HIGH-THROUGHPUT MICROFLUIDIC CAPTURE
OF RARE CELLS FROM LARGE VOLUMES OF
BLOOD

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A DISSERTATION
PRESENTED TO THE FACULTY
OF PRINCETON UNIVERSITY
IN CANDIDACY FOR THE DEGREE
OF DOCTOR OF PHILOSOPHY

RECOMMENDED FOR ACCEPTANCE
BY THE DEPARTMENT OF
ELECTRICAL ENGINEERING
ADVISER: PROFESSOR JAMES C. STURM

MAY 2016
Abstract

Deterministic lateral displacement (DLD) arrays are microfluidic devices capable of high-resolution separation of particles based on size. DLD arrays have been applied to separation of large cells from blood for a wide variety of diagnostic and analytical purposes. The volume of blood processed in these applications has been limited by volume-dependent performance degradation and throughput (volume/time) limitations. We address these issues in three ways in this thesis. First, we develop fabrication methods that increase the density of DLD arrays on a chip of a given area, resulting in an increase in the volumetric flow rate for a given pressure by a factor of ten. Second, we identify conventional platelet-driven clot formation as the source of the volume-dependent performance degradation and develop a method to completely inhibit clot formation in the DLD array, resulting in a 1000-fold increase in the volume of blood processed without device performance degradation. Third, we characterize the effect of post shape on the behavior of cells at high flow rates, corresponding to moderate Reynolds numbers (Re), by showing how post shape can be used to minimize shear-induced compression that reduces the target cell yield and to minimize hydrodynamic asymmetry that results in undesirable displacement of erythrocytes. Lastly, we finish by showing how post shape and row spacing can be used to minimize anisotropic conduction in DLD arrays that leads to non-ideal behavior of particles even at low Re.
Acknowledgements

I would like to thank the many people who have contributed to the work in this thesis. First, I would like to thank my advisor, Prof. James C. Sturm, for being an excellent mentor and for his unwavering support, encouragement, and enthusiasm. I would also like to thank Prof. Robert H. Austin for giving me the opportunity to do research in his lab and for his guidance in thinking about scientific problems and setting up experiments.

I would like to thank all of my colleagues and friends in the Sturm and Austin groups. Special thanks to Kevin Loutherback for introducing me to the equipment and facilities when I first joined the group and to Jason Puchalla for reading this thesis and providing helpful feedback. I am grateful to Amy Wu, Yu Chen, Qiucen Zhang, Saurabh Vyawahare, Ke-Chih Lin, George Liu, Karen Malatesta, and Brandon Comella for many interesting discussions.

I would like to thank my research collaborators for their contributions. I am grateful to the staff at GPB Scientific LLC for providing equipment for experiments and guidance in developing my experimental setup. Thanks to Dr. Curt Civin and his lab at the University of Maryland Medical School for helpful discussions and collaboration on early experiments. I would also like to thank Dr. Ken Pienta and his lab at the Johns Hopkins University School of Medicine for providing the PC3 cancer cells used in this thesis.

Devices used in the experiments in this thesis were fabricated at the Princeton Institute for the Science and Technology of Materials (PRISM) Micro/Nano Fabrication Laboratory. I appreciate the assistance from the cleanroom staff, especially Pat Watson, Bert Harrop, Mikhail Gaevski, and Joe Palmer. I would also like to thank the Physics Department Machine Shop staff, especially Bill Dix, for making the manifolds and other pieces of equipment used in the experiments in this thesis.
The research described was funded by GPB Scientific LLC, an NIH STTR Award (Project 1R41HL110574) via GPB Scientific LLC, and by the National Cancer Institute (Grant U54CA143803) through the Physical Sciences in Oncology Centers (PSOC) Program.

I would like to thank all of my friends at Princeton and elsewhere. Finally, thanks to my family for their constant encouragement and support.
To my parents.
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Chapter 1

Introduction

1.1 Continuous Flow Separations

Separation of cells, functionalized beads, or other particles based on size, density, compressibility, or other properties is a critical step in many analytical and preparative techniques in medicine and biology [61]. These separations have traditionally been performed using bulk processes, such as filtration or centrifuge, which require more preparative steps and larger quantities of reagents. Microfluidic techniques offer a method of performing these separations more quickly, with better resolution, and using smaller quantities of reagents. These devices are fabricated using techniques borrowed from integrated circuit technology, and devices have been patterned in a variety of bulk materials, including silicon, glass, polymer, and plastics [61].

A wide variety of microfluidic techniques have been developed to perform such separations. All of these techniques rely on an effective force applied to the desired particles in a direction perpendicular to the flow, causing lateral displacement of a particle when the flow direction is defined as vertical. The magnitude of this displacement is dependent upon properties of the particle, such as size, density, compressibility, or charge. This spatial displacement of particles as the particles move
through the channel allows for both on-chip analysis of the spatial distribution of the particles and direction into separate outlet channels for off-chip analysis. For most applications involving high-throughput separations of cells from blood, three microfluidic techniques have been used: acoustic standing waves, inertial lift and Dean drag forces, and deterministic lateral displacement, which is the focus of this thesis.

Microfluidic separation techniques rely on laminar flow, which is the condition under which a particle will follow a streamline unless acted upon by an external force. The flow is characterized by the Reynolds number Re, which is a dimensionless number used to characterize the ratio of inertial to viscous forces in the flow. It is defined as \( \text{Re} = \frac{\rho v L}{\mu} \), where \( v \) and \( L \) are the characteristic velocity and length scale that describe the flow, \( \rho \) is the density of the fluid, and \( \mu \) is the dynamic viscosity of the fluid. The flow is purely laminar up to Re of approximately 40, beyond which laminar flow is locally disrupted in certain regions due to vortex shedding. More relevant for the work in this thesis is Re~1, when inertial effects become important, causing changes in the flow pattern, up to Re of 20.

### 1.2 Continuous Flow Separation of Blood

The ability to separate cells in blood into distinct populations based on physical properties such as size or compressibility has numerous diagnostic and analytical applications. In the case of epithelial tumor cells, it has been shown that increased deformability of the epithelial tumor cells correlates with increased metastasis. Erythrocytes infected with malaria are approximately 50 times stiffer than erythrocytes that are not infected with malaria and have been isolated on this basis using microfluidic techniques. Two major classes of microfluidic techniques have been used in separating cells in blood based on physical properties: active, which require an energy source input, and passive, which do not. We briefly discuss the major
achievements using techniques in each class in sorting cells in blood based on physical properties.

1.2.1 Passive Microfluidic Separation Techniques

Passive microfluidic separation techniques do not require an energy input to enable sorting and typically rely on properties of the flow within the microfluidic channel. Most applications involving high-throughput separations of cells from blood utilize three techniques: cross-flow filtration, inertial lift and Dean drag forces, and deterministic lateral displacement (DLD). Perfusion in continuous microfluidic cross-flow has been used to separate leukocytes from whole human blood [82]. Inertial lift forces, specifically the shear-gradient lift force and the wall lift force, cause certain cells to migrate to a fixed equilibrium distance away from the channel walls and have been used to separate cancer cells from blood on the basis of size and deformability [33]. High-throughput separation of cancer cells from blood has also been achieved using size-selective vortex technology and slanted-spiral microfluidics, both of which also rely on inertial lift and Dean drag forces to achieve separation (Dean drag forces relate to particle behavior in vortices which form perpendicular to the flow direction in sufficiently curved micro-channels.) [74, 83]. DLD has been used to separate blood into leukocytes, erythrocytes, and platelets and plasma, as well as for separation of specific cell types from whole blood for a wide variety of diagnostic applications, which we discuss later in this chapter [14].

1.2.2 Active Microfluidic Separation Techniques

We define active microfluidic separation techniques as those which require an energy input to enable sorting or require the attachment of a functionalized label to the desired particles. Some active microfluidic separation techniques, such as dielectrophoresis, magnetic-force-based sorting, and optical trapping, require the attach-
ment of functionalized beads with specific properties to enable the separation of cells from blood [72]. Micro-fabricated magnetic stripes in the bottom of a micro-channel have been used to separate leukocytes, which had been selectively labelled with magnetic nanoparticles, from whole blood [38]. In contrast, separation using acoustic standing waves is typically label-free and depends only on the relative densities and compressibilities of the particles and the carrier medium. Acoustic standing waves have been used to transfer erythrocytes from contaminated plasma to clean plasma in a microfluidic channel [64].

1.3 Deterministic Lateral Displacement: Principles of Operation

Deterministic lateral displacement (DLD) arrays are microfluidic devices that offer continuous-flow separation of particles suspended in a fluid based on size. The mechanism of action is that suspended particles in a fluid that are larger than a critical size experience sequential displacement (bumping) from one streamtube to an adjacent one in a direction perpendicular to the flow by micro-posts that are arranged in a tilted rectangular or parallelogram array [30]. Particles smaller than this critical size travel in the average flow direction, while particles larger than this critical size travel at the array tilt angle.

Figure 1.1 is a schematic showing the relevant parameters and typical particle behavior. The critical size is a function of the gap, $G$, and the array tilt angle $\epsilon = d/\lambda$ [35]. The tilt angle, $\epsilon$, is typically chosen such that $1/\epsilon$ is an integer. For circular posts, the diameter of the post is typically set equal to the gap, $G$. The array shown in Figure 1.1 uses a parallelogram unit cell, although the tilt can also be achieved using an array with a tilted rectangular unit cell. Since there can be no
Figure 1.1: Schematic showing the separation of a large particle from a small particle via a DLD array with a row-shift fraction of one-third. The row-shift fraction ($\epsilon = d/\lambda$) divides the flux in the gap (G) between the posts into $1/\epsilon = 3$ streamtubes, which are divided by streamlines. Large particles, defined as particles larger than the width of the first streamtube next to the post on the side of the post that displaces the particle, are displaced by the post with each subsequent row in the direction of the flow, traveling at the array tilt angle over the course of many rows of posts. Small particles, defined as particles smaller than the width of the first streamtube next to the post on the side of the post that displaces the particle, follow the streamtube, traveling in the average fluid flow direction over the course of many rows of posts (illustrated by the black arrow). Image courtesy of J.A. Davis et al. [14].

fluid flux into the walls, the fluid flow is on average strictly vertical, as shown by the black arrow in Figure 1.1.

The device works by asymmetric bifurcation of the fluid streams around the posts. The array tilt angle, $\epsilon$, divides the flux in the gap, G, between the posts into $1/\epsilon$ streamtubes. $1/\epsilon - 1$ of these streamtubes pass through the next gap, while the remaining streamtube passes around the opposite side of the next post. These streamtubes are separated by stagnation streamlines that begin and end on posts and show how the fluid divides around the posts.
Particles in the flow behave in one of two ways depending upon the size of the particle relative to the size of the first streamtube next to the post on the side of the post that displaces the particle. If the radius of the particle is larger than the width of the streamtube next to the post, the particle will be bumped by the post into the adjacent streamtube. If the radius of the particle is smaller than the width of the streamtube next to the post, the particle will remain within the current streamtube and follow a zigzag trajectory as it travels through the array. Because of the cyclical way the streamtubes move through the gaps, large particles travel at the tilt angle of the array while small particles follow the average flow direction, which is vertical.

The critical size separating large particle behavior from small particle behavior can be predicted using a method developed by D. Inglis et al. The flux in the gap is divided into $1/\epsilon$ streamtubes of equal volume flux, and the critical particle radius is the width of the streamtube next to the post on the side of the post which bumps the particle. This can be found by simple integration over the velocity profile in the gap.

Post geometry can be used to alter the velocity profile in the gap therefore also the critical size. K. Loutherback et al. showed that asymmetric triangular posts allow for a 50% larger gap for a given critical size and tilt angle. Asymmetric triangular posts can also be used to achieve a different critical size in the same array for opposite flow directions. The effect of post shape is investigated in detail in Chapter 4.

Because deterministic lateral displacement relies only on particles following streamlines unless displaced by posts, the separation is independent of flow velocity as long as the flow remains laminar to first order. At Re>1, even if the flow is laminar, several second order effects can affect DLD array performance. These are discussed in Chapter 4 and include inertial effects affecting the fluid flow and particle paths and effects of the fluid flow on the particle shape and orientation. To first
order, the separation resolution is enhanced by increased flow velocity, since increased flow velocity allows less time for diffusion \[30\]. Compared to acoustic and inertial microfluidic cell sorting techniques, deterministic lateral displacement does not have a minimum channel length that depends on flow velocity.

The capabilities of DLD arrays have been expanded through use in combination with active microfluidic separation techniques. Dielectrophoresis has been used to allow tunability of the critical size in DLD arrays, which is typically determined purely by hydrodynamic conditions \[5\]. An electrode-based virtual pillar array which combined DLD with dielectrophoretic effects has been used to achieve 99% separation of leukocytes from erythrocytes \[8\]. The DLD concept has also been extended using interdigital transducers to create acoustic standing waves at an angle to the average flow direction, which allows size-based separation of particles \[12\].

1.4 Deterministic Lateral Displacement: Applications toward Separation of Nucleated Cells from Blood

Table 1.1: Physical characteristics and concentrations of the major components in whole adult human blood.

<table>
<thead>
<tr>
<th>Cell Type</th>
<th>Cell Shape</th>
<th>Diameter ($\mu$m)</th>
<th>Concentration ($\text{mL}^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelet</td>
<td>Various</td>
<td>1-5</td>
<td>5.0-10.0x10^7</td>
</tr>
<tr>
<td>Erythrocyte</td>
<td>Biconcave Discoid</td>
<td>2x7.5</td>
<td>4.0-5.5x10^9</td>
</tr>
<tr>
<td>Leukocyte</td>
<td>Spheroid</td>
<td>5-20</td>
<td>4.0-10.0x10^6</td>
</tr>
</tbody>
</table>

Blood contains cells that range in size from 1$\mu$m to 20$\mu$m, with the size of a cell often being related to its biological function. With a critical size that is set by design parameters, DLD arrays are well suited to separating blood into its main components.
of leukocytes, erythrocytes, and platelets [14]. The sizes and concentrations of these components in whole adult human blood are shown in Table 1.1.

DLD arrays have been applied to the capture of variety of rare cells of biological interest from whole blood. Compared with other microfluidic techniques, the major advantage of DLD arrays in these applications is the ability to capture these rare cells without the background of erythrocytes present in whole blood [40]. D.W. Inglis et al. demonstrated that DLD arrays can be used to separate malignant lymphocytes from healthy lymphocytes [36]. L.R. Huang et al. have used DLD arrays to capture nucleated red blood cells from the peripheral blood of pregnant women for applications in prenatal diagnostics [32]. S.H. Holm et al. have used DLD arrays to separate parasites from human blood [26]. B. Zhang et al. have used DLD arrays to separate cardiomyocytes from blood [91]. Typical volumes of blood processed for such applications have been limited to 100 µL per DLD array due to volume-dependent performance degradation.

While the capture efficiencies achievable with DLD arrays are sufficiently high to be useful in rare cell capture (>85%), the volumes of blood processed in the previous examples were small (<100 µL). This is insufficient for many diagnostic purposes. To process large volumes of blood, high-throughput operation of DLD arrays for capture of cancer cells from blood has recently been demonstrated. K. Loutherback et al. have demonstrated capture of greater than 85% of cancer cells from diluted whole blood at flow rates as high as 10 mL/min (Figure 1.2) [50]. Z. Liu et al. demonstrated capture of cancer cells from diluted whole blood at similar flow rates and showed that capture rates remain high at these high flow rates with DLD arrays using triangular posts but decrease with increasing flow rate for DLD arrays using circular posts [47]. However, both of these experiments still only processed small volumes of blood (<200 µL per DLD array).
1.5 High-Throughput, High-Volume Separation of Cells from Blood using DLD Arrays

This thesis aims to set the foundation for using DLD arrays to process large volumes of blood (10-100 mL). Two potential applications are the capture of circulating tumor cells (CTCs) and harvesting of stem cells. First, CTCs have been identified in the blood of patients with metastatic disease and can be used to assess the severity of disease and the effectiveness of various treatment strategies [13]. CTCs can be isolated from blood using DLD arrays since these cells typically have diameters of 15 to 30 µm, with only slight overlap in the size distribution with leukocytes [58]. Second, there are a number of applications involving isolation of leukocytes and cells similar in size to leukocytes from whole blood. One particularly interesting application is the

Figure 1.2: Concentrator DLD array used to achieve high-throughput (10 mL/min) enrichment of cancer cells from blood. The path of the MDAMB231 cancer cells are shown as green streaks. The cells are evenly distributed across the entire width of the array at the inlet and are concentrated into the center of the array at the outlet. (Figure courtesy of K. Loutherback.)
isolation of hematopoietic stem cells (HSCs), which are similar in size to leukocytes, from umbilical cord blood (UCB) [75]. Currently, isolation of HSCs from UCB is achieved using centrifugation technology, which has limited resolution resulting in loss of HSCs and contamination by erythrocytes and involves large amounts of reagents. We examine separation of leukocytes from whole blood as a model system for isolation of HSCs from UCB.

Both of these applications require processing 10-100 mL of blood. CTC concentrations in the blood of patients with metastatic cancer can be as low as 1-10 mL$^{-1}$, typically requiring processing of $\sim$10 mL of blood for diagnostic applications [71]. UCB is collected in volumes of 50-100 mL [78], and maximizing the stem cell yield requires processing this entire volume.

Applications of DLD arrays toward separation of cells from large volumes of blood (10-100 mL) face two major limitations. First, while the flow rates at which DLD arrays can be operated has increased by a factor of $10^5$, the volume of blood that can be processed with DLD arrays has not increased by a similar factor due to clot formation in the array. Increasing the volume of blood that can be processed with DLD arrays requires identifying and inhibiting the mechanisms driving this clot formation. Inhibition of clot formation in DLD arrays will also allow for collection of target cells in a buffer stream without the background of erythrocytes and platelets present in blood. This is not possible with the concentrator design that has been used for high-throughput enrichment of cancer cells from blood with DLD arrays [50, 47].

Second, the behavior of cancer cells, leukocytes, and erythrocytes at high flow rates has not been well characterized in two ways. The behavior of erythrocytes at high flow rates in DLD arrays is important because a major advantage of DLD arrays over other microfluidic technologies is the ability to separate nucleated cells without the background of erythrocytes present in whole blood. For nucleated cells, such as cancer cells and leukocytes, the main concern with high throughput operation of DLD
arrays is minimizing shear-induced reduction in size of the cell, so that a high capture efficiency based on displacement of cells above the critical size of the DLD array can be maintained. Overcoming these two limitations will allow for high efficiency separation of nucleated cells from large volumes of whole blood using DLD arrays.

1.6 Thesis Outline

Chapter 2 details the experimental methods and setup for high-throughput operation of DLD arrays. We introduce three fabrication processes that increase the density of DLD arrays in a given chip area by a factor of 4. Two of the three fabrication processes increase the maximum depth of the DLD arrays by a factor of 2.5, yielding a 10-fold increase in volumetric flow rate for a given pressure over that which was achievable using previous fabrication processes. We also introduce a bubble-free manifold design for operating the high-throughput manifold over long times to process large volumes. Lastly, we introduce the imaging setup and cell labelling techniques utilized for the experiments in Chapter 3.

Chapter 3 demonstrates the inhibition of clot formation, removing a key limitation to processing large volumes of blood using DLD arrays. We identify that clot formation occurs dynamically at the beginning of the array and is not due to clots that form in the tubing leading to the chip. We then show that platelets, not leukocytes or erythrocytes, are the driver of this clot formation, and we examine the biological and physical mechanisms that contribute to clot formation. We then show that the activity of calcium ions and thrombin are the dominant mechanisms driven clot formation. We inhibit these mechanisms and show that this method of inhibition is effective at both high and low flow rates and correspondingly high and low shear rates.
Chapter 4 discusses the effects of post geometry on the behavior of erythrocytes and leukocytes in DLD arrays at high flow velocities (moderate Re: 1<Re<20). First, we examine the effect of post geometry on the extent of cell deformation against the micro-posts at high flow rates. We show that the extent of cell deformation is directly related to the shear at the post surface against which the cell can be deformed resulting from fluid bending around the post. Second, we examine the flow velocity dependent erythrocyte contamination of the product that occurs with asymmetric triangular posts but not with circular posts in DLD arrays with shallow tilt angles (tilt < 1/20). Using eight different post geometries, we show that the extent of erythrocyte contamination of the product at high flow velocities is directly related to the asymmetry of the fluid centripetal acceleration distribution in the gap. Finally, we combine the results of these two sets of experiments to achieve high yield separation of leukocytes from blood with minimal erythrocyte contamination using a DLD array with diamond-shaped posts.

Chapter 5 examines the effect of post geometry on anisotropic conduction in the DLD array at low Re. Anisotropic conduction in DLD arrays, which causes a misalignment of the pressure gradient to the average flow direction, can be caused by either a rhombic (not tilted square) unit cell array or by post shapes that are asymmetric. We first examine how the tilt of the array and post geometry contribute to anisotropic conduction. We then show that anisotropic conduction can be minimized by increasing the spacing between the rows in the direction of the flow.

Chapter 6 summarizes our contributions and makes suggestions for future work in this area.
Chapter 2

High-Throughput Deterministic Lateral Displacement Array
Experimental Setup

2.1 Introduction

In this chapter, we cover the major components of the experimental setup for operating high-throughput deterministic lateral displacement arrays. First, we develop fabrication techniques to increase the channel depth and density of DLD arrays on a chip of a given size. We then develop a manifold through which fluidic connections are made to the chip that can be operated at high pressures and is resistant to the formation of bubbles that can form when operating the DLD arrays for long times to process large volumes. We then present the imaging setup used to visualize beads, leukocytes, and cancer cells in the DLD array.
2.2 Parallelization of Deterministic Lateral Displacement Arrays for High-Throughput Processing of Blood

Figure 2.1: (a) SEM image of a DLD array etched into silicon using deep reactive ion etching (DRIE). Image courtesy of Keith Morton. (b) Setup for sandblasting through-holes, through which fluid enters and exits the chip. Limitation in the precision of the through-hole size and positioning of the nozzle limit the number of DLD arrays that can fit in a given area. Image courtesy of David Inglis.

Figure 2.2: The through-holes of the DLD array chip mate via O-rings to an acrylate manifold, through which fluidic connections are made to the DLD array chip. Image courtesy of David Inglis.

Deterministic lateral displacement (DLD) arrays with deep channels (depth > 10 µm) have traditionally been fabricated in silicon using deep reactive ion etching.
(DRIE) (Figure 2.1(a)). Through-holes through which fluid enters and exits the chip have traditionally been created via sand-blasting (Figure 2.1(b)). The through-holes of the DLD array chip mate via O-rings to an acrylate manifold, through which fluidic connections to the DLD array chip are made(Figure 2.2). The lack of precision in size and placement of the through-holes made via sand-blasting limits the number of DLD arrays that can be placed in a given area on a silicon chip. For this reason, a typical 1.5-mm wide DLD array requires approximately 8-mm width on the silicon chip. We present three techniques that overcome this limitation: (i) a technique using an oxide layer for pattern transfer that allows two patterns to be etched into a single side of a silicon wafer, one of which is for precisely placed through-holes, (ii) a technique using backside alignment that allows etching of two patterns, one of which is for precisely placed through-holes, into a silicon wafer by first performing photolithography and etching one side of the wafer and then performing the same on the opposite side of the wafer, and (iii) a technique that utilizes a powerful laser with alignment capability to within 10 µm and size control to within 25 µm to blast the through-holes. These three techniques allow for fitting a typical 1.5-mm wide DLD array in approximately 1.6-mm width on a silicon chip.

2.2.1 Oxide Mask Technique: All Front-Side Processing

The sandblasted through-holes, with the required spacing to compensate for the lack of precision in size and placement, cause a typical 1.5-mm wide DLD array to require approximately 8-mm width on a silicon chip. In principle, with anisotropic deep reactive ion etching, we could etch through-holes patterned via photolithography such that a typical 1.5-mm wide DLD array would require only 1.6-mm width on the silicon chip, a reduction in area of a factor of five. A challenge however occurs when the micro-post array is etched first. In trying to spin a 7 µm thick layer of AZ4330 photoresist over the micro-post array, which is typically etched to a depth greater than
Figure 2.3: Cross-sections of major steps in the fabrication process that uses an oxide mask to achieve etching of two aligned patterns on the front side of a silicon wafer.

