Isolation of cells from whole blood using shear-induced diffusion

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Extraction of cells of interest directly from whole blood is in high demand, yet extraordinary challenging due to the complex hemodynamics and hemorheology of the sample. Herein, we describe a new microfluidic platform that exploits the intrinsic complex properties of blood for continuous size-selective focusing and separation of cells directly from unprocessed whole blood. The novel system only requires routinely accessible saline solution to form a sandwiched fluid configuration and to initiate a strong effect of shear-induced diffusion of cells, which is coupled with fluid inertia for effective separation. Separations of beads and cells from whole blood have been successfully demonstrated with high efficiency (89.8%) at throughput of 6.75 mL/hr (10^6–10^7 cells/s) of whole blood. Rapid isolation of circulating tumor cells (CTCs) from peripheral blood sample of hepatocarcinoma patients is also shown as a proof of principle.

Isolation of cells directly from whole blood with minimal pretreatment is of high demand in liquid biopsy and cytopathology. Minimizing sample preparation not only reduces user intervention and increases reproducibility, but also diminishes labor involved and minimizes process time, as well as lowers testing cost\(^3,4\). This is especially vital in isolation of rare cells, such as circulating tumor cells (CTCs) from patient peripheral blood\(^8,9\), where loss of even a single cell can lead to substantial inaccuracies due to rarity of these cells\(^7,8\). However, direct isolation of target cells from whole blood is prohibitively challenging due to complex hemodynamics and hemorheology.

Many types of microfluidic cell sorting devices have been reported to tackle the challenge of rare cell isolation from blood\(^9\). External forces, including magnetic\(^10\), electric\(^11,12\), acoustic\(^13\) and optical\(^14\), have been used in active microfluidic systems for focusing and extraction of target cells from suspensions\(^15\). Meanwhile, passive systems that rely purely on channel geometry, carrier fluid and cell properties have received attention due to their simplicity and high throughput\(^15,16\). These include deterministic lateral displacement (DLD)\(^17,18\), pinched flow fractionation (PFF)\(^19,20\), hydrodynamic filtration\(^21,22\), inertial migration\(^23,24\), viscoelastic focusing\(^25,26\) and their combinations\(^27,28\). Additionally, biological affinity has been widely used to target specific cell surface markers and improve selectivity of microfluidic cell sorting\(^8,29\). While tremendous progress has been achieved, these platforms are not able to work with unprocessed whole blood and generally require a number of sample preparation steps, including lysis of red blood cells (RBCs), immunoselection, or sample dilution. Direct separation of cells from whole blood remains largely unexplored despite of the persistent interest.

The handful of microfluidic devices that can handle whole blood are based on principles of cell margination\(^30,31\), cross-flow filtration\(^32,33\), deterministic lateral displacement\(^34,35\) and immunoselection\(^36,37\). Additionally, cell deformability coupled with tapered post array\(^38\) and incorporation of ridges on the top wall of a rectangular channel\(^39\) have also been exploited to differentiate cell populations passively. However, these approaches suffer from low throughput (0.3–16.7 µL/min) or mediocre separation efficiency (e.g., 27% in continuous\(^39\) and 72% in discontinuous\(^40\) cross-flow devices), yet require sophisticated design (e.g., DLD\(^34,35\) and ridged channel\(^39\)), operational complexity\(^36,38\), or large device footprint. Hence, these existing approaches are far from practical, and the need for a simple device with high-performance (in terms of efficiency and throughput) still exists.

Herein, we report on a new passive approach for continuous separation from unprocessed whole blood. Our novel separation technique is based on shear-induced diffusion of particles in concentrated suspensions, and

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is for the first time applied to cell separation from whole blood in a straight, rectangular microfluidic channel (Fig. 1). With a flow of saline solution flanked by sample streams, bioparticles rapidly migrate out of side streams and focus into the cell-free center under the influence of shear-induced diffusion and fluid inertia. Such lateral migration is strongly dependent on cell size. We have successfully demonstrated focusing of polystyrene particles in whole blood within 10 mm downstream length, offering ~90% efficiency. More intriguingly, our throughput remains extremely high (10⁶-10⁷ cells/s or 6.75 mL/h), which surpasses the ultra-fast spiral inertial devices. As a proof-of-concept, we successfully separated HepG2 cells spiked in human blood (>89% efficiency) and also isolated CTCs directly from patient blood in our device.

