






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Adhesion-based high-throughput label-free cell sorting using ridged microfluidic channels

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Numerous applications in medical diagnostics, cell engineering therapy, and biotechnology require the identification and sorting of cells that express desired molecular surface markers. We developed a microfluidic method for high-throughput and label-free sorting of biological cells by their affinity of molecular surface markers to target ligands. Our approach consists of a microfluidic channel decorated with periodic skewed ridges and coated with adhesive molecules. The periodic ridges form gaps with the opposing channel wall that are smaller than the cell diameter, thereby ensuring cell contact with the adhesive surfaces. Using three-dimensional computer simulations, we examine trajectories of adhesive cells in the ridged microchannels. The simulations reveal that cell trajectories are sensitive to the cell adhesion strength. Thus, the differential cell trajectories can be leveraged for adhesion-based cell separation. We probe the effect of cell elasticity on the adhesion-based sorting and show that cell elasticity can be utilized to enhance the resolution of the sorting. Furthermore, we investigate how the microchannel ridge angle can be tuned to achieve an efficient adhesion-based sorting of cells with different compliance.

1 Introduction

Identification and isolation of cells that express desired molecular surface markers are required in a variety of applications in the biological sciences, cell therapy, and medical diagnostics.^{1–4} For example, adhesion-based cell sorting is needed in the manufacturing workflow of chimeric antigen receptor (CAR) T cell therapies. A major challenge to improving these therapies is a lack of efficient methods to evaluate CAR expression efficiency in conjunction with CD4 and CD8 expression, required as part of the release criteria. In addition, there is a lack of separation methods that are sensitive to cell surface expression, while not inducing changes in cell morphology, viability, molecular content, activation state, and phenotype.

Labeling methods (for example: FACS and MACS) can be used for adhesion-based cell sorting and offer high purity with high enrichment. However, these techniques provide a binary output for the receptor expression and are not yet able to fractionate into multiple outputs for finer sensitivity to the molecule of interest. Another major drawback of these methods is the changes they induce by labeling tags or tag removal, such as the risk of tag-induced activation of sorted cells, which restrains their downstream use.^{5,6} To solve this issue, negative selection of target cells can be used. Yet, even this can

contaminate therapy products with magnetic beads coated with foreign antibodies.

To overcome these difficulties, label-free approaches can be employed leveraging differences in cells adhesive properties as the mechanism of separation.^{7–9} As an example of cell adhesion sorting using micro/nano structures, Kwon *et al.*¹⁰ used patterned microchannels to separate and enrich human breast cancer cells from epithelial cells. Detachment assays were used to measure cells adhesion. Specific adhesion of cells using proteins, peptides, and other molecular mediators can be employed to capture desired cells. Antibody coated surfaces in microfluidic channels were demonstrated to be effective for capturing circulating endothelial progenitor cells (EPCs).¹¹ Hashimoto, Kaji, and Nishizawa¹² combined dielectrophoresis with adhesion-based microfluidics to selectively capture neutrophils from a mixture of leukocytes. Zhang *et al.*¹³ used a microchannel coated with basement membrane extract to separate heterogeneous breast cancer cell lines based on their adhesive capacities. Yang *et al.*¹⁴ achieved circulating tumor cell (CTCs) capture from blood cells using a TA-functionalized film. Negative selection is also an option for enriching rare target cells.^{15,16} A tandem microfluidic systems was developed to isolate in a high-throughput, label free manner CTCs that exhibit high compliance and low adhesion.¹⁷ Singh *et al.*¹⁸ were able to isolate fully reprogrammed induced human pluripotent stem cells (hPSCs) from heterogeneous reprogramming cultures by capturing target cells on a substrate in a microchannel under shear flow. Microfluidic assay was developed to separate genetically-related yeast strains based on adhesion

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strength enabling the rapid screening and fractionation of yeast based on adhesive properties.¹⁹ However, these methods are not continuous since they capture and arrest cells from the flow. Chang, Lee, and Liepmann²⁰ used microstructured fluidic channels with array of pillars coated with E-selectin to capture and arrest cells from the free flow. Bose *et al.*²¹ demonstrated affinity flow fractionation where interactions with asymmetric molecular patterns laterally displace cells in a continuous manner. Similarly, Choi, Karp, and Karnik²² described a separation process called “deterministic cell rolling”, which separates cells based on transient molecular interactions with slanted ridges on the channel floor in a continuous process. However, these approaches are limited to low flow rates to maintain rolling interactions with the adhesive surface and to prevent hydrodynamic forces from detaching the cells without any separation.

