeper than that in the typical protein IEF setting (between 0.001 and 0.2 unit per 100 µm³). The anode pH decreased from 7.4 to 3.8 after the first pulse and further to 2.0 after the 10th pulse. Similar trend was observed at the cathode where the pH became 11.6 after the first pulse, and increased to 12.2 after the 10th pulse. At the steady state (after 10th pulse), a pH range of 3.8 to 10.5 was covered in a narrow 50 µm distance along x.

The establishment and dynamics of the pH gradient could be observed experimentally by having fluorescein dissolved in the buffer. Fluorescein is pH-sensitive and its fluorescence reduces significantly in acidic environments (e.g. the fluorescence intensity of fluorescein decreases by about 70% as the pH changes from 10 to 5, due to its protonation). ESI Fig. S3† shows that the fluorescence on the right side of the chamber close to the anode turned acidic (indicated by the loss of fluorescein fluorescence). This led to a change in the moving direction for these cells, causing them to move towards the cathode. It is worth noting that cells stabilized when they crossed 10–20 µm over the fluorescein boundary line to the right, indicating that the neutralization in the cell surface charge occurred under a low pH. For the cells which were at the far left of the boundary line, they experienced only movement toward the right (anode) due to the fact that they were always under the basic pH. The effect of electroosmotic flow in here can be ignored as the chamber was closed during the process. The similar movement trajectories of cells can also be seen in ESI Video S2.†

It is interesting to track the dynamics in the electrophoretic mobility of a cell under the pulse sequence and the local pH at the moment. The pH varied with the location inside the chamber and the number of pulses that have been exerted. We calculated the local pH values for cells at a particular moment based on our COMSOL modeling and tracked the corresponding velocity, and therefore the electrophoretic mobility, based on the videometric data. Fig. 4 shows that the local pH strongly affected both the direction and magnitude of the electrophoretic mobility. First of all, all cells moved from the cathode to the anode (shown with negative sign) when pH was higher than 6.5 (with an average µ of −19.86 mm² kV⁻¹ s⁻¹ and a standard deviation of 8.01 mm² kV⁻¹ s⁻¹), whereas the vast majority of the mobilities were positive when the pH was lower than 4.5 (the average value of µ was 1.68 mm² kV⁻¹ s⁻¹ and the standard deviation was 4.20 mm² kV⁻¹ s⁻¹). This confirms that cells were negatively charged at high pH and positively charged at low pH.

Fig. 3  Time-lapse images that show the movement of cells inside the chamber in relation to the boundary formed by fluorescein. The cells were stained by Hoechst and suspended in the buffer containing fluorescein. A buffer system containing 4.8 mM Na₂HPO₄, 1.2 mM KH₂PO₄, and 250 mM sucrose was used. The pulses (1000 V cm⁻¹) were applied once every 10 s and lasted for the first 30 ms of each 10 s cycle.
with a threshold range of pH 4.5–6.5 separating the two regimes. This fairly wide range for pI may be related to the heterogeneity in the cell population. Other than pH, mechanisms such as cell polarization in the electric field may also have effect on the distribution of the cell surface charge and the electrophoretic mobility.24

We also tested various buffer concentrations (2.4 mM Na2HPO4–0.6 mM KH2PO4, 4.8 mM Na2HPO4–1.2 mM KH2PO4, and 8 mM Na2HPO4–2 mM KH2PO4) experimentally (ESI Fig. S4†). The results indicate that the mobility of cells was not significantly affected by the buffer concentration in this range. This confirms that the buffering power of the solution was not enough to counteract the generation of protons and hydroxides by the electrochemical reactions.

Our device and setup allow IEF of mammalian cells of various types and various concentrations (we show IEF of Jurkat cells at high and low concentrations in ESI Fig. S5†), owing to the ultrahigh pH gradient formed and the free solution arrangement. The focusing process involves a complex interplay among the spatiotemporally varying parameters of local pH, cell surface charge, and cell electrophoretic mobility (shown in the schematic of Fig. 5). It is also worth noting that electroporation of cells may occur when the field intensity was higher than a threshold of 300–400 V cm−1.25,26

To summarize, we demonstrate the IEF of mammalian cells in a microfluidic chamber by creating an ultrahigh pH gradient in free solution via electrolysis of water. The use of a closed chamber facilitates formation of significant pH gradient based on microscale electrochemical reactions and also eliminates potential undesired flow (e.g. electroosmotic flow) which is detrimental to pH gradient establishment. The free solution scheme allows the easy movement of cells. The focusing of cells requires high field intensities for the dc pulses, because of the relative low mobility of cells due to pH-induced surface charge. The ultrahigh and nonlinear pH gradient close to the center of the chamber permitted focusing of cells with minor difference in their pI. Our approach works universally for different mammalian cell types and concentrations.