**Results and Discussion**

**Human blood and passive focusing.** Before we discuss details of our device operation and results, we briefly review blood rheology and its implications on cell focusing. Human blood is a two-phase fluid with
various formed elements, exhibiting complex rheological properties. Approximately 45% volume of whole blood is comprised of blood cells, while the rest is plasma which is the aqueous solution with numerous proteins.40 The majority (~95%) of the suspended blood cells are RBCs. Density of normal whole blood is about 1056 kg/m³, which is close to that of water (1000 kg/m³) and is primarily determined by plasma and cells.40 Although plasma shows Newtonian behavior40,41, fluid dynamics of whole blood is non-Newtonian, mainly attributed to the dominant population of RBCs (10⁹ cells/mL) and their deformability40-43. The mutual interactions of RBCs and interplays with plasma give rise to the viscoelastic dynamics of whole blood. Viscoelasticity can be characterized in terms of Weissenberg number, described as \( \text{Wi} = \frac{\gamma}{\lambda} \), where \( \lambda \) is the characteristic relaxation time and \( \gamma \) is the shear rate.46,47 In a microchannel with height \( h \), \( \gamma = \frac{2U_f}{h} \), where \( U_f \) is the average flow velocity. Both viscosity and elasticity of blood response to fluid shear. At 37°C, its viscosity is about \( 4 \times 10^{-3} \text{Pa} \cdot \text{s} (4 \text{cP}) \) at high shear rate \( (\gamma > 100 \text{s}^{-1}) \) and lowering shear rate increases viscosity significantly which is known as the shear thinning effect.40

Viscoelasticity of whole blood suggests a possibility of preferential migration of cells within the complex fluid and subsequent cell separation. Particle (and cell) migration in viscoelastic fluids has been investigated both analytically48,49 and experimentally50-52. The migration in such flow is primarily subjected to elastic force \( (F_e) \) and inertial forces in sheared flows (Fig. 1a). The former is described in first and second normal stress differences \( (N_1 \text{ and } N_2) \)50,52 and the latter includes mainly shear-induced lift force \( (F_{w}) \), wall-induced lift force \( (F_{w}) \), and rotation-induced lift force \( (F_{r}) \). As illustrated in Fig. 1a, specific focusing positions emerge when inertial forces are dominant at moderate Reynolds number \( (Re = \rho U_f D_h/\mu) \), where \( \rho, D_h, \text{and } \mu \) are fluid density, channel hydraulic diameter and dynamic viscosity. On the other hand, particles migrate away from the high to low shear rate region undergoing elastic force (mainly \( N_1 \) since \( N_2 \) is significantly smaller)50,54. Recent works using viscoelastic fluid have shown successful focusing and separation of particulates including polystyrene spheres50,54, blood cells50,52,55 and even DNAs56 in microchannels.

Despite demonstrations in inertial57, elastic58, elasto-inertial59 or inertio-elastic60 systems, whole blood has rarely been directly used in these platforms due to its complex composition and nonlinear rheological properties. Inertial separation is only applicable to Newtonian fluid \((\text{Wi} = 0)\) and hardly working in whole blood. Focusing of cells using elastic force is ostensibly feasible considering the viscoelasticity of blood. However, the operational condition of negligible inertia \((Re \approx 0)\) imposes minimal flow rate \((\sim \mu l/\text{hr})\) and thus reduced shear rate50-52,56,57, which could completely ruin device performance. In whole blood, the RBCs aggregate in large numbers and form rouleaux at low shear rate, especially when \( \gamma < 100 \text{s}^{-1} \). The aggregation not only significantly increases the fluid viscosity but also minimizes the intercellular spacings40,45,58,59. Escalated viscosity and consequently larger drag force necessitates stronger elastic force for driving cells and without free space cells are difficult to move laterally. Thus, neither of the two approaches alone work for separation of whole blood.

Recent investigations on synergetic interaction of fluid inertia and viscoelasticity \((\text{Wi} > 0, Re > 0.1)\) suggest potential focusing of bioparticles within blood flow, considering the distinctly intriguing properties of RBCs and blood. Mildly increased inertial force (typically \( 0.1 < Re < 10 \)) could effectively eliminate the focusing positions near four corners in a square channel46,50. As shown in Fig. 1b, wall induced lift force \( (F_w) \) becomes sufficiently strong to repel particles inward and they subsequently focus in the channel axis under influence of the elastic force \( (F_e) \). The elevated shear rate \((>10^5)\) as indicated by \( Re \) helps disaggregate RBC rouleaux and thus reduce blood viscosity (complete dispersion of RBC aggregate occurs when \( \gamma > 200 \text{s}^{-1} \))59,60. Furthermore, deformation of RBCs and formation of RBC layers at high shear rate diminish effective volume fraction. This creates additional free space among cells40,45,58,60, reduces fluid viscosity and results in a Newtonian-like behavior, which leads to a possibility of lateral migration of cells subjected to both inertial and elastic forces.

