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Microfluidic label-free bioprocessing of human reticulocytes from erythroid culture†

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In vitro erythroid cultures from human hematopoietic stem cells produce immature red blood cells (RBCs) called reticulocytes, which are important for RBCs production, and are widely used in scientific studies of malaria pathology, hematological diseases and protein translation. However, *in vitro* reticulocyte cultures contain expelled cell nuclei and erythroblasts as undesirable by-products and current purification methods such as density gradient centrifugation and fluorescence-activated cell sorting (FACS) are not optimal for integrated bioprocessing and downstream therapeutic applications. Developments in Dean flow fractionation (DFF) and deterministic lateral displacement (DLD) microfluidic sorting methods are ideal alternatives due to label-free size sorting, throughput scalability and low manufacturing cost. DFF sorting of reticulocytes from whole erythroid culture showed a 2.4-fold increase in cell recovery compared to FACS albeit with a lower purity; DLD sorting showed comparable cell recovery and purity with FACS using an inverse-L pillar structure to emphasize size and deformability sorting of reticulocytes. The viability and functional assurance of purified reticulocytes showed conserved cell deformability and supported the propagation of malaria parasites. Collectively, our study on label-free RBCs isolation represents a significant technical advancement towards developing *in vitro* generated viable human RBCs, opening opportunities for close-loop cell manufacturing, downstream therapeutic and research purposes.

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1. Introduction

Reticulocytes are nucleus-free, immature red blood cells (RBCs) originating from self-renewing hematopoietic stem and progenitor cells (HSPCs) within the bone marrow. Reticulocytes are important in various clinical prognosis and diagnosis applications,^{1–4} are commercially used to produce cell-free protein translation systems as they contain high levels of ribosome and are free from deoxyribonucleic acid (DNA),⁵ and are potential precursors for *in vitro* RBCs production.⁶ Reticulocytes are also the preferable invasion targets for malaria parasites, including *Plasmodium vivax* and *Plasmodium falciparum*,^{7–10} making them important for investigating malaria pathology. However, biological and medical advancements in these areas are limited by the availability of reticulocytes as they are transient in the human body and comprise only 0.5–2.5% of blood cells.^{11,12} The ability to manufacture reticulocytes *in vitro* will circumvent the need for large volume blood draws and laborious cell isolation processes.

In vitro erythroid cultures of reticulocytes have progressed significantly achieving more than 60% purity with >300 000 fold expansion.¹³ However, this process requires erythroblasts, to eject their nuclei resulting in a viscous



culture medium due to the accumulation of the expelled nuclei contaminants.⁶ Downstream applications such as cell-free protein translation, RBCs production and malaria invasion of reticulocytes require purified reticulocytes free from extra-cellular DNA. Thus, the expelled nuclei as well as erythroblasts must be removed from the culture, so as not to interfere with the downstream applications of reticulocytes.

Current reticulocyte purification protocols are mainly developed for cord blood and peripheral blood using, leukocyte reduction filters,^{14–16} density gradient centrifugation,^{17,18} and magnetic- and fluorescence-activated cell sorting (MACS; FACS).^{19–22} These methods have limited yield (ranging from 64–80% purity), throughput and scalability due to the need for discontinuous sample handling processes, cell-labeling and cost. Density gradient methods though well-established are challenged to purify these cultured cells as the expelled nuclei have varying density depending on the degree of degradation. While MACS has been employed to purify reticulocytes from blood samples based on the CD71 surface marker, this cannot be used in an erythroid culture where all the cells express CD71. Thus, an alternative sorting method that is continuous, efficient and low-cost is required to process erythroid cultures for the bulk production of reticulocytes.

Recent advances in microfluidic cell sorting systems have enabled label-free continuous sorting of cells based on biophysical parameters, such as size, deformability, electrical and acoustic properties.^{23,24} Given that erythroid cultures contain mainly reticulocytes, nucleated erythroblasts and expelled nuclei, which are different in size, shape and deformability,²⁵ we investigate here two microfluidic sorting methods for the isolation of human reticulocytes generated from HSPCs; namely Dean flow fractionation (DFF)^{26,27} and deterministic lateral displacement (DLD).²⁸ Both DFF and DLD are passive cell size dependent sorting methods and have shown high efficiency in blood cell sorting.^{27,29–31} These methods do not require cell labelling, are low cost and are scalable using modular and parallel device setups.^{32–35} DFF systems dominate in ultra-high processing rates of mL min⁻¹,³⁶ while DLD systems enable precision in cell shape, size and deformability discrimination.^{30,37}

In this study, we employed a trapezoidal cross-section design in DFF sorting to enable circulating purification of reticulocytes integrated with an automated continuous culture platform. For DLD sorting, high reticulocyte purity was achieved using the sorting properties of an inverse L-shape pillar structure that allows selective isolation of the larger and deformable reticulocytes from the smaller and rigid nuclei in the erythroid culture. We show that both DFF and DLD could sort 100 million cells significantly faster at 1.5 to 2 hours compared to FACS, which is considered as the gold-standard method that requires 4 hours. DFF sorting of reticulocytes yields close to 72.3% recovery with 80.0% purity, while DLD sorting has a lower yield of 27.6% recovery with 97.4% purity. The DLD sorting is comparable to FACS system for which the sorting recovery is 30.6% with more than 99.7% purity.