A high throughput adhesion based sorting method was recently reported²³ that consists of a microchannel decorated with diagonal ridges and coated with adhesion molecules. Flow cytometry data for cells collected at the microchannel outlets showed different cell distributions correlated with the differences in cell adhesiveness indicating sorting dependence on adhesion levels. A unique aspect of this approach is the ability to operate at high throughput. The experiments demonstrated flow of cells at rates up to 0.2 m s^{-1} , which is over two orders of magnitude higher than previous methods. Results indicated that sorting did not activate the cells due to the brief interaction time of cells with adhesion ligands beneficial for downstream cell analysis. We note that ridged microfluidic channels have been previously used to sort cells and microparticles by size and elasticity.^{24–27}

Computational modeling has been used to probe adhesive behavior of cells, especially in the context of cell rolling.^{28–30} These models are usually based on theoretical frameworks which describe adhesive interactions using chemical reaction kinetics and relate the kinetic rates to the applied force. Bell³¹ and Dembo *et al.*³² adapted this kinetic theory and proposed an exponential constitutive law between the dissociation rate and the force. Evans, Berk, and Leung³³ and Evans and Ritchie³⁴ employed a power law instead of the exponential law used in the Bell model.³¹ Jadhav, Eggleton, and Konstantopoulos³⁵ developed a three-dimensional computational model to predict receptor-mediated rolling of deformable cells in shear flow. Hammer and Apte³⁶ simulated the rolling and adhesion of selectin-mediated neutrophil using a kinetics framework. Khismatullin and Truskey³⁷ used a modified Dembo model and evaluated the effect of cell deformability and viscoelasticity on receptor-mediated leukocyte adhesion to ligand coated surface. Interactions of compliant adhesive capsules and cells in patterned microfluidic channels have been studied using computer simulations with applications to the sorting and separation.^{38–41}

In this work, we first examine the rolling of deformable cells represented by a fluid-filled elastic particle on a flat substrate patterned with diagonal adhesive stripes. We show that the cells follow different trajectories based on the adhesiveness and explain the sorting mechanism. We further show that this separation method is limited to low flow rates due to the

hydrodynamic lift detaching cells from the substrate. We then demonstrate high-throughput sorting of adhesive cells in ridged microfluidic channels. The periodic diagonal ridges compress cells propelled by the flow, thereby ensuring cell contact with adhesive surfaces even at high flow rates. We systematically investigate the effects of cell adhesion and elasticity on cell trajectories in the ridged microchannel and demonstrate that cells with different adhesiveness follow different trajectories. The simulations reveal that cell elasticity can be used to enhance sorting of adhesive cells. Furthermore, we show that the resolution of the adhesion-based cell sorting can be increased by adjusting the angle of the microchannel ridges. Together these studies suggest ways to improve the sensitivity of separation to adhesion in the limit of transient interactions characterized by fast timescales.

2 Methods

We employ a three-dimensional computational approach^{42–45} to model the motion of compliant neutrally-buoyant adhesive cells in a microfluidic channel filled with a viscous fluid. The simulations are performed using an inhouse three-dimensional fluid-structure interaction solver based on the lattice Boltzmann method (LBM) for fluid flow coupled in a two-way manner with a lattice spring model (LSM) for solid mechanics.

LBM uses time integration of the discretized Boltzmann equation $f_i(\mathbf{r} + \mathbf{c}_i \Delta t, t + \Delta t) = f_i(\mathbf{r}, t) + \Delta_i[f(\mathbf{r}, \Delta t)]$ for a distribution function $f_i(\mathbf{r}, t)$ describing the mass density of fluid particles at a lattice node \mathbf{r} and time t propagating along a lattice direction i with a constant velocity \mathbf{c}_i .^{46,47} Here, Δt is the time step. The collision operator Δ_i describes the change in f_i due to collisions at lattice nodes. In three dimensions, we use a cubic lattice D3Q19 with 19 discrete velocities.

The compliant cell is modelled as a fluid-filled elastic membrane. LSM uses a triangulated network of harmonic springs that connect regularly spaced mass nodes on the cell surface to represent an elastic membrane.^{48,49} The stretching

elastic energy associated with a node at a position \mathbf{r}_i is $U_s(\mathbf{r}_i) =$

$$0.5 \sum_j k_s (r_{ij} - r_{ij}^{\text{eq}})^2 \text{ where } r_{ij} = |\mathbf{r}_i - \mathbf{r}_j| \text{ and } r_{ij}^{\text{eq}} \text{ are respectively}$$

the length and the equilibrium length of a spring between two nodes with positions \mathbf{r}_i and \mathbf{r}_j , and k_s is the spring constant. Bending rigidity of cell membrane is included by considering the change of dihedral angle of adjacent pairs of triangles. The bending energy is given by $U_b = \sum_i k_b [1 - \cos(\theta_i - \theta_{\text{eq}})]$, where k_b is the bending spring constant, θ_i is the instantaneous angle between two adjacent triangles with a common edge i , and θ_{eq} is the equilibrium angle.