Experimental section

Cell cultures

Jurkat cells (ATCC, Manassas, VA) were grown in a humidified incubator containing 5% CO2 at 37 °C in RPMI-1640 growth medium supplemented with 10% fetal bovine serum. The cell concentration was measured every day using a hemocytometer, and fresh medium was added to the flask when needed, to keep the cell concentration below 3 × 106 cells per ml. CHO-K1 (ATCC, Manassas, VA) cells were grown in F-12K growth medium supplemented with 10% fetal bovine serum (Atlanta Biologicals, Flowery Branch, GA) and 100 U per ml penicillin–100 mg mL−1 streptomycin (Life Technologies Corporation, Grand Island, NY) in a 25 cm2 flask, and in a humidified incubator containing 5% CO2 at 37 °C. Every 2 days they were trypsinized and subcultured at a ratio of 1 : 10 in order to be maintained in the exponential growth phase.

After the cells were harvested, they were centrifuged at 300 g for 5 min at room temperature and washed with phosphate buffer saline (PBS, Fisher Scientific, Suwanee, GA). They were then, resuspended in a low-conductivity phosphate buffer (4.8 mM Na2HPO4, 1.2 mM KH2PO4, and 250 mM sucrose, pH = 7.4, unless otherwise specified) at a concentration of 3 × 106 cells per ml as determined by a hemocytometer. In the case where they were fluorescently labeled, cells were kept in medium at 3 × 106 cells per ml, and Hoechst 33342 (Life
Microfluidic device fabrication

Multilayer soft lithography based on polydimethylsiloxane (PDMS, R. S. Hughes Company, Sunnyvale, CA) was used for the fabrication of the device in our experiments. The masks were designed using Macromedia Freehand and printed on a 4000 dpi film by Infinity Graphics (Okemos, MI). SU-8 2025 (Microchem Corporation, Newton, MA) or AZ 9260 (Capitol Microfabrication Materials, Carrollton, TX) photo-resist was spin-coated on a 3-inch silicon wafer (University Wafer, South Boston, MA) for the control channel (~60 μm deep) and the fluidic channel (~13 μm deep), respectively. A final hard bake of the AZ 9260 features at 130 °C for 1 min created a circular cross-section needed for the fluidic channel. Degassed prepolymer mixture consisting of monomer (RTV 615 A) and curing agent (RTV 615 B) at a mass ratio 10 : 1 was poured in a Petri dish on top of the control layer master (~5 mm in the PDMS thickness) and spun at 1100 rpm (~110 μm in the PDMS thickness formed) on the fluidic layer mask followed by bake at 80 °C for 0.5 h. The two layers were then aligned and bonded after oxygen plasma treatment (Harrick Plasma, Ithaca, NY). Another bake followed for 45 min to ensure strong bonding between the layers. The access holes for the inlets and the outlets were punched.

Pre-cleaned glass slides were used for the fabrication of the surface gold electrodes. On top of each slide a 20 nm-thick layer of titanium (Kurt J. Lesker Company, Clairton, PA) was deposited, followed by the deposition of a 150 nm-thick layer of gold (Kurt J. Lesker Company, Clairton, PA) using an e-beam evaporator (PVD 250; Kurt J. Lesker Company, Clairton, PA). The metal layer was then patterned using AZ 9260 before being wet-etched using standard gold etchant (Aldrich, Milwaukee, WI) and titanium etchant (HF : H2SO4 : DI water = 1 : 1 : 10). The remaining photoresist was stripped using acetone, leaving 2 μm thick gold films to support the electrodes. On top of each slide a 20 nm-thick layer of titanium (Kurt J. Lesker Company, Clairton, PA) was deposited, and titanium etchant (HF : H2SO4 : DI water = 1 : 1 : 10) was used to etch the remaining photoresist. The remaining gold films were then patterned using AZ 9260 before being wet-etched using standard titanium etchant (HF : H2SO4 : DI water = 1 : 1 : 10). The remaining photoresist was stripped using acetone, leaving 2 μm thick titanium films to support the cell culture supports. The PDMS device and the glass slide were bonded using oxygen plasma followed by 0.5 h bake at 80 °C to create the assembled microfluidic device.