Finally, we demonstrated the retention of the biological function of DFF and DLD sorted reticulocytes by assessing the cell deformability by micropipette assay and overall viability by malaria parasite infectivity, which require purified and healthy host cells for their growth. Collectively, our results represent technical advancements towards human RBCs manufacturing and downstream label-free applications for therapeutic, diagnostic and research purposes.

2. Results and discussion

2.1. Cell composition of the *in vitro* erythroid culture from CD34⁺ HSPCs

Stem cells have the potential to multiply continuously and are the key to achieving large quantities for the manufacturing of blood.^{38,39} Large-scale synchronized axenic culture system based on peripheral CD34⁺ HSPCs has the highest yield and enucleation rate reported so far.¹³ The entire erythroid culture consists of an expansion of HSPCs, followed by 5 differentiation phases with varying cell density, cytokines and growth factors resulting in a 14 700-fold expansion over 23 days (Fig. S1, ESI[†]). A day 12 intermediary cryogenic storage protocol was developed to reduce the culture time for terminal differentiation down to 11 days post cell thaw. This eliminates the need for continuous 23 day blood manufacturing. Throughout the culture, synchronous differentiation of erythroid cells was monitored by flow cytometry-based detection of surface markers (Fig. S1a, ESI[†]) and by Wright–Giemsa staining to observe the cell morphology (Fig. S1f–k, ESI[†]). The final *in vitro* culture contains approximately 66.4% reticulocytes, 19.2% nuclei and 13.6% erythroblasts (Fig. 1a and b). The corresponding enucleation rate is more than 80%.

To evaluate the biological functionality of the culture-derived reticulocytes, the entire crude erythroid culture was tested for its ability to sustain *Plasmodium falciparum* infection. *P. falciparum* is the most virulent of human malaria species that can propagate in reticulocytes and mature RBCs.⁴⁰ However, using the crude erythroid culture, we observed that the parasitemia failed to increase (Fig. S2a, ESI[†]). Giemsa-stained thin smears of the *P. falciparum* culture revealed intact expelled nuclei (Fig. S2b, ESI[†]) that eventually became degraded (Fig. S2c, ESI[†]). There was no apparent parasite expansion in the crude culture as compared to a parallel infection using RBCs from the peripheral blood. A likely explanation is that the high density of degrading extracellular DNA in the erythroid culture trap and interfere with invasion of the newly released parasites. Therefore, the elimination of nuclei and erythroblasts are essential for the downstream usage of the erythroid culture and in this study we have compared reticulocyte purification by FACS and two innovative microfluidics sorting using DFF and DLD techniques.

2.2. FACS sorting of reticulocytes

FACS is the gold standard for cell sorting due to well-established protocols and high specificity. The reticulocytes