To model adhesion between the cell and microchannel surfaces, we use the Morse potential that has a repulsive and attractive components, depending on the separation distance. The Morse potential associated with two nodes separated by a distance r is given by $U_A(r) = D_e \left(1 - e^{-\frac{r-r_e}{K_e}}\right)^2$, where D_e is the



depth of the potential well, K_e is a parameter controlling the width of the potential, and r_e is the equilibrium distance. The force derived from the Morse potential is given by $F_A(r) = 2\frac{D_c}{K_e}e^{-\frac{r-r_e}{K_e}}\left(1 - e^{-\frac{r-r_e}{K_e}}\right)$. Each LSM node on the particle surface can adhesively interact with nodes on the microchannel surfaces located within a cut-off distance $r_{\text{cut}} = 1.5r_e$.

The fluid and solid models are coupled *via* the boundary conditions at the fluid–solid interface that utilize the interpolated bounce back method^{50,51} and the momentum exchange method.⁵² The computational methodology for fluid structure interactions has been previously extensively validated by simulating flows with different rigid and deformable particles including mesh independence study, and found good agreement with relevant theories and experiments.^{24,26,53–56}

3 Computational setup

The computational setup for simulating cells propelled by a fluid flow in a microchannel where the substrate is patterned with adhesive patches is illustrated in Fig. 1a, whereas the setup for simulating cell motion in a microfluidic channel with diagonal ridges is presented in Fig. 1b. The following dimensions are used for the computational setups: height $H = 1.5d$, width $W = 5d$, and length $L = 10d$. The adhesive stripe and ridge have a width of $W_r = 1.5d$ and are inclined at an angle ϕ with respect to the channel axis. The ridge forms a gap h with the bottom wall of the channel. The magnitude of the gap $\lambda = h/d$ is normalized by the particle diameter $d = 20$. Note that we express the dimensional quantities in LB units unless indicated otherwise.

To characterize the flow within the channel, we define a channel Reynolds number $\text{Re} = \frac{\rho H U_0}{\mu}$, which represents the ratio between the fluid inertia and viscous forces. The Reynolds number is based on the channel height H , the fluid density ρ and viscosity μ , and the characteristic flow velocity in the channel U_0 due to a pressure gradient $\Delta P/L$, defined as

$U_0 = \frac{\Delta P H^2}{12 L \mu}$. The compliant particle is characterized by a capillary number $\text{Ca} = \frac{\mu \dot{\gamma}_0 d}{2 G_s}$. Here, $\dot{\gamma}_0 = \frac{6 U_0}{H}$ is the characteristic shear rate, and $G_s = \frac{E t_s}{2(1 + \nu)}$ is the two-dimensional shear modulus of the particle elastic shell, where E is the particle Young's modulus, t_s is the shell thickness, and ν is the Poisson's ratio. Thus, the capillary number represents the ratio between viscous stresses on the particle and elastic stiffness of the particle shell. The adhesion level is controlled by varying the dimensionless adhesion strength $\beta = \frac{D_c/K_e}{3\pi\mu d U_0}$.

In our simulations the particles are initially positioned at the channel centerline. The particle trajectory is represented using dimensionless particle's center of mass coordinates $C_x = c_x/d$ and $C_z = c_z/d$. The particle motion is quantified using the dimensionless particle displacement per ridge $\delta = \frac{c_{z2} - c_{z1}}{d}$ where c_{z1} and c_{z2} are the particle's center of mass z -coordinate at $x = L/3$ and $x = L$, respectively.