**Particle migration in blood flow.** When particles are spiked into whole blood, no discernable migration takes place, despite the lateral migration expected due to the synergetic interaction of the inertial and elastic forces. Figure 2 illustrates a straight rectangular microchannel with 100 \( \mu \text{m} \times 50 \mu \text{m} \times 50 \mu \text{m} \) cross-section. Whole blood spiked with 18.7 \( \mu \text{m} \) diameter fluorescent particles was pumped at 225 \( \mu \text{l/}\text{min} \) (corresponding to \( Re = 50 \) in terms of Newtonian water flow54,61). Considering the Fåhræus -Lindqvist effect, the apparent viscosity of the blood streams and lower in the saline stream. These sharp gradients coupled with shear rate in the channel...
flow can lead to a strong shear-induced diffusion of particles and cells\(^{64}\), which may result in an effective and fast mixing of blood and saline. We first hypothesized that, if the flow rates of whole blood and saline buffer equals each other, such mixing may help to establish a similar rheological and flow conditions to those in the 2× diluted blood, permitting lateral migration and focusing of larger cells and particles. Indeed, our preliminary results in Fig. 1b show both rapid mixing of blood cells and successful focusing of 18.7 µm particles from whole blood (Hct = 45%) within 10 mm downstream length, which is even shorter than the \(L_f\) in 2× diluted blood.

Migration due to shear-induced diffusion. The observed fast and size-dependent focusing of particles can primarily be attributed to the collective effect of shear-induced diffusion and fluid inertia. Since blood cells were not fully mixed with saline solution while larger particles were focusing, simple dilution mixing (aforementioned hypothesis) cannot explain the observed phenomenon. In fact, particles in concentrated suspensions are known to undergo complex interactions that give rise to net lateral migration in shear flows\(^{64,65}\). This is termed shear-induced diffusion. In our device, the migration observed in the side streams of whole blood could be dominated by the shear-induced diffusion due to the unmatched concentrations and viscosities between adjacent fluid layers in the sheared flow. Leighton and Acrivos\(^{64}\) have shown that lateral drift of diffusion is mainly as a result of irreversible interaction among cells in a concentrated suspension. In the direction normal to the plane of shear, the effective diffusivity (\(D\)) of particles relates to shear rate, particle diameter, concentration (\(\phi\)) and viscosity and can be expressed as\(^{64}\),

\[
D = K \frac{\phi^2 \mu}{\mu} \frac{d\phi}{d\gamma} a^2
\]

where \(K\) is a dimensionless coefficient showing weak dependence on particle concentration (\(\phi\)). This is equivalent to volume fraction for particle suspension or hematocrit (Hct) for blood sample. The diffusion directs from blood layer (high concentration, viscosity and shear rate) to saline buffer (low concentration, viscosity and shear rate). As indicated by equation (1), the diffusivity (and thus lateral diffusion velocity) is strongly dependent on concentration and particle size as well as shear rate. Essentially, this size-selective shear-induced diffusion promotes particle lateral migration in our co-flow device, leading to a faster focusing as compared to single flow of two-fold diluted blood (Fig. 2).

We observed that the shear-induced diffusion did not effectively promote particle focusing in whole-blood only flow (Fig. 2). This could be mainly attributed to two factors that suppress the hydrodynamic diffusion. First, the velocity profile of whole blood flow in a microchannel is blunter than normal Poiseuille flow due to the shear thinning effect\(^{64}\). Such blunt profile produces relatively flat shear rate across the most of channel width. Secondly, no established viscosity and concentration gradients exist to amplify the effect of shear-induced diffusion in single whole-blood flow.

While concentration gradient enhances shear-induced diffusion, it also facilitates the general diffusion of cells into saline layer, which is known as Brownian motion. However, Brownian motion cannot be the driving factor in our device. On one hand, migration dynamics observed in our device differs from Brownian motion. According to Stokes-Einstein Equation (\(D_0 = k_BT/(6\pi\mu a)\), where \(k_B\) is Boltzmann’s constant and \(T\) is absolute temperature),
diffusivity $D_0$ of spherical particles due to Brownian motion is inversely proportional to particle diameter. Hence larger particles shall migrate slower than smaller ones, which contradicts our observation where 18.7 μm particles reached the centerline much faster than RBCs (6–8 μm). Further, particle motion in our flow configuration is in fact dominated hydrodynamically. Brownian motion is negligible as compared to shear-induced drift, as indicated by the dimensionless Péclet number ($Pe = \frac{\gamma a^2}{D_0}$). Brownian motion predominates particle behavior when $Pe < 1$. Péclet number within our device is on the order of $10^5$–$10^6$ depending on particle/cell size, which is excessively larger. As a result, the effect of the general diffusion exerts minimal effect on particle lateral migration in our case.