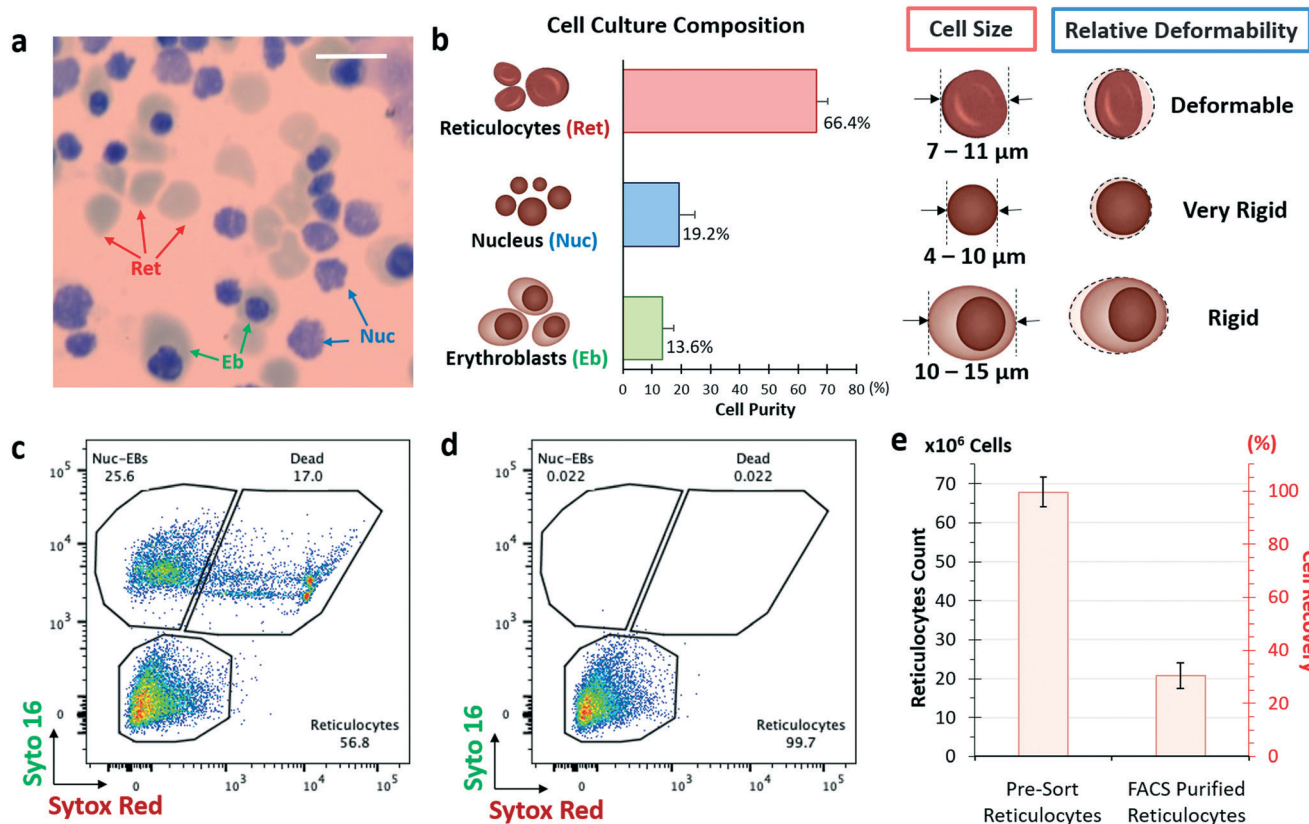


Fig. 1 Large-scale *in vitro* erythroid culture from CD34⁺ HSPCs. a) Schematic of the 6 phases *in vitro* erythroid culture with a representative image of Giemsa stain of day 23 erythroid culture. Indicated with arrows are reticulocytes (Ret), expelled nuclei (Nuc) and nucleated erythroblasts (Eb). Scale bar denotes 20 μm . b) Shows day 23 cell culture composition of reticulocytes (Ret), nuclei (Nuc) and erythroblasts (Eb) ($n = 3$; \pm SD) with schematics of the biophysical properties of each cell types, which are size and deformability. c) Shows representative FACS pseudo-color density plots of day 23-erythroid culture. Staining of SYTO 16 vs. SYTOX red dyes, where indication corresponds to reticulocytes, nuclei and erythroblasts. Double positive cells are dying and dead cells. d) Shows the re-analysis of reticulocytes purified by FACS sorting. e) both reticulocyte count and recovery were analysed for pre-sorted reticulocytes and FACS purified samples ($n = 3$ and error bars of mean represent SD).

isolation from the erythroid culture was conducted by SYTO16 (permeable DNA stain) and SYTOX red (non-permeable to live intact cells) staining to differentiate reticulocytes, nucleated particles and dead cells (Fig. 1c).^{41,42} It is noted that the CD71 marker, which is used to isolate reticulocytes from whole blood, is not used as a label here as all erythroblasts and reticulocytes express CD71 (Fig. S1e, ESI[†]). Nuclei and erythroblasts were further differentiated based on granularity and cell size (Fig. S3, ESI[†]).

100 million cells from the day 23 erythroid culture with reticulocyte count of 67.9 ± 3.7 million cells at a purity of 66.4% were subjected to purification by FACS sorting (BD ARIA II FACS machine). After FACS sorting, the reticulocyte count dropped to 20.8 ± 1.8 million, which accounted to a cell recovery of 30.6% (Fig. 1e) and reticulocyte purity of 99.7% (Fig. 1d). The low cell recovery may be due to cell death from high-shear stress sorting events by squeezing 6000–10000 cells per second in a 100 μm nozzle. Sample preparation including centrifugation, staining and washing steps could also contribute to the significant cell loss. Additionally, to ensure >99% reticulocyte purity, a tight gating is required to exclude overlapping events which may

also remove reticulocytes to the waste channel (Fig. S4[†]). The total time taken to sort 100 million cells was more than 4 hours (excluding the time required for pre-sorting processing).