4 Results and discussion

4.1 Cell motion over adhesive stripes

We first consider the case of compliant cells rolling on a substrate patterned with diagonal adhesive stripes, as shown in Fig. 1a. Here, only the diagonal patch shown in blue is adhesive, whereas the remaining channel surfaces are non-sticky. The motion of adhesive cells over microfluidic surfaces patterned with stripes was previously examined computationally and experimentally.^{39,57,58}

Fig. 2a presents the trajectories of the particle's center of mass in the x - z plane for particles with different adhesion strengths β and $\text{Ca} = 3.5 \times 10^{-3}$. As the adhesion strength β increases, the deflection δ increases. In Fig. 2b, we show a schematic illustration of the particle trajectory across an adhesive diagonal stripe and the main forces acting on the particle. Before encountering the stripe, the particle moves along the channel centerline due to a

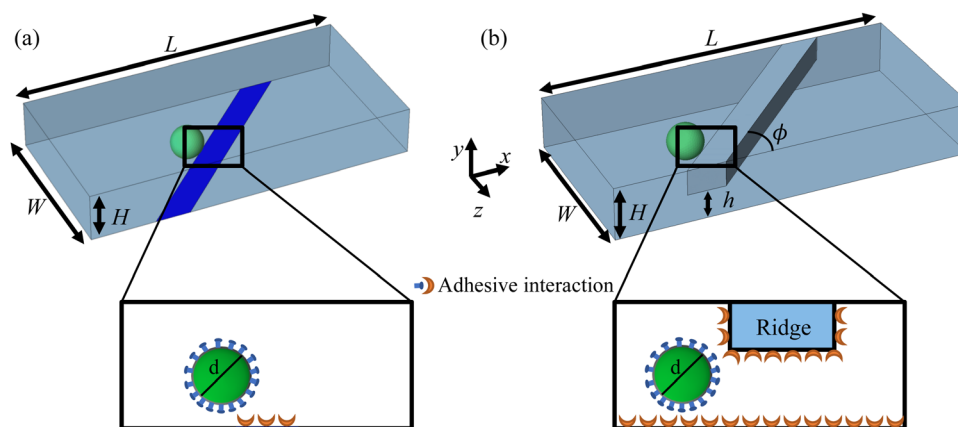


Fig. 1 (a) Microfluidic channel with substrate patterned with diagonal adhesive stripes. Only the patch shown in blue is adhesive. (b) Microfluidic channel with diagonal ridges for adhesion based sorting. The channel internal surfaces are coated with adhesion molecules that attach to the particle. Fluid flows in the x -direction.



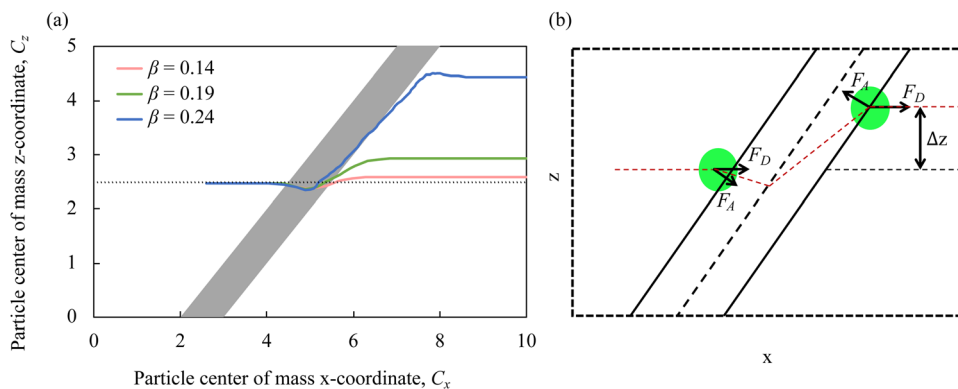


Fig. 2 (a) Particle trajectories in the x - z plane in a microchannel with an adhesive stripe for $Ca = 3.5 \times 10^{-3}$, $\phi = 45^\circ$, and $Re = 3$. (b) Schematic illustrating the trajectory of a particle due to the hydrodynamic drag force F_D and the adhesion force F_A as the particle crosses an adhesive stripe resulting in a transverse displacement Δz .

drag force F_D . When the particle approaches the adhesive stripe, it is pulled towards the centerline of the stripe by an adhesive force F_A . The adhesive force deflects the particle trajectory in the negative x direction. When the particle crosses the middle of the stripe, the adhesive force flips its direction and acts to keep the particle on the stripe to minimize the energy.³⁹ The combined action of F_A and F_D propels the particle along the stripe until it reaches the back edge of the stripe where the action of the adhesive force diminishes. After that the particle moves along the flow direction due to the drag force F_D . As a result, the particle experiences a net positive lateral displacement Δz with increasing adhesion.

Fig. 3a shows how the dimensionless particle displacement $\delta = \Delta z/d$ depends on the adhesion strength β for particles with different elasticity Ca . We find that δ increases with increasing β . Furthermore, δ increases with increasing Ca . Thus, softer particles experience larger lateral displacement as they cross the adhesive stripe, which can be explained by a larger contact area between softer particles and the adhesive stripe increasing the free energy change and, therefore, F_A .³⁹

Fig. 3b shows δ as function of β for different channel flow rates expressed in terms of the Reynolds number Re . We find that the deflection decreases as Re increases due to a greater contribution of the hydrodynamic drag force F_D . Furthermore, the increase of

Re above the values reported in Fig. 1b results in the particle failure to interact with the adhesive stripe due to an increased lift force translating the particle away from the substrate. Thus, the method while can be efficient in separating cells is limited to relatively low flow rates^{57,58} and, therefore, low cell throughput.