The other main driving force stems from fluid inertia. Particles flowing in Newtonian dilute suspensions are subjected to inertial forces when Reynolds number is not vanishing. These forces are strongly size-dependent: $F_i \sim a^3$, $F_s \sim a^2$, and $F_0 \sim a$. Wall-induced lift force ($F_w$) always deflects particles toward channel center and sheared-induced lift force ($F_s$) drives particles in the opposite direction. Rotational force ($F_r$) drives particles to their equilibrium positions when they are in proximity of walls. Inertial forces are the dominant driving factors that entrain particles and cells once they enter the Newtonian saline stream in our sandwiched flow. Although inertial forces also exist in the whole blood streams ($Re \approx 15$), they contribute little on particle lateral migration. While the high concentration of RBCs is favored for shear-induced diffusion, the strong cell-cell interaction hampers the effect of inertial forces, which is evidenced by the observation of no discernible migration in whole blood (Fig. 2).

In fact, cell-cell interaction is generally avoided in inertial focusing devices by limiting cell/particle volume fraction ($\phi$) to be less than 2%. Inertial effect diminishes sharply at higher $\phi$. Little effect of inertia in concentrated suspension was also pointed out by Madanshetty and Nadim in their experimental investigation of shear-induced diffusion ($\phi \approx 25\%$). Nevertheless rotational force might have some effect on migration due to the strong shear thinning effect displacing particles to sidewalls and to the formation of RBC layers acting as transient walls in whole blood. This is partly evidenced by the particle migration within 2×-diluted blood streams (Fig. 2).

Since whole blood is essentially non-Newtonian, it is possible that elastic force may affect the migration. Although blood plasma is believed to be Newtonian, the addition of deformable RBCs gives rise to non-linear behavior of whole blood. The elastic component of blood is too small to be measured when Hct < 20%. At higher hematocrit the elasticity of blood increases sharply with the third power of Hct. However, this viscoelasticity could still be very weak for whole blood considering typical value of Hct = 45%. Should the elastic effect influence particle migration, it could only contribute positively. On one hand, the elastic force is also strongly size-dependent ($F_e \sim a^3$). On the other hand, while shear-induced lift force counteracts both elastic and wall forces vertically in our low AR channel, forces in horizontal direction are essentially in harmony, directing from sidewalls to channel center (Fig. 1c).

In summary, particle migration within side streams of whole blood is dominated by shear-induced diffusion and inertial force is the primarily driving force in shear stream for particle focusing from whole blood. Owing to the strong size dependence of both shear-induced diffusion and inertial effect, larger particles in whole blood migrate much faster than smaller RBCs and achieve tight focusing in the middle of channel width before being significantly contaminated by the RBCs. These two effects work in tandem to transport larger particles out of the blood stream.

**Device optimization.** Successful demonstration of particle focusing and separation from whole blood led us to explore the dynamics of biparticle motion within our co-flow system to improve focusing quality and isolation performance (Fig. 3). According to the merged images and the intensity profiles in Fig. 1b, particles collected from the inner outlet (target outlet) accompanied considerable number of RBCs, downgrading the separation purity. Since RBC contamination primarily arises from strong shear-induced diffusion, careful tuning of RBC cross-stream diffusion can be an effective way to improve the separation quality. According to equation (1), we could reduce the RBC diffusion by decreasing either cell volume fraction (Hct) or shear rate as the diffusion coefficient is strongly dependent on cell concentration ($D = \phi \cdot \tau$) and scales with shear rate ($\tau = \gamma \cdot \mu$). While reduction of shear rate helps inhibit cell diffusion, it can significantly suppress inertial forces ($F_e \sim a^3$) which is one of the main driving force. In addition, both approaches could adversely affect the throughput. Thus, the former is preferred since it is more effective.

We first examined the lateral diffusion of RBCs downstream the channel at three cell volume fractions (Fig. 3). At Hct = 45%, the two blood streams expanded into the central saline layer drastically, leaving only ~8 μm spacing between them at channel end (10 mm downstream) when the total flow rate was 225 μL/min and flow rates were equal for blood ($Q_b$) and saline solution ($Q_s$). Two-fold dilution of blood in saline exhibited remarkable improvement in terms of confinement of RBCs as the spacing tripled (~25 μm). In both cases, blood streams expanded nonlinearly with respect to downstream length, fast in the first half and slowly in the other half as shown in Fig. 3a–c. The observed reduction of lateral diffusion is indeed implied in equation (1) since cell concentration continuously decreases as diffusion progressing downstream the channel and cells experience larger shear rate when they are closer to side walls at the first half channel. Excessive dilution of blood (200×) led to the minimized lateral diffusion and utmost confinement of RBCs within side streams indicated by the unchanged stream widths throughout the channel. The variations of initial stream fractions for the three cases were mainly resulting from different fluid viscosities of blood streams.