2.3. DFF microfluidic design for the enrichment of reticulocytes

Others and we have previously reported various applications of DFF microfluidics for size-based cell separation.^{26,27,43–47} Cells flowing in spiral DFF microchannel experience lift force and centrifugal acceleration-induced Dean drag force, which results in the formation of two Dean vortices on the cross-sectional plane of the channel. Interplay of the lift and Dean drag forces cause cells to migrate laterally while flowing through the microchannel, and stay at a lateral position where the net force is zero (Fig. 2a and b). The magnitudes of the lift and Dean drag forces experienced by a cell correlate with the cell size, thus cells with different sizes focus at different lateral positions, enabling size-based cell separation.^{26,46} A trapezoidal cross-section spiral microfluidic chip with one inlet and two outlets was employed for DFF-



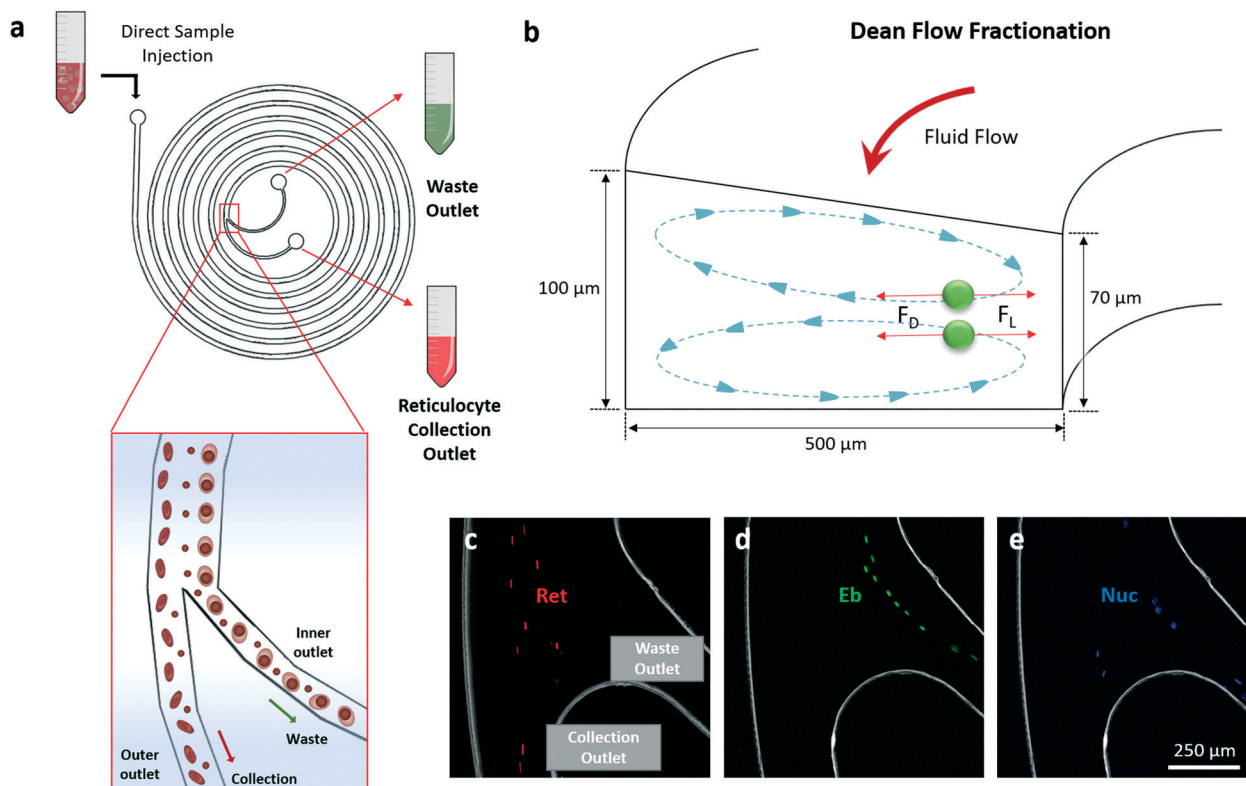


Fig. 2 DFF microfluidics for reticulocytes sorting. a) Design of a DFF microfluidic chip with a single inlet injection of erythroid culture samples and two bifurcated outlets for cells sorting to collection and waste outlets. b) Schematics of secondary fluid flow on the cross-sectional plane of the DFF device. The Dean vortices created inside the spiral channel would align the particles based on size by the balance of drag force (F_D) and lift force (F_L). The schematics are not drawn to scale. Purified samples of reticulocytes (Ret), erythroblasts (Eb) and nuclei (Nuc) are used to characterise the sorting dynamics in DFF microfluidics at 0.8 mL min^{-1} sorting flow rate as shown in (c)–(e) respectively. Scale bar shows length of $250 \mu\text{m}$.