4.2 Cell motion in ridged microchannel

To ensure the contact of cells with adhesive microchannel surfaces independently of the channel flow rate, we use diagonal ridges that form a gap h between the ridge surface and the channel wall that is smaller than the cell diameter d , as shown in Fig. 1b. Furthermore, we consider that all internal surfaces of the ridged microchannel are coated with adhesion molecules, thereby eliminating the need for adhesive patterning.

Fig. 4a shows the trajectories of the particle's center of mass in the x - z plane for particles with different adhesion strength β . The figure also includes a typical streamline in the fluid flow passing through the microchannel middle plane. We find that the particles follow different trajectories depending on the magnitude of the adhesion β . The less adhesive particles deflect more along the ridge, whereas more adhesive particles more closely follows the flow streamline. Thus, the adhesion facilitates the particle crossing the ridge and, as a result, the negative transverse z -displacement increases with increasing β .

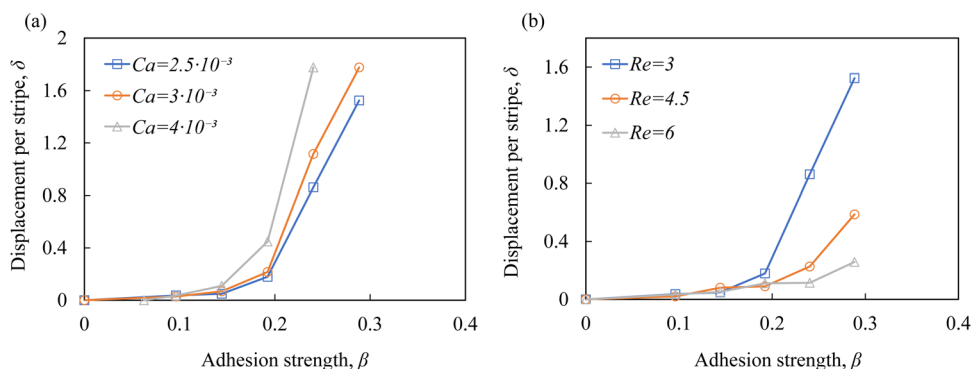


Fig. 3 Particle dimensionless lateral displacement per stripe $\delta = \Delta z/d$ as a function of the adhesion strength β in microchannels with adhesive stripes for (a) $Re = 3$ and (b) for $Ca = 2.5 \times 10^{-3}$, $Ca = 3.75 \times 10^{-3}$, and $Ca = 5 \times 10^{-3}$, respectively. The stripe angle is $\phi = 45^\circ$.



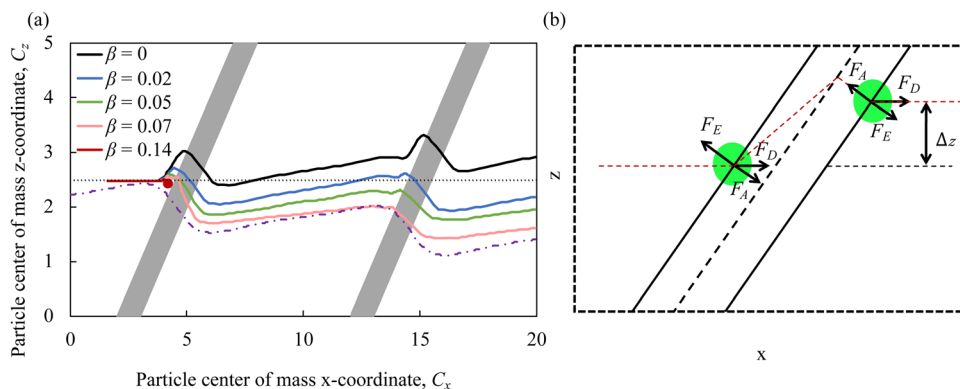


Fig. 4 (a) Particle trajectories in the x - z plane in a ridged microchannel for $Ca = 2.5 \times 10^{-3}$, $Re = 3$, $\lambda = 0.75$, and $\phi = 45^\circ$. Dash-dotted line represents a fluid streamline. Red line refers to a case of firm adhesion where the particle is arrested due to a large adhesion force. (b) Schematic illustrating the trajectory of a particle in a ridged microchannel due to the hydrodynamic force F_D , elastic force F_E and adhesion force F_A as the particle crosses a ridge resulting in a transverse displacement Δz .