60 \mu m, the photoresist coverage is non-uniform. When we etch the through-holes, the regions that are not covered by photoresist due to the non-uniform coverage are also etched through along with the through-holes, resulting in randomly placed holes through the array.

Oxide masks have been used for pattern transfer in the fabrication of micro-electro-mechanical systems (MEMS) devices [65]. We grow an oxide with thickness of 1.2 \mu m using a wet oxidation process. Using a standard deep reactive ion etch (DRIE) Bosch process, the selectivity between the oxide and silicon is such that 80 \mu m of Si can be etched for each 1 \mu m thickness of the oxide mask. For comparison, 100 \mu m of Si can be etched for each 1 \mu m of thickness in AZ4330 photoresist mask using the same etch recipe.

The major steps involved in this fabrication process are shown in Figure 2.3. The micro-post array is first patterned in a 1.4 \mu m thick layer of AZ5214 photoresist. The pattern is then transferred to the oxide using a buffered oxide etch (BOE). A 7 \mu m thick layer of photoresist is then spun on top of the patterned oxide layer. The photoresist coverage is uniform now because we are spinning a 7 \mu m thick layer of photoresist on top of a patterned surface with 1.2 \mu m depth instead of 60 \mu m.
depth (Figure 2.3), as we attempted to do in the process without the oxide mask. The through-holes are then etched using the 7 μm thick photoresist as a mask. This photoresist mask is then removed, and the micro-post array is etched through the oxide mask to a depth of 80 μm.

With this process, we can etch through-holes as small as 100 μm by 100 μm squares through a standard 550 μm thick 4-inch wafer. Minimum spacing between through-holes is 500 μm. We successfully fabricate a 70-mm by 36-mm chip containing 18 DLD arrays with 108 through-holes of 100 μm by 100 μm size using this process.

The major drawback of this process is that the 1.2-μm oxide mask was etched isotropically using BOE. This results in the size of the micro-posts being reduced in size by an amount equal to the thickness of the oxide. While this can be easily compensated for in the case of circular posts, it can also affect the sharpness of post geometries with corners, such as triangles. This reduction in sharpness can have a significant effect on critical size [49]. Possible future work could use an anisotropic dry etch of the oxide layer.

### 2.2.2 Backside Alignment Technique

![Diagram of backside alignment process](image)

Figure 2.4: Cross-sections of the major steps in the fabrication process that uses backside alignment to pattern and etch the micro-post array on one side of the wafer while through-holes are patterned and etched from the other side of the wafer.
Silicon is nearly transparent for near infrared light. This allows for a mask alignment technique in which a pattern in photoresist on one side of a wafer can be aligned to features that have been etched into the opposite side of the wafer. Alignment is easier if the depth to which features have been etched on the opposite side of the wafer is large, since contrast is achieved through thickness-dependent absorption of the infrared light by the silicon wafer. Here, we use this backside alignment technique to pattern through-holes in photoresist aligned to a micro-post array that has been etched into the opposite side of the wafer.

The major steps involved in this fabrication process are shown in Figure 2.4. A 3 \( \mu \text{m} \) thick layer of AZ4330 photoresist is spun onto one side of a double-side polished silicon wafer. The micro-post array is patterned and etched to a depth of 160 \( \mu \text{m} \). A 7 \( \mu \text{m} \) thick layer of AZ4330 photoresist is then spun onto the opposite side of the wafer. A specially designed chuck that elevates the wafer above the surface of the chuck so as not to scratch the features that have been etched into the opposite side of the wafer is used in spinning on the photoresist. The pattern for the through-holes is then aligned to the micro-post array etched on the opposite side of the wafer using a mask aligner equipped with an infrared light source and camera. The through-holes are then etched using a standard anisotropic etch (Bosch) process. Details of this fabrication process can be found in Appendix B.

This fabrication process overcomes two major limitations imposed by the previous process. First, the limitation on etch depth achievable for the micro-post array is limited by the maximum aspect ratio at which the micro-posts can remain standing, which is approximately 10:1, not by the thickness of the mask through which the micro-post is etched. Second, the dimensions of the micro-posts are identical to the mask pattern, whereas with the oxide mask process, the dimensions of the micro-posts were affected by the isotropic etching of the oxide mask. These are the two main practical differences between the two techniques.
2.2.3 Laser Cutting Technique

Patterning through-holes using the backside alignment technique and the oxide mask technique can be challenging due to non-uniformities in the photoresist layer caused by particles in the photoresist. Instead of using a Bosch process to etch through the wafer, we use a powerful YAG laser to cut through-holes as small as circles with 100 µm diameter. Although our system is not set up for alignment to automate the positions at which through-holes are cut, commercial systems can be programmed to achieve this with positional precision as high as 10 µm. We use this technique to rapidly fabricate chips with a single channel, requiring only six through-holes, for testing the effects of different post geometries in Chapters 3 and 4 of this thesis.

2.3 Sealing of the Micro-Post Array

Sealing of the micro-post array is important in high-throughput applications for two reasons. First, any irregularities in the sealing layer cause distortions in the flow pattern, which can cause displacement of particles below the critical size. Second, the sealing layer needs to be able to withstand high pressure. It has also been shown that a sealing layer with a high Young’s modulus, such as the Pyrex glass slides used in anodic bonding, makes the flow less sensitive to fluctuations caused by the syringe pump [90].

2.3.1 Pressure-Sealing Tape

Before sealing with the pressure-sealing tape, the chip is cleaned using a standard Piranha clean and then exposed to an oxygen plasma so that the surface is made hydrophilic. The Piranha clean begins with immersing the chip in a 1:1 mixture of 96% sulfuric acid and 30% hydrogen peroxide for 15 minutes. The chip is then
rinsed with DI water and dipped in 2% hydrofluoric acid for 1 minute. The chip is rinsed with DI water and then dried under nitrogen. In order to make the surface hydrophilic, the chip is exposed to a 1kW oxygen plasma for 90s.

Compared to a thin PDMS layer backed by a glass slide, which is the method previously used to seal DLD arrays, the pressure-sealing tape (9795R, 3M Co.) contains fewer defects that can cause deviations in the flow pattern in the chip. The backing from the pressure-sealing tape is removed, and the tape is positioned on top of the chip with no pressure applied. A rubber rolling pin is then used to seal the tape to the chip without any bubbles in the array region. Compared to the depth to which the chip is etched, there is negligible caving of the tape into the array (Figure 2.5).

One advantage of the pressure-sealing tape is that it can be easily removed to enable cleaning and re-use of the chip. To remove the tape, the sealed chip is first soaked for roughly two days in acetone. A razor is then used to lift the tape from
the corner of the chip, creating a tab. Using this tab, the tape can be easily peeled from the chip before the acetone evaporates. If the acetone evaporates before peeling is completed, the chip can be simply re-dipped in acetone before peeling of the tape is continued.

This method has been used at pressures as high as 50 psi without leaking. However, at pressures above 10 psi, the tape can detach from the chip in certain regions over the array, causing irregular behavior of particles in the array. Therefore, anodic bonding of Pyrex lids was investigated (Section 2.3.2). All of the experiments involving imaging were conducted using DLD arrays sealed with pressure-sealing tape.

2.3.2 Anodic Bonding

We use air anodic bonding to attach a Pyrex lid to the silicon chip into which the DLD array has been etched. The surface properties of the Pyrex slide and silicon chip are extremely important in successfully achieving anodic bonding since the bonding is due to the forces between opposite charges that are driven towards the interface between the Pyrex slide and silicon chip [45]. The Pyrex slide is cleaned by immersion for 15 minutes in a 1:1 mixture of 96% sulfuric acid and 30% hydrogen peroxide. The silicon chip is cleaned in the same way as the Pyrex slide with an additional step. To remove the native oxide from the surface of the silicon chip, the chip is immersed for 90s in 2% hydrofluoric acid. The Pyrex slide and silicon chip are pushed together without being blown dry and mounted in the holder shown in Figure 2.6, which is used to align the two pieces for bonding. The holder with the Pyrex slide and silicon chip mounted are kept under water until anodic bonding is performed to prevent oxidation of the surface of the silicon chip.

The setup for anodic bonding is shown in Figure 2.6. The holder with the Pyrex slide and silicon chip are removed from under water and placed on the hot plate. A piece of graphite is placed on top of the Pyrex slide, and the voltage probe is placed
Figure 2.6: Setup for anodically bonding a Pyrex glass slide to a silicon chip into which a DLD array has been etched. A voltage of -1000V is applied on top of the Pyrex slide through a probe connected to the Pyrex slide through a piece of graphite, which is simply placed on top of the Pyrex slide. The base of the hot plate is grounded. The Pyrex slide and silicon chip are placed on top of a stainless steel holder to align the two pieces. Anodic bonding is performed at a temperature above 500°C in air.

on top of this piece of graphite. Alignment between the silicon chip and Pyrex in the holder is checked. Both the Pyrex slide and silicon chip are still wet at this point. The hot plate is then set to 550°C, and the temperature is allowed to stabilize over 30 minutes. The voltage, set at -1000V, is then turned on. As bonding proceeds, the dark region shown in Figure 2.6 expands to eventually cover the entire area of the chip. Once bonding is complete, the hot plate is turned off, while the voltage is left on as the chip cools to prevent detachment of the bond due to uneven thermal shrinking of the Pyrex slide and silicon chip. The voltage is turned off once the hot
plate reaches a temperature of 50°C. For safety, the voltage supply is operated with a current trip set to 5 mA.

### 2.4 Bubble-Free, High-Pressure Manifold Design

![Acrylic manifold for fluidic connections to a lidded silicon chip into which the DLD arrays have been etched. The through-holes on the silicon chip align to reservoirs in the manifold. Each reservoir is connected on one side to a fluidic connection and on the other side to a hydrophobic filter, which is used to prevent bubbles from entering the chip. Spring-loaded screws evenly distribute the pressure onto the O-rings surrounding the reservoirs, allowing for operation at high pressures.](image)

Figure 2.7: Acrylic manifold for fluidic connections to a lidded silicon chip into which the DLD arrays have been etched. The through-holes on the silicon chip align to reservoirs in the manifold. Each reservoir is connected on one side to a fluidic connection and on the other side to a hydrophobic filter, which is used to prevent bubbles from entering the chip. Spring-loaded screws evenly distribute the pressure onto the O-rings surrounding the reservoirs, allowing for operation at high pressures.

The silicon chip is mounted into an acrylic manifold to which fluidic connections are made (Figure 2.7). This manifold has two advantages over manifolds that have been used previously. First, it is made resistant to bubbles forming in the reservoirs in the manifold by connecting each reservoir to both a fluidic connection and a hydrophobic filter, through which bubbles can exit. Second, it uses spring-loaded screws to evenly distribute the pressure to the O-rings that surround each reservoir, making it possible for the manifold to be used at pressures as high as 50 psi.

Prior to the experiment, the manifold is thoroughly rinsed with DI water and blown dry with nitrogen. The manifold is then heated overnight at 40°C to drive
off any remaining moisture. The O-rings are then placed around the reservoirs, the sealed chip is mounted, and the top plate is screwed onto the manifold, pressing the chip onto the O-rings. As shown in Figure 2.7, hydrophobic filters are attached to the white luer fittings extending from the manifold, and barbed-luer fittings are used to connect peroxide-cured silicone tubing to the remaining white luer fittings extending from the manifold.

Figure 2.8: A dual-input syringe pump is used to drive the sample and buffer inputs of the bubble-free, high-pressure manifold. T-valves are used to release bubbles in the tubing connecting the syringes to the manifold.

A dual-input syringe pump is used to drive the sample and buffer inputs of the bubble-free, high-pressure manifold (Figure 2.8). T-valves are used to release bubbles that may have formed in the tubing connecting the syringes to the manifold and to
enable changing of the syringes driving the inputs to the manifold. In order to ensure complete wetting of the tubing, manifold reservoirs, and DLD array chip, wetting is performed using a single syringe containing 2% Pluronic F108 surfactant in DI water. 60 mL syringes containing the sample and buffer are loaded following wetting, making use of the T-valves to avoid the introduction of bubbles into the reservoirs in the manifold. A detailed list of the parts required to assemble the setup shown in Figure 2.8 can be found in Appendix C.

2.5 Imaging Setup and Fluorescent Labelling of Cells

In this section, we describe the setup we use to image cells in the DLD array and the procedures we use to fluorescently label the cells for imaging.

2.5.1 Imaging Setup

A green fluorescent protein (GFP) filter set is used with a blue LED source (470 nm) to image appropriately labelled particles or cells. Blue light from the LED source is reflected off the dichroic mirror onto the DLD array chip. The emitted fluorescence from the cells or particles in the DLD array then passes through the dichroic mirror and then through a lens to the camera (Figure 2.9). The filter cube, lens, and camera are mounted on a xyz-stage so that different regions of the chip can be focused on.

In order to image the behavior of particles or cells in the DLD array, we use GFP-fluorescent polymer microspheres in place of cells or GFP-fluorescently labelled cells. Fluorescent beads above the critical size of the DLD array being displaced from the outer sides of a DLD array toward the center in parallel DLD arrays are shown in Figure 2.10. The image was taken using the imaging setup described above and shown in Figure 2.9.
2.5.2 Fluorescent Beads

Fluoresbrite YG microspheres (Polysciences, Inc.) with sizes of approximately 3 µm, 6 µm, and 10 µm were used to verify the critical sizes of the DLD arrays used in this thesis.

2.5.3 Leukocytes

Leukocytes are labelled with the green fluorescent nucleic acid stain, SYTO13. SYTO13 is added at concentrations of 6-10 µL per mL of whole blood. An incubation period of 45 minutes is observed after labelling before the experiment is run. The label becomes brighter with increasing incubation time up to at least 90 minutes, so regulation of the incubation time is important for experiments involving
Figure 2.10: Image of 10 µm green fluorescent beads in 1.5 mm wide parallel DLD arrays with 18 µm circular posts, 18 µm gaps, and 1/42 tilt (critical size ~4 µm). Adjacent DLD arrays are separated by horizontal blue lines in the image. Flow is from left to right, and the 10 µm beads, which are above the critical size of the DLD array, are directed from the outer sides of each DLD array toward the center.

measurement of the intensity of the fluorescence. No bleaching of SYTO13 from the blue LED in the imaging setup has been observed for time periods up to 90 minutes.

2.5.4 Cancer Cells

PC3 cancer cells that have been transfected with a green fluorescent protein (GFP) vector are used to enable imaging. The PC3 cancer cells were obtained from the lab of Dr. Ken Pienta at Johns Hopkins University School of Medicine.
2.6 Standard Blood Preparation and Disposal Protocols

Blood used in the experiments in this thesis is shipped overnight from Interstate Blood Bank (Memphis, TN), and experiments are performed within 36 hours of the time that the blood is drawn. All blood handling, including loading of the blood into the syringes, is performed in a Biosafety Level 2 (BSL-2) hood. Blood is diluted in a buffer containing anti-coagulants to prevent clot formation as discussed in Chapter 3 and filtered through a Steriflip 20 μm nylon net filter prior to loading into a syringe to remove any clots that may have formed during transit.

Disposal involves adding bleach to the blood to be disposed such that the final concentration is 10% bleach and disposing down the drain after 20 minutes have passed since the addition of the bleach. Syringes, tubing, hydrophobic filters, and barbed-luer fittings are disposed in a standard bio-hazard box. The manifold is disassembled and cleaned with ethanol. A 1% Contrad 70 solution can be used after this to remove any cells or other debris that remains on the manifold, and the manifold is soaked in a 1% Contrad 70 solution until next use.

Detailed information about ordering blood, preparing blood for experiments, and counting of cells in blood can be found in Appendix D.
Chapter 3

Inhibition of Clot Formation in DLD Arrays

3.1 Introduction

Previous work has focused on using DLD arrays to selectively capture rare cells of biological interest. D.W. Inglis et al. demonstrated that DLD arrays can be used to separate malignant lymphocytes from healthy lymphocytes [36]. L.R. Huang et al. have used DLD arrays to capture nucleated red blood cells from the peripheral blood of pregnant women for applications in prenatal diagnostics [32]. S.H. Holm et al. have used DLD arrays to separate parasites from human blood [26]. B. Zhang et al. have used DLD arrays to separate cardiomyocytes from blood [91]. Typical volumes of blood processed for such applications have been limited to 100 μL per DLD array.

While the capture efficiencies achievable with DLD arrays are sufficiently high to be useful in rare cell capture (>85%), capturing biologically useful quantities of rare cells requires processing of large volumes of blood. Recently, K. Loutherback et al. operated DLD arrays at flow rates as high as 10 mL/min, removing one key barrier to processing large volumes of blood [50]. However, even at this high flow rate, the
volume of blood processed was limited to less than 200 $\mu$L per DLD array due to clogging in the array. In this chapter, we demonstrate that this clogging process is due to the formation of blood clots and identify and inhibit the underlying physical and biological mechanisms driving this process.

Clot formation in DLD arrays imposes three significant limitations on device performance. First, the clot increases the fluidic resistance of the array, limiting the flow rate for a given pressure. Second, the clot formation can alter the flow pattern in a way that affects the critical size or simply displace cells below the critical size, making it appear that these cells behave similar to cells above the critical size and thus decreasing the enrichment. Third, the clot formation captures target cells, thus decreasing the yield of a separation process.

The limitations imposed by clot formation in the DLD array have been addressed in recent work. S. Zheng et al. showed that clogging occurred where the cells entered the array and explore the effects of dilution and age of the blood on clogging [93]. S.H. Holm et al. reported no clot formation with coagulation of blood being prevented by EDTA at a concentration of 6 mM. However, the volumes of blood being processed were still very small ($\sim$10 $\mu$L), the dilution was high (20x), and the flow rate was low ($\sim$3 $\mu$L/min) [26]. D.W. Inglis et al. mentioned the regular observation of blockages arising in the array from large clot-like structures in the blood despite the removal of such clot-like structures via pre-filtration before blood is loaded onto the array [37].

The effect of surface properties on clot formation in silicon microfluidic devices has been studied extensively [81]. Clot formation in DLD arrays has been observed in both silicon devices and PDMS devices, suggesting that activation of the contact activation coagulation pathway due to contact with silicon dioxide (glass) is not the dominant mechanism driving clot formation in DLD arrays [32, 93, 37]. In both silicon and PDMS DLD arrays, bovine serum albumin (BSA) has been used at a concentration of 1% (1g per 100 mL) to reduce non-specific adhesion [36, 32, 26, 91, 50, 37].
use BSA in this work at this concentration as a control in order to allow comparison with previous work.

Our experiments are conducted using DLD arrays reactively-etched in silicon to a depth of 160 µm by an anisotropic deep-etching process. The arrays are lidded with a pressure-sealing tape consisting of a thin silicone layer with a polyolefin backing. The lidded device is mounted in an acrylic manifold to which fluidic connections are made via O-rings. A dual-syringe pump is used to drive buffer and sample (diluted blood) through the chip. Flow velocities in the array are reported as the syringe volume flow rate divided by the cross section formed by the depth of the channel (160 µm) and the period of the array (post width + gap width = 100 µm) times the number of gaps through which the fluid is flowing (14 for a typical array) for a total effective width of 1.4 mm.

Whole adult human blood from healthy donors was purchased from Interstate Blood Bank (Memphis, TN) for use in our experiments. Blood samples were shipped overnight in acid citrate dextrose anti-coagulated tubes at a temperature of 4°C, and experiments were performed within 36 hours of the draw time. Blood was obtained, processed, and disposed of in accordance with policies and procedures set by the Princeton University Institutional Biosafety Committee (IBC) as described in Section 2.6 in the previous chapter.

3.2 Origins of Clot Formation

In order to capture rare cells in a product free from non-target cells present in blood, we use a three-input mirrored DLD array (Figure 3.1). Blood enters the chip through two through-holes (not shown) that each connect to four injector channels (visible via fluorescently tagged leukocytes) that lead to the top and bottom sides of the array. Buffer enters the chip through a through-hole (visible due to light from back of chip)
Figure 3.1: Fluorescent image of stained leukocytes in whole blood entering the DLD micro-post array through injector channels. Leukocytes are stained with SYTO13 at a concentration of 10 µL per mL of whole blood. Blood is diluted 1:3 in a running buffer composed of PBS, 1% BSA (1% = 1g per 100 mL), and 1 mM EDTA. The DLD array is a mirrored design in which sample enters at the outer sides of the array through four 40-µm-wide injector channels each on top and bottom, and cells larger than the critical size are displaced toward the center of the array, in which buffer alone is input through four 40-µm-wide injector channels. The array parameters are 60 µm right isosceles triangles, 40 µm gaps, and 1/50 tilt angle, yielding a critical size of approximately 6 µm. The array is 40 mm long, and each half of the mirrored array is 0.7 mm wide. The diluted blood input flow rate (total over both the top and bottom halves of the mirrored array) is 0.10 mL/min and the buffer input flow rate is 0.16 mL/min, giving a flow velocity of 1.8 cm/s. This image was taken with an exposure time of 2.5 seconds at t=30 minutes, after 0.7 mL equivalent volume of undiluted whole blood (2.8 mL of diluted blood in running buffer) had flowed through the array.

that leads to four injector channels (not visible, but which connects the through-hole to the beginning of the array) that lead to the middle of the array. There is a large amount of stuck fluorescent leukocytes at the beginning of the array. Experimentally, this is indicative of a clogging phenomenon that will cause an increase in the pressure for a given flow rate.

From the image in Figure 3.1 alone, we do not know if capture of the leukocytes at this location is caused by their direct adhesion to the posts or if they are caught due to clogging caused by a different mechanism. Further, note that the clogging does not occur in the injector channels and only happens at the beginning of the array. Since the injector channels are the same width as the gap in the micro-post array,
this is significant because it demonstrates that clogging happens dynamically at the beginning of the array and is not due to large clumps that have formed off-chip.

Blood contains three main components: leukocytes, erythrocytes, and a platelet-rich plasma. By experimentally removing or adding these different blood components before the sample enters the chip, we determine the relative contributions of the different components of blood to clogging. Leukocytes, because of their role in wound healing, have some adhesive properties, so we first isolate leukocytes via erythrocyte lysis buffer and centrifuge and re-suspend in a volume of running buffer equal to four times the original volume of blood from which the leukocytes were obtained.
Using the isolated leukocytes as the input to the chip, we see the leukocytes alone do not result in significant clogging (Figure 3.2). Note that the x-axis in Figure 3.2 represents the corresponding volume of the original blood from which the leukocytes were obtained, not the input volume to the chip which was four times greater. We use such a metric throughout the paper so one can directly relate results to the clinically relevant amount of blood processed. A very approximate but experimentally useful number for comparison is that the pressure begins to increase for a brightness of stuck leukocytes corresponding to a value of 1 on the y-axis in Figure 2. While the pressure increase alone might not be a problem, we note that when it occurs, the fluid flow in parts of the array is no longer parallel to the walls due to local clogging, which detrimentally affects the bumping mechanism.

Since erythrocytes do not have biologically significant adhesive properties, we then obtain a combination of erythrocytes and leukocytes via centrifuge by removing platelet-rich plasma via pipette and then re-suspending the erythrocytes and leukocytes in a volume of running buffer equal to four times the original volume of blood from which the leukocytes and erythrocytes were obtained. Similarly, we see that the combination of erythrocytes and leukocytes do not contribute significantly to clogging. However, when we run a combination of leukocytes and platelet-rich plasma diluted 4x in running buffer relative to the original volume of blood from which these were obtained (without erythrocytes), we see a substantial increase in clogging. Following this, we run a combination of erythrocytes, leukocytes, and platelet-rich plasma diluted 4x in running buffer relative to the original volume of blood from which these were obtained, and we observe a slight further increase in clogging, which is consistent with platelets being able to adhere to not just other platelets and leukocytes but also erythrocytes. These results suggest a conventional clotting process driven by platelets is the primary cause of clogging in the DLD array.
The surface of our chips is a thin coating of silicon dioxide from the oxygen plasma treatment and from reaction with air. Since it is well-known that contact with hydrophilic silicon dioxide causes clot formation through contact-activation coagulation [2], we examine two methods of preventing this: direct inhibition of the contact activation coagulation pathway and surface coatings. We used corn trypsin inhibitor to break the coagulation cascade due to contact activation by the surface by preventing the conversion of coagulation factor XII into XIIa [2]. Our results show that a 3.5 μM concentration of corn trypsin inhibitor results in a 40% decrease in clot formation, but not enough to enable 10X higher throughputs. We also examined the effect of a fluorocarbon coating to reduce the hydrophilic nature of the surface. The coating increased angle of 112° compared to a contact angle of 4° for our control surface. However, we found that this results in a two-fold increase in clot formation. This is consistent with prior work that showed with a wettability gradient on polyethylene that platelets under shear conditions adhere more strongly to hydrophobic surfaces compared to hydrophilic surfaces [76]. In order to further examine the contribution of protein adsorption on the silicon dioxide surface to clot formation, we use Pluronic F108, which is a block copolymer consisting of ethylene oxide and propylene oxide, in the running buffer at a concentration of 2 mg/mL instead of BSA [46]. Pluronic F108 reduces clot formation by less than 10% compared with BSA, which is consistent with protein adsorption onto the silicon surface not playing a major role in clot formation. The combination of these three results lead us to conclude that it is not surface effects but rather biological mechanisms that are involved in clot formation in DLD arrays.