based purification of the reticulocytes from erythroblasts and free nuclei (Fig. 2a and b). This DFF microfluidic chip design was developed from a similar design described in our previous study on the separation of blood cells.²⁷ The widths of the two outlets were modified to be equal ($250 \mu\text{m}$) in order to accommodate the different composition of cell population in this study. The use of trapezoidal cross-section geometry could improve separation resolution as compared to conventional rectangular shape.⁴⁸ It also removes the need of sheath buffer at the inlets to synchronize the initial position of cells,^{27,47} which greatly enhances the volumetric sorting rate and enables low concentration sorting. To demonstrate the performance of the chip, each population of reticulocytes, erythroblasts and nuclei were first prepared by FACS sorting. The FACS-purified reticulocytes, erythroblasts and nuclei were then pumped into the DFF microfluidic chip respectively at the same flow rate (0.8 mL min^{-1}), and their focusing positions at the outlet branching point were recorded using a high-speed camera (Movie S1, ESI†). Results showed that majority of reticulocytes were sorted to the collection outlet (Fig. 2c). Almost all erythroblasts were sorted to the waste outlet (Fig. 2d), demonstrating efficient separation between the erythroblasts and reticulocytes. Nuclei had approximately equal chance to be sorted into the two outlets due to wider size distribution compared to erythroblasts and reticulocytes (Fig. 2e).

2.3.1. Circulated sorting of reticulocytes using DFF. A circulation system setup for DFF-based cell sorting to enhance reticulocyte purity was designed (Fig. 3a). Fig. 3a shows the use of a spinner flask for homogeneous suspension of the erythroid culture and infusion of the cells into a DFF microfluidic chip for continuous sorting. The waste outlet is connected to a waste reservoir and the purified reticulocytes are diverted back to the bulk cell culture in the spinner flask through the collection outlet (Fig. 3b and c).

Employing real-time high-speed video recording, we observed that large amounts of cells were sorted into waste outlet at the beginning of the circulation, and drastically decreased after an hour (Fig. 3d–f). The circulated sorting was completed after 2 hours, when no more cells were sorted into the waste outlet (Fig. 3f). FACS analysis of the erythroid culture before (pre-sort) and after (purified) sorting, and the cells collected from the waste outlet (waste) was performed to quantify the cell compositions (Fig. S5, ESI†). In the purified cell population, reticulocytes were enriched from 66.4% to 80.0% after 2 hour recirculated sorting, erythroblast was reduced from 13.6% to 2.7% (Fig. 3g). From an initial 100 million cells (containing 67 million reticulocytes), 49 ± 1.1 million reticulocytes were recovered after purification. This corresponded to a recovery of 72.3% of the reticulocytes with a purity of 80.0% (Fig. 3h).