Furthermore, we find that when the adhesion strength β exceeds a critical value β_{cr} , the particles are unable to cross the ridge as they get immobilized by the adhesive interaction with the ridge.

The motion of compliant adhesive particles in a ridged microchannel is governed by three main forces, as illustrated in Fig. 4b. The fluid drag force F_D acts along the flow direction due to the relative velocity between the fluid and the particle. The elastic force F_E arises due to the particle elastic deformation by the ridge and is directed away from the ridge. The adhesive force F_A is proportional to the gradient of the adhesive energy and is directed towards the ridge. Thus, the elastic force F_E and the adhesive force F_A have opposing effect on the particle trajectory and effectively counteract each other. Indeed, when F_A is weak, F_E causes the particle to move along the ridge leading to particle cross-channel displacement in the positive z direction. On the other hand, when F_A is comparable or exceeds F_E , it cancels out the effect of F_E on the particle trajectory and the particle follows the flow streamline. When F_A is too strong, F_D is unable to translate an adhered particle and the particle becomes immobilized by the ridge. We note that the idealized particle trajectory shown in Fig. 4b somewhat deviates from the trajectories obtained in the simulations reported in Fig. 4a. This can be related to the non-linearity associated with particle deformation by the ridge. The particle deformation gradually grows until approximately a half of the particle enters in the gap formed by the ridge. After that the elastic resistance due to particle deformation rapidly weakens and the particle can readily traverse the ridge under the action of the hydrodynamic and adhesion forces. Furthermore, diagonal ridges in the microchannel give rise to a circulatory secondary flow that transport fluid in the cross stream direction, thereby deflection particle trajectory in the positive z direction.^{24–26}

In Fig. 5, we plot the particle displacement per ridge δ as function of the adhesion strength β for particles with different elasticity Ca . The figure shows that at higher adhesion levels with $\beta > 0.05$, δ only slightly changes with Ca indicating that the motion under the ridge is dominated by the adhesive interactions. However, for weaker adhesion the particle displacement

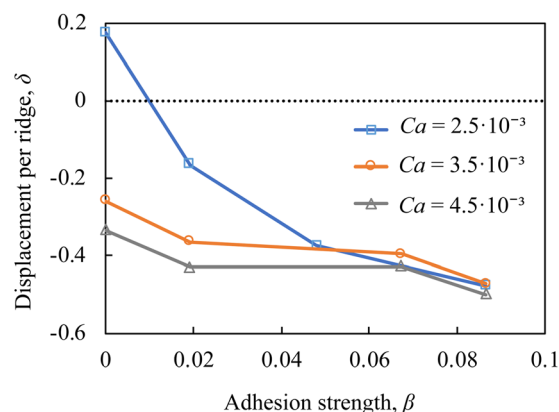


Fig. 5 Particle lateral displacement per ridge $\delta = \Delta z/d$ as a function of the adhesion strength β in a ridged microchannel with $Re = 3$, $\lambda = 0.75$, and $\phi = 45^\circ$.

increases as adhesion decreases. The difference in the displacement between particles with weak and high adhesiveness is the most significant for stiffer particles with low Ca . Thus, the sorting resolution increases as particle stiffness increases.

Fig. 6a and b compare the trajectories of particles with $Ca = 3.5 \times 10^{-3}$ and different adhesiveness in microfluidic channels with ridge angles $\phi = 30^\circ$ and $\phi = 15^\circ$, respectively. Note that the two figures have different scales of the x -axes, so the ridge angles are not shown to scale. We find that the separation between particles with different β is greater using ridges with $\phi = 15^\circ$ compared to $\phi = 30^\circ$. Furthermore, the trajectories of particles with $\beta = 0.03$ and $\beta = 0.05$ practically coincide when the ridge angle is $\phi = 30^\circ$, whereas for $\phi = 15^\circ$, the trajectories of such particles are well separated. Thus, the use of lower ridge angle not only increases the separation resolution, but also enables separation of particles with greater adhesiveness.