3.3 Clot Formation Mechanisms

Clot formation is a very extensive field, and the processes driving clot formation in vivo are well understood. However, clot formation in our devices occurs in a shear-
dependent manner in a silicon device environment at shear rates higher than typical physiological levels (>1500 s$^{-1}$) and in a relatively short time (<1 s in the array). Thus, while the conventional understanding of clot formation in vivo may not be directly applicable, we use it as a framework to determine the mechanisms driving clot formation in DLD arrays.

Clot formation is driven by two complementary, mutually dependent processes shown schematically in Figure 3.3: platelet activation and coagulation. During platelet activation, the contents of stored granules are released into the blood plasma, triggering a clotting cascade, and the platelets change shape from spherical to stellate. Platelet activation can be triggered by a variety of factors, with thrombin-induced
platelet activation and mechanical stress-induced platelet activation being the two most relevant to DLD arrays. Coagulation involves the conversion of the soluble protein fibrinogen into insoluble strands of fibrin. In the coagulation cascade, pro-thrombin is converted into thrombin, which catalyzes the conversion of fibrinogen into fibrin in the presence of calcium to form a clot [24].

From this diagram, we notice two mutual positive feedback interactions between platelet activation and coagulation. First, thrombin, which is a product of the coagulation cascade, induces platelet activation by activating protease-activated receptors on the cell membrane of platelets [60]. Second, platelet activation causes the release of calcium ions into the blood plasma, which promotes coagulation. Therefore, both inhibiting the action of thrombin and chelating calcium ions should result in a significant reduction in clot formation due to the breaking of this feedback loop [24].

3.4 Reduction of Clogging

In this section, we examine the effectiveness of biochemical and physical methods for reducing clot formation in DLD arrays.

3.4.1 Chelation of Calcium and Inhibition of Thrombin

In order to inhibit the effect of calcium ions in the blood plasma, either ethylenediaminetetraacetic acid (EDTA) or acid citrate dextrose (ACD) can be used. While both work by chelating calcium, EDTA binds calcium more strongly and irreversibly than ACD. Recent studies have also found EDTA to be more effective than ACD at inhibiting the activity of the glycoprotein IIb/IIIa receptor, which is a key calcium-dependent integrin involved in platelet aggregation [11]. For these reasons, our work focuses primarily on EDTA. Our results (Figure 3.4) show that increasing the concentration of EDTA from 1 mM to 5 mM results in a 10-fold reduction in clot formation.
Figure 3.4: Effect of the calcium-chelating anticoagulant EDTA and the direct thrombin inhibitor PPACK on clot formation in the DLD array at flow velocities of 1.8 cm/s, 4.5 cm/s, and 20.3 cm/s. Clot formation is measured as the fluorescent brightness of stuck leukocytes as a function of the equivalent volume of undiluted whole blood that has passed through a single DLD array. Leukocytes are stained with SYTO13 at a concentration of 6 µL per mL of whole blood. Blood is diluted 1:3 in a running buffer composed of PBS, 1% BSA (1% = 1g per 100 mL), and either 1 mM or 5 mM EDTA. PPACK is then added so that the final concentration in the diluted blood is 40 µM. DLD array parameters are the same as in Figure 3.1.

as measured by the brightness of stuck leukocytes for up to ~0.7 mL equivalent volume of undiluted whole blood processed over a time of 30 minutes. While this is a significant reduction in clot formation, an even larger reduction is necessary to prevent an increase in pressure due to clot formation since pressure scales nonlinearly with the shrinkage of the gap caused by clot formation.

In order to inhibit thrombin-induced platelet activation, we use the direct thrombin inhibitor D-Phenylalanyl-L-prolyl-L-arginine chloromethyl ketone (PPACK). Compared to heparin, PPACK is more effective since it is able to act against both free thrombin and clot-bound thrombin, while heparin can only act against free thrombin [84]. We do not examine the effect of the direct thrombin inhibitor hirudin as previous studies have found PPACK to be a similarly effective anticoagulant.
for blood when used at a concentration of at least 40 \( \mu M \). Adding PPACK at a concentration of 40 \( \mu M \) on top of the 5 mM EDTA results in a further 5-fold reduction in clot formation (Figure 3.4). At a flow rate of 1.8 cm/s, after 30 minutes, during which time 0.7 mL of undiluted blood has been processed, only a barely discernible increase in stuck leukocytes due to clot formation is observed.

![Figure 3.5: Effect of the calcium-chelating anticoagulant EDTA and the direct thrombin inhibitor PPACK on clot formation in a DLD array with circular posts at flow velocities of 0.4 cm/s and 0.9 cm/s. Except for DLD array parameters of 18 \( \mu m \) circular posts, 18 \( \mu m \) gaps, and 1/42 tilt angle, experimental conditions are as in Figure 3.4.](image)

Figure 3.5: Effect of the calcium-chelating anticoagulant EDTA and the direct thrombin inhibitor PPACK on clot formation in a DLD array with circular posts at flow velocities of 0.4 cm/s and 0.9 cm/s. Except for DLD array parameters of 18 \( \mu m \) circular posts, 18 \( \mu m \) gaps, and 1/42 tilt angle, experimental conditions are as in Figure 3.4.

We also examined the effect of EDTA and PPACK on reducing clogging in arrays with circular posts. The results (Figure 3.5) were qualitatively similar. As with triangular posts, we observe a significant decrease in clot formation with the combination of 5 mM EDTA and 40 \( \mu M \) PPACK compared to the control of 1 mM EDTA.

In order to keep processing times under one hour even when processing large volumes of blood, we next examine the effect of flow rate on clot formation under conditions of optimal calcium chelation and thrombin inhibition described here.
3.4.2 Clot Formation, High Flow Velocity, and Processing Large Volumes of Blood

In addition to keeping processing times short even for large volumes of blood, there are two further reasons for increasing the flow velocity. First, increasing the flow velocity reduces the time in which platelets can form an aggregate within the array and reduces the time in which leukocytes can adhere to the micro-posts. Second, the higher shear stress from the higher flow velocity has been shown to cause disaggregation of platelet aggregates that have already formed [31]. On the other hand, a higher velocity gives a higher shear, which could cause more shear-induced platelet activation and clogging. Thus we performed experiments increasing the flow velocity first by a factor of 2.5 and then by a factor of 11.3 from the velocity of the previous section for triangular posts.

When we increase the flow velocity by a factor of 2.5 to 4.5 cm/s to achieve a higher processed volume in 30 minutes, we observe an additional two-fold reduction in clot formation over the first 0.7 mL and were able to successfully process 2.2 mL equivalent volume of undiluted whole blood over 35 minutes. We then increase the flow velocity to 20.3 cm/s to measure the extent of clot formation as 14 mL of blood flows through a single DLD array in 45 minutes. We again observe a similarly slow linear increase in the fluorescent brightness of stuck leukocytes over the entire 14 mL of blood (Figure 3.4).

At all three flow velocities, over the 30-45 minute experimental time, the stuck leukocytes do not appear primarily near the input end of the chip, but are rather uniformly distributed throughout the array. This leads us to conclude that no platelet-driven plug formation is observed with 5 mM EDTA and 40 µM PPACK, and that the slow linear increase in fluorescent brightness of stuck leukocytes is due to isolated leukocyte adhesion to the posts. Furthermore, the average shear rate at the flow velocity of 20.3 cm/s is 25,000 s\(^{-1}\), which is more than 10 times typical physiological
shear rates of 50 s$^{-1}$ to 2000 s$^{-1}$ \cite{20} and well above the 10,000 s$^{-1}$ at which shear-induced platelet activation has been reported \cite{27}. Even under these conditions, the combination of 5 mM EDTA and 40 $\mu$M PPACK prevents platelet plug formation in DLD arrays. We note that the high shear may also have a positive benefit of detaching adherent cells from the posts \cite{53}. High flow rates were also tested with circular posts (Figure 3.5), and again large volumes could be processed with minimal clogging.

### 3.4.3 Effects of Mechanical Stress and Gap Scaling on Clot Formation

We seek the source which initiates the clogging process in the array. Mechanical stress, particularly in the form of shear stress in a microfluidic channel, is a well-known cause of platelet activation, which could lead to clot formation \cite{69, 27, 25}. From Figure 3.1 and similar experiments, we do not observe any clogging in the injector channels leading to the array. This implies that shear at sidewall alone is not a major source of clogging. Figure 3.6 shows a 2-D simulation of the flow speeds in the injector channels and DLD array with triangular posts. For 150 $\mu$m depth channels, the simulation condition represents a flow of 110 $\mu$L/min per injector channel, which is comparable to an input flow rate of 1.30 mL/min for the chip of Figure 3.1. Shear stresses in the injector channels have a peak of 16,000 s$^{-1}$, which is comparable to the peak shear stress in the gap in the array. However, experimentally, even at shear rates exceeding this, only negligible clogging is observed in the injector channels.

As seen in Figure 3.1 the clogging begins at the start of the DLD array. From the simulations, the peak shear stress in the injector channels does not differ significantly from the peak shear stress in the array, but the shear stress increases and decreases rapidly vs. time as the fluid accelerates and decelerates in the array. It has previously been shown that for a single constriction, similar to a single gap in the array, in a
blood vessel, a clot expands from a small aggregate of platelets through adhesion of platelets in the low-shear zones at the downstream face of the clot [59]. While no such low-shear zones exist in the injector channels, the acceleration and deceleration of the fluid in the gaps provide a number of such low-shear zones for a clot to form.

Shear-induced platelet activation requires platelet membrane glycoproteins Ib and IIb-IIIa as well as the presence of von Willebrand factor [63]. We measure the effect of the glycoprotein IIb-IIIa inhibitor tirofiban at a concentration of 100 ng/mL of blood to test the contribution of shear-induced platelet aggregation to clot formation [73]. Our results (with 1 mM EDTA) show that tirofiban at this concentration reduces clot
formation by 40% at a flow velocity of 1.8 cm/s for the first 0.7 mL of blood through the array. This provides further support that mechanical shear stress is an important factor in initiating the clogging process in the array.

We examined the effect of scaling the gap size by comparing the clogging in two triangular arrays — one with post and gap of 40 μm and 27 μm, respectively, and the second with post and gap of 60 μm and 40 μm, respectively, with only 1 mM EDTA added to the input mixture (Figure 3.7). A qualitatively different clogging pattern was observed as the gap was reduced; there was more clogging immediately after the beginning of the array (much more intense fluorescence), and less farther downstream. This qualitative difference was consistently observed over different flow speeds. In order for a larger gap to become clogged by a clot, an aggregate of platelets at least as large as the gap needs to form. Thus, for a given concentration of platelets, it takes longer for such an aggregate to form since it would require a greater number of platelet-platelet interactions. Furthermore, if the inlets are held at a fixed pressure, the flow velocity in the DLD array with the smaller gap will be slower than in the DLD
array with the larger gap, leading to more time for clots to form. Rapid clogging near the array inlet would prevent the transport of leukocytes (and platelets and other clotting agents) downstream to the farther end of the array. The combination of these effects explains the more rapid clogging right at the beginning of the array with smaller gaps.

3.4.4 Effects of Acid Citrate Dextrose and Heparin on Clot Formation

![Graph showing effects of anticoagulants on clot formation.](image)

Figure 3.8: Effect of the calcium-chelating anticoagulants EDTA and ACD and the indirect thrombin inhibitor heparin, on clot formation in the DLD array at a flow velocity of 1.8 cm/s. Clot formation is measured as the fluorescent brightness of stuck leukocytes as a function of the equivalent volume of undiluted whole blood that has passed through a single DLD array. Leukocytes are stained with SYTO13 at a concentration of 6 µL per mL of whole blood. Blood is diluted 1:3 in a running buffer composed of PBS, 1% BSA (1% = 1g per 100 mL), and either 1 mM EDTA, 5 mM EDTA, 1:9 diluted ACD, or 40 units/mL heparin. DLD array parameters are the same as in Figure 3.1.

We examine the effect of two other commonly used anticoagulants, acid citrate dextrose (ACD) and heparin, on clot formation in DLD arrays. Like EDTA, ACD also inhibits clot formation by chelating free calcium ions, although less strongly and irre-
versibly than EDTA. Similar to PPACK, heparin inhibits clot formation through inhibi-
tion of thrombin, although the inhibitory mechanism of heparin is indirect through the
activation of the enzyme inhibitor antithrombin III rather than directly inhibiting the active site of thrombin as PPACK does.

ACD binds calcium less strongly and irreversibly than EDTA. We use the standard
ACD solution A, which is composed of 22.0 g/L C3434 (Citric Acid, trisodium salt, 
dihydrate); 7.3 g/L C0759 (Citric Acid, anhydrous); and 24.5 g/L G7528 (D-(-)-Glucose), added to the buffer at the recommended dilution ratio of 1:9. The difference in clot formation as observed from the fluorescent brightness of stuck leukocytes between 1:9 ACD and 5 mM EDTA is very small (Figure 3.8), further demonstrating the importance of calcium ions in the clot formation process in DLD arrays.

Heparin activates the enzyme inhibitor antithrombin III, which inactivates throm-
bin and other proteases involved in clot formation. We add heparin sodium salt to
the buffer such that the final concentration of heparin is 40 units/mL. There is sig-
nificantly more clot formation as observed from the fluorescent brightness of stuck leukocytes with heparin than with either 5 mM EDTA or 1:9 ACD (Figure 3.8). This is consistent with previous work by J.I. Weitz et al. showing that clot-bound throm-
bin (thrombin bound to fibrin) is not inhibited by heparin but is inhibited by PPACK
and hirugen, which is another synthetic direct thrombin inhibitor [85].

3.4.5 Effects of Anti-Platelet Drugs on Clot Formation

Anti-platelet drugs are used to inhibit clot formation by preventing platelet aggrega-
tion. Tirofiban, which we discuss in Section 3.4.3 above, is one such drug that works by inhibiting platelet membrane glycoproteins IIb-IIIa, which are involved in shear-induced platelet aggregation. Recently, combinations of anti-platelet drugs have been used to improve effectiveness of platelet aggregation inhibition through synergistic ef-
ficts between different anti-platelet drugs.
Figure 3.9: Effect of anti-platelet drug cocktails on clot formation in the DLD array at a flow velocity of 1.8 cm/s. Clot formation is measured as the fluorescent brightness of stuck leukocytes as a function of the equivalent volume of undiluted whole blood that has passed through a single DLD array. Leukocytes are stained with SYTO13 at a concentration of 6 µL per mL of whole blood. Blood is diluted 1:3 in a running buffer composed of PBS, 1% BSA (1% = 1g per 100 mL), and 1 mM EDTA. In addition, Cocktail 1 contains 100 µM aspirin, 10 µM clopidogrel, and 100 µM dipyridamole, and Cocktail 2 contains the same as Cocktail 1 plus 30 µM cilostazol. DLD array parameters are the same as in Figure 3.1.

One particularly powerful combination of anti-platelet drugs that has been examined recently is the combination of aspirin, clopidogrel, and dipyridamole [28]. We add these drugs to the diluted blood sample such that the final concentrations are 100 µM aspirin, 10 µM clopidogrel, and 100 µM dipyridamole. The decrease in clot formation is a modest 15% over the first 0.7 mL of blood through the DLD array (Cocktail 1, Figure 3.9). The inhibition of shear-induced platelet aggregation by another anti-platelet drug, cilostazol, has been shown to be enhanced by the presence of dipyridamole [29]. We run the experiment with the four anti-platelet drugs added to the diluted blood sample such that the final concentrations are 100 µM aspirin, 10 µM clopidogrel, 100 µM dipyridamole, and 30 µM cilostazol. The decrease in clot...
formation is 75% over the first 0.7 mL of blood through the DLD array (Cocktail 2, Figure 3.9), which is a significant improvement from the addition of 30 µM cilostazol.

Even the powerful anti-platelet drug cocktails we examine here are significantly less effective in inhibiting clot formation in DLD arrays than the combination of EDTA and PPACK we describe in Section 3.4.1. Furthermore, the cost of inhibiting clot formation in DLD arrays with the anti-platelet drug cocktails we describe here is approximately ten times higher per unit volume of blood compared with the method using EDTA and PPACK we describe in Section 3.4.1. Thus, inhibition of clot formation in DLD arrays via calcium chelation (EDTA) and direct thrombin inhibition (PPACK) is both the most effective and the most cost effective.

3.4.6 Effects of Flow Rate and Blood Dilution on Clot Formation

While chemical and biological factors affect the affinity of platelets towards aggregation, the actual formation of a clot is a physical process affected by physical factors. Interactions between platelets are necessary in order for platelet aggregates to form, and the interaction frequency is determined by the concentration of platelets. A second factor that affects whether platelets form aggregates is the duration the platelets are in the chip. This is controlled by the flow velocity.

Under the control conditions of 1 mM EDTA in the buffer, we examine how these two factors affect clot formation in the DLD array (Figure 3.10). Assuming an interaction between two platelets is necessary for aggregation, increasing the dilution will reduce the interaction frequency by the square of the increase of the dilution. We observe that a 3.3-fold increase in the dilution and a corresponding 10-fold decrease in interaction frequency significantly slows the rate of clot formation for the same flow velocity.
Figure 3.10: Effect of flow rate and blood dilution on clot formation in DLD arrays. Clot formation is measured as fluorescent brightness of stuck leukocytes as a function of the equivalent volume of undiluted whole blood that has passed through a single DLD array. Leukocytes are stained with SYTO13 at a concentration of $10 \mu$L per mL of whole blood. Blood is diluted either 1:3 or 1:10 in a running buffer composed of PBS, 1% BSA (1% = 1g per 100 mL), and 1 mM EDTA. DLD array parameters are 60 $\mu$m right isosceles triangles, 40 $\mu$m gap, and 1/50 tilt angle, and the flow velocities used are either 1.8 cm/s or 18 cm/s.

Increasing the flow velocity by a factor of 10 reduces the time in which platelets can form an aggregate within the array by a factor of 10. The higher flow velocity also has the added benefit that higher shear stress has been shown to cause disaggregation of platelet aggregates that have already formed [31]. We observe that this 10-fold reduction in the time for aggregation formation significantly slows that rate of clot formation. Combining the 10-fold decrease in collision efficiency with the 10-fold decrease in aggregate formation time, we show that clot formation can be reduced by a factor of 10 as measured by the fluorescent brightness of stuck leukocytes.
3.5 DLD Array Performance for High Volume Separation at High Flow Rates

Table 3.1: DLD Array Performance for High Volume Separation at High Flow Rates for Different Array Parameters

<table>
<thead>
<tr>
<th>Post: 60µm, Gap: 40µm, Tilt: 1/50, Triangular Posts, Center Wall</th>
<th>PC3 Cancer Cells</th>
<th>Leukocytes</th>
<th>Erythrocytes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample: $3.1 \times 10^6$/mL</td>
<td>Sample: $1.7 \times 10^6$/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product: $8.2 \times 10^6$/mL</td>
<td>Product: $5.0 \times 10^6$/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample/Product Volume: 3.4</td>
<td>Sample/Product Volume: 3.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yield: 77%</td>
<td>Yield: 8.6%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample/Product Volume: 3.4</td>
<td>Sample/Product Volume: 3.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yield: 4.5%</td>
<td>Yield: 4.5%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Post: 64µm, Gap: 34µm, Tilt: 1/26, Circular Posts, No Center Wall</th>
<th>PC3 Cancer Cells</th>
<th>Leukocytes</th>
<th>Erythrocytes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample: $3.2 \times 10^6$/mL</td>
<td>Sample: $1.9 \times 10^6$/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Product: $8.0 \times 10^6$/mL</td>
<td>Product: $7.1 \times 10^6$/mL</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample/Product Volume: 2.9</td>
<td>Sample/Product Volume: 2.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yield: 86%</td>
<td>Yield: 1.3%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample/Product Volume: 2.9</td>
<td>Sample/Product Volume: 2.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yield: 0.03%</td>
<td>Yield: 0.03%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

We then tested the performance of three-input mirrored DLD arrays, as in Figure 3.1, to harvest prostate cancer cells (PC3) from blood using the conditions for minimum clogging from Section 3.4.1. The fractions of the total number of cancer cells, leukocytes, and erythrocytes, respectively that appear in the product compared to that processed through the input are measured to characterize the device performance. The fraction of cancer cells, the diameters of which are above the critical size of the array, is taken as a measure of the collection efficiency. The fraction of undesired erythrocytes (RBCs) appearing in the product is a measure of the depletion efficiency, which is another important parameter in characterizing three-input mirrored DLD arrays (a low yield is desired). The fraction of leukocytes (WBCs) appearing in the product is also measured so that the enrichment of cancer cells and purity of the product can be calculated. The target PC3 prostate cancer cells have an average diameter of 17 µm [52]. The results of our experiments are summarized in Table 3.1.

As a first experiment, the PC3 cells are collected along a central wall in the middle of the array with triangular posts, as in Figure 3.1 [34]. With array parameters of 60
\( \mu m \) triangular posts, 40 \( \mu m \) gaps, and 1/50 tilt angle, the critical size is approximately 6.1 \( \mu m \). The array length was 40.00 mm, the total width of the array region including the center wall was 1.44 mm, with 0.62 mm width of post array in both the top half and bottom half of the array. The product is collected from a region spanning the central 0.22 mm of the array including the center wall. GFP-labelled PC3 cells were spiked in 14 mL of whole blood diluted 1:3 in a running buffer composed of PBS, 1% BSA, 5 mM EDTA, and 40 \( \mu M \) PPACK at an approximate concentration of 3x10^4 per mL. After running through the array for 45 minutes at a flow velocity of 20.3 cm/s (processing an input volume of 56 mL), the collection efficiency of PC3 cells into the product outlet was 77%. However, erythrocyte contamination of the product was significant with nearly 4.5% of the input erythrocytes appearing in the product. At the high Reynolds number used in these flows (Re \( \sim \) 10), it has recently been reported that erythrocytes, which have a discoid shape, behave in a DLD array as if they have a much larger size than they do at low Reynolds number, an effect not observed for circular posts [70]. Thus more are collected as large cells than would otherwise occur.

We then created an array with 64 \( \mu m \) circular posts, 34 \( \mu m \) gaps, and 1/26 tilt angle, for which the critical size is approximately 10 \( \mu m \). The central collection wall in the mirrored array was also removed to present fewer surface for leukocyte sticking – the fluidic balance between the top and bottom regions kept the output buffer stream confined to this central region [34]. The array length was 38.22 mm, the total width of the array region including the central collection region was 2.24 mm, with 1.06 mm width of post array in both the top half and bottom half of the array. The product is collected from a region spanning the central 0.35 mm of the array including the collection region. The experiment is run under the same conditions as the first experiment except that the duration is now 38 minutes and the average flow velocity is 21.7 cm/s. The collection efficiency of PC3 cells is 86%, slightly higher than in
the triangle post chip, but the erythrocyte yield has dropped over 100X, from 4.5% to 0.03%, and now consists of small clumps of 3-5 erythrocytes. This surprising very large effect of the post shape on undesired erythrocyte collection will be one of the main topics of Chapter 4.

This represents the highest collection efficiency of cancer cells for a large volume of diluted whole blood (>7.5 mL) processed in a short time (<1 hour). By comparison, E. Sollier et al. used size-selective vortex technology to process 7.5 mL of blood in 20 minutes but achieved a maximum collection efficiency of less than 10% [74]. Using another inertial cell separation technique, slanted spiral microfluidics, M.E. Warkiani et al. processed 7.5 mL of blood in 8 minutes achieving a capture efficiency greater than 80%, but use of this technology requires lysis of red blood cells before processing [83].