Device operation and electropulsation

The microfluidic device was mounted on an inverted fluorescence microscope (IX-71, Olympus, Melville, NY) equipped with a 10× dry objective and a CCD camera (ORCA-285; Hamamatsu, Bridgewater, NJ). The fluidic layer was flushed by the phosphate buffer supplemented with Tween 20 at 0.02% before the experiments, using a syringe pump (Fusion 400, Chemxy, Stafford, TX). The surface electrodes were connected to a high voltage reed relay (5501, Coto Technology, North Dingtown, RI) and a dc power supply (PS350, Stanford Research Systems, Sunnyvale, CA) for application of electric pulses of milliseconds. A schematic of the system setup is illustrated in ESI Fig. S1† and the setup was similar to that in our previous publication.23

Image analysis

Image analysis was performed using a software package Fiji in order to extract the velocity of cells in the fluorescence image series. A MATLAB (Mathworks, Natick, MA) code was written for analyzing the standard deviations in the cell location.

COMSOL modeling

COMSOL Multiphysics 4.3 (Burlington, MA) was used to simulate the pH profiles in the microfluidic chamber. The “transport of diluted species” module was used to simulate the H+, OH–, HPO42−, H2PO4– concentration profiles due to diffusion and generation/consumption for 500 seconds (the duration of applying 50 pulses). A 1-D geometry was determined to be sufficient to accurately solve the concentration profiles. Eqn (1) was used to compute the concentrations of the ions.

\[
\frac{\partial c_i}{\partial t} + \nabla \cdot (-D_i \nabla c_i) = R_i
\]

where \( c_i \) is the concentration of species \( i \), \( D_i \) is the diffusion coefficient, and \( R_i \) is the generation or consumption of ions due to electrochemical reactions (\( R_i \)), the neutralization reaction between \( H^+ \) and \( OH^- \) (\( R_{\text{neut}} \)), and the reaction between the conjugate acid–base pair in the buffer \( HPO_4^{2-} \) and \( H_2PO_4^- \) (\( R_{\text{buf}} \)). The diffusion coefficients are \( D_{H^+} = 9.31 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \), \( D_{OH^-} = 5.32 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \), \( D_{HPO_4^{2-}} = 0.759 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \), \( D_{H_2PO_4^-} = 0.959 \times 10^{-9} \text{ m}^2 \text{ s}^{-1} \).

Generation of ions \( H^+ \) and \( OH^- \) (\( R_{\text{neut}} \)) takes place at the anode (\( R_{H^+} \)) and cathode (\( R_{OH^-} \)) during each pulse (Eqn (2)), assuming a homogeneous profile over the surface of the electrodes:

\[
R_{\text{neut}} = R_{H^+} + R_{OH^-} = \frac{i}{nFV}
\]

where \( i \) is the current (~0.34 μA as observed in our experiments), \( n \) is the number of electrons involved in the reaction, and \( V \) is the volume.

Neutralization reaction occurs when protons meet hydroxides:

\[
H^+ + OH^- \leftrightarrow H_2O
\]

\[
R_{\text{neut}} = R_{H^+} = R_{OH^-} = -k_1[H][OH] + k_2 \times 55.65
\]

We consider one equilibrium between \( HPO_4^{2-} \) and \( H_2PO_4^- \):

\[
HPO_4^{2-} + H^+ \leftrightarrow H_2PO_4^-
\]

\[
R_{\text{buf}} = R_{H^+} = R_{H_2PO_4^-} = -k_3[HPO_4^{2-}][H^+] + k_4[H_2PO_4^-]
\]
Acknowledgements

We thank NSF CBET 1016547 and NCI R21CA174577 for financial support of this research.

References