To explain the effect of ridge angle ϕ on the trajectories of adhesive particles, we consider how relative direction of forces acting on the particle change with ϕ (Fig. 4b). The elastic force F_E and the adhesive force F_A are oriented normal to the ridge. Thus the direction of F_E and F_A changes with ϕ , whereas their



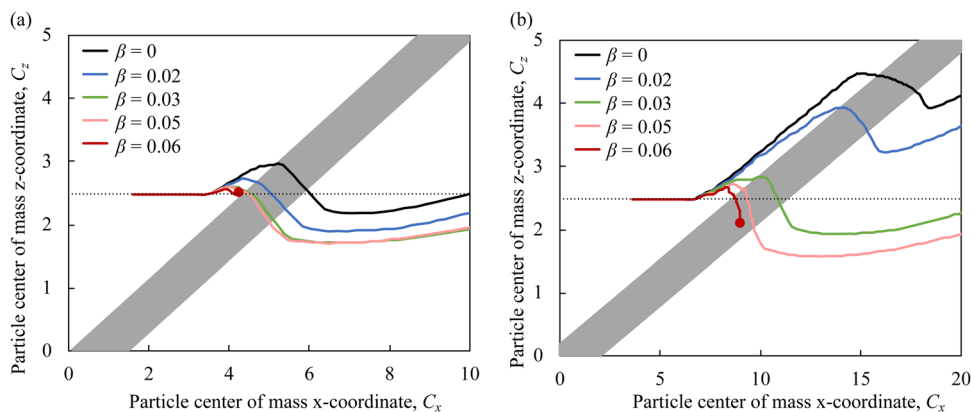


Fig. 6 Particle trajectories in the x - z plane in ridged microchannels with ridge angles (a) $\phi = 30^\circ$ and (b) $\phi = 15^\circ$. Other parameters are $Ca = 3.5 \times 10^{-3}$, $\lambda = 0.75$, and $Re = 3$. Red lines present the trajectories for particles that are arrested at the ridge due to a large adhesion force.

magnitudes are independent of ϕ . The drag force F_D acts along the flow direction independently of ϕ . Since the component of F_D acting normal to the ridge balances F_E and F_A , the magnitude of this component of F_D does not change with ϕ . This requires that the magnitude of F_D increases with decreasing ϕ , and therefore, the component of F_D directed along the ridge increases for smaller ϕ . This, in turn, results in a greater translation of the particles along the ridge enhancing the trajectory separation for particles with different adhesiveness.

Fig. 7 presents the particle displacement per ridge δ in microchannels with $\phi = 15^\circ$ and $\phi = 30^\circ$ as a function of the adhesion strength β for particles with $Ca = 2.5 \times 10^{-3}$ and $Ca = 3.5 \times 10^{-3}$. Note that the displacement $\delta = 2$ corresponds to the condition where particles deflect along the ridge all the way until they reach the channel sidewall. Furthermore, the dashed lines in Fig. 7 indicate the conditions where particle motion is arrested at the ridge due to the large adhesion force. For stiffer particles with $Ca = 2.5 \times 10^{-3}$, the microchannel with $\phi = 30^\circ$ provides significant separation among nonadhesive particles and particles with the adhesiveness up to $\beta = 0.05$. The same particles do not separate in the microchannel with $\phi = 15^\circ$, where they are deflected by the ridge to the channel side wall. On

the one hand, softer particles with $Ca = 3.5 \times 10^{-3}$ and different β display minor differences in δ in channels with $\phi = 30^\circ$ and, therefore, cannot be separated in such channels. The use of ridges with $\phi = 15^\circ$ significantly enhances the separation of softer particles. In fact, the channel with $\phi = 15^\circ$ yields separation of softer particles with $Ca = 3.5 \times 10^{-3}$ comparable to the separation of stiffer particles with $Ca = 2.5 \times 10^{-3}$ in microchannels with $\phi = 30^\circ$. This result suggests that adjustment of the ridge angle can be used to achieve an efficient adhesion-based sorting of particles with specific stiffness.

Finally, we assess the impact of higher Re on adhesion-based particle separation in the ridged microchannels. We modify Re by changing the fluid viscosity, in which case Ca does not change with Re . Fig. 8a and b show the trajectories of particles with $Ca = 2.5 \times 10^{-3}$ and different adhesiveness in microfluidic channels with $Re = 60$ and ridge angles $\phi = 30^\circ$ and $\phi = 15^\circ$, respectively. Note that the two figures are not shown to scale. We find that the separation between particles with different β occurs even at high Re that significantly exceeds the maximum Re that can be used in microchannels with patterned stripes (Fig. 1a). Similarly to lower Re (Fig. 6), ridges with smaller angle ϕ result in greater separation of adhesive particles. Thus, our simulations show that microfluidic channels with diagonal ridges can be effectively used for high-throughput sorting of adhesive particles.