### 3.6 Conclusions

Clogging in DLD arrays processing blood has limited their applicability for capture of rare cells to blood volumes well under 1 mL for each array. In this chapter, we have first shown that clogging of the array is caused by a conventional platelet-driven clot formation process. Next, we identified the presence of calcium ions and thrombin as drivers of platelet-driven clot formation process and disabled these two drivers using the calcium-chelating anticoagulant EDTA and the direct thrombin inhibitor PPACK.

Using these approaches, we have demonstrated that 14 mL of blood can be processed using a single DLD array in less than 45 minutes at flow velocities as high as 20 cm/s with no significant clot formation by the successful harvesting of PC3 prostate cancer cells from blood. Future work could include using a second DLD array in series to further concentrate the harvested cells [30]. Standard micro-fabrication techniques...
have been previously used to produce a chip containing 15 DLD arrays in parallel, which would require an area of 36 mm by 70 mm for the array presented in this paper [91, 40]. Combining a chip containing 15 of our DLD arrays in parallel with the advances presented in this chapter should allow for processing of well over 100 mL of blood for rare cell capture in less than one hour.

The work in this chapter was largely described in [18].
Chapter 4

Post Geometry Design for High-Throughput Harvesting of Nucleated Cells from Blood with Minimal Erythrocyte Contamination Using DLD Arrays

4.1 Introduction

Separation of rare cells from whole blood using DLD arrays requires processing large volumes of blood in short times. In the previous chapter, we demonstrated an approach to inhibit clot formation in DLD arrays, allowing for separation of PC3 cancer cells from 14 mL of blood in less than one hour using a single DLD array [18]. In this chapter, we examine how post geometry affects capture efficiency of target cells above the critical size of the array and rejection efficiency of cells below the critical size of the array, specifically at high flow rates corresponding to Re>1.
As the flow rate increases, the shear the cells experience passing through the gap between the micro-posts increases, resulting in compression of the cells against the micro-posts. This compression can be sufficient to reduce the size of these cells below the critical size of the array, causing these cells not to be displaced. Z. Liu et al. have shown that significantly greater compression of cancer cells occurs with circular posts than with asymmetric triangular posts, resulting in a 4-fold decrease in capture efficiency at mL/min flow rates for the same array critical size \([47]\). However, in the previous chapter, we also demonstrated that erythrocyte contamination of the product increases significantly at high flow rates with asymmetric triangular posts, which should indicate that the erythrocytes effectively acted larger at higher flow rates, not smaller. Further puzzling is that this last effect does not occur with circular posts.

The mechanism that causes this increase in erythrocyte contamination of the product with asymmetric triangular posts but not circular posts at high flow rates, and correspondingly higher (moderate) Re, has not been previously studied. As background, inertial effects in DLD arrays at moderate Re \((5<Re<40)\) have been studied, with an observed increase in separation efficiency with increasing Re due to a decrease in critical size resulting from asymmetry of the array \([54]\). It has also been suggested that the increasing separation efficiency results partly from shear-gradient lift forces arising from vortices that form behind the micro-posts at \(5<Re<40\) \([56]\). The behavior of erythrocytes in DLD arrays has also been examined, since the shape of erythrocytes is discoid and not spherical. J. Beech et al. showed that channel depth can be used to change the orientation of the erythrocytes such that the erythrocyte behaves as a spherical particle with size corresponding to either the long or short axis of the discoid \([4]\). More recently, post geometry was used to induce changes in the orientation (flipping) of the erythrocytes to achieve a similar effect in DLD arrays.
with deep channels, in which the erythrocytes would normally be expected to behave as spherical particles with size corresponding to the short axis of the discoid \[88\text{, }70\].

We begin in Section 4.2 with an overview of the fundamental mechanisms that govern particle (cell) behavior in microfluidic devices at high flow rates (moderate Re). In Section 4.3, we explain the experimental structures and modelling of streamlines. We then experimentally determine and fundamentally explain (i) (Section 4.4) how post geometry affects cell compression due to shear resulting in reduced yield of leukocytes and (ii) (Section 4.5) undesired erythrocyte displacement at high flow rates. Finally, in Section 4.6, we show that with optimal post shape, high yield (\(\sim 85\%\)) of leukocytes with efficient rejection of erythrocytes (\(>99.98\%\)) can be achieved at up to Re of 2.3, corresponding to processing of nearly 4mL of blood in less than 45 minutes in a single mirrored DLD array.

4.2 Fundamental Mechanisms at High Flow Rates

At these high flow rates, corresponding to Re of \(\sim 15\), we have found that the capture efficiencies of the large cells above the critical size of the array can degrade and the rejection of the small cells below the critical size of the array decreases. In this chapter, we examine the fundamentals behind this performance degradation and show how both can be affected by post shape. With optimal post shape, high yield of leukocytes (\(\sim 85\%\)) with efficient rejection of erythrocytes (\(>99.98\%\)) can be achieved up to Re of 2.3, corresponding to processing of nearly 4mL of blood in less than 45 minutes.
4.2.1 Changes in Fluid Flow at High Re

As Re increases, terms that could be neglected in the Navier-Stokes equation at low Re become significant. Assuming no external body forces, the full Navier-Stokes equation is:

\[ \rho \cdot \left( \frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \cdot \nabla)\mathbf{u} \right) = -\nabla P + \mu \nabla^2 \mathbf{u} \] (4.1)

where \( \rho \) is the density of the fluid, \( \mathbf{u} \) is the fluid velocity, \( P \) is the pressure, and \( \mu \) is the dynamic viscosity of the fluid. At low Re, the inertia terms, which are the two terms on the left-hand side of equation 4.1 are neglected. These terms are called "inertial" because they depend on changing the momentum (or the "inertia") of the particle. Note that these terms depend on the mass density times the velocity or the time derivative of the velocity. The first term is always neglected in analysis of DLD arrays, since we do not consider the time-evolution of the flow; we only consider steady-state. The second term results from considering the transport of the velocity field in the flow (material derivative of the velocity). As we increase Re, this term becomes increasingly significant, because it is proportional to the square of the velocity, leading to the effects we consider below.

The Reynolds number, Re, in the DLD array is defined as

\[ Re = \frac{\rho u G}{\mu} \] (4.2)

where \( \rho \) is the density of water at room temperature, \( u \) is the average velocity of the fluid in the gap, \( G \) is the width of the gap between the posts, and \( \mu \) is the dynamic viscosity of water at room temperature. A particle Reynolds number, \( Re_p \), has also been used to characterize hydrodynamic lift forces and can be calculated from the DLD array Re in equation 4.2 as:
\[ \text{Re}_p = \text{Re} \frac{a^2}{G^2} \]  

where \( a \) is the radius of the particle \[77\]. \( \text{Re}_p \) is less than 1 for all experiments and models in this chapter, and we state all results in terms of the DLD array \( \text{Re} \).

It has previously been shown using rigid, non-deformable particles (polyethylene beads) that the critical size decreases by about 10% for circular posts as \( \text{Re} \) increases from 1 to 30 \[56, 54\]. At high \( \text{Re} \), for particles that are denser than the fluid, such as erythrocytes and the polymer microsphere beads in our experiments, it has been shown previously that micro-post geometry becomes important due to the possibility of impaction of particles on the micro-posts as well as particles not following the streamlines due to rapid acceleration of the fluid around certain micro-post geometries \[23, 3, 6\].

We now examine inertial and shear effects in fluid flow and how these effects could affect the behavior of cells suspended in the fluid flowing through the DLD array at moderate \( \text{Re} \) (\( \text{Re} > 1 \)). The effects include inertial lift and Dean drag forces (Section 4.2.2), lift forces from hydrodynamic asymmetry (Section 4.2.3), centripetal acceleration effects (Section 4.2.4) and shear effects on cell shape (Section 4.2.5).

### 4.2.2 Inertial Lift and Dean’s Flow Forces

In a straight micro-channel, suspended particles experience two oppositely directed lift forces as \( \text{Re} \) increases (Figure 4.1(a)). The shear-gradient lift force is directed from the center of the channel toward the walls and is a result of the parabolic flow profile. The wall effect lift force is directed away from the channel wall and decreases with distance from the wall. Both of these lift forces are caused by the effect the particle itself has on the fluid flow distribution. In a straight micro-channel, a minimum flow rate has been determined for the particles to reach stable equilibrium positions as
Figure 4.1: (a) Two oppositely directed inertial forces in straight micro-channels arise at high Re: a shear-gradient lift force directed away from the center of the channel and a wall-effect lift force that decreases with distance from the wall. Based on a balance between these two oppositely directed forces, particles can reach stable equilibrium positions. (b) In curved micro-channels, force from Dean’s flow, which results from two counter-rotating vortices perpendicular to the flow direction, combines with the inertial lift forces to create stable and unstable equilibrium positions in the channel for particles suspended in the fluid. Image taken from [17].

As a result of these two oppositely directed lift forces in a given length along the flow direction [17].

In curved micro-channels, Dean’s flow is present in addition to these two lift forces. Dean’s flow is caused by two counter-rotating vortices in the direction perpendicular to the flow, which result from the faster moving fluid near the channel center causing the more stagnant fluid near the walls to re-circulate in order to conserve mass (Figure 4.1(b)). These vortices result in a drag force on particles suspended in the fluid. The minimum flow rate for particles to reach equilibrium positions in this case is the
same condition as for the straight microchannel with an additional condition on the curvature of the channel relative to the channel width [17].

Theory to characterize the effect of inertial lift forces, specifically the shear gradient and wall lift forces, in DLD arrays has been developed and characterized by Lubbersen et al. [56]. Under the theory developed by Lubbersen et al., the flow rates we examine here are less than 0.01 the minimum flow rate required for particles to achieve stable equilibrium positions in the gap due to these inertial lift forces [56]. This makes it significantly less likely in our experiments that the shear-gradient and wall effect lift forces are causing particles below the critical size of the array to cross one or more streamlines and behave as particles above the critical size of the array compared to in the experiments of Lubbersen et al. [56], which were conducted at more than 0.2 times the minimum flow rate required for particles to achieve stable equilibrium positions due to these inertial lift forces. Note that the theory developed by Lubbersen et al. [56] is based on the length of the post in the direction of the flow, considering the channel length as the length of a single gap.

4.2.3 Lift Effects from Asymmetry

Less well-known than inertial (shear-gradient and wall effect) lift forces is the non-inertial hydrodynamic lift force for deformable particles, such as erythrocytes. Even in the regime of low Re, the symmetry of the Stokes regime (Re<1) can be broken in a way that enhances suspended deformable particles (vesicles) to undergo cross-streamline migration. This can occur due to asymmetry of the walls or interactions between suspended particles [19]. Non-inertial hydrodynamic lift has been used to separate circulating tumor cells from erythrocytes [21]. Hydrodynamic lift forces arising from asymmetry of the walls in this way have not been studied at Re above 1.
Figure 4.2: (a) Non-inertial hydrodynamic lift induced separation, which relies on asymmetry of the walls to enhance particles to cross streamlines, is used to separate MV3 melanoma cells from erythrocytes at Re of 0.37. Image taken from [21]. (b) Pinched flow fractionation, which relies on hydrodynamic effects resulting from liquid with particles and liquid without particles being injected into a pinched region that then expands into a wider region, is used to separate polymer microspheres of different sizes. For the geometry shown, separation is dependent upon the relative flow rates of the liquid with particles and the liquid without particles. Image taken from [86].

Hydrodynamic asymmetry has also been used to separate particles based on size. In pinched flow fractionation, a fluid with particles and a fluid without particles are injected into inlets that lead to a pinched channel. Based on the relative flow rates of the two fluids, particles are focused along one sidewall in the pinched channel. When the pinched region broadens into wider channel, large particles experience a force from the expanding flow toward the center of the channel while smaller particles are directed toward the channel walls [86]. Geometric asymmetry has been shown to enhance this effect [80], and pinched flow fractionation has also been applied to separation of cancer cells from leukocytes [66].
4.2.4 Centripetal Acceleration and the “Slingshot” Effect

As a particle follows a straight streamline, it is pushed by a viscous drag force, $F_D$, parallel to its motion, so that it moves at a velocity, $v_p$, equal to that of the fluid, $v_f$.

$$F_D = 6\pi \mu v_p R_p$$ (4.4)

where $\mu$ is the dynamic viscosity of the fluid and $R_p$ is the radius of the particle.

Figure 4.3: Qualitative picture of a particle moving to an adjacent streamline when the fluid flow undergoes curvature.

When the streamline curves, if the particle were to follow the streamline, a centripetal force, $F_C$, would be required to cause the needed centripetal acceleration $a_C$:

$$F_C = (\rho_p - \rho_f) V_p a_c$$ (4.5)

where $\rho_p$ and $\rho_f$ are the particle and fluid mass densities, respectively, and $V_p$ is the volume of the particle, and

$$a_c = \frac{v_p^2}{R_c}$$ (4.6)
where $R_c$ is the radius of curvature.

Because of its inertia, the lack of an actual centripetal force will cause a particle denser than the fluid to move to an adjacent stream of greater radius, where the drag force will then cause the particle to accelerate along the new streamline (Figure 4.3) until the new streamline curves and the process repeats. Thus the particle will repeatedly be moving to outside streamlines when the fluid streamlines curve. We refer to this informally as the “slingshot” effect. Because the required centripetal force is proportional to the square of the velocity, this effect of moving particles across streamlines at curves is expected to increase as the flow rate increases.

![Figure 4.4: (a) Plot of streamlines in a DLD array with 60 \(\mu\)m circular posts, 40 \(\mu\)m gaps, and 1/20 tilt, and flow velocity corresponding to Re of 20. (b) Plot of streamlines in a DLD array with 60 \(\mu\)m asymmetric isosceles triangular posts, 40 \(\mu\)m gaps, and 1/20 tilt, and flow velocity corresponding to Re of 20. Note the sharp curvatures in the gap region with asymmetric triangular posts is circled in red in (b) compared to the curvatures with circular posts in (a).](image)

We expect this process to depend on the post shape because different post shapes result in different curvatures of the streamlines, especially in the gap. Note in Figure 4.4 that the curvatures of streamlines in the gap can be much sharper for posts with sharp points in the gap (Figure 4.4(b)) than those for posts with smooth edges, such as circular posts (Figure 4.4(a)). If the velocities in the two gaps were the
same, the required centripetal forces for particles to follow streamlines and not move
to adjacent streamlines would be much larger for the posts with sharp points, as in
Figure 4.4(b).

We now describe an approach to quantitatively estimate how much this effect
might be expected to move a particle laterally across streamlines. We assume the
effect is “small, so that the particle does approximately follow a streamline over short
distances, even with curves. At each point along its path, there is some small motion
perpendicular to the streamline at curves, because of the lack of a centripetal force.
This we can estimate by assuming the small motion perpendicular to the streamline
leads to a drag force on the particle in that direction, equal to the centripetal force
which would be needed to keep the particle on the streamline. We define $v_L$ as the
lateral velocity (perpendicular to the streamline). Starting with:

$$ F_C = F_D \quad (4.7) $$

$$(\rho_p - \rho_f)V_p a_c = 6\pi \mu v_L R_p \quad (4.8)$$

and solving for the lateral velocity, $v_L$, yields:

$$ v_L = \frac{(\rho_p - \rho_f)V_p a_c}{6\pi \mu R_p} \quad (4.9) $$

From the lateral velocity, we can find the distance perpendicular to the stream-
line that the particle has traveled due to the centripetal acceleration force, $x_L$, by
integrating over the streamline path approximately traced by the particle:

$$ x_L = \int v_L d\left(\frac{y}{v_y}\right) \quad (4.10) $$
where \( y \) is the distance in the average flow direction and \( v_y \) is the velocity in the average flow direction. These calculations are carried out numerically for erythrocytes for different post shapes at high Reynolds number in Section 4.5.3.

### 4.2.5 Effects on Cell Shape

Two effects become important as the flow rates, and correspondingly Re, at which DLD arrays are operated increases. First, the shear on the cells as the cells pass between the micro-posts increases, causing deformation of the cells that reduces the effective size of the cells [4]. This reduction of the effective size of the cells can be sufficient to change the behavior of the cells in the DLD array [68, 87]. Second, at high flow rates, certain post geometries displace erythrocytes, which are well below the critical sizes of the arrays in these cases [88, 70].

**Shear Stress on Spheroid Cells (Leukocytes and Cancer Cells)**

Deformable spheroid cells have been shown to assume an elongated shape at high shear rates (Figure 4.5), which leads to cells navigating through the DLD array as if the cells have smaller effective radii [4]. Thus, target cells to be harvested are not displaced into the product if the effective radii of these cells falls below the critical size of the array. Compared with cancer cells, leukocytes are deformed more for the same shear in a DLD array than cancer cells [41], and the deformability of leukocytes is a critical parameter in scaling of DLD arrays for high-throughput separation of leukocytes from large volumes of blood [37].

As the flow rate increases, the shear the cells experience passing between the micro-posts increases. Shear-induced cell deformation typically results from fluid bending around the up-stream face of the micro-post on the side in the direction of the tilt [15]. Z. Liu et al. showed that shear-induced deformation of cancer cells is significantly less with triangular posts compared with circular posts at high flow...
rates [47]. Recently, D. Holmes et al. demonstrated that leukocytes can be separated into subpopulations based on a combination of size and deformability in DLD arrays using circular posts [28]. M. Al-Fandi et al. developed a theoretical model for shear-induced compression of cells against the micro-posts in a DLD array and proposed diamond and airfoil shaped micro-posts to minimize this compression [1]. However, the effect of post geometry on shear-induced cell deformation has not been examined experimentally.

**High Flow Rate Effects on Discoid Cells (Erythrocytes)**

Hydrodynamic effects on the behavior of erythrocytes in DLD arrays are particularly interesting because erythrocytes are discoid in shape. J. Beech et al. have shown
that in deep channel DLD arrays (e.g. >20 µm deep), such as the ones we use in our experiments, the shear causes the erythrocytes to align such that the long axis is parallel to the flow, and the erythrocytes behave as spherical particles with a diameter equal to the smallest dimension [4]. This alignment occurs to minimize the cross-section of the erythrocyte that is exposed to viscous drag from the flow. Recently, K.K. Zeming et al. and S. Ranjan et al. have shown that post geometry can be used to induce flipping (90° rotation) of the aligned erythrocytes, causing the erythrocytes to behave as particles larger than the critical size of the array even in deep channel DLD arrays [88, 70]. Because the major advantage of DLD arrays over other microfluidic technologies is the ability to separate nucleated cells from whole blood without the background of erythrocytes, characterizing the behavior of erythrocytes in DLD arrays with respect to post geometry at high flow rates is very important.

With this as background, we now go on to describe experiments and modelling to understand the behavior of erythrocytes and leukocytes in DLD arrays with different post shapes at high Re. These results are quite unexpected, and we show with optimal post shape that we can harvest leukocytes efficiently without substantial erythrocyte contamination.

4.3 Array Structure, Post Shapes, Experimental Conditions, and Modelling Approach

In this section, we discuss the array structure, post shapes, experimental conditions, and modelling approaches that we use in this chapter.
4.3.1 DLD Arrays for Separation of Nucleated Cells from Whole Blood

Figure 4.6: Schematic of the DLD array design used for separation of nucleated cells, such as leukocytes or PC3 cancer cells, from whole blood. The DLD array is a mirrored design in which sample enters at the outer sides of the array, and cells larger than the critical size are displaced toward the center of the array, in which buffer alone is input. The buffer input spans 9/19 the total width of the array, and each of the sample inputs span 5/19 the total width of the array. The product is collected at the end of the array from a width spanning 3/19 of the array width, and each of the waste outlets span 8/19 the total width of the array.

In order to capture leukocytes or cancer cells in a product free from non-target cells present in blood, we use a three-input mirrored DLD array (Figure 4.6). Blood
sample enters the array through inlets connected to the right and left sides of the array, while buffer enters the array through inlets connected to the center of the array. Target cells larger than the critical size of the array are displaced toward the center of the array into the buffer stream and are then collected through the product outlet, which spans $3/19$ the total width of the array.

Figure 4.7: Schematic of mirrored array with circular posts for collection of leukocytes. The tilt on the left side of the array is to the right, and the tilt on the right side of the array is to the left. This displaces cells above the critical size of the array toward the center from each side. Instead of a wall in the middle, we use a fluidically balanced central collection region.

Mirrored DLD arrays have typically been designed to collect cells above the critical size of the array using a central wall [47, 50] that is fluidically balanced to maintain vertical flow even near the wall using the method developed by D. Inglis [34]. However,
adhesion of cells, particularly leukocytes, along this central wall is problematic in maintaining high capture efficiency for large volumes of blood. We instead use a fluidically balanced central collection region instead of a wall to capture displaced cells in the DLD arrays used in our experiments to avoid this problem (Figure 4.7).

Our experiments are conducted using DLD arrays reactively-etched in silicon to a depth of 160 $\mu$m by an anisotropic deep reaction ion etching (DRIE) process. The array is lidded with a 1-mm-thick Pyrex glass slide anodically bonded to the silicon chip into which the DLD array has been etched. The lidded device is mounted in an acrylic manifold to which fluidic connections are made via O-rings. A dual-syringe pump is used to drive buffer and sample (diluted blood) through the chip. Flow velocities in the array are reported as the syringe volumetric flow rate divided by the cross section formed by the depth of the channel (160 $\mu$m) and the gap between the posts (36 $\mu$m for cancer cell experiments, 17 $\mu$m for leukocyte experiments) times the number of gaps through which the fluid is flowing (23 for cancer cell experiments, 41 for leukocyte experiments) for a total effective width of 2.3 mm for cancer cell experiments and 1.5 mm for leukocyte experiments. This gives the average flow velocity in the gap, which is used along with the gap size as the characteristic length to calculate the Reynolds number.

Blood used in the experiments is obtained from Interstate Blood Bank (Memphis, TN), and experiments are conducted within 36 hours of the time the blood is drawn. Blood is diluted 1:3 in a buffer of PBS, 1% BSA (1%=1g BSA per 100 mL PBS), 5mM EDTA, and 40$\mu$M PPACK to inhibit clot formation [18]. The running buffer into which the leukocytes are collected is composed of PBS and 1% BSA.

4.3.2 Post Shapes and Array Parameters

The post shapes and critical sizes for the leukocyte yield and erythrocyte product contamination experiments and modeling are shown in Tables 4.1 and 4.2 below.
The critical sizes for circular posts are computed using the well-known experimental fit for critical size in circular post arrays with shallow tilt (<1/10) angles:

\[ D_c = 1.4G\epsilon^{0.48} \]  

(4.11)

where G is the gap size and \( \epsilon \) is the tilt angle of the array [15].

For arrays with asymmetric isosceles triangular posts with shallow tilt (<1/10) angles, based on theory by K. Loutherback et al. [49], we propose a similar fit for critical size:

\[ D_c = 1.0G\epsilon^{0.48} \]  

(4.12)

This fit agrees well with experimental results with polymer microsphere beads (3 \( \mu \)m, 6 \( \mu \)m, and 10 \( \mu \)m), leukocytes, and PC3 cancer cells for asymmetric isosceles triangular post arrays with tilts of 1/20, 1/26, 1/36, and 1/42.

The critical sizes for all post geometries are first computed at the stated Re by the method proposed by D. Inglis et al. [35] using the velocity profile generated by the 2D laminar flow module in COMSOL. Fundamentally, this method examines the width of the streamline next to the post, as shown in Figure 1.1. This is done with an array with posts that can fit within a 60 \( \mu \)m by 60 \( \mu \)m square, 40 \( \mu \) gaps, and 1/20 tilt. The critical sizes for symmetric post shapes (including quarter circle and semi-circle) are then scaled to the appropriate gap and tilt using the experimental fit for circular posts. The critical sizes for asymmetric post shapes are scaled to the appropriate gap and tilt using the experimental fit for asymmetric triangular posts.