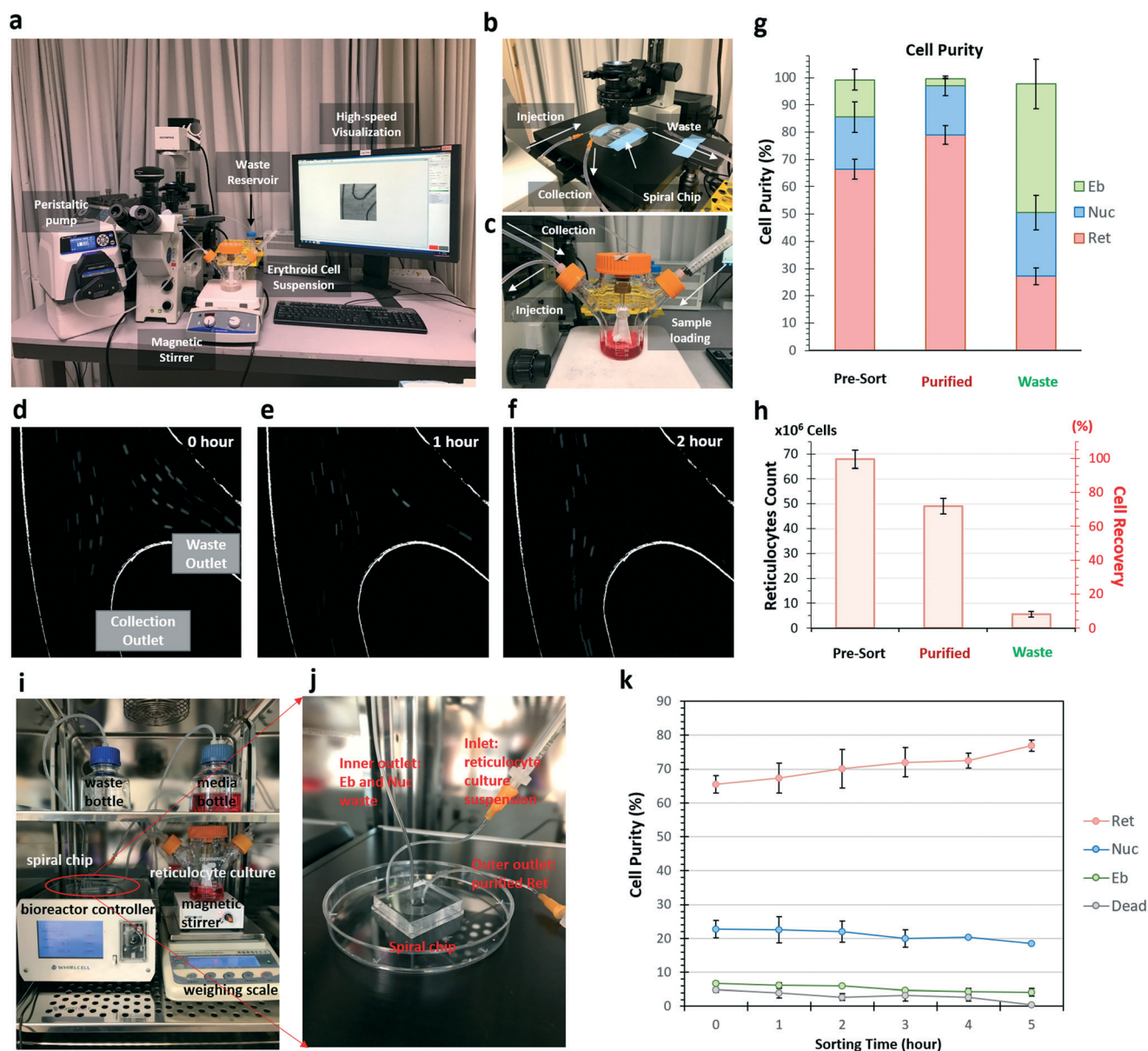


Fig. 3 Circulated sorting of reticulocytes with DFF and application in perfusion bioreactor. a) Setup of a benchtop circulated sorting system. b) DFF microfluidics on the microscope stage showing the fluid flow of samples injection, collection and waste outlets. c) A spinner flask containing the erythroid cell suspension. The sample injection inlet and the collection outlet, which redirects purified reticulocytes back to the spinner flask, are indicated. Representative images of sorting paths at outer outlet captured by high-speed camera during the circulated sorting. Real-time monitoring of cell collection at waste outlet at 0 hour d), 1 hour e) and 2 hours f) of circulation. g) Percentage of reticulocytes (red), nuclei (blue) and erythroblasts (green) from g) in the pre-sort, purified and waste populations. Results represent mean values (\pm SD) from three independent experiments ($n = 3$). h) Absolute number of reticulocytes and recovery percentages from outer (purified) and inner (waste) outlets are counted after 2 hours of DFF microfluidic sorting ($n = 3$; bars represent mean, and error bars represent SD). i) A continuous perfusion bioreactor setup for reticulocyte purification inside CO_2 incubator. j) A close view of the DFF microfluidic chip coupled with the bioreactor system for perfusion and purification. k) Change of various cell compositions in the culture system during continuous perfusion.

2.3.2. DFF for continuous purification of reticulocytes in culture. It is worth noting that the high-throughput feature of DFF sorting can be achieved at a sorting concentration, which resembles the culture concentration (2–4 million cells per mL), suggesting the potential of coupling this technology with a perfusion bioreactor for continuous processing. To demonstrate this concept, we set up a lab-scale perfusion culture system with a 100 mL working volume and 200

million unpurified cells inside a standard CO_2 incubator (Fig. 3i). Similar to the circulating sorting system, the cell suspension in the culture vessel was pumped into the DFF chip to remove erythroblasts and nuclei to a waste bottle, and recycle purified reticulocytes to the culture vessel (Fig. 3j). A perfusion bioreactor controller machine (WhirlCell Inc., Singapore) was used to automatically feed fresh media into the culture vessel to maintain the working volume. We



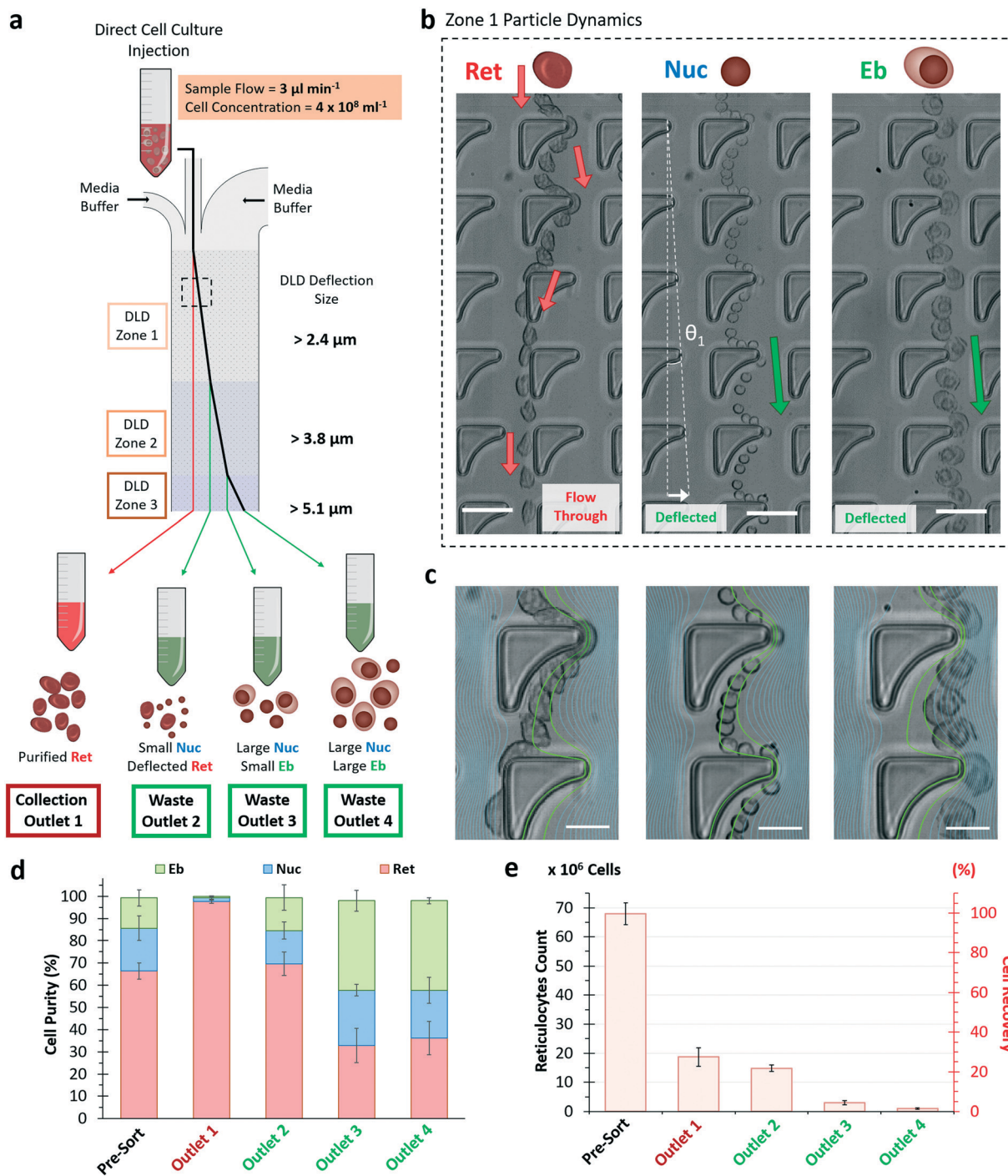


Fig. 4 Design of DLD device for microfluidic sorting of *in vitro* erythroid culture. **a**) Schematics of the sorting performance and expected outlet distribution of cell types from the erythroid culture. The concentrated sample (400 million cells per mL) with a volume of 250 μ L is injected to the device at a rate of 3 μ L per min. The four device outlets are designated as reticulocyte collection outlet 1, waste outlet 2, 3 and 4 respectively. **b**) Particle dynamic motions of Ret, Nuc and Eb within the inverse-L pillar structure in zone 1 of the DLD device shown by the image sequence overlay. The angle of deflection is denoted as $\theta_1 = 1.0^\circ$ and red (Ret) and green (Nuc and Eb) arrows indicated the paths of each cells. The white arrow denotes the lateral displacement length for the given section. Scale bar, 20 μ m. **c**) Simulated fluid streamlines are overlaid on the magnified image for comparison of the particle migration trajectories within inverse-L DLD structures for Ret, Nuc and Eb respectively (from left to right). The green streamlines denote the two closest streamline boundaries to the pillar protruded edge. The scale bar shows 10 μ m. **d**) Percentage of the cell distribution for pre-sort, FACS sorted and cells collected from DLD outlets 1–4 for Ret (red), Nuc (grey) and Eb (white). DLD outlet 1 for reticulocyte collection is highlighted in blue. **e**) Reticulocyte number from pre-sort sample and collected after FACS and DLD device with $n = 3$ and error bars represent SD.



operated the perfusion culture for 5 hours, and sampled cell suspensions from the bioreactor vessel every hour for FACS analysis. Results showed continuous increase of reticulocyte purity and decrease of erythroblasts, nuclei, as well as dead cells in the culture (Fig. 3k). At the end of the 5 hour perfusion culture, 76% reticulocyte purity and 82% recovery were achieved from 200 million unpurified cells (Fig. S5, ESI†). This performance was similar to results obtained with circulating sorting of 100 million cells for 2 hours (Fig. 3g and h). This lab-scale system indicated the possibility of processing large volume (liters) of cell culture using multiplexed DFF modules.^{49–51}

2.4. High purity reticulocyte isolation using the DLD device with inverse-L pillars

Purification of reticulocytes using size alone is challenging due to the large over-lapping size distribution of the expelled nuclei and reticulocytes (Fig. 1b). This is due to the fact that degradation of the expelled nuclei results in their size expansion from its compact 4 μm to a size similar to reticulocytes (Fig. 1a). Guźniczak *et al.* have investigated both mechanical and size properties of HSPCs and showed that deformability is a vital parameter to differentiate between reticulocytes and nuclei.²⁵

To sort reticulocytes from the erythroid culture, we utilized an inverse-L DLD microfluidic technique to rapidly isolate reticulocytes with high purity using size and deformability sorting (Fig. 4a and S6, ESI†). Previously, inverse-L DLD structures showed distinct sorting profiles of 3.0 μm rigid spherical beads relative to bi-concave shaped RBCs in a fixed DLD device critical cut-off size (D_c) of 3.3 μm (see eqn (1)).³⁷ The earlier study suggests effective sorting based on bi-concave RBC shape however the current DLD study on erythroid culture eliminates the shape effect as reticulocytes are deformable with a spherical morphology.

Thus, the designed DLD microfluidic chip has 3 inlet ports with a narrow sample injection stream sandwiched by 2 media buffers (Fig. S6, ESI†). As the sample enters the device, the biological particles flow through zones 1 to 3 of D_c 2.4, 3.8 and 5.1 μm respectively. Large particles (more than specified D_c) deflect laterally into waste outlets 2 to 4, while small particles (less than all D_c) are collected from outlet 1. This design was based on the hypothesis that reticulocytes deform significantly and have an apparent size smaller than the smallest D_c of 2.4 μm . It is also important to note that the D_c is a DLD apparent cut-off cell size, a parameter of the dynamic cell size and does not reflect the actual physical size under static measurement.

To understand and visualize the individual sorting dynamics of the sample components of Ret, Nuc and Eb, a flow of 50 million cells per mL sample was observed in zone 1 ($D_c = 2.4 \mu\text{m}$) (Fig. 4b). Reticulocytes were more deformable and interacted with the surrounding fluids *via* compression, transverse and rotational motion (Movie S2, ESI†). This complex hydrodynamic interaction resulted in reduced

deflection in DLD. They slipped past the pillar and flowed towards the collection outlet as though its overall net flow direction was unhindered as shown in the red arrows (Fig. 4b). Meanwhile, nuclei and erythroblasts both experienced a deflected flow path. Thus, the inverse L-shape amplifies differences between rigid bodies and deformable bodies of similar size.

To elucidate the effects of deformability in cell migration within the inverse-L structures, streamline profiles of the inverse-L structure was overlaid onto a magnified view of the cell trajectory (Fig. 4c). The green streamlines denote the closest two simulated streamline boundaries to the protrusion of the inverse-L structure where sorting of DLD occurs. These green streamlines cut through the center of the Ret and Nuc as they flow past the inverse-L protrusion. The Ret being deformable and pliant, wraps around the apex of the protrusion with a horizontal width comparable to the nuclei diameter (see ESI† Fig. S7). As the cells flow past the protrusion of the inverse-L, reticulocytes deform, relax back into an elliptical shape and slip out of the streamline boundaries while nuclei maintain their spherical shape and trajectory within the green streamline boundaries. Erythroblasts being larger and more rigid follow the streamline trajectory and DLD pillar gradient.

We directly injected the crude erythroid culture samples at a flow velocity of 3 μL per minute using 250 μL of cells at 400 million cells per mL (equivalent to ~3–4% hematocrit) into the DLD device, which processed 100 million cells in 1.5 hours. This concentration is approximately 15 \times blood dilution and various DLD works have shown effective sorting performance even up to whole blood density.^{52,53} Increasing the cell concentration to 400 million cells per mL show that the cells are still relatively sparse ensuring DLD pillar interactions for effective cell sorting (Movie S3, ESI†).

The final purity of reticulocytes at outlet 1 was 97.4% with 27.6% cell recovery with little variation from independent culture samples (Fig. 4d and e). Outlet 2, 3 and 4 showed significant reduction in reticulocyte purity at 64.9%, 23.4% and 22.3% respectively (Fig. S8†). Outlet 2 consisted of a relatively large number of reticulocytes, which suggest that some of these cells could have an apparent size larger than 2.4 μm . Most of erythroblasts were sorted into outlets 3 and 4.

2.4.1. Size versus deformability sorting performance of DLD pillars. Reticulocyte deformability-based sorting is attributed to fluid flow properties induced by inverse-L shaped DLD pillars. Fig. 5 shows the sorting comparison of conventional circle DLD pillars and inverse-L DLD pillars using FACS purified populations of reticulocytes and nuclei (Fig. S9, ESI†) at a sample processing flow velocity of 3 $\mu\text{L min}^{-1}$. Conventionally, a circle DLD pillar structure enables precise size-based sorting of particles and have shown deformability based sorting of RBCs.³⁰ However, the influence of deformability-based reticulocyte sorting from rigid nuclei is less pronounced for the circle DLD post compared to the inverse-L pillars (Fig. 5) at this flow velocity.