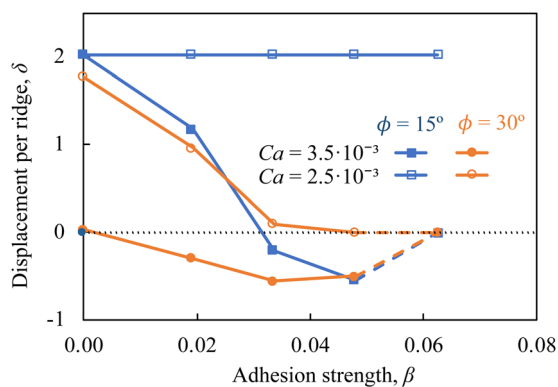


Fig. 7 Particle lateral displacement per ridge $\delta = \Delta z/d$ as a function of the adhesion strength β in microchannels with ridge angles $\phi = 15^\circ$ and $\phi = 30^\circ$. Other parameters are $\lambda = 0.75$ and $Re = 3$. Dashed lines indicate the conditions where the particle motion is arrested due to a large adhesion force.

5 Conclusions

Using three-dimensional computer simulations, we examine a microfluidic method for label-free separation and sorting of compliant cells by their adhesiveness. The approach consists of flowing cells in a microchannel decorated with skewed periodic ridges coated with adhesive molecules. The ridges form a gap smaller than the cell diameter thereby ensuring cell contact with adhesive surfaces independently of the channel flow rate. Thus the method opens up separation opportunities at high flowrate and high cell throughput.

Our simulations identified three modes of adhesive particle transport through the ridged microchannels. In the case of small



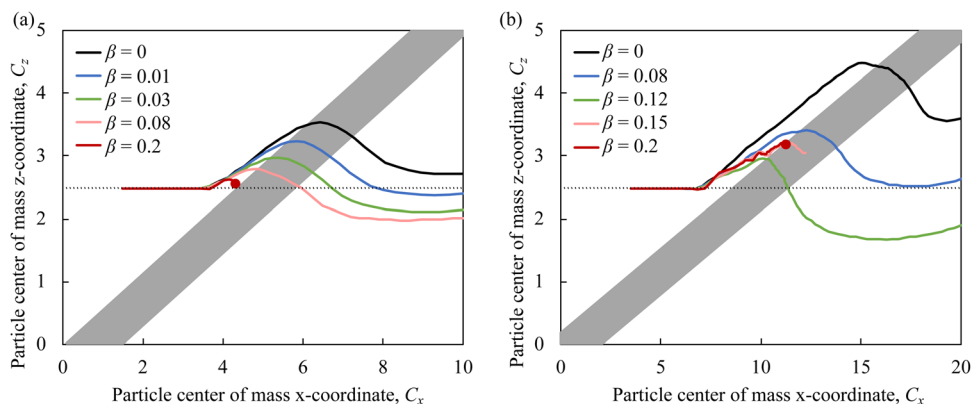


Fig. 8 Particle trajectories in the x - z plane in ridged microchannels for $Re = 60$ with ridge angles (a) $\phi = 30^\circ$ and (b) $\phi = 15^\circ$. Other parameters are $Ca = 2.5 \times 10^{-3}$, $\lambda = 0.75$. Red lines present the trajectories for particles that are arrested at the ridge due to a large adhesion force.

ridge angle, cell rolling can occur along the diagonal ridges in which a threshold of adhesion eliminates cell sliding. In the case of high adhesion, cells can occlude the ridge leading to extended contact times and irreversible binding. Finally, cell trajectories can be modified by adhesive ridges in a manner that greater displacement differences were observed for stiffer particles with low Ca number.

Our results demonstrate that cell trajectories within ridged microchannels are sensitive to the adhesion strength, which can be used to continuously sort cells based on their adhesiveness by collecting cells at different microchannel outlets. The simulations reveal that particle elasticity is essential for the adhesion-based sorting in ridged microchannels and can be used to enhance the sorting resolution by counteracting the adhesion force on the cells. We also show that the ridge angle can be tuned to achieve efficient adhesion-based sorting for cells with different mechanical properties. Furthermore, our simulations for $Re = 60$ show that ridged microfluidic channels can be effectively used for high-throughput adhesion-based particle sorting.

It is further interesting to examine the role of adhesion type on cell-ridge interactions. In these studies, we considered interactions represented by the Morse potential in which bond strength develops instantaneously as in the case of electrostatic interactions. In the case of specific bond adhesion, which is contact time dependent such as described by the Bell model, additional parameter space emerges affecting the cell transport that can be explored.

Data availability

All the data for the current study are available from the corresponding authors on reasonable request.

Conflicts of interest

There are no conflicts to declare.

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