In addition to the evolution of stream fractions, blood dilution also modified the patterns of velocity and shear rate profiles at identical flow rates (Fig. 3b). For excessive dilution (e.g., 200×), the viscosity of side streams closely matched with saline solution and both the velocity and shear rate profiles was continuously and smoothly distributed within channel cross-section, forming a large rectangular region (approximately 65 μm × 15 μm) of low shear rate. While such shear rate pattern is preferred in inertial focusing where particles equilibrate in the two positions...
near walls, it is not desired for migration due to shear-induced diffusion, as particles would focus into a broad band instead of a tight streak. For less diluted blood, the increased viscosity difference between adjacent fluids rendered to the distorted velocity profile with elevated velocity in saline stream and broke the previous low shear rate region into three sections (Figs 3 and S2). For undiluted whole blood, the two side regions tended to vanish, with the lowest shear rate region restricted to a small blue spot (~13 µm in diameter) in the center. Such a small region is ideally preferred for tight focusing of bioparticles dominated by shear-induced diffusion which drives particles to region of low shear rate. Nevertheless, this focusing spot was bounded by a high shear rate blockade (the green annulus in Fig. 3b), which could otherwise prevent migration of bioparticles from the side streams theoretically. Considering the remarked concentration gradient across the blockade and the development of flow downstream, cells could still migrate into the saline stream. Thereafter, blood cells along with plasma rapidly diffuse into the saline layer as evidenced by the evolution of intensity profile of blood streams in Fig. 1b(v–vii). The fast-diffusive migration increases the viscosity of center layer, lowers the viscosity gradient across the fluid layers and thus reshapes the velocity and shear rate profiles downstream. Subsequently, inertial force, potentially also with elastic force, drives the bioparticles toward the channel center.

**Figure 3.** Dynamics of whole blood flow in the co-flow microfluidic systems. (a) Shear induced diffusion of blood cells leads to expansion of blood streams. (b) Modified velocity and shear rate profiles within channel cross-section at start position obtained from numerical simulation (ESI Group, ACE+). Dashed lines represent the interfaces of the two adjacent fluids. (c) Interface between blood and saline streams. Channel length was 10 mm. Each micrographs was generated from the standard deviation of 500 bright field images.
We aimed to achieve both tight focusing stream and broad central buffer region for high quality separation. The buffer region is indicated by the spacing between the two sharp boundary lines in Fig. 3c. While particles could achieve single-file focusing in whole blood, the separation spacing was minimal. The $2 \times$ dilution serves to significantly enlarge the spacing for easy separation. As dilution factor increased, the separation spacing expanded and meanwhile the focusing quality improved, as evidenced in the decrease of particle stream width (Fig. 4a). The measurements were taken right at the output trifurcation (end of expansion with width $\approx 192 \mu m$). Particle stream achieved tight focusing at 5 times dilution ($Hct = 9\%$) but the fractions of blood streams remained unstabilized until dilution factor reached $40 \times$ ($Hct = 1.1\%$, Figure S3), which is consistent with our previous measurements. Despite better focusing quality and larger separation spacing, the throughput markedly decreased at larger dilution factor. While longer channel may help to achieve full focus in this case, the throughput was reduced considerably. In accordance with the intensity profile, balanced flow rates ($\alpha = 1$) were beneficial as tight focusing is achieved in the channel center. Further increase of flow ratio only showed slight decrease in the spacing between the two blood streams, and the focusing quality slightly downgraded as well. We also tried to enhance the input flow rate to $\sim 312 \mu L/min$ and set $\alpha = 0.56$. We found the separation spacing was comparable to that when $Q = 225 \mu L/min$ and $\alpha = 1$, since higher shear rate at larger flow rate strengthened cell diffusion according to equation (1), and the particles were focused but not completely as shown in Fig. 4c. As a result, equal flow rates shall be the optimal in our short channel.

Separation of cells from whole blood. As a model system, we evaluated the performance of our co-flow system using polystyrene particles as surrogates for cells. We firstly spiked fluorescent particles into blood sample and measured the outcomes (Fig. 5). The optimal flow condition was chosen ($Q = 225 \mu L/min$ and $\alpha = 1$). First, 18.7 $\mu m$ diameter particles were spiked into a $2 \times$ diluted blood and injected into our channel. The central sharp stream (green) at output shows well focused particles (stream width = 24 $\mu m$) and the blood cells (RBCs) were confined within the side streams, giving $13 \mu m$ gap at each side of the particle stream and permitting an easy separation of cells from whole blood.
separation of larger targets. The corresponding intensity profiles also reveal a considerable improvement in both focusing quality and separation spacing for \(2 \times\) dilution as compared to undiluted whole blood (Fig. 1e). We noticed in both cases particles were rapidly entrained into two confluent streams at half channel length, and then merged into one stream centered channel width. The adverse effect of shear-induced diffusion on RBC contamination was well remedied, as uncovered by the intensity profiles of blood streaks (Fig. 5a). The micrographs of
the collected samples from inner and side outlets imply high separation efficiency and effective removal of RBCs in the target outlet.

Quantitative results of collected samples confirmed the high-profile performance (Fig. 5b). 94.4% of fluorescent particles were collected from the inner inlet and 97.6% of RBCs were removed. Blood cells in Fig. 5b refer to RBCs as the quantity of WBCs was thousand times lower than RBCs. Our system was highly reliable as indicated by the small error bars. Since separation efficiency is defined as the ratio of target count in central outlet over the total count of target in both outlets, it equals to the normalized count in Fig. 5b. Hence, the separation efficiency was 94.4% for 18.7 μm particles in a blood diluted two times. To compare the separation efficiency of whole blood processing, we also introduced the undiluted whole blood spiked with the same number of particles into the device. It turned out that the separation efficiency remained highly promising (~90%). But the rejection rate of RBCs markedly decreased (64.8%) for inner outlet (target outlet). This was expected according to the strong influence of shear-induced diffusion in whole blood. While these separation efficiencies are comparable to recently work by Geislinger et al.78 using pinched flow fractionation in viscoelastic fluid, our throughput overpowers their system (20 μl/hr) by a factor of 337. Our separation scheme also outperforms existing systems based on cell margination90,91 and cross-flow filtration32,33 and is simpler than DLD34,35 and other36,37 devices. While DLD device from Austin' group79,80 showed comparable efficiency (~86%) with very high flow rate, their system requires 5–20 times dilution and multiple additives to prevent clogging. Similar dilution factor is also necessary for inertial microfluidic devices regardless of various geometries78-80. Recently, Lee et al.36 reported separation of MCF-7 cells from whole blood in their contraction-expansion device, showing a slightly better efficiency. Nevertheless, their sample throughput (5μL/min) was much lower than ours (112.5μL/min).

Isolation of cells from clinical sample. To validate the fidelity of our system for cell separation from blood, we utilized human blood spiked with Hep G2 cells to demonstrate the extraction of circulating tumor cells which are typically larger than blood cells. Hep G2 cell line is a perpetual cell line which was derived from the liver tissue of a male with hepatocarcinoma. The size distribution was measured and plotted in Fig. 5b(iv). ~ 90% of Hep G2 cells were found larger than 15 μm. We spiked 1 ml Hep G2 cell suspension (initial concentration 1.7 × 10^6 cells/ml in PBS solution) into 1 ml human whole blood from venipuncture and carefully stirred to allow adequate mixing. The mixture was processed through our device at room temperature for ~14 min. The Hep G2 cells were pre-stained before mixing for easy counting. Our results confirmed the high separation efficiency for cells as well (89.1%), which is well-matched with size distribution of cells. The slightly decreased rejection rate of RBCs was primarily due to the variation of fluid resistances of outlet branch among different fabrication batches. Based on the concentrations measured in Fig. 5b(vi) and the collected volumes, we estimated the recovery rate of 82% (the ratio of target cells collected in central outlet over total cells injected). Considering the broad spectrum of Hep G2 cell size (10–45 μm) and the cutoff size (~15 μm) of our channel (in the case of inertial focusing)84, this recovery rate could still be very high and was comparable to that (85%) in the spiral devices38,39, where blood lysis was required and the separation efficiency was lower.

As a proof of concept, we also demonstrated isolation and identification of circulating tumor cells (CTCs) from peripheral blood donated by patients with hepatocarcinoma (TNM staging: T3N0M0). 1 ml blood was obtained through venipuncture and collected in Vacutainer tube coated with EDTA. Hepatocarcinoma cells (HCC) are typically larger than most of blood cells and the CTCs in these patient bloodstreams can be even fewer compared to other cancer types (e.g., breast cancer)35. Hence it is more challenging to separate CTCs for these patients. We utilized a post array right after the target outlet to immobilize suspected CTCs (Figure S3) and to minimize the chance of cell loss due to otherwise off-chip cell collection. After a single run, we successfully extracted and identified one CTC from 1 ml blood samples by onchip immunostaining. While white blood cells were CD45+ (Red), CTC was identified by pan-cytokine (CK, green). The identified cell was 23 μm in diameter (Fig. 5c). Two additional tests using the same protocol were carried out in two days and each detected one CK-CD45- cell, with diameters of 18 and 26 μm, respectively. Some leukocytes were also observed in the target outlet (Fig. 5c) due to their broad size variation. The cutoff size of our chip was about 15 μm and thus WBCs sized larger than that also entered the immobilization chamber. Observing these results, we believe our approach could be a powerful alternative in isolation or depletion of larger targets from highly concentrated sample.

