Continuous microfluidic immunomagnetic cell separation

David W. Inglis a) 
Princeton Institute for the Science and Technology of Materials (PRISM), Department of Electrical Engineering, Princeton University, Princeton, New Jersey 08544

R. Riehn and R. H. Austin 
Department of Physics, Princeton University, Princeton, New Jersey 08544

J. C. Sturm 
Princeton Institute for the Science and Technology of Materials (PRISM), Department of Electrical Engineering, Princeton University, Princeton, New Jersey 08544

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We present a continuous-flow microfluidic device that enables cell by cell separation of cells selectively tagged with magnetic nanoparticles. The cells flow over an array of microfabricated magnetic stripes, which create a series of high magnetic field gradients that trap the magnetically labeled cells and alter their flow direction. The process was observed in real time using a low power microscope. The device has been demonstrated by the separation of leukocytes from whole human blood. © 2004 American Institute of Physics. [DOI: 10.1063/1.1823015]

Immunomagnetic cell separation, in which magnetic particles are selectively attached to cells, has become a common technique for biological cell isolations. Typically dextran beads impregnated with iron oxide (Fe₂O₃) and coated with antibodies are bound to a particular antibody receptor on the surface of a cell. In previous work, cells are either trapped on ferromagnetic fibers which induce locally high magnetic field gradients from a relatively uniform externally applied field and then washed out after the field is removed, or are deflected in a continuous flow by an externally applied gradient of magnetic field in a macroscopic device. In this letter, we present a magnetic cell separator that has two distinct advantages over current methods. First, it is planar and capable of being integrated with other components into a microfluidic total analysis system (μTAS). Further, because it is operating with continuous flow it can separate a large range of volumes without modification. The magnetic field gradients are created by micropatterns of a magnetic material. We also present a numerical analysis of the forces on cells labeled with superparamagnetic beads in such a device, and demonstrate the selective cell by cell separation of leukocytes from blood.

In our device, cells flow over a region of integrated microfabricated ferromagnetic stripes [Fig. 1(a)]. The magnetic field pattern from each stripe creates a magnetic trap that alters the movement of only those cells coated with superparamagnetic beads. Cells with sufficient quantity of beads become trapped over the magnetic stripes and flow only along the stripe direction, not parallel to the fluid flow (Fig. 2). Quantitatively the magnetic force \( \mathbf{F}_M \) on a particle with a magnetic dipole \( \mathbf{m} \) in a magnetic field \( \mathbf{H} \) is

\[
\mathbf{F}_M = \mu_0 \mu_B \nabla (\mathbf{m} \cdot \mathbf{H}).
\]

The beads used in our case are superparamagnetic, leading to a saturation of the magnetization at relatively low magnetic fields of 0.02 T, and negligible remanence. Thus \( \mathbf{m} \), for fields greater than 0.02 T, can be treated as a vector quantity of constant magnitude, but parallel to \( \mathbf{H} \), and the force becomes

\[
\mathbf{F}_M = S \mu_0 \mu_B \nabla H.
\]

where \( S \) is the number of bohr magnetons \( \mu_B \) per bead and \( H \) is the magnitude of the magnetic field. The magnetically labeled cell is subject to both a force \( \mathbf{F}_{Mz} \) in the vertical direction \([z \text{ as defined in Fig. 1(a)}]\) and a force \( \mathbf{F}_{Mx} \), which acts in the plane of the device \([x \text{ as defined in Fig. 1(a)}]\). This in plane force, \( \mathbf{F}_{M\perp} \) in Fig. 2., will be perpendicular to the stripe. \( \mathbf{F}_{Mz} \) pulls the magnetic bead towards the substrate while \( \mathbf{F}_{M\perp} \) acts to trap the cell in the area above the stripe. The force on a whole cell can be found by integrating the contributions of all magnetic beads on it.

For low Reynolds-number flow, the velocity \( \mathbf{v} \) of a tagged cell in the plane of the device for a magnetically labeled object with viscous drag factor \( \xi \) in a flow with bulk flow velocity \( \mathbf{v}_{\text{flow}} \) is \( \mathbf{v} = \mathbf{v}_{\text{flow}} + \mathbf{F}_M / \xi \). We can identify \( \xi \mathbf{v}_{\text{flow}} \) with a drag force \( \mathbf{F}_D \). If the component of the drag force perpendicular to the magnetic stripes \( \mathbf{F}_{D\perp} \) becomes less than the maximum of the magnetic force \( \mathbf{F}_{M\perp} \) (Fig. 2), then the cell will be trapped above the stripe because the drag force is not sufficient to push the particle past the stripe. When this happens, the vector component of the drag parallel to the stripe will push the cell along the stripe and it will flow at an angle \( \theta \) compared to the unlabeled particles. For stripes aligned at an angle \( \theta \) to the fluid flow the condition for trapping can be expressed as

\[
\xi \mathbf{v}_{\text{flow}} \sin(\theta) < \mathbf{F}_{M\perp} / \xi.
\]

The magnetic field gradient needed for trapping at a given bulk flow velocity becomes progressively smaller as \( \theta \) approaches 0°, but the net displacement then also approaches 0. On the other hand, for \( \theta \) approaching 90° the displacement becomes infinite but the magnetic force \( \mathbf{F}_{M\perp} \) has to fully balance \( \xi \mathbf{v}_{\text{flow}} \). (Note that even if Eq. (3) is not fulfilled, cells are slightly deflected as they travel over the stripe, but by an order of magnitude less than if they are trapped. A complete solution of the problem will be given elsewhere.)
The magnetic force on a cell in the lateral and vertical directions was modeled assuming 10-\mu m-wide, 2-\mu m-thick nickel stripes at a period of 35 \mu m with a uniform out-of-plane magnetization \( M = 0.08 \) T, and a uniform external field of 0.08 T. The nickel magnetization was chosen to be set at the level because it was measured experimentally in our laboratory.