**Leukocyte Yield Experiments**

The post shapes and critical sizes used for the leukocyte yield experiments are shown in Table 4.1.
Table 4.1: Post shapes used in the leukocyte yield experiments. Experiments are conducted at a flow velocity corresponding to Re of 2.3, with the average velocity in the gap between the posts being 13.5 cm/s. Array parameters are posts that can fit within a 19 \( \mu \text{m} \) by 19 \( \mu \text{m} \) square, 17 \( \mu \text{m} \) gaps, and 1/42 tilt, except for the asymmetric isosceles triangle, which has posts with 34 \( \mu \text{m} \) height and width, 22.5 \( \mu \text{m} \) gaps, and 1/42 tilt.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>Critical Size (( \mu \text{m} )) at Re = 2.3</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \downarrow )</td>
<td>( \uparrow )</td>
<td>( \downarrow )</td>
<td>Asymmetric Triangle</td>
<td>3.8</td>
</tr>
<tr>
<td>( \downarrow )</td>
<td>( \Uparrow )</td>
<td>( \downarrow )</td>
<td>Circle</td>
<td>4.0</td>
</tr>
<tr>
<td>( \downarrow )</td>
<td>( \Uparrow )</td>
<td>( \downarrow )</td>
<td>Quarter Circle</td>
<td>4.0</td>
</tr>
<tr>
<td>( \downarrow )</td>
<td>( \Uparrow )</td>
<td>( \downarrow )</td>
<td>Rhombus</td>
<td>4.2</td>
</tr>
<tr>
<td>( \downarrow )</td>
<td>( \Uparrow )</td>
<td>( \downarrow )</td>
<td>Symmetric Triangle</td>
<td>4.0</td>
</tr>
<tr>
<td>( \downarrow )</td>
<td>( \Uparrow )</td>
<td>( \downarrow )</td>
<td>Diamond</td>
<td>3.5</td>
</tr>
</tbody>
</table>

The first column shows the post shape as oriented with respect to the flow, which is vertical. For the orientation of posts shown in Table 4.1, the tilt of the array is to the right, meaning each subsequent row is offset 1/42 of a horizontal period (period = post + gap = 36 \( \mu \text{m} \)). Thus, the unit cell of the array is rhombic with interior angles of 90° +/- 1.36°. In the actual mirrored array (Figure 4.6 and Figure 4.7), the right half of the array, which has tilt to the left, has posts that are flipped horizontally with respect to those shown in Table 4.1.

We define symmetry about the center of the gap with respect to an axis parallel to the average flow direction. Thus, the asymmetric triangle is asymmetric by this definition, just as the symmetric triangle is symmetric. Also, note that the rhombus
has interior angles of $90^\circ \pm 2.72^\circ$, with two sides perpendicular to the average flow direction and the other two sides at an angle of $1.36^\circ$ relative to the tilt angle.

The variation in the critical size with post shape in Table 4.1 is mainly a result of the effect of the post shape on the velocity profile in the gap and to a lesser extent an effect of increasing Re on the velocity profile in the gap. The critical size is still significantly larger than the expected 2.5 $\mu$m size of erythrocytes for a deep-channel (depth $\geq 20 \mu$m). The maximum flow velocity and Re at which experiments can be conducted is limited by the maximum force which the syringe pump can apply, and average flow velocity in the gap of 13.5 cm/s and Re of 2.3 are currently the maximum achievable for an array with 17 $\mu$m gaps. The fluidic resistance of the array scales with the square of the gap width, so significantly higher flow velocities and Re can be achieved in arrays with a larger gap width.

**Erythrocyte Product Contamination Experiments**

The post shapes and critical sizes used for the erythrocyte product contamination experiments are shown in Table 4.2 which is similar in layout to Table 4.1. We seek to examine post-shape-dependent erythrocyte contamination of the product over a wider range of flow rates (velocities) and Re than we examine for post-shape-dependent compression of leukocytes. The limitation on the maximum flow rate we can operate the DLD array is determined by the maximum pressure that can be supplied by the syringe pump divided by the fluidic resistance of the array. The fluidic resistance of the array scales inversely with the square of the width of the gap between the posts. Thus, we use arrays with the array parameters that were used in the cancer cell harvesting experiments in Chapter 3 to examine the dependence of erythrocyte contamination of the product on post shape at Re up to 15, corresponding to an average flow velocity in the gap of 42.9 cm/s.
Table 4.2: Post shapes used in the erythrocyte product contamination experiments. Experiments are conducted at a flow velocity corresponding to Re of 15, with the average flow velocity in the gap between the posts being 42.9 cm/s. Array parameters are posts that can fit within a 62 µm by 62 µm square, 36 µm gaps, and 1/26 tilt for all post geometries.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>Critical Size (µm) at Re = 15</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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<td>Asymmetric Triangle</td>
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<td></td>
<td>Circle</td>
<td>10.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rounded Triangle</td>
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<td></td>
<td></td>
<td>Extended Semi-Circle</td>
<td>9.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Upright Triangle</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Quarter Circle</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Square</td>
<td>11.4</td>
</tr>
</tbody>
</table>

4.3.3 Modelling of Flow Streamlines

Modelling is performed in the numerical simulator package, COMSOL, using the 2D laminar flow module. All simulations are performed with array parameters of 60 µm posts, 40 µm gaps, and 1/20 tilt. The array size is 700-µm width (perpendicular to the flow direction) and 2-mm length (parallel to the flow direction). The walls of the array are designed to preserve vertical flow near the edge of the array [34]. Boundary conditions are "no slip" at the walls of the array and the edges of the posts. The flow rate is controlled by the normal inflow velocity across the top boundary of the array.
Figure 4.8 shows surface plots of the fluid velocity and streamlines in a DLD array with 60 \( \mu \text{m} \) circular posts, 40 \( \mu \text{m} \) gaps, and 1/20 tilt at Re of 1 and 20. Some qualitative conclusions can be drawn from Figure 4.8. As Re increases, the high velocity region in the gap becomes elongated because of the fluid’s inertia in the high velocity gap region. The fluid does not like to spread out in the broadening region...
between the posts at high Re, and at Re>10, vortices behind the posts form for this reason. The increased flow velocity increases the fluid shear as the fluid bends around the bumping side of the post into the gap. In section 4.4, we discuss how this contributes to cell compression against the posts and how certain post shapes minimize this effect. It has been suggested that the vortices that form at Re∼10 prevent particles below the critical size from behaving in zig-zag mode, thus making these particles behave as particles above the critical size of the array [56, 54]. We use the streamlines and detailed flow pattern extensively in explaining the behavior of erythrocytes as Re increases beyond 1 in section 4.5.

4.4 Leukocyte Yield and Post Shapes at High Flow Rates

In this section, we experimentally determine and fundamentally explain how post shape affects leukocyte yield at high flow rates.

4.4.1 Experiment and Results

We use arrays with parameters described in Table 4.1 to separate leukocytes from whole blood. The length of the array is 39.3 mm, and the total array width is 1.5 mm. The experimental conditions at which leukocyte capture is measured are (i) the average flow velocity in the gap is 13.5 cm/s, the Reynolds number (gap width * average flow velocity in the gap / kinematic viscosity of water at room temperature) is 2.3, and (ii) the average shear rate (average flow velocity in the gap / half the width of the gap) is 15000 s$^{-1}$. The results for the six post geometries tested are shown in Table 4.3 below.

Although the critical sizes for all of the post shapes are similar (Table 4.1), the leukocyte yields vary strongly based on post shape. The leukocyte yields vary from
Table 4.3: Leukocyte yield (harvested into the central product outlet) for different post geometries under the same experimental conditions. Array parameters are 19 µm posts, 17 µm gaps, and 1/42 tilt for all post geometries except the asymmetric triangle, which has 34 µm posts, 22.5 µm gaps, and 1/42 tilt. The average flow velocity in the gap is 13.5 cm/s, the Reynolds number is 2.3, and the average shear rate is 15000 s⁻¹ for all post geometries. Volume of blood was greater than 3 mL for each post shape, and blood was diluted 1:4. The running time ranged from 30 to 40 minutes for each post shape.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>Leukocyte Yield (%)</th>
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<td>18.7</td>
</tr>
<tr>
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<td><img src="image" alt="Arrow" /></td>
<td>Rhombus</td>
<td>32.9</td>
</tr>
<tr>
<td></td>
<td><img src="image" alt="Symmetric Triangle" /></td>
<td><img src="image" alt="Arrow" /></td>
<td>Symmetric Triangle</td>
<td>64.8</td>
</tr>
<tr>
<td></td>
<td><img src="image" alt="Diamond" /></td>
<td><img src="image" alt="Arrow" /></td>
<td>Diamond</td>
<td>83.6</td>
</tr>
</tbody>
</table>

18.7% to 83.6%. Leukocyte yields for post shapes with relatively flat surfaces parallel to the flow in the gap are significantly lower than the leukocyte yields for post shapes with vertices facing into the gap. The circle and quarter circle (quarter circle with rounded edge facing upstream and on the bumping edge) post geometries resulted in less than 30% of leukocytes being displaced into the product. This low yield is a high Re effect as the leukocyte yield was 88% using a nearly identical DLD array with circular posts (18 µm posts, 18 µm gaps, 1/42 tilt, with critical size 5% greater than the circular post array used in the experiment in Table 4.3) operated at Re of 0.07, corresponding to an average flow velocity in the gap of 0.4 cm/s.
Although offering less surface area against which the leukocytes could be compressed compared to circular and quarter-circular posts, rhombus-shaped posts displaced only 32% of the leukocytes into the product. Posts with sharp corners facing into the gap had much higher leukocyte yields, with symmetric isosceles triangles displacing 64% of erythrocytes into the product and asymmetric isosceles triangular posts and diamond posts displacing more than 80% of leukocytes into the product.

4.4.2 Modelling

Figure 4.9: Surface plot of the shear rate around a circular post (left) and a diamond post (right) in a DLD array with the flow having a Re of 15. Flow direction is from top to bottom, and the array tilt is to the right. Note that there is a significantly larger high-shear region on the right and left sides of the circular post compared to the diamond post. Cell compression occurs in the high-shear region on the upstream facing half of the right (bumping) side of the post, as a cell in the first streamline next to the post follows the streamline around the post. The white circles qualitatively represent a leukocyte (WBC) path following the first streamline around the bumping side of the post.

We hypothesize that the low yield of leukocytes is due to shear stress elongating the cells, so the cells act “smaller” than expected and below the critical size for bumping (See Section 4.2.5 and Figure 4.5). The shear rate is the rate at which a progressive shearing deformation is applied to the fluid, in this case, by the walls of the micro-posts and can be represented in a 2D coordinate system by:
\[
\gamma_{ij} = \frac{\partial v_i}{\partial x_j} + \frac{\partial v_j}{\partial x_i}
\]  

(4.13)

where \(v_i\) and \(v_j\) are the velocities in the i and j directions, respectively, and \(x_i\) and \(x_j\) are the displacements in the i and j directions, respectively.

We now examine the shear rate around micro-posts of various shapes in a DLD array with micro-posts with height and width of 60 \(\mu\)m, gaps of 40 \(\mu\)m, and tilt of 1/20 in a flow having Re of 15 using the 2D laminar flow module in COMSOL (Figure 4.9). Qualitatively, the shear rate is a measure of the extent to which the fluid bends around the post. There is a significantly larger high shear rate region on the right (bumping) and left sides of the circular post compared with the diamond post (Figure 4.9). This means that the fluid, in bending around the post, interacts far more with the circular post than with the diamond post, and a cell following this fluid will correspondingly interact (be deformed) more by a circular post than by a diamond post, as can be seen from the path followed by the leukocytes in Figure 4.9. The cell interaction with the diamond post is more of a glancing blow, whereas with a circular post, the interaction is a more sustained deformation.

The shear rate near a micro-post in a DLD array is highly dependent on post geometry. For shallow tilt angles (tilt < 1/10), the shear rate is sensitive primarily to post shape, and there is little qualitative dependence on post size, gap size, or Re (for Re>1), as long as these are maintained to fixed values when comparing among post shapes. There is a significantly larger high shear region on the sides parallel to the flow direction of a circular post compared with a diamond post (Figure 4.9) or a triangular post. This high-shear region results from fluid bending around the micro-post, leading us to examine the relationship between the shear from the fluid bending around the micro-post to the extent of cell deformation.

The shear rate of the fluid at the surface of the micro-post comes from two sources. The first is that the fluid velocity is constrained to zero at the surface of the micro-
post, while it has some finite velocity in the gap. An upper bound for the shear rate from this source at a given flow rate can be found by taking the average value of the shear rate in the region highlighted in green for a square post as shown in Figure 4.10(a). The second source, which results in significantly higher shear rates, is from the fluid bending around the micro-post. The region in which the shear rate from this source contributes to cell compression is highlighted in red in Figure 4.10(b) and can more generally be described as the region at the surface on the bumping side of the post that has an outward normal that points between 0° and 90° on a standard unit circle, where 0° points from the bumping side of the post into the gap perpendicular to the average flow direction. In other words, it is the upstream edge on the side where the particle will “bump” against the post.

We integrate the shear rate from the fluid bending around the micro-post over the region at the surface of the micro-post against which cell compression can occur. We define the shear rate from the first source described above as $\gamma_0$ and the total shear rate as $\gamma$. For each micro-post shape, we then compute:
Figure 4.11: Fraction of leukocytes displaced into the product as a function of the shear rate from fluid bending around the micro-post integrated over the surface of the micro-post at which cell compression can occur for six different post geometries. The shear rate from fluid bending around a square micro-post integrated over the surface of the micro-post against which a cell can be compressed is shown for reference. The shape of the posts is schematically indicated, assuming the flow direction is vertical from top to bottom. The tilt of the array is to the right as one moves down through the array, and the flow is from top to bottom.

\[
\int_{L} (\gamma(s) > \gamma_0) \cdot (\gamma(s) - \gamma_0) \cdot ds
\]  

(4.14)

where \( L \) is along the edge of the post against which a cell can be compressed and integration only occurs for \( \gamma(s) > \gamma_0 \). The integral is computed for each of the six post geometries using the model described in Section 4.3.3 and compared to the experimental results from Section 4.4.1 in Figure 4.11.

From Figure 4.11 we see that leukocyte yield is inversely related to the shear rate at the surface of the post integrated over the surface against which the cell can be compressed. Since we are only integrating along the surface of the post, this means that the area of the "bumping" surface of the post that contributes to fluid bending around the post is significant. Posts with vertices (sharp corners) facing into the gap result in cells experiencing a glancing blow against the post, resulting
in less deformation. Cells with surfaces nearly parallel to the average flow direction interact with the post over a greater area and longer time, resulting in more significant deformation.

We now turn to examine the effect of post shape on erythrocyte displacement into the product at high flow velocities in the next section. In Section 4.6, we discuss which post shapes give the best combination of high leukocyte yield and low erythrocyte contamination in the product output.

4.5 Dependence of the Erythrocyte Contamination of the Product Output on Post Shape at High Flow Rates

In this section, we experimentally determine and fundamentally explain how post geometry contributes to erythrocyte contamination of the product at high flow rates.

4.5.1 Experimental Dependence on Post Shape at High Flow Rate

We use DLD arrays with parameters described in Table 4.2 to examine the effect of post geometry on erythrocyte contamination of the product at Re greater than 1. These array parameters were originally used for harvesting PC3 cancer cells from blood, which have a much larger critical size of $\sim 17 \text{ µm}$ than one would have for harvesting leukocytes ($\sim 4.5 \text{ µm}$). Nevertheless, the lessons we learn here about undesired contamination of erythrocytes into the product output applies to designs with smaller critical sizes, as will be shown in Section 4.6. In the PC3 cancer cell harvesting project, which is described in Section 3.3 of Chapter 3, we originally used asymmetric triangular posts because they allow for a wider gap for a fixed critical size.
Table 4.4: Erythrocyte contamination of the product and displacement of polymer microspheres (beads) below the critical size of the DLD array into the product for different post geometries under the same experimental conditions. Array parameters are 62 µm posts, 36 µm gaps, and 1/26 tilt for all post geometries. The average flow velocity in the gap is 42.9 cm/s, the Reynolds number is 15.4, and the average shear rate is 24000 s⁻¹ for all post geometries.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>% RBCs in Product</th>
<th>% 5.7 µm Beads in Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Asymmetric Triangle</td>
<td>4.9</td>
<td>4.6</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Circle</td>
<td>0.03</td>
<td>0.01</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Rounded Triangle</td>
<td>52.3</td>
<td>33.7</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Extended Semi-Circle</td>
<td>0.04</td>
<td>0.03</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Upright Triangle</td>
<td>3.1</td>
<td>--</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Quarter Circle</td>
<td>0.03</td>
<td>--</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Downturned Triangle</td>
<td>65.2</td>
<td>39.6</td>
</tr>
<tr>
<td>↓</td>
<td></td>
<td></td>
<td>Square</td>
<td>0.43</td>
<td>--</td>
</tr>
</tbody>
</table>

size and tilt angle, leading to higher flow rates for a fixed applied pressure and less clogging due to cell adhesion to the posts. However, we noticed surprisingly high erythrocyte contamination of the product that did not decrease even as we increased the critical size of the asymmetric triangular post array from 6 µm to 7.5 µm by changing the array tilt while keeping the flow Re constant, implying that the cause of the erythrocyte displacement was hydrodynamic as opposed to bumping. When we changed the asymmetric triangular posts to conventional circular posts of the same size, the erythrocyte contamination of the product diminished by a factor of 150 for the same flow Re. This motivated us to investigate the effect of post shape on erythrocyte displacement in DLD arrays as Re increases above 1.

The length of the array is 38.2 mm, and the width is 2.3 mm. The experimental conditions at which erythrocyte contamination of the product is measured are that
the average flow velocity in the gap is 42.9 cm/s, the Reynolds number (gap width * average flow velocity in the gap / kinematic viscosity of water) is 15.4, and the average shear rate (average flow velocity in the gap / half the width of the gap) is 24000 s$^{-1}$. The results for the eight post geometries tested are shown in Table 4.4.

Reorientation of erythrocytes by post shapes can change their apparent size and behavior in the DLD array [88, 70]. The shapes and motion of erythrocytes in microfluidic devices at various flow rates, including the effects of inertial lift forces at moderate Reynolds numbers, has been studied extensively [22]. Recently, the behavior of erythrocytes in DLD arrays has been examined through simulation studies at low Reynolds numbers [42, 92]. In order to determine the extent to which qualitative types of erythrocyte behavior contribute to the effects we observe here, we run the same experiments with 5.7+/−0.38 µm non-deformable polymer microsphere beads, which are well below the critical size of the array for all post shapes considered. The displacement of the polymer microsphere beads into the product very closely tracks the displacement of erythrocytes into the product (Table 4.4), which shows that effect of post shape on erythrocyte displacement is not due to reorientation of erythrocytes by the posts or other specific flow-velocity-dependent behavior, such as shape changes, of erythrocytes.

The main observation from these experimental results (Table 4.4) is that the fraction of erythrocytes displaced into the product (vs. the erythrocytes input into the DLD array) varies by a factor of over 2000, from 0.03% to 65.2%. In analyzing these results, we consider symmetry about an axis parallel to the average flow direction, and the terms “symmetric” and “asymmetric” are used to refer to the existence or non-existence, respectively, of symmetry about this axis (Symmetry about an axis perpendicular to the flow direction is referred to as “up-down symmetric”). Note that erythrocyte contamination of the product is significantly higher for asymmetric triangular post geometries than for symmetric and asymmetric but very rounded post
geometries. Even though square posts have sharp corners, the erythrocyte displacement into the product with square posts is at least an order of magnitude less than with any of the asymmetric triangular post geometries, indicating that hydrodynamic asymmetry is a significant driver of erythrocyte contamination of the product. By hydrodynamic asymmetry, we mean that the post shape is asymmetric such that the curvature of the streamlines entering the gap around the bumping side of the post at one side of the gap is very different than the curvature of the streamlines entering the gap around the non-bumping side of the post at the opposite side of the gap.

4.5.2 Experimental Dependence on Flow Rate and Post Symmetry

We now examine the dependence of the undesired erythrocyte collection in the product output on flow velocity. For reference, Re of 1 corresponds to an average flow velocity in the gap of 2.9 cm/s, and Re of 20 corresponds to an average flow velocity in the gap of 58.8 cm/s.

The fraction of erythrocytes displaced into the product is shown as a function of Re for DLD arrays with asymmetric triangular and circular posts with the same post size, gap size, and tilt angle (Figure 4.12). First, we note that for conventional circular posts there is little erythrocyte contamination of the product at all Re. For asymmetric triangular posts, even at Re of 1, there is measureable (concentration > 1x10^4 mL^-1) erythrocyte contamination of the product. With asymmetric triangular posts, the erythrocyte contamination increases significantly with Re before leveling off at Re~10, with approximately 5% of erythrocytes in the product. With circular posts, the fraction of erythrocytes in the product increases very slowly with Re, never exceeding 0.03% of erythrocytes (concentration: 1.1x10^5 mL^-1) even at Re of 20, which is an order of magnitude less than the fraction of erythrocytes appearing in
Figure 4.12: Fraction of erythrocytes in the product vs. Re for asymmetric triangular posts and circular posts. The fraction of erythrocytes in the product increases significantly with Re for asymmetric triangular posts but not for circular posts. The asymmetric triangular posts are isosceles triangles with height (in the direction of the flow) and width (perpendicular to the direction of the flow) of 62 $\mu$m, and the gap is 36 $\mu$m. The circular post array has 62 $\mu$m diameter circular posts with 36 $\mu$m gaps. The tilt is 1/26 for both asymmetric triangular and circular post arrays.

the product with asymmetric triangular posts at Re of 1 and is mainly comprised of small clumps of 3-5 erythrocytes.

This confirms that the high undesirable lateral displacement of erythrocytes into the product with asymmetric triangular posts is a high flow velocity ($1<$Re$<20$) effect. From this section and the previous section, we have learned that erythrocyte displacement into the product occurs in a flow-velocity-dependent manner for post shapes that lack symmetry about an axis parallel to the average flow direction. Further, this effect is not due to the reorientation of erythrocytes by certain post shapes or the specific velocity-dependent behavior of erythrocytes, as similar behavior is observed for non-deformable polymer microspheres.
4.5.3 Modelling

Model Definition

In Section 4.2.4, we showed that a particle following a streamline is acted upon by a viscous drag force, which keeps the particle near the streamline, and a centripetal acceleration force, which causes the particle to travel perpendicular to the streamline with lateral velocity, $v_L$. For an erythrocyte, this lateral velocity, $v_L$ is given by:

$$v_L = \frac{(\rho_{RBC} - \rho_f)V_{RBC}a_c}{6\pi\mu R_{RBC}}$$  \hspace{1cm} (4.15)

where $\rho_{RBC}$ and $\rho_f$ are the densities of the erythrocyte ($1.13\times10^3$ kg/m$^3$) and fluid ($1.03\times10^3$ kg/m$^3$), respectively, $V_{RBC}$ is the volume of the erythrocyte ($1.10\times10^{-16}$ m$^3$), $a_c$ is the centripetal acceleration, $\mu$ is the dynamic viscosity of the fluid (1x$10^{-3}$ kg/(m·s)), and $R_{RBC}$ is the radius of the erythrocyte ($5\times10^{-6}$ µm).

Using the 2-D velocity surface in COMSOL (without any particles), which is described in Section 4.3.3 as the input, we numerically evaluate $a_c$ by tracing fluid paths starting across the width of one gap into the subsequent gap in the direction of the flow (Figure 4.13). The 2-D velocity surface encodes the x and y components of the velocity at 1 µm intervals in both the x and y directions. The fluid path is traced by multiplying the x and y components of the velocity at the current point by the time to travel 1 µm in the y-direction (which is the flow direction), which is equal to 1 µm divided by the y-component of the velocity at the current point. Although the grid spacing is 1 µ, weighted averages of the two nearest points on either side of the current point along the x-axis are used in tracing the fluid to the next point in the y-direction. The total acceleration vector is then computed by the vector difference between the velocity at the next point and the velocity at the current point divided by the time to travel between the two points, which is (1 µm)/$v_y$, where $v_y$ is the average of the y-velocities at the current and next points. The centripetal acceleration
Figure 4.13: A DLD array with zero tilt is shown to illustrate tracing of fluid paths from one gap to the next in the direction of the flow to enable calculation of the centripetal acceleration. Flow is along the y-axis. Fluid paths, shown in blue, are traced at equally spaced points along the x-axis from one gap to the subsequent gap in the direction of the flow. The fluid paths are shown as straight lines for illustrative purposes. The actual fluid paths are traced numerically using data from the 2-D velocity surfaces computed in COMSOL, which is described in Section 4.3.3.

can then be computed from the dot product between the total acceleration and the vector average of \( n_1 \), which is the unit normal to the velocity vector at the current point, and \( n_2 \), which is the unit normal to the velocity vector at the next point.

We have now calculated the centripetal acceleration, \( a_c \), at 1 µm intervals along each of the paths that we seek to evaluate. From equation [4.15], we can calculate the lateral velocity of the erythrocyte induced by the centripetal acceleration at each point. We now seek the total lateral displacement of the erythrocyte over each fluid path we trace from one gap to the subsequent gap in the direction of the flow. In order to calculate this, we have to evaluate the integral (See Section 4.2.4):
\[ x_L = \int v_L d\left( \frac{y}{v_y} \right) \]  

(4.16)

over each path from one gap to the subsequent gap in the direction of the flow. Here, distance in the direction of the flow (vertical) and \( v_y \) is the velocity in the direction of the flow (vertical).

Note that our model defines positive centripetal acceleration \((a_c > 0)\) for fluid with increasing velocity curving to the left as the fluid moves through the gap and negative centripetal acceleration \((a_c < 0)\) for fluid with increasing velocity curving to the right as the fluid moves through the gap.

**Model Application**

Figure 4.14: Surface plots of the magnitude of the fluid velocity in a DLD array with 60 \( \mu \text{m} \) posts, 40 \( \mu \text{m} \) gaps, and 1/20 tilt at Re of 20 for (a) circular posts and (b) asymmetric isosceles triangular posts. The erythrocyte shown in red should take the zig-zag path, which is shown in silver, leading it to the waste output. At this Re, it often instead takes the black path with asymmetric triangular posts but not with circular posts, leading it to the product output. Note that the velocity vectors across the narrowest width of the gap shown in red are oriented nearly entirely vertically for circular posts but have a significant horizontal component for asymmetric triangular posts.
Asymmetric triangular posts are often used in DLD array because compared with circular posts, asymmetric triangular posts offer a larger gap for a given critical size and tilt angle [49]. Because both asymmetric triangular posts and circular posts are popular and because there is a very large (>100X) difference in undesired erythrocyte displacement into the product at Re of 15 between arrays with these two post shapes, we first focus our attention on these two post shapes.

Surface plots of the fluid velocity in DLD arrays with circular and asymmetric triangular posts at Re of 20 are shown in Figure 4.14. At Re of 20, the fluid bends around the two post shapes into the gap in different ways, as shown by the red arrows. With circular posts, the red arrows are relatively vertical across the width of the gap. However, with asymmetric triangular posts, the red arrows closest to the vertex of the triangle (the bumping side of the post) indicate that the velocity here has a significant horizontal component. Below the gap, the vectors point back to the left. We seek to evaluate if the centripetal acceleration from the fluid bending around the asymmetric triangular post in this way contributes to the higher displacement of erythrocytes into the product.

In order to make this evaluation, we evaluate, for both circular and asymmetric triangular post arrays, the integral in Equation 4.16 over each of 39 fluid paths that we trace from one gap to the next in the flow direction starting at 1 µm intervals across the width of the first gap using the 2-D surface plot of the fluid velocity in Figure 4.14 (Re = 20). This method is schematically illustrated in Figure 4.13. The values of $x_L$ for each of the 39 paths that we trace starting at 1 µm intervals across the width of the first gap for circular and asymmetric triangular posts are shown below in Figure 4.15.

For displacement of a particle from the “slingshot” effect to be large enough for the particle to reach the product output, in each unit cell it would have to be displaced as much as if it were a large particle bumping normally. $x_L$ over one period would have
Figure 4.15: Total lateral displacement, $x_L$, of the erythrocyte due to centripetal acceleration as fluid is traced from a point $x \, \mu m$ into the gap from the bumping side of a post to the subsequent gap in the direction of the flow. The flow velocity corresponds to Re of 20. Negative $x_L$ for small values of $x$ results from tracing fluid that zig-zags to the next gap opposite the tilt direction instead of flowing into the next gap following the tilt.

to equal, on average, the tilt angle times the gap size, which is $1/20 \times 40 \, \mu m = 2 \, \mu m$ in this case. For both asymmetric triangular posts and circular posts at Re of 20, $x_L$ does not exceed $0.02 \, \mu m$ for any of the 39 paths that we trace from one gap into the next in the direction of the flow (Figure 4.15). This is two orders of magnitude too small for displacement of erythrocytes into the product to result from the “slingshot” effect. Thus, we conclude that this conceptually simple mechanism is not responsible for the undesired erythrocyte contamination, assuming our assumption that the particle does not disturb the flow distribution is correct.

4.5.4 Other Post-Shape-Dependent High Flow Rate Effects

Although the flow rate is too small by two orders of magnitude for inertial (shear-gradient and wall effect) lift forces to affect particle trajectories for the array parameters considered here, hydrodynamic effects due to asymmetry, such as those involved in non-inertial lift cell sorting [19, 21] and pinched flow fractionation [86, 80, 66],
are still possible. Since the tilt angle is shallow (tilt angle < 1/20) in our work, for circular posts, fluid enters the gap at one side of the gap with centripetal acceleration of similar magnitude but opposite sign as at the opposite side of the gap. However, for asymmetric triangular posts, the fluid has a significantly higher centripetal acceleration around the vertex pointing into the gap from the bumping side of the post compared to fluid entering around the flat side of the asymmetric triangular post at the opposite side of the gap. This is similar to the conditions for pinched flow fractionation described in Section 4.2.3.

Note that “centripetal acceleration” is in some sense a proxy for “curvature” of the streamlines and hydrodynamic effects in general, as would cause pinched flow effects. Therefore, we examine how centripetal acceleration varies in the array with different post geometries. For simplicity, we examine the centripetal acceleration of the fluid across the narrowest point of the gap, not the entire unit cell as in the previous section.

We first examine the flow velocity dependence of erythrocyte contamination of the product with circular and asymmetric triangular posts and how this relates to the centripetal acceleration distribution in the gap between the posts for each of these two post shapes. We then alter the degree of asymmetry of fluid entering the gap by altering the up-down symmetry (symmetry about an axis perpendicular to the flow direction) of the asymmetric triangular posts and by altering the roundedness of the triangular posts. Finally, we show that it is the combination of sharp corners and post asymmetry that causes significant erythrocyte contamination of the product at high flow velocities corresponding to Re>1.
Flow Velocity Dependence of Erythrocyte Contamination of the Product with Circular and Asymmetric Triangular Posts

We showed in Section 4.5.2 that erythrocyte contamination of the product increases significantly with Re in DLD arrays with asymmetric triangular posts but not in DLD arrays with circular posts. In order to better understand this effect, we begin by examining surface plots of the fluid velocity in DLD arrays with asymmetric triangular and circular posts at Re of 1 and 20 (Figure 4.16). For circular posts, the main difference between the surface plots of the fluid velocity at Re of 1 (Figure 4.16(a)) and Re of 20 (Figure 4.16(c)) is that at Re of 20, the high velocity region in the gap between the posts is elongated compared to the high velocity region in the gap...
between the posts at Re of 1. This occurs because the inertia of the fluid prevents it from spreading out in the subsequent region following the gap in the direction of the flow (vertical). With asymmetric triangular posts, we can see from the surface plots of the fluid velocity at Re of 1 (Figure 4.16(b)) and at Re of 20 (Figure 4.16(d)) that this inertia of the fluid has an additional effect, which causes the high velocity region in the gap between the posts at Re of 20 to be asymmetric. At Re of 20, the fluid coming around the vertex of the post on the left side of the gap has substantial curvature compared to fluid entering the gap around the flat side of the post at the right side of the gap (Figure 4.17). This leads us to examine how the centripetal acceleration distribution changes with Re for circular and asymmetric triangular post shapes (Figure 4.18).

With the symmetric circular posts, the centripetal acceleration distribution in the gap is similar at Re of 1 and Re of 20, with the only significant difference being in the
Figure 4.18: The acceleration of the fluid perpendicular to the streamlines (centripetal acceleration) in the gap between the posts is plotted as a function of distance from the bumping side of the post for circular posts and asymmetric triangular posts for (c) Re=1 and (d) Re=20. For the asymmetric triangular posts, note that the centripetal acceleration distribution is broadly spread across the gap at Re of 1, but is mostly confined to a region extending 7 µm into the gap from the bumping side of the post at Re of 20. In contrast, the centripetal acceleration distribution in the gap for circular posts is similar at Re of 1 and Re of 20, with the main difference being the magnitude of the centripetal acceleration.

magnitude of the centripetal acceleration. The centripetal acceleration of the fluid around the bumping side of the post at one side of the gap is approximately equal in magnitude to the centripetal acceleration of the fluid around the non-bumping side of the post at the opposite side of the gap at both Re of 1 and Re of 20. With the asymmetric triangular posts, the centripetal acceleration distribution is spread broadly across the gap at Re of 1, but is mostly confined to a region extending 7 µm into the gap from the bumping side of the post at Re of 20, which represents
about 1/20 of the total flow through the gap by volume. This results from the centripetal acceleration around the vertex that comprises the bumping side of the post for the asymmetric triangular post increasing rapidly with Re while the centripetal acceleration around the flat side of the asymmetric triangular post at the opposite side of the gap remains at approximately zero. Because the shape of the centripetal acceleration distribution across the gap qualitatively matches the observed effect of erythrocyte contamination of the product vs. flow rate (Section 4.5.2) for circular vs. asymmetric triangular posts, we strongly suspect that it is related to the underlying mechanism. In the next two sections, we examine, using other post shapes, how geometry-induced changes in the centripetal acceleration around the bumping side of the post at a fixed Re affect erythrocyte contamination of the product.

**Effect of Up-Down Asymmetry of Left-Right Asymmetric Triangular Posts on Erythrocyte Contamination of the Product**

![Figure 4.19: Definition of fluid “angle of attack” illustrated in a DLD array with (a) downturned triangular posts, which have a $\theta_{\text{Attack}}$ of 90°, and (b) quarter circular posts, which have a $\theta_{\text{Attack}}$ of ~0°.](image)

In this section, we examine the effect of up-down symmetry (symmetry about an axis perpendicular to the flow) of left-right asymmetric triangular posts on erythrocyte contamination of the product. We begin with the up-down symmetric left-right asymmetric triangular post (Figure 4.20(a)) that we have used so far in this chapter, and break the up-down symmetry in two ways to create downturned triangular posts(Figure 4.20(b)) and upright triangular posts(Figure 4.20(c)). In order to exam-
Figure 4.20: Surface plots of fluid velocity in a DLD array with 60 µm posts, 40 µm gaps, and 1/20 tilt at Re of 20, the experimentally observed fraction of erythrocytes displaced into the product at Re of 15, and the fluid “angle of attack” for (a) asymmetric triangular posts, (b) downturned triangular posts, (c) upright triangular posts, and (d) square posts.

The effect of sharp corners on erythrocyte contamination of the product, we then compare erythrocyte contamination of the product with these three post shapes to erythrocyte contamination of the product with a square post (Figure 4.20(d)) instead of a circular post.

Analysis of the surface plots of fluid velocity in DLD arrays with these four post shapes yields qualitative insights into the physical mechanism driving erythrocyte contamination of the product. We begin by looking at the angle between the average flow direction (vertical) and the tangent to the upstream face of each of these four
post shapes, which we call the “angle of attack” and how this angle relates to undesired erythrocyte displacement. With downturned triangles, this angle is 90°, the largest for these four post shapes, and the undesired erythrocyte displacement is similarly the largest at 65.2%. For the up-down symmetric left-right asymmetric triangle and the upright triangle, these angles are 60° and 45°, respectively, and the undesired erythrocyte displacements are 4.9% and 3.1%, showing a clear, monotonic relationship between the angle of attack and undesired erythrocyte displacement. For square posts, however, the angle of attack is 90°, and the undesired erythrocyte displacement is only 0.43%. Thus, we conclude that, qualitatively, there is a strong, direct relationship between the angle of attack and undesired erythrocyte displacement for left-right asymmetric (asymmetric about an axis parallel to the flow direction) post shapes. Comparing erythrocyte displacement with square posts to erythrocyte displacement with circular posts, we conclude that this relationship between angle of attack and erythrocyte displacement is much weaker for left-right symmetric post shapes, and as we saw in the previous section with circular posts, post shapes with left-right symmetry have low erythrocyte contamination of the product output.

From these results, it is clear that left-right post shape asymmetry is driving undesired erythrocyte displacement, and we now examine the centripetal acceleration distribution in the gap to understand how hydrodynamic effects resulting from “curvature” of the streamlines may be contributing to this effect (Figure 4.21).

For the three left-right asymmetric triangular post shapes we consider, the angle of attack is a proxy for the amount of fluid that experiences high centripetal acceleration in entering the gap around the vertex that comprises the bumping side of the post. Since the vertical velocity profile across the width of the gap is parabolic, most of the fluid flows through the middle of the gap, with very little flowing at the sides of the gap. Thus, one way to determine how much fluid experiences high centripetal acceleration entering the gap around the bumping side of the post is to look at how
Figure 4.21: The acceleration of the fluid perpendicular to the streamlines (centripetal acceleration) in the gap is plotted as a function of distance from the bumping side of the post for asymmetric triangular, downturned triangular, upright triangular, and square post geometries at Re=20. Note that for the square post, the centripetal acceleration is computed across the top of the gap, where the black line is shown in Figure 4.20(d).

far into the gap from the bumping side of the post the high fluid centripetal acceleration extends. From Figure 4.21 we see that, qualitatively, the high fluid centripetal acceleration region extends much farther into the gap for the downturned triangle than for the up-down symmetric asymmetric triangle and the upright triangle, which have centripetal acceleration distributions in the gap that are much more similar. In section 4.5.5, we quantify this measure of the volume of fluid experiencing high centripetal acceleration by integrating the centripetal acceleration multiplied by the vertical velocity (how much fluid flows in that region) over the width of the gap and show that this value correlates well with undesired erythrocyte displacement.

The relationship between the fluid centripetal acceleration distribution in the gap and undesired erythrocyte displacement can also be viewed in terms of symmetry of the fluid centripetal acceleration distribution in the gap. In each of the three left-right asymmetric triangular post shapes we examine in this section, the centripetal acceleration at the right side of the gap is limited to small values due to the “no-
slip” condition at the flat side of the asymmetric triangle bounding that side of the gap. Therefore, how far into the gap from the bumping side of the post the high fluid centripetal acceleration region extends is also a measure of the asymmetry of the fluid centripetal acceleration distribution with respect to an axis parallel to the flow direction and passing through the center of the gap ($x = 20 \mu m$ in Figure 4.21). In contrast to the three left-right asymmetric triangular post shapes, the square post has a nearly symmetric fluid centripetal acceleration distribution in the gap by this definition.

The relatively low (0.43%) undesired erythrocyte displacement for square posts is interesting for three reasons. First, it shows that the relationship between angle of attack and undesired erythrocyte displacement is much weaker for left-right symmetric post shapes than for left-right asymmetric post shapes. Second, it had been suggested that sharp corners on post shapes can induce flipping of erythrocytes, resulting in undesired displacement [70, 88]. This result shows that this is not a major contributor to undesired erythrocyte displacement. Third, the relatively low erythrocyte displacement into the product with square posts provides further evidence that lift effects arising from interactions with the walls of the posts do not play a significant role in erythrocyte displacement into the product at high flow rates [92, 67]. Compared to other post shapes, square posts provide the most wall area (wall area along the entire length of the gap) that can exert a force on erythrocytes perpendicular to the flow direction, causing the erythrocytes to cross streamlines, yet this results in displacement of only 0.43% of erythrocytes.

**Effect of Roundedness of Asymmetric Triangular Posts on Erythrocyte Contamination of the Product**

The “roundedness” of the asymmetric triangular posts is important because it is often determined by the process used to fabricate DLD arrays and has been shown to
Figure 4.22: (a) As the roundedness of the asymmetric triangle increases from 0.5 μm, which is the experimental fabrication limit, to 30 μm, which renders the asymmetric triangle effectively a combination of half of a square and half of a circle ("extended semi-circle"), the erythrocyte contamination increases by a factor of 10 before decreasing to an amount comparable to that with circular posts. This effect is not unique to erythrocytes and also occurs for spherical beads with a diameter of 5.7 +/- 0.38 μm, which is well below the critical size of the array for all post geometries examined here. (b) Centripetal acceleration of the fluid versus distance into the gap from the bumping side of the post for each of the four post geometries in (a) at Re of 20.

Increasing the roundedness of 60 μm asymmetric isosceles triangular posts from 0.5 μm to 7.5 μm results in a more than 10-fold increase in the fraction of erythrocytes displaced into the product (Figure 4.22(a)). Further increasing the roundedness to
create an extended semi-circle (Figure 4.22(a)) causes the fraction of erythrocytes displaced into the product to decrease to a level similar to that with circular posts. The effect of the roundedness of asymmetric isosceles triangular posts on the fraction of erythrocytes displaced into the product correlates well with the asymmetry in the distribution of the centripetal acceleration of the fluid in the gap (Figure 4.22(b)). As the roundedness of the asymmetric isosceles triangular post increases from 0.5 \( \mu \text{m} \) to 7.5 \( \mu \text{m} \), the fluid accelerating around the bumping side of the post sweeps out a larger radius, causing the region of high centripetal acceleration from fluid accelerating around the bumping side of the post to extend further into the gap, while the centripetal acceleration around the non-bumping side of the post on the opposite side of the gap is limited to approximately zero. The further the region of high centripetal acceleration from fluid accelerating around the bumping side of the post extends into the gap, the more fluid is affected due to the vertical velocity profile in the gap being parabolic, which results in most of the fluid flowing near the center of the gap. We show in section 4.5.5 that the erythrocyte contamination of the product is directly related to the integral of the centripetal acceleration multiplied by the vertical velocity over the width of the gap.

**Effect of Post Asymmetry and Post Roundedness on Erythrocyte Contamination of the Product**

Right triangular posts and quarter-circular posts both lack left-right (about an axis parallel to the flow direction) and up-down (about an axis perpendicular to the flow direction) symmetry (Figure 4.23). However, at Re of 15, right triangular posts displace 100 times more erythrocytes into the product than quarter circular posts. Similarly, asymmetric isosceles triangular posts displace more than 100 times more erythrocytes into the product than extended semi-circular posts at Re of 15, as shown in section 4.5.4. This shows that the post asymmetry does not affect erythrocyte
contamination of the product at high flow rates (Re>1) when the radius of curvature of the post on the upstream edge is at least half the size of the post itself. This effect can be seen in the symmetry (about an axis parallel to the flow direction through the center of the gap) of the high velocity region in the gap at Re of 20, which is asymmetric for right triangular posts but symmetric for quarter circular posts (Figure 4.23).

\[ \text{4.5.5 Working Model of Dependence of Undesired Erythrocyte Displacement on Post Shape at High Flow Rates (Moderate Re) and Physical Interpretation} \]

In this section we present a method, supported by our experimental results, to predict undesirable erythrocyte displacement into the product based on post shape at high flow rates (moderate Re), a qualitative physical model that explains undesired
erythrocyte displacement based on post shape at high flow rates, and a discussion of the possible underlying physical high flow rate mechanisms that could be causing undesired erythrocyte displacement based on our experimental results.

**Predictive Capability of Undesired Erythrocyte Displacement into the Product Based on Post Shape at High Flow Rates**

![Image](image)

Figure 4.24: Experimentally measured fraction of erythrocytes displaced into the product (Table 4.4) as a function of integrated centripetal acceleration times vertical velocity across the width of the gap at Re of 20. Note that the y-axis (“Fraction of RBCs in Product”) is log-scale.

In Section 4.5.4 for various post shapes, we have seen that asymmetry in the fluid centripetal acceleration distribution in the gap correlates well with erythrocyte contamination of the product at high flow rates (Re > 1). For symmetric post geometries, we have seen that the centripetal acceleration distribution is symmetric about the center of the gap, with a difference in sign indicating opposite curvature. For asymmetric post geometries, we have seen that the extent of erythrocyte contamination of the product is directly related to how far into the gap the high centripetal acceleration around the bumping side of the post at one side of the gap extends relative to how
Table 4.5: Experimentally measured fraction of erythrocytes displaced into the product at Re of 15 and integrated curvature times vertical velocity across the width of the gap ($\chi/G$) at Re of 20 for the post shapes in Table 4.4.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>% RBCs in Product</th>
<th>$\chi/G$ ($m^2/s^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
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<td>Asymmetric Triangle</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Circle</td>
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<td>13216.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rounded Triangle</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Extended Semi-Circle</td>
<td>0.04</td>
<td>74500.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Upright Triangle</td>
<td>3.1</td>
<td>86345.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Quarter Circle</td>
<td>0.03</td>
<td>62372.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Downturned Triangle</td>
<td>65.2</td>
<td>137814.9</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>Square</td>
<td>0.43</td>
<td>9481.3</td>
</tr>
</tbody>
</table>

far into the gap the high centripetal acceleration around the non-bumping side of the post at the opposite side of the gap extends. The “how far into the gap” matters because little fluid travels next to the walls; most of the fluid travels near the middle of the gap, as can be seen in the vertical velocity ($v_y$) profiles in Figure 4.25. In order to quantify this phenomenon, we compute $\chi$, which we define as the integral of the centripetal acceleration, $a_c$, times the vertical velocity through the gap, $v_y$ across the width of the gap, $G$:

$$\chi = \int_0^G v_y(x) \cdot a_c(x) dx$$  \hspace{1cm} (4.17)

Numerically, our simulations are divided into 39 streamlines through the gap. The streamlines have a spacing of 1 $\mu$m, with the first one separated from the post by 1 $\mu$m. Thus, we compute:
Figure 4.25: Magnitude of the vertical velocity across the width of the gap in DLD arrays with circular and asymmetric triangular posts at Re of 20. Note that the peak vertical velocity is shifted toward the bumping side of the post ($x=0 \, \mu m$) for asymmetric triangular posts compared to circular posts, as characterized by Loutherback et al. [51].

$$\frac{\chi}{G} = \sum_{k=1}^{39} v_y(k) \cdot a_c(k)$$  \hspace{1cm} (4.18)

For each of the eight post geometries in Table 4.4 we calculate $\chi/G$ at Re of 20 and compare to the erythrocyte contamination of the product observed experimentally (Figure 4.24 and Table 4.5). The results are striking. Except for a small deviation for the square posts, there is a monotonic relationship between $\chi/G$ and the fraction of undesired erythrocytes displaced into the product. Thus, we feel we have a strong prediction capability for undesired erythrocyte collection at high flow rates that we can use for designing new post shapes.
Figure 4.26: As Re increases, the “attack angle” of the leading edge of the post into the gap becomes important with respect to inertial effects. We show the attack angle of the leading edge of the post into the gap relative to the average flow direction for (a) the downturned triangle and (b) the quarter circle. (c) The “attack angle” strongly correlates with undesired erythrocyte displacement into the product at Re of 15 for asymmetric post shapes.

### Qualitative Physical Model Explains Dependence of Erythrocyte Displacement at High Flow Rates on Post Shapes

The above subsection has shown a strong correlation between the asymmetry of flow curvature in the gap and undesired erythrocyte collection. We now qualitatively discuss why certain post shapes have this property.

(i) **Asymmetry**: Gaps which are symmetric do not have hydrodynamic asymmetry. Any undesired inertial effect pushing a particle one way at one side of the gap will be countered by an opposite vector at the other side of the gap. As expected, the circular and square posts both have relatively low undesired erythrocyte collection.

(ii) **Attack Angle of Leading Edge of Post Shape into the Gap**: At low Re (no inertial effects), fluid flow is reversible. Reversing the sign of the pressure gradient
Figure 4.27: As Re increases, the “attack angle” of the leading edge of the post into the gap becomes important with respect to inertial effects. We show the attack angle of the leading edge of the post into the gap relative to the average flow direction correlates strongly with undesired erythrocyte displacement into the product for left-right asymmetric post shapes (shown in gray) and significantly less strongly with undesired erythrocyte displacement into the product for left-right symmetric post shapes (shown in black). Note that the y-axis (“Fraction of RBCs in Product”) is log-scale.

reverses the flow direction, but streamlines are unchanged. This changes at high Re. Namely, we may expect a post with a large “attack angle” going into the gap to lead to more inertial effects, since the fluid and especially particles trying to follow the fluid must undergo a larger change in direction of velocity (centripetal acceleration), as shown in Figure 4.26(a) and (b) for the cases of large and small “attack angle”, respectively.

Undesired erythrocyte displacement into the product at Re of 15 correlates well with the attack angle of the leading edge of the post into the gap for left-right asymmetric post shapes (Figure 4.26(c)). The post shape with the largest attack angle, the downturned triangle, has the largest erythrocyte displacement into the product, while the two post shapes with nearly 0° angle, the quarter circle and the extended semi-circle, have very little erythrocyte displacement into the product. For left-right
symmetric post shapes, the correlation between the attack angle of the leading edge of the post into the gap and undesired erythrocyte displacement into the product at Re of 15 is significantly weaker compared to that for left-right asymmetric post shapes (Figure 4.27). While the qualitative model gives us good physical intuition, it unfortunately cannot explain the high erythrocyte displacement by the rounded asymmetric triangle (7.5 μm radius), even though the quantitative numerical model of the previous section can predict it.

Possible Physical Mechanisms for Undesired Erythrocyte Collection at High Flow Rates

So far, we know that the amount of fluid undergoing high centripetal acceleration into an asymmetric gap correlates well with undesired erythrocyte collection at high flow rates. We hypothesize that it is actually responsible for it.

We now discuss which of the possible mechanisms that we describe in detail in Section 4.2 could be responsible for this:

- **Inertial Lift and Dean’s Flow Forces:** The undesired erythrocyte collection is unlikely to be due to these mechanisms because the minimum flow rate required for particles (or cells) to reach stable equilibrium positions due to these forces is two orders of magnitude greater than the highest flow rates used in our experiments. This makes it highly unlikely that these forces are sufficient to cause particles to cross streamlines, resulting in displacement into the product.

- **Lift Effects from Asymmetry:** Undesired erythrocyte collection in the product output cannot be due to lift effects specific to vesicles (erythrocytes) arising from asymmetry, since the effect we observe also occurs for rigid polymer microspheres. However, effects similar to those involved in pinched-flow fractionation could be responsible for the effect we observe.
• **Centripetal Acceleration and the “Slingshot” Effect:** Undesired erythrocyte collection in the product output cannot be due to this mechanism as the displacement resulting from this mechanism is at least two orders of magnitude too small to cause erythrocyte displacement into the product, as calculated in Section 4.5.3.

• **Erythrocyte Reorientation (Flipping):** It cannot be due to this mechanism because the flow-rate dependent displacement into the product is observed for rigid polymer microsphere beads below the critical size of the array as well as erythrocytes (Section 4.5.1), indicating that this effect is not due to reorientation of the erythrocytes or other specific, flow-dependent behavior of erythrocytes.

Thus, we conclude that effects similar to those involved in pinched-flow fractionation are responsible for erythrocyte displacement into the product at high flow rates for asymmetric post shapes. Pinched flow fractionation has been extensively characterized experimentally [66, 80, 86], but the fundamental mechanisms are not well understood.

This work has focused on arrays with shallow tilt angles (tilt < 1/20), which are common in applications involving separation of nucleated cells from blood due to the large size range of the target cells. In arrays with steeper tilt angles, symmetric post geometries can cause an asymmetric centripetal acceleration distribution in the gap, resulting in displacement of particles below the critical size of the array at high flow rates. Results by Y.S. Lubbersen et al. [56, 54] show that this occurs for circular and quadrilateral posts with tilt angles of 1/6 and 1/4, respectively, as Re is increased from 2 to 30. One possible solution to this undesirable effect is to use asymmetric posts to compensate for hydrodynamic asymmetry arising from the tilt angle of the array in these cases.

Future work will probably need combined fluid and particle modelling of the flow. Calculation of the flow pattern with no particles and predicting particle behavior
from this flow profile is likely not sufficient for further work, since high flow velocity hydrodynamic lift effects depend on the effect of the particle on the flow pattern.

4.6 Optimal Post Geometry for High-Throughput Separation of Nucleated Cells from Blood with Minimal Erythrocyte Contamination

The optimal post geometry for high-throughput separation of nucleated cells from blood has two characteristics: (1) high nucleated cell (leukocyte) collection efficiency and (2) low collection efficiency of undesired erythrocytes. Section 4.4 showed (1) requires minimum shear at the post surface against which the cell can be compressed and is achieved using post shapes that have a vertex pointing into the gap in the bumping direction. Section 4.5 showed (2) requires that the flow in the gap have a symmetric centripetal acceleration distribution and is satisfied by using post shapes with symmetry about the axis parallel to the flow direction.

Section 4.4 described leukocyte collection at Re of 2.3, corresponding to an undiluted blood flow rate of 100 µL/min and processing of an undiluted blood volume of 3 to 4 mL for different post shapes. We now report the undesired erythrocyte collection at the product output in these experiments. The results are shown in Table 4.6 and plotted in Figure 4.28 with leukocyte yield (harvested into the central product outlet) on the y-axis and erythrocyte fraction in the product output on the x-axis. Consistent with our results in the previous section, post shapes that are symmetric about an axis parallel to the flow direction result in relatively low erythrocyte displacement into the product compared to post shapes that are asymmetric about this axis (Figure 4.28). There is a 130-fold increase in erythrocyte displacement into the product between asymmetric triangular posts and circular posts, whereas the difference in erythrocyte
Table 4.6: Leukocyte yield (harvested into the central product outlet) and erythrocyte fraction in the product output for different post geometries under the same experimental conditions. Array parameters are 19 µm posts, 17 µm gaps, and 1/42 tilt for all post geometries except the asymmetric triangle, which has 34 µm posts, 22.5 µm gaps, and 1/42 tilt. The average flow velocity in the gap is 13.5 cm/s, the Reynolds number is 2.3, and the average shear rate is 15000 s\(^{-1}\) for all post geometries. Volume of blood was greater than 3 mL for each post shape, and blood was diluted 1:4. The running time ranged from 30 to 40 minutes for each post shape.

<table>
<thead>
<tr>
<th>Tilt</th>
<th>Post</th>
<th>Flow</th>
<th>Geometry</th>
<th>% WBCs in Product</th>
<th>% RBCs in Product</th>
</tr>
</thead>
<tbody>
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<td></td>
<td></td>
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<td>Asymmetric Triangle</td>
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<td></td>
<td></td>
<td></td>
<td>Circle</td>
<td>25.2</td>
<td>0.009</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Quarter Circle</td>
<td>18.7</td>
<td>0.011</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rhombus</td>
<td>32.9</td>
<td>0.015</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Symmetric Triangle</td>
<td>64.8</td>
<td>0.038</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Diamond</td>
<td>83.6</td>
<td>0.009</td>
</tr>
</tbody>
</table>

displacement into the product between symmetric triangular posts and circular posts is only 4-fold.

For most post shapes, one can achieve high leukocyte yield or low erythrocyte contamination, but not both. Fortuitously, diamond posts simultaneously achieve a high (>80%) fraction of leukocytes displaced into the product with a low level of erythrocyte displacement into the product comparable to that achieved with circular posts (<0.02%).

Combining high-throughput operation of DLD arrays with inhibition of clot formation, we separate leukocytes from 3.7 mL of whole blood in less than 45 minutes using a single DLD array with diamond posts. The leukocyte capture rate is 83%, which is the highest reported in a DLD array in which the average shear rate exceeds 10,000 s\(^{-1}\). Erythrocyte concentration in the product is \(3 \times 10^4\) mL\(^{-1}\), representing a depletion by a factor of more than 10,000 relative to the starting sample. Altogether,
Figure 4.28: Fraction of leukocytes in the product versus fraction of erythrocytes in the product for six different DLD array post geometries at Re of 2.3. Note that the x-axis (Fraction of RBCs in the Product) is log-scale. The shape of the posts is schematically indicated, assuming the flow direction is vertical. The tilt of the array is to the right as one moves down through the array, and the flow direction is from top to bottom.

This represents the highest efficiency separation of leukocytes from the largest volume of blood in the shortest time ever achieved using DLD arrays. For comparison, in the largest scale separation of leukocytes from whole blood previously reported, D. Inglis et al. processed undiluted blood at a rate of 4 \( \mu \text{L/min} \) (1/20 as fast as we report in terms of undiluted blood flow rate) per DLD array, separating 98% of leukocytes from 80 \( \mu \text{L} \) of blood with more than 100 times the erythrocyte contamination of the product that we report here [37].

4.7 Conclusion

Shear-induced cell deformation and potential inertial effects arising from Re increasing above 1 present two major limitations to high-throughput separation of nucleated cells from whole blood using DLD arrays. In this chapter, we show:
(i) The extent of shear-induced cell deformation is proportional to the shear rate of the fluid bending around the surface of the micro-post against which the cell can be compressed. It can be minimized by using post geometries with vertices pointing into the gap in the bumping direction.

(ii) The displacement of erythrocytes into the product at high flow rates for certain micro-post geometries correlates well with asymmetry in the centripetal acceleration distribution in the gap between the micro-posts and can be minimized by using post shapes with symmetry about an axis parallel to the flow direction.

We then show that diamond micro-posts overcome both these limitations, simultaneously achieving maximum displacement of leukocytes into the product while minimizing erythrocyte displacement into the product.

Using this approach, we have demonstrated that 83% of leukocytes can be separated from ∼4 mL of whole blood in less than 45 minutes with a single DLD array with diamond posts, resulting in a product with 10,000-fold depletion of erythrocytes relative to the starting blood sample. Standard micro-fabrication techniques have been previously used to produce a chip containing 15 DLD arrays in parallel, which would require an area of 36 mm by 70 mm for the array presented in this paper [40, 36]. Combining a chip containing 15 of our DLD arrays in parallel with the advances presented in this chapter should allow for high-efficiency separation of leukocytes from over 50 mL of blood in less than one hour.
Chapter 5

Controlling Anisotropic Conduction in DLD Arrays with Post Geometry at Low Reynolds Number

5.1 Introduction

The performance of deterministic lateral displacement (DLD) arrays in separating particles based on size can be enhanced or diminished by misalignment of the pressure gradient to the average flow direction due to anisotropic conduction. This misalignment of the pressure gradient to the average flow direction has been shown to create a third type of particle behavior, mixed motion, which results in particles in an intermediate size range traveling at an angle between the average flow direction and the array tilt angle for an array with circular posts. In this chapter, we show (i) how post geometry and tilt angle affect this misalignment of the pressure gradient and (ii) how increasing the spacing between rows in the direction of the flow minimizes
the effect of post geometry on anisotropic conduction. We then present a practical application of using the misaligned pressure gradient to adjust the critical size of the DLD array.

Figure 5.1: (a) A tilted square unit cell array with gap size $L$ and tilt angle ($\epsilon$) of 1/5. (b) A rhombic unit cell array with gap size $L$ and tilt angle ($\epsilon$) of 1/5. Array is oriented such that the average flow direction is vertical, from top to bottom.

Although the original DLD array designed by L.R. Huang et al. used an array with a tilted square unit cell \[30\], rhombic unit cell arrays have become more widely used since rhombic unit cell arrays allow for placing multiple arrays in series with varying critical sizes (Figure 5.1). In a rhombic array, the rows are perpendicular to the average flow direction while the columns are parallel to the tilt angle. This results in anisotropic conduction, in which the conductivity in the direction of the flow is less than the conductivity in the direction perpendicular to the flow by an amount dependent upon the tilt angle of the array. In contrast, a DLD array with a tilted square unit cell is expected to be an isotropic conductor with fixed conductivity at any angle. However, we show in Section 5.2 that this is only the case for certain post shapes.

A fundamental assumption of the theory that characterizes the critical size for DLD arrays is that the average fluid flow is at a fixed angle to the post lattice that forms the array \[35\]. Previous work in minimizing the discrepancy between the
anisotropic conduction of rhombic DLD arrays with this fundamental assumption has focused primarily on two areas. First, D. Inglis devised a method for designing the boundaries of DLD arrays to ensure vertical flow by causing one streamline to shift opposite the tilt direction with each subsequent row in the direction of the flow [34]. Second, T. Kulrattanararak et al. showed that anisotropy of the array causes particles in an intermediate size range to travel at an angle in between the average flow direction and the tilt angle [43, 44]. This effect is similar to the multidirectional sorting modes with DLD arrays achieved by B.R. Long et al. by using rational number (N/M, where N and M are both integers) tilt angles [48].

Recently, post shape engineering in DLD arrays has emerged as a way to increase throughput [49, 50], sort non-spherical particles [88, 70, 39], and reduce the shear experienced by cells traveling through the array [1]. However, this recent work has focused on operation of DLD arrays at Re > 1. In this chapter, we focus on the effect of post shape on anisotropic conduction in the array at Re ~0.01 and how post geometry can be used to achieve a different effective critical size for the same array parameters.

More recently, anisotropic conduction in the array has been used to enhance separation efficiency and throughput. K.K. Zeming et al. have adjusted the ratio of the spacing between the rows in the direction of the flow to the gap size in order to enhance the throughput and separation efficiency of erythrocytes [89]. This work built on previous work showing that a single line of obstacles tilted at an angle to the average flow direction is sufficient to concentrate particles based on size from sufficiently dilute suspensions of particles [55, 16]. This allows for scaling of the fluidic resistance of the array based on the starting concentration (particles/volume) of particles in the suspension from which concentration is desired.
5.2 Anisotropic Conduction

The difference between isotropic fluid conduction and anisotropic fluid conduction in an array of square posts is demonstrated in Figure 5.2. Figure 5.2(a) shows a square array of circular posts, which has been verified in simulation (details below) to be an isotropic conductor. This means that a flow along axis $\mathbf{u}_1$ is driven by a pressure differential $P_1$ in the $\mathbf{u}_1$ direction and will not generate a pressure differential $P_2$ in the direction of axis $\mathbf{u}_2$. Figure 5.2(b) shows a square array of upright asymmetric triangular posts, which has been observed in simulation as an anisotropic conductor. This means that a flow along axis $\mathbf{u}_1$ driven by a pressure differential $P_1$ in the $\mathbf{u}_1$ direction will generate a pressure differential $P_2$ in the direction of axis $\mathbf{u}_2$. For a flow along axis $\mathbf{u}_1$, the ratio $P_2/P_1$ can be used as a measure of anisotropic conduction.

In a standard DLD array, a pressure is applied to drive the flow in a given direction (vertical in this case). Since there can be no flux through the horizontal side walls, if the array design is that of an anisotropic conductor, an internal pressure gradient will develop to keep the flow vertical. This internal pressure gradient can be used to
characterize the anisotropic conduction of the array. This is analogous to the Hall effect in semiconductors.

Figure 5.3: Schematic of simulation used to characterize anisotropic conduction in the DLD array with rhombic unit cell. A uniform pressure, $P$, is applied across the top of the array, and the bottom of the array is maintained at a uniform pressure of 0. This pressure differential drives the flow through the array, which is 20 rows long and 15 columns wide. The transverse pressure gradient, $\Delta P = P_2 - P_1$, is calculated from the difference between the pressures measured at the right and left walls of the array along the line through the middle of the array above. The array shown here has 60 $\mu$m triangular posts, 40 $\mu$m gaps, and 1/20 tilt. The walls are designed to ensure vertical flow at the sides of the array [34].

The array used in our simulations is 20 rows long and 15 columns wide (Figure 5.3). It is a “rhombus” array (not a tilted square array), and the rhombus shape alone introduces a fluid flow anisotropy, which is dependent upon the tilt angle of the array. In the next section, we investigate the effect of post shape on anisotropic conduction at different array tilt angles. The posts we use can be inscribed within a square with
Table 5.1: Post geometries for which the transverse pressure generated by a pressure gradient in the direction of the flow is measured. All post shapes can be fit within a square with side length 60 $\mu$m. Flow direction is from top to bottom with respect to post shape in the table.

<table>
<thead>
<tr>
<th>Post Shape</th>
<th>Geometry</th>
</tr>
</thead>
<tbody>
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<td>Circle</td>
</tr>
<tr>
<td><img src="image" alt="Right Up-Down Symmetric Triangle" /></td>
<td>Right Up-Down Symmetric Triangle</td>
</tr>
<tr>
<td><img src="image" alt="Left Up-Down Symmetric Triangle" /></td>
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<tr>
<td><img src="image" alt="Upright Asymmetric Triangle" /></td>
<td>Upright Asymmetric Triangle</td>
</tr>
<tr>
<td><img src="image" alt="Downturned Asymmetric Triangle" /></td>
<td>Downturned Asymmetric Triangle</td>
</tr>
<tr>
<td><img src="image" alt="Diamond" /></td>
<td>Diamond</td>
</tr>
<tr>
<td><img src="image" alt="Square" /></td>
<td>Square</td>
</tr>
<tr>
<td><img src="image" alt="Left-Right Symmetric Triangle" /></td>
<td>Left-Right Symmetric Triangle</td>
</tr>
</tbody>
</table>
side 60 µm, the gap is 40 µm, and the tilt is varied between 1/5 and 1/50. The array unit cell is rhombic and the walls are designed using the method by D. Inglis to ensure vertical flow near the edges of the array [34]. Simulations are performed using the COMSOL 2D laminar flow module at Re ∼0.01 by applying a pressure P of 1 Pa at the top boundary of the array. No-slip boundary conditions along the walls of the array and the edges of the posts are applied. The transverse pressure, ΔP=P_{2} - P_{1}, is then calculated in the simulation to characterize the anisotropic conduction in the DLD array. The post geometries we examine in this chapter are shown in Table 5.1.

5.3 Effect of Post Shape and Tilt Angle on Transverse Pressure Gradient

Figure 5.4: Relative transverse pressure gradient (ΔP/P) as a function of tilt angle (tilt angle is 1/N) for six different post geometries in a DLD array with a rhombic unit cell. The simulation is performed at Re ∼0.01 for all post geometries tested. Note that the relative transverse pressure gradient remains significant even for large N for the upright asymmetric triangle but decreases to very small values at large N for the other five post geometries.
The effect of tilt angle on the transverse pressure gradient was measured for six different post shapes (Figure 5.4). All of the post shapes are sized such that the post shapes can fit within a square with side length of 60 µm.

The transverse pressure gradient decreases with increasing N (increasingly shallow tilt angles) for all six post shapes. Note that at large N, there is very little tilt, the rhombic unit cell is approximately square, and the differences in the relative transverse pressure gradient (measure of anisotropic conduction in the DLD array) are primarily a function of post shape. Circular posts at large N approach the square array of Figure 5.2(a) and thus have no anisotropic effects.

We expect post shapes with symmetry about axes parallel and perpendicular to the average flow direction (circle, diamond, square) to have little anisotropic effects based on shape. A single qualitative view might expect a triangular post with a vertex pointing into the gap on one side to cause anisotropic conduction, but this is contradicted by the up-down symmetric triangle. At present, there is no intuitive understanding about why the upright asymmetric triangular post shape causes such a large anisotropic effect.

In order to better understand why the upright asymmetric triangular post shape causes such a large anisotropic effect, we examine four triangular post shapes that lack symmetry about at least one axis (Figure 5.5). Two of these post shapes have up-down symmetry but lack left-right symmetry (right and left up-down symmetric triangles), while the other two lack both left-right and up-down symmetry (upright and downturned asymmetric triangles). Both the right and left up-down symmetric triangular post shapes have transverse pressure gradients that decrease with increasing N, and the direction of the transverse pressure gradient is not sensitive to the orientation of the post shape in these two cases. For the upright and downturned asymmetric triangular post shapes, the transverse pressure gradient remains significant even at large N, and the direction of the transverse pressure gradient is sensitive.
Figure 5.5: Relative transverse pressure gradient ($\Delta P/P$) as a function of tilt angle (tilt angle is $1/N$) for four asymmetric triangular post geometries in a DLD array with a rhombic unit cell. The simulation is performed at $Re \sim 0.01$ for all post geometries tested. Note that the relative transverse pressure gradient remains significant even for large $N$ for the upright and downturned asymmetric triangular post shapes but decreases to very small values at large $N$ for the up-down symmetric triangular post shapes. Also, note that the direction of the transverse pressure gradient is sensitive to the orientation of the up-down asymmetric post shapes but not the up-down symmetric post shapes.

to the orientation of the post shape, as indicated by the opposite sign of the transverse pressure gradient for the downturned asymmetric triangle compared to the upright asymmetric triangle. We next examine the effect of spacing of the rows in the direction of the flow on the transverse pressure gradient with upright and downturned asymmetric triangular post shapes in order to better understand why these two post shapes cause such a large anisotropic effect.
Figure 5.6: (a) Square array of upright asymmetric triangular posts is an anisotropic conductor. It is observed in simulation that a flow along axis $\mathbf{u}_y$, driven by a pressure difference $P_y$, generates a pressure difference $P_x$ along axis $\mathbf{u}_x$. (b) We examine the effect of doubling (and tripling, not shown) the spacing between rows, $D_y/D_x$, on the pressure difference $P_x$ along the axis $\mathbf{u}_x$ generated by a flow along the axis $\mathbf{u}_y$, which is driven by a pressure difference $P_y$.

5.4 Effect of Spacing between Rows on Transverse Pressure Gradient for Different Post Geometries

We have established that post shape contributes to anisotropic conduction in DLD arrays even if the tilt angle is shallow. In this section, we examine how spacing within the array can be used to minimize anisotropic conduction. We cannot modify the gap size without changing the critical size of the array \[35\]. Therefore, we examine how we can change the spacing between rows in the direction of the flow to minimize anisotropic conduction.

In Figure 5.6(a), we show the standard array, with unit size gap and $D_y/D_x$ spacing between the rows. The fluidic resistance $R_y$ for fluid flowing along $\mathbf{u}_y$ is fixed by the gap. However, the fluidic resistance $R_x$ for fluid flowing along $\mathbf{u}_x$ can be significantly
reduced by increasing the spacing of the $u_x$ rows along the $u_y$ axis. Figure 5.6(b) shows the array with this spacing doubled compared to the array in Figure 5.6(a).

Thus, if applying a vertical (along $u_y$ axis) pressure gradient causes a flow component in the $u_x$ direction, only a very small pressure gradient in the $u_x$ direction is needed to induce a flow to counterbalance this, leading to a small $P_x/P_y$.

![Graph](image)

Figure 5.7: The relative transverse pressure gradient ($\Delta P/P$) was measured as the spacing between rows in the direction of the flow ($D_y$) was increased relative to the gap ($D_x$) for three post geometries in a DLD array with tilt $1/10$. Although upright and downturned asymmetric triangles have significantly larger relative transverse pressure gradients compared to up-down symmetric triangles when the row spacing in the direction of the flow is equal to the gap, this difference decreases rapidly as the spacing between rows of posts in the direction of the flow increases relative to the gap size.

Thus, the effect of post shape on the transverse pressure gradient is highly dependent on the spacing of the rows of posts in the direction of the flow ($D_y$) relative to the gap between posts in the direction perpendicular to the flow ($D_x$). We simulate the transverse pressure gradient as a function of $D_y/D_x$ for the upright asymmetric triangle, downturned asymmetric triangle, and up-down symmetric triangle (Figure 5.7). Simply doubling $D_y/D_x$ causes the transverse pressure gradient to decrease by more than a factor of 7 for the upright and downturned asymmetric triangular posts. Increasing $D_y/D_x$ to 3 almost completely removes all anisotropic conduction effects.
This shows that the transverse pressure gradient is sensitive to the transverse fluidic resistance ($R_x$), since $R_x$ decreases by a factor of 4 when $D_y/D_x$ is doubled and by a factor of 9 when $D_y/D_x$ is tripled.

5.5 Conclusion

Anisotropic conduction in DLD arrays, which causes a misalignment of the pressure gradient to the average flow direction, can be caused by either a rhombic (not tilted square) unit cell array or by post shapes which are asymmetric. For a grid with approximately equal spacing in the x and y directions, this misaligned pressure gradient is minimized by post shapes that are symmetric about axes parallel and perpendicular to the average flow direction. For arrays with a tilt angle achieved with a rhombic unit cell array, as is commonly used in DLD arrays, a transverse (perpendicular to the average flow direction) pressure develops between the sidewalls of the array in order to keep the average flow direction straight, since there can be no fluid flux into the sidewalls. The magnitude of this transverse pressure is determined by the steepness of the tilt of the array and the asymmetry of the post shape, and a significant transverse pressure can cause deviations in particle behavior in the array from what would be expected by conventional DLD array theory. The magnitude of this transverse pressure is highly sensitive to the spacing of the rows in the average flow direction, and increasing (doubling or tripling) this spacing for arrays with steep tilt angles or asymmetric post shapes can reduce the magnitude of this transverse pressure to very small values.
Chapter 6

Conclusion

6.1 Summary

Deterministic lateral displacement (DLD) arrays have been used to fractionate blood into leukocytes, erythrocytes, and platelets, based on hydrodynamic size [14]. Compared to other microfluidic cell sorting technologies, DLD arrays allow for separation of nucleated cells without the background of erythrocytes, platelets, and smaller non-target cells and without requiring significant (>5X) dilution of the blood for operation. DLD arrays have been applied to separation of a variety of cancer cells [50, 47], malignant lymphocytes [36], and fetal nucleated erythrocytes [32] from blood. However, the volume involved in these applications has been limited to less than 100 µL due to volume-dependent performance degradation and processing time constraints. In this thesis, we presented three approaches toward improving the volume of blood that can be processed in a given time within a fixed chip area.

In Chapter 2, we made improvements to the device fabrication processes and experimental setup to allow processing of large volumes of fluid in shorter time periods. We presented three fabrication processes that increased the number of DLD arrays that could fit in a given area on a silicon chip by a factor of four. Two of these three
fabrication processes allowed a 2.5-fold increase in the maximum depth to which the DLD array could be etched, resulting in a 10-fold increase in the volumetric flow rate for a given pressure differential over that which was achievable with previous fabrication processes. We also introduced a bubble-free design of the high-throughput manifold, through which fluidic connections are made to the DLD array, removing another barrier to reliable operation of DLD arrays over longer time periods.

In Chapter 3, we identified that volume-dependent performance degradation in processing blood with DLD arrays occurs due to conventional platelet-driven clot formation. We showed the activity of calcium ions and thrombin are the dominant mechanisms driving clot formation in the DLD array and presented a method using the calcium-chelating anti-coagulant, EDTA, and the direct thrombin inhibitor, PPACK, to completely inhibit clot formation in silicon DLD arrays. We then demonstrated, using this method, capture of 86% of PC3 cancer cells from 14 mL of blood in less than 38 minutes with only 0.03% of erythrocytes contaminating the product using a single DLD array. Putting 10 DLD arrays in parallel on a chip, which is easily technically achievable [40], would allow processing of >100 mL of blood using a single chip. Previously, single DLD arrays were limited to processing less than 250 µL of blood.

In Chapter 4, we explored the effects of post shape on the behavior of leukocytes and erythrocytes at high flow rates (moderate Re). We showed that the extent of cell deformation against the micro-posts is directly related to the shear at the surface of the post resulting from fluid bending around the post and can be minimized by using post shapes that have vertices pointing into the gap. We also identified flow velocity-dependent erythrocyte contamination of the product with asymmetric triangular posts but not circular posts. We showed that the extent of this flow velocity-dependent erythrocyte displacement is directly related to asymmetry in the fluid centripetal acceleration distribution in the gap and that this flow velocity-dependent
erythrocyte displacement does not occur for post shapes for which the fluid centripetal acceleration distribution in the gap is sufficiently symmetric. This enabled, for the first time, harvesting of leukocytes from blood at a high flow rate (100 µL/minute undiluted blood flow rate), requiring shear rate greater than 10,000 s⁻¹ and Re of 2.3, with both high leukocyte yield (83%) and low erythrocyte contamination (<0.02%).

In Chapter 5, we examine the effect of post shape on anisotropic conduction at low Reynolds number. Anisotropic conduction occurs in DLD arrays with a parallelogram unit cell instead of a rotated square unit cell and leads to a pressure gradient that is misaligned to the average flow direction. This causes particles in a certain size range to migrate at an angle between the average flow direction and the tilt angle of the array and can potentially be used to adjust the critical size of the array for a given set of array parameters. We examine how the tilt angle of the array contributes to anisotropic conduction, show that the lateral (perpendicular to the average flow direction) velocity induced by different post shapes is the dominant contributor to anisotropic conduction at a given tilt angle, and demonstrate that increasing the spacing of the rows in the flow direction significantly reduces the misaligned pressure gradient for a given post shape.

6.2 Future Work

Widespread use of DLD arrays for therapeutic and diagnostic applications in medicine will require fabrication of DLD arrays in low-cost, disposable materials. Recently, R.S. Powell et al. have devised a method to fabricate thermoplastic DLD arrays using low-cost, regulatory-approved materials and biocompatible methods [62]. Future work could include combining the method we presented for inhibition of clot formation with use of these thermoplastic DLD arrays to experimentally measure thermoplastic DLD array performance for large volumes of blood. Decreases in device performance
compared to silicon DLD arrays could be compensated through stacking and parallelization of DLD arrays, which should be easier with thermoplastics than with silicon due to the ease with which thermoplastics can be bonded together.

The advances presented in this thesis overcome two major limitations toward separation of large cells from large volumes of blood in short times. These advances may allow pursuit of two interesting applications. First, umbilical cord blood, which is typically collected in volumes of approximately 100 mL, provides a rich source of hematopoietic (blood-forming) stem cells (HSCs) that can be used for treatment of a wide variety of diseases, including a variety of cancers, bone marrow failure syndromes, immunodeficiencies, and metabolic disorders. DLD arrays provide greater separation efficiencies and superior erythrocyte depletion compared to current state-of-the-art technology [75]. Future work would include experiments to characterize DLD array performance for separation of HSCs from cord blood samples instead of using leukocyte separation from adult peripheral blood as a model system. Second, we demonstrate that DLD arrays can be used to separate cancer cells from larger volumes and with greater separation efficiency than the U.S. Food and Drug Administration-approved CellSearch system [71]. Future work would include experiments using primary cancer cells, instead of cell lines, and development of a system for on-chip detection of small numbers of cancer cells for high-sensitivity diagnostic applications.

On-chip processing of blood offers the advantages of more precise control of reaction time and lower required quantities of reagents. Recently, high-quality on-chip washing of leukocytes for diagnostic applications [9] and on-chip lysis of leukocytes for analytical applications [10] using DLD arrays were demonstrated. Future work that may be enabled by the advances presented in this thesis would include extensions of such applications to rare cells, which would require processing large volumes of blood in short times.
Appendix A

Publications and Presentations

A.1 Peer-Reviewed Publications

- J D’Silva, K Loutherback, RH Austin, and JC Sturm. Controlling Anisotropic Conduction in DLD Arrays with Post Geometry at Low Reynolds Number. (in preparation)


### A.2 Conference Presentations

- **CYTO 2016**, Seattle, WA. – Microfluidic Post Geometry Design for High-Throughput (100 µL/min) Harvesting of Leukocytes from Blood using Deterministic Lateral Displacement Arrays. (talk, accepted)


- **MRS Spring Meeting 2013**, San Francisco, CA. – High-Throughput, High-Enrichment Microfluidic Cell Sorters for Capture of Circulating Tumor Cells from Whole Blood. (poster)

- **APS March Meeting 2013**, Baltimore, MD. – Minimizing Platelet Activation-Induced Clogging in Deterministic Lateral Displacement Arrays for High-Throughput Capture of Circulating Tumor Cells. (talk)
Appendix B

Backside Alignment Fabrication Process

This appendix details the steps in the backside alignment fabrication process described in Chapter 2 Section 2.2.2. Devices are fabricated on double-side-polished 100-mm-diameter, 550-µm-thick silicon wafers.

Front-Side Processing

1. Dehydration bake the wafer for 2 minutes at 95°C.

2. Spray on HMDS and spin immediately using recipe 3.

3. Apply ∼3 mL AZ 4330 photoresist using a plastic transfer pipette and spin using recipe 3.

4. Soft bake at 95°C for 3 minutes.

5. Expose for 1 minute using Channel 2 on the MA6 mask aligner with hard contact setting.

6. Wait 10 minutes.
7. Develop using AZ 300 MIF developer. Development usually takes 2.5 minutes, but this can vary depending on conditions such as how long the photoresist has been outside the refrigerator and room humidity.

8. Inspect under microscope to ensure features are completely developed.

9. Etch with recipe 3 (200 µm trench etch) on the Samco 800 for 120 cycles (1 cycle ≈ 1.3 µm if different etch depth is desired).

**Back-Side Processing**

1. To prevent scratching of features that have been etched into the front side of wafer from photoresist on the hot plate, place a clean stainless steel plate on top of the hot plate and allow temperature to reach 95°C. Flat stainless steel plates can be ordered from McMaster Carr.

2. Dehydration bake the wafer for 2 minutes at 95°C.

3. Mount in specially designed spinner chuck that prevents the front side of the wafer from touching the surface of the chuck. This specially designed chuck consists of a circular stainless steel plate that has been machined on both sides. On the front side, three metal rods protrude from the surface near the edge of the plate, spaced 120° apart, at a radius that allows for a standard 4” wafer to be placed with a flat against one metal rod and the edges of the wafer flush against the other two metal rods. This allows for rotation of the wafer with the chuck. O-rings are placed on the metal rods against the surface of the chuck, and the wafer is placed on top of the O-rings such that the front side of the wafer does not touch the chuck. The bottom of the stainless steel plate has a region cut out that allows the plate to fit on top of a standard spinner chuck, and the plate is held on top of the standard spinner chuck via vacuum. An O-ring around the recessed region may be necessary to achieve vacuum seal to the
standard spinner chuck such that the plate rotates with the standard spinner chuck.

4. Soft bake at 95°C for 4 minutes and 15 seconds.

5. Use backside alignment capability (infrared (IR) light probes beneath wafer, special IR transparent chuck, IR-sensitive camera) on the MJB4 to align through-hole pattern to features etched on the front side of the wafer.

6. Expose for 6 cycles, 12 seconds cycle time, 10 seconds wait time in between cycles. This is overexposure of the pattern even for this thickness of photoresist, but this ensures that the through-hole pattern fully develops.

7. Post-exposure bake for 2 minutes at 95°C.

8. Wait 10 minutes.

9. Develop using AZ 300 MIF developer. Development usually takes 10 minutes, but can vary significantly.

10. Inspect under microscope to ensure through-holes are fully developed. Re-align, re-expose, and re-develop if necessary.

11. Use crystal bond to mount on carrier wafer for etching. Crystal bond can be applied by placing carrier wafer on hot plate set to 100°C and then applying the crystal bond as one would apply glue using a glue stick. Avoid applying crystal bond in areas of the carrier wafer that would interact with features etched into the front side of the fabrication wafer to avoid scratching these features.

12. Cover any pin-holes in the photoresist or other non-uniformities in the photore sist coverage with Kapton tape.

13. Etch with recipe 3 (200 µm trench etch) on the Samco 800 for 400 cycles.
14. Dismount from carrier wafer and visually inspect (hold up to the light) to ensure all through-holes have gone through the wafer. Dismount is achieved by placing on a hot plate at 100°C and using a razor blade to pry the two wafers apart. If all the through-holes have not gone through, re-mount and etch another 50 to 100 cycles of recipe 3 on the Samco 800.

15. Re-mount onto carrier wafer and remove the Kapton tape carefully to avoid breaking the fabrication wafer.

16. Dismount from carrier wafer.


19. Load wafer on ring with dicing tape.

20. Dice using appropriate recipe on ADT Dicing Saw with blade designed for cutting silicon.

21. Carefully remove chip from dicing tape after exposing tape to ultraviolet light to decrease adhesiveness of the tape.


Due to the long etch times and use of Kapton tape and crystal bond, the full Piranha clean is required before running experiments with chips fabricated using this method. The full Piranha clean process is detailed below.

**Piranha Clean**

1. Ensure no liquid solvent (acetone, isopropanol, ethanol) is present on chip. This process is performed with the chip loaded into a Fluoroware dipper basket.
2. Mix 96% sulfuric acid and 30% hydrogen peroxide in a 1:1 ratio in a glass beaker in sufficient quantity to completely immerse the chip.

3. Immerse the chip in solution for 15 minutes.

4. Rinse the chip under DI water and immerse the chip in large (>500 mL) Fluoroware beaker filled with DI water.

5. In a second large Fluoroware beaker, mix solution of 2% hydrofluoric (HF) acid by diluting 49% HF.

6. Remove the chip from the DI water beaker and immerse it in the 2% HF beaker for 1 minute.

7. Remove the chip from the 2% HF beaker and rinse under DI water. Immerse the chip in the Fluoroware beaker containing DI water.

8. Remove the dipper basket from the beaker and remove the chip from the dipper basket. Blow the chip dry under nitrogen.
Appendix C

Parts for Experimental Setup

This appendix details the parts necessary to set up and run the experiments described in Chapter 3 and Chapter 4.

C.1 Syringe Pump and Syringes

- Syringe pump: Fusion 200 High Precision Syringe Pump, Chemyx, Item Number 07200
- Syringes: 60 mL Disposable Syringe with Luer-Lok Tip; Fisher Scientific, Catalog No. 13-689-8

C.2 Parts for Manifold

O-Rings

- Inlet reservoirs: Metric Buna-N O-Ring 1 mm Width, 20 mm ID, McMaster Carr, Catalog No. 9262K618
- Outlet reservoirs: Metric Buna-N O-Ring 1 mm Width, 19 mm ID, McMaster Carr, Catalog No. 9262K617
Spring-Loaded Screws

- Spring: Type 302 Stainless Steel Compression Spring, 1.0” Length, .313” OD, .029” Wire Diameter, McMaster Carr, Catalog No. 1986K4
- Screw: 18-8 Stainless Steel Truss Head Slotted Machine Screw, 10-32 Thread, 2” Length, McMaster Carr, Catalog No. 19785A838
- Washer: Type 316 Stainless Steel Flat Washer, Number 10 Screw Size, 0.203” ID, 0.438” OD, McMaster Carr, Catalog No. 90107A011

Hydrophobic Filters

- Millex-FG Filter Unit, 0.2 μm, Hydrophobic PTFE, 25 mm, PVC, Ethylene Oxide Sterilized, Millipore, Catalog No. SLFG025LS

Fluidic Connections to Manifold

- Female Luer x 10-32 UNF Thread, Nylon, Cole Parmer, Catalog No. 45502-60

C.3 Tubing and Connectors

- Tubing: Peroxide-Cured Silicone Tubing, 1/8” o.d. x 1/16” i.d., Cole Parmer, Catalog No. EW-06411-62
- T-Valves: High-Pressure Stopcock, 1050 psi Max, 3-Way, Cole Parmer, Catalog No. EW-30526-34
- Female Barbed Luer: Female Luer x 1/16” Hose Barb Adapter, PP, Cole Parmer, Catalog No. EW-45508-00
- Male Barbed Luer: Male Luer x 1/16” Hose Barb, PP, Cole Parmer, Catalog No. EW-45518-22
Appendix D

Blood Protocols

This appendix details protocols used to obtain and handle blood used in the experiments in Chapter 3 and Chapter 4.

D.1 Blood Source

Blood is obtained in 8 mL ACD-coated tubes that are shipped overnight from Interstate Blood Bank (Memphis, TN). Blood is obtained from registered donors whose blood has cleared tests for common blood-borne viruses (HIV, Hepatitis B, etc.) within the previous six weeks, and only preliminary tests for these common blood-borne viruses are performed before the blood is shipped. Blood is shipped in an insulated container containing ice packs and is stored from the time of arrival to the time of experiment in a 4°C refrigerator.

D.2 Blood Preparation

Buffer for experiments involving blood is prepared as follows. We begin with phosphate buffered saline (PBS) from which the calcium and magnesium ions have been removed (Fisher Scientific, Catalog No. SH3002802). 1 g of bovine serum albumin
(BSA, Fraction V, Heat Shock Treated, Fisher Scientific, Catalog No. BP1600-100) is added for every 100 mL of PBS and is dissolved with gentle swirling. EDTA (Anhydrous, Crystalline, Sigma Aldrich, Catalog No. E6759-100G) is added to PBS such that the final concentration is at least 5 mM and not more than 7.5 mM. With a magnetic stir bar in the beaker, the EDTA typically takes at least 1 hour to dissolve. During this time, the buffer is also degassed using a vacuum pump. The solution is then filtered using a disposable filter unit with 0.2 µm pore size (Fisher Scientific, Catalog No. 09-741-04).

The water soluble form of PPACK dihydrochloride (Santa Cruz Biotechnology Inc., Catalog No. SC-201291A) is dissolved in water such that the final concentration is 0.2 mg/mL. The powder dissolves easily with gentle rocking of the centrifuge tube, and no magnetic stir bar is required. The solution is filtered through a Steriflip filter with 20 µm pore size (Millipore, Catalog No. SCNY00020) so that it does not need to be filtered before adding to the diluted blood before running the experiment.

Blood is diluted 1:3 in the buffer and filtered through a Steriflip filter with 20 µm pore size (Millipore, Catalog No. SCNY00020) to remove any clots that may have formed during transport. PPACK solution is then added such that the final concentration of PPACK is at least 40 µM and not more than 100 µM.

D.3 Counting Cells

Cells (and fluorescent polymer microspheres) are counted via haemocytometer (Figure D.1). The top of the haemocytometer is covered with a thin glass cover-slip to create a channel over a counting grid that is 100 µm deep. Approximately 10 µL of solution is injected into each of the loading ports such that the region over the counting grid is covered by fluid. A microscope is then used with either a 10X, 20X, or 60X objective lens to count cells over the counting grid region, which is enclosed
Figure D.1: The haemocytometer used to count cells in all of the experiments in Chapter 3 and Chapter 4 is shown above. The top of the haemocytometer is covered with a glass cover-slip as shown. Approximately 10 $\mu$L of solution is injected via micro-pipette into each of the loading ports. The haemocytometer is then placed on a microscope, and the cells are counted above a grid in the region inside the red rectangle. There are two counting grid regions inside the red rectangle, one for each loading port.

by the red rectangle in Figure D.1. In order to view fluorescently labelled cells, a microscope equipped with a mercury lamp is used.

Leukocytes that have been labelled with the fluorescent nucleic acid label, SYTO13, are shown above the central counting grid of the haemocytometer (enclosed by dotted red lines) in Figure D.2. A mercury lamp is used to excite fluorescence in the labelled leukocytes. The concentration of leukocytes in the solution being measured can be determined by counting the number of cells inside the dotted red square in Figure D.2, which has dimensions in the plane of 1 mm by 1 mm and a depth of 100 $\mu$m. The concentration of leukocytes in the solution is this number multiplied by $10^4$ per mL.
Leukocytes labelled with SYTO13 (fluorescent nucleic acid label) are shown above the central counting grid (enclosed by the dotted red square), which is illuminated using a mercury lamp. The central counting grid is 1 mm by 1 mm in the plane, and the depth of the channel bounded above by the glass cover-slip is 100 $\mu$m. This means the concentration of leukocytes in the solution that is being measured here is the number of leukocytes in the square enclosed by the dotted red line multiplied by $10^4$ per mL. Image courtesy of Yu Chen.

In measuring the concentration of leukocytes and erythrocytes in diluted whole blood, fluorescent labelling of leukocytes is used to distinguish between leukocytes and erythrocytes. Fluorescent labelling is also used to distinguish cancer cells from unlabelled leukocytes and erythrocytes. Significant dilution of the sample before it is loaded onto the haemocytometer is often required for measuring the concentration of erythrocytes. The concentration of erythrocytes can also be measured using counts within the smaller square grids within the large square enclosed by dotted red lines and multiplying by the appropriate number to get the total concentration. This is best performed with the 60X objective on the microscope. With the 60X objective, the erythrocytes can also be distinguished from the leukocytes by a very visible dot in the middle of the cell, resulting from the biconcave morphology of the erythrocytes.
Counting of leukocytes or cancer cells in blood can be made significantly easier with the removal of erythrocytes. In order to achieve this, we use erythrocyte lysis buffer (1X RBC Lysis Buffer, eBiosciences, Catalog No. 00-4333). 10 mL of lysis buffer is added to 1 mL of blood and incubated at room temperature for 10 minutes. The reaction is then stopped by adding 30 mL of 1X PBS. The solution is spun at 400g for 8 minutes, and the supernatant is removed via pipette. The pellet is then resuspended in PBS, and the cell count is performed via haemocytometer.
Appendix E

COMSOL Simulation Setup

In this appendix, we detail the COMSOL simulation setup used to perform the simulations in Chapter 4. All simulations are performed using the 2D laminar flow module. The following parameters should be set by clicking on the laminar flow tab in the model builder menu. The stationary form of the Navier-Stokes equation for incompressible fluids is used. The option to neglect the inertial term and the option to use the shallow channel approximation are both de-selected (not checked).

E.1 Schematic

The DLD array used in the simulations is shown in Figure E.1. The DLD array is 19 rows long (parallel to the flow direction) and 6 columns wide (perpendicular to the flow direction). Boundary conditions are no-slip at the walls and at the edges of the posts. Flow rate is set by uniform inflow velocity across the top of the array, and the bottom of the array is held uniformly at zero pressure. Array parameters are 60 μm asymmetric triangular posts, 40 μm gaps, and 1/20 tilt.

The uniform inflow velocity was related to the Reynolds number, Re, using a Bezier polygon across a gap in the middle of the array (the gap at the top of the red square shown in Figure E.1). By adding a 1D plot line graph and selecting this...
Figure E.1: The DLD array simulated using the 2D laminar flow module in COMSOL is 19 rows long (parallel to the flow direction) and 6 columns wide (perpendicular to the flow direction). Boundary conditions are no-slip at the walls and at the edges of the posts. Flow rate is set by uniform inflow velocity across the top of the array, and the bottom of the array is held uniformly at zero pressure. Array parameters are 60 µm asymmetric triangular posts, 40 µm gaps, and 1/20 tilt. The red square shows the 100 µm by 100 µm unit cell that is analyzed in Section E.3.

Bezier polygon, the velocity across the gap could be simulated for different normal inflow velocities. The default expression and units for the y-axis of the 1D plot line graph should be velocity magnitude (spf.U) and m/s, respectively. The velocity used in the Reynolds number calculation is 2/3 times the peak velocity in the gap. For Re of 20, the normal inflow velocity used is 0.224 m/s. The data from the 1D plot line
graph of the velocity in the gap can also be exported and opened in Matlab or Excel to determine the critical size using the method of D. Inglis et al. [35].

E.2 Shear Rate Simulations

Figure E.2: The shear rate at the surface of the post against which a cell can be compressed can be obtained using the 1D plot line graph function and selecting the line shown in red for the asymmetric triangular post. The y-axis data and expression must be changed to spf.sr and 1/s, respectively. The data from the 1D plot can then be exported as a text file that can be opened in Matlab or Excel for analysis.

The shear rate at the surface of the post against which the cell can be compressed can be calculated for a simulation that has already been run as follows. Add a 1D plot group to the results, and select line graph. The expression and units for the y-data should be set to spf.sr and 1/s, respectively. The edge or multiple edges of
the post can be selected simply by clicking on the edge, as shown in Figure E.2 and added to the selection section of the line graph options. Data from the 1D plot can be exported as a text file for analysis in Matlab or Excel by right-clicking on the line graph tab and adding plot data to export.

### E.3 Centripetal Acceleration Simulations

![Figure E.3: Magnified image of the region within and around the red square in Figure E.1 that is used in the fluid centripetal acceleration simulations in Chapter 4. The green arrows are from the arrow surface that allows extraction of the x and y components of the velocity at 1 µm intervals within the red square, which has dimensions of 100 µm by 100 µm. Data from the arrow surface is exported as a text file, and the centripetal acceleration of the fluid as it travels from one gap to the next in the direction of the flow is computed numerically in Matlab.](image)
We seek to evaluate the fluid centripetal acceleration (acceleration perpendicular to the streamlines) as fluid travels from one gap to the next in the direction of the flow. To do this, we need to know the \( x \) and \( y \) components of the velocity at evenly-spaced points in a grid covering a unit cell of the DLD array (100 \( \mu \text{m} \) by 100 \( \mu \text{m} \) in this case). One way to obtain this data is to add an arrow surface on top of the 2D velocity surface plot in COMSOL. This can be done by right-clicking on the velocity tab under the results section in the model builder menu, and clicking on arrow surface. The range of coordinates and the spacing within the range at which arrows will be placed can be specified in the menu for the arrow surface. We specify a 100 \( \mu \text{m} \) by 100 \( \mu \text{m} \) area with 1 \( \mu \text{m} \) spacing along each axis that stretches vertically from one gap to the next in the direction of the flow and is centered horizontally about the center of the first gap. This arrow surface is bounded by the red square in Figure E.3, and the larger arrows, indicating higher fluid velocity, are visible in green in the gap. The velocity data from this arrow surface is then exported as a text file for analysis in Matlab by right-clicking on the arrow surface tab and adding the plot data to export.
Bibliography