Conclusions
In conclusion, we have successfully demonstrated a new scheme for continuous focusing and separation of bioparticles directly from human whole blood. Our system takes advantages of shear-induced diffusion that utilizes the intrinsic complex nature of blood, and couples it with inertial force for size-based separation. A separation efficiency of ~90% was achieved in whole blood spiked with microparticles at blood flow rate of 6.75 ml/hr, with a throughput up to 10^7 cells per second which is even higher than the ultra-fast spiral systems. In a 2 × diluted sample, higher separation efficiency (94.4%) for particles and 89.1% for Hep G2 cells were achieved. The majority (>96.6%) of RBCs were removed simultaneously at either case. Since our system only need routine saline solution as buffer, we could completely eliminate any sample preparation steps in our system even at 2 × dilution mode, using a saline-preloaded Vacutainer tube to collect clinical sample. No additives will be necessary and thus little contamination. Furthermore, we have successfully demonstrated the isolation of CTCs from clinical sample as a proof of prototype. We observed that the purity of CTCs obtained from our device needs to be further improved to meet the need of various applications. Our simple system permits size-selective isolation without external forces, easy setup and operation, and fast processing. We envision our approach could be a promising powerful technique for a wide range of diagnostic or prognostic applications.
Materials and Methods

Device fabrication. Microchannels were fabricated using standard soft lithography. Briefly, we utilized negative resist of SU-8 3025 (MicroChem Corp.) to pattern the microchannels on a 4" polished silicon wafer by conventional photolithography. Polydimethylsiloxane (PDMS, Dow Corning®) was casted on the wafer and peeled after 2 hour curing on 80 °C hotplate. Replicated channels in PDMS were bonded to 1" × 3" glass slides (Citotest Labware Manufacturing Co., Ltd) using surface plasma treatment (Harrick Plasma PDC-002). The inlet and outlet ports were punched manually using stainless flat head needles.

Experimental setup. Samples and buffer solutions were injected into the PDMS device with a syringe pump (NE-4002 ×, New Era Pump Systems, Inc.) to sustain stable flow rate. The loaded syringe was connected to 1/16" Peek tubings (IDEX Health & Science LLC) using proper fittings (IDEX Health & Science LLC) and then secured to the device inlets. Output of each outlet were collected in 1.5 mL centrifuge tubes. The outlet resistances were carefully tuned for different applications; the resistance ratios of central outlet over single side outlet were 0.597 and 1.29 for whole blood and 2 × diluted blood sample, respectively.

The flows of the fluorescently-labeled beads and cells in microchannels were visualized at successive downstream positions using an inverted fluorescence microscope (Leica DMI 4000) equipped with a high-speed EM-CCD camera (iXon ultra 897, Andor Technology Ltd). Analogous to microparticle streak velocimetry (µ-PSV), flowing particles generated streaks across each frame, and we analyzed fluorescent intensities and locations of these particle streaks. Fluorescent, bright-field and phase-contrast images were acquired during experiments. At least 500 frames were obtained and stacked using ImageJ® at each downstream position to improve image contrast. Fluorescence intensity linescans were used in quantitative analyses of focusing quality. Full width at half maximum (FWHM) was used to determine the widths of particle and blood streams in Figs 3 and 4.

Numerical simulations. Numerical model (Figure S1) was created in CFD-GEOM (ESI Group) with the same layout to the actual device. To reduce the calculation load, we ended the channel at 2 mm downstream length. Unstructured triangle mesh was used and the mesh cell size spanning from 0.02 to 2 µm, resulting in a total of 250,000 cells. Flow module in CFD-ACE+ (ESI Group) was used to model steady state flow. Total flow rate at 225 µL/min and flow rate ratio α = 1 were set, which were identical to our optimal experimental conditions. To model blood stream at different dilution factors, we adjusted dynamic viscosity from 4 to 1 mPa-s for the side sample streams and kept the central saline stream unchanged at 1 mPa-s. Densities of the fluids were rate at 225

Blood sample collection and preparation. Human peripheral blood was obtained from healthy and patient donors after informed consent and according to experimental protocols approved by institutional review board of Sir Run Run Shaw Hospital. Blood was collected in 2 mL Vacutainer (BD). Whole blood was diluted in saline solution to reach different hematocrit/cell concentrations (1 × 10^4, 5 × 10^4, 10 × 10^4, 20 × 10^4, 40 × 10^4, 80 × 10^4, and 200 × 10^4 dilution). Original hematocrit was determined as 45% from blood test report carried out in hospital. Fluorescent 18.7 µm diameter polystyrene particles (Polyscience, Inc.) were spiked into each sample at concentration of ~5.4 × 10^6 particles/mL.

Cell culture and staining. HepG2 cells provided by Professor Xiujun Cai at Zhejiang University were cultured in complete growth medium comprised of DMEM (Gibco Cat. No. 11995-065) with 10% (vol/vol) FBS (Gibco Cat. No. 10099-133) and 1% (vol/vol) Penicillin-Streptomycin (Gibco Cat. No. 15140-148) in 25 cm² flask. Cell passages were carried out when the confluence reached 70–80%. Briefly, cells were rinsed twice with PBS (phosphate-buffered saline) at room temperature after removal of culture media; then 1 mL Trypsin-EDTA solution (Gibco Cat. No. 25300-054) was added and kept in incubator for 4 min to digest the cell layer. Trypsinization was quenched by adding 1 mL complete growth media; cell suspension was transferred into 15 mL conical tube and centrifuged at 1000 rpm for 5 min; cell pellet was re-suspended in saline and cell concentration determined by haemocytometry. A 5 mL of diluted cell suspension at concentration of 5 × 10^4 cells/mL was aspirated into a new flask for new culture. Culture flask was kept in 37 °C incubator with humidified atmosphere of 5% CO₂ after spreading the cells evenly by rocking the flask back and forth.

Cells were stained using Hoechst 33258 stain (Molecular Probes) at a final concentration of 2 µg/mL. Cells were incubated in a humidified chamber at 37 °C and 5% CO₂ for 20–30 min before re-suspending to desired concentration. Blood cells and suspected CTCs were collected from the target outlet and stained immunoselectively. Cells were fixed using 4% formaldehyde for 30 min in exhaust hood and permeabilized by 0.2% Triton X-100 (Sigma-Aldrich) for 5 min. The permeabilized cells were washed in PBS buffer supplemented with 0.5% BSA. FITC-conjugated pan-cytokeratin (CK) antibody (1:100) and PE-Texas Red conjugated CD45 (1:100) antibody were used successively and allowed to incubate for 30 min. used DAPI for nuclei staining. The stained cells were washed with 1–2 mL of PBS buffer supplemented with 0.5% (wt/vol) BSA for observation under microscope.

Cell counting. Cell counting was implemented using hemocytometry. Samples collected from each outlet were gently stirred to allow random dispersion. Due to the high concentration of RBCs, we used fluorescently-activated counting, which counted target cells by fluorescent spots under microscope. A 100 × dilution of samples was performed before loading sample into hemocytometry for RBC counting due to the high concentration of erythrocytes. This dilution was not necessary for counting fluorescently labeled cells or particles. Images of the counting chamber were taken by CCD camera (bright field for RBCs and fluorescent field for labeled targets) and the counting was performed automatically using ImageJ® “Analyze particle” module with a proper threshold set to discriminated cells. Each sample was counted at least three times to improve accuracy.

Device performance was quantified using the following well-established terms,
### Data availability.
Most data are presented in the main text or the supplementary information. All data are available from the corresponding author on reasonable request.

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### Efficiency Calculation

\[
\text{Efficiency} = \frac{\text{Number of target cells from target outlet}}{\text{Total number of target cells from both outlets}}
\]

### RBC Rejection Rate Calculation

\[
\text{RBC rejection rate} = \frac{\text{Number of RBCs from waste outlet}}{\text{Total number of RBCs from both outlet}}
\]

### Recovery Rate Calculation

\[
\text{Recovery rate} = \frac{\text{Number of target cells from target outlet}}{\text{Number of target cells injected}}
\]
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Author Contributions
J.Z. developed the separation concept and designed the device. C.T. fabricated the devices. J.Z. and C.T. conducted the microfluidic experiments. C.T. and B.H. fulfilled the cell immunostaining steps. Y.L. cultured the cancer cell lines and prepared cell samples. Y.F. and X.L. collected the samples from patients and healthy donors. J.Z. and C.T. analyzed the data. J.Z. implemented the numerical modeling and wrote the manuscript. C.T., Y.L. and B.H. contributed to the method section. I.P. edited the manuscript. X.Y. provided lab resources.

Additional Information
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