The cells are attracted to the center of each magnetic stripe and 2.2 \mu m above the wires. The magnetic moment of 35 \mu m with a uniform out-of-plane 

FIG. 1. (a) Cross section of the magnetic separation device showing qualitative field lines from magnetized nickel structures embedded into a silicon substrate. For simulations, the nickel is 2 \mu m thick, 10 \mu m wide at period of 35 \mu m. (b) Calculated force in the \( x \) (lateral) direction on 10-\mu m-diam cells A (touching surface) and B (center is 10 \mu m above surface). Nickel magnetization \( M = 0.08 \) T, external field of \( H = 0.08 \) T, beads of dipole moment \( m = 1.8 \times 10^{-5} \mu_B \) and \( N = 5000 \) beads per cell was assumed. (c) Calculated force in the \( z \) (vertical) direction on cells A and B. The net force per period is negative for both cells.

The device consists of a silicon substrate containing recessed magnetic stripes. To fabricate the magnetic stripes, 10-\mu m-wide channels were etched at a period of 35 \mu m into a silicon substrate and then sputter coated with nickel. The wafer was then chemically mechanically polished to achieve a smooth surface with recessed nickel stripes, 2 \mu m thick, with their surfaces planar with the silicon surface [Fig. 1(a)]. A 75 \nm silicon dioxide layer was then deposited on the wafer surface by plasma-enhanced chemical vapor deposition. A polydimethylsiloxane layer containing 15-\mu m-deep microfluidic channels was then sealed to the substrate to direct the flow of cells. A uniform flow of fluid in one direction (at an angle \( \theta \) to the stripes) was imposed over the substrate by positive pressure combined with a row of microfluidic resistors at the top and bottom of the substrate. The cells were injected through a 40-\mu m-wide aperture in the middle of the microfluidic resistors. An external field \( H \) of up to 0.1 T was applied using a NbFeB magnet of dimensions 2.5 cm diameter by 1.9 cm.

To demonstrate operation of the device we labeled cells with CD45 Microbeads from Miltenyi Biotech (Auburn, CA). The beads are 20–100 nm in diameter and were estimated in the previous calculations to have an average magnetic moment of \( 1.8 \times 10^{-5} \mu_B \). The CD45 antigen is expressed to varying degrees on all leukocytes. Whole, undiluted human blood less than 24 h old was incubated in a heparin-coated container with a vital nucleic stain, Hoechst 33342 (Hoechst, Frankfurt, Germany) at 100 \mu g/ml, for 15 min at 37 °C, and then incubated with the microbeads for 15 min at 9°C. After this preparation we expect all leukocytes to fluoresce when illuminated with 488 nm light, and to be coated with at least 4000 beads. To reduce adhesion of the cells to the walls of the device, the device was soaked in buffer containing 1 g/ml F108 for at least 1 h. During operation, phosphate buffered saline containing 2 mM ethylenediaminetetraacetic acid (EDTA) and 2 \mu g/ml F108 was used.

In Fig. 3, we present a sequence of images that illustrates the difference in the path of a tagged leukocyte from that of untagged red blood cells. The fluid layer is 15 \mu m thick, the external field is 0.1 T, and the average flow speed is 110 \mu m/s, inferred from the red blood cells, which are not deflected. The path of the red blood cells can be seen from...
the single image which comprises the background of Fig. 3. Superimposed is the fluorescent image of a single leukocyte at intervals of 0.33 s. It has become trapped over the magnetic stripe and follows it, diverging from the untagged cells.

Figure 4 shows histograms of multiple leukocyte separations under three different conditions. The lateral positions of the cells are measured with respect to the center of the red blood cell stream (defined as $x=0$ μm) after traveling 1.15 mm in the device. The magnetic stripes are tilted at $\theta = 11^\circ$ to the fluid flow. In Fig. 4(a), the device has been demagnetized, no external magnetic field was applied, and the flow speed was 180 μm/s. No separation occurred, and all cells followed the fluid flow direction to exit the camera’s field of view at approximately $x=0$ μm. The width of the peak is identical to the width of the red blood cell stream.

In Fig. 4(b) an external magnetic field of 0.08 T was applied, and the flow speed was 240 μm/s. There is a peak in the histogram at $x=220$ μm, representing 40% of the distribution, which corresponds to cells becoming trapped and following magnetic stripes. Presumably the separation condition given in Eq. (3) has been met for these cells. We calculate the separation condition to be met for cells traveling slower than 240 μm/s for $\theta=11^\circ$, assuming 5000 beads on a 10 μm spherical cell on the bottom of the channel. The peak at approximately $x=30$ μm contains 60% of all leukocytes; these have been deflected only slightly. The bimodal distribution may be caused by variations in the number of beads attached to cells.

In Fig. 4(c) the external magnetic field is still at 0.08 T, but the flow speed has been increased fivefold to 1200 μm/s. In this case no separation has occurred and again the width of the distribution is equal to that of the red blood cell stream. The failure to separate occurs because the fluidic drag force which pushes the cells through each magnetic trap is now five times larger than in Fig. 4(b) while the magnetic force is unchanged.

We believe that with minor modifications such as increasing the thickness of the stripes and increasing the external field, the separation force can be increased by an order of magnitude, allowing for higher throughput and higher recovery.

In summary, we have presented a microfluidic implementation of a device which allows continuous cell by cell separation from a flow stream by selectively tagging with magnetic beads. The device uses microfabricated magnetic features to induce a lateral force on streaming tagged cells. The device has been used to separate leukocytes from whole blood and should be useful for integration into a microfluidic total analysis system ($\mu$TAS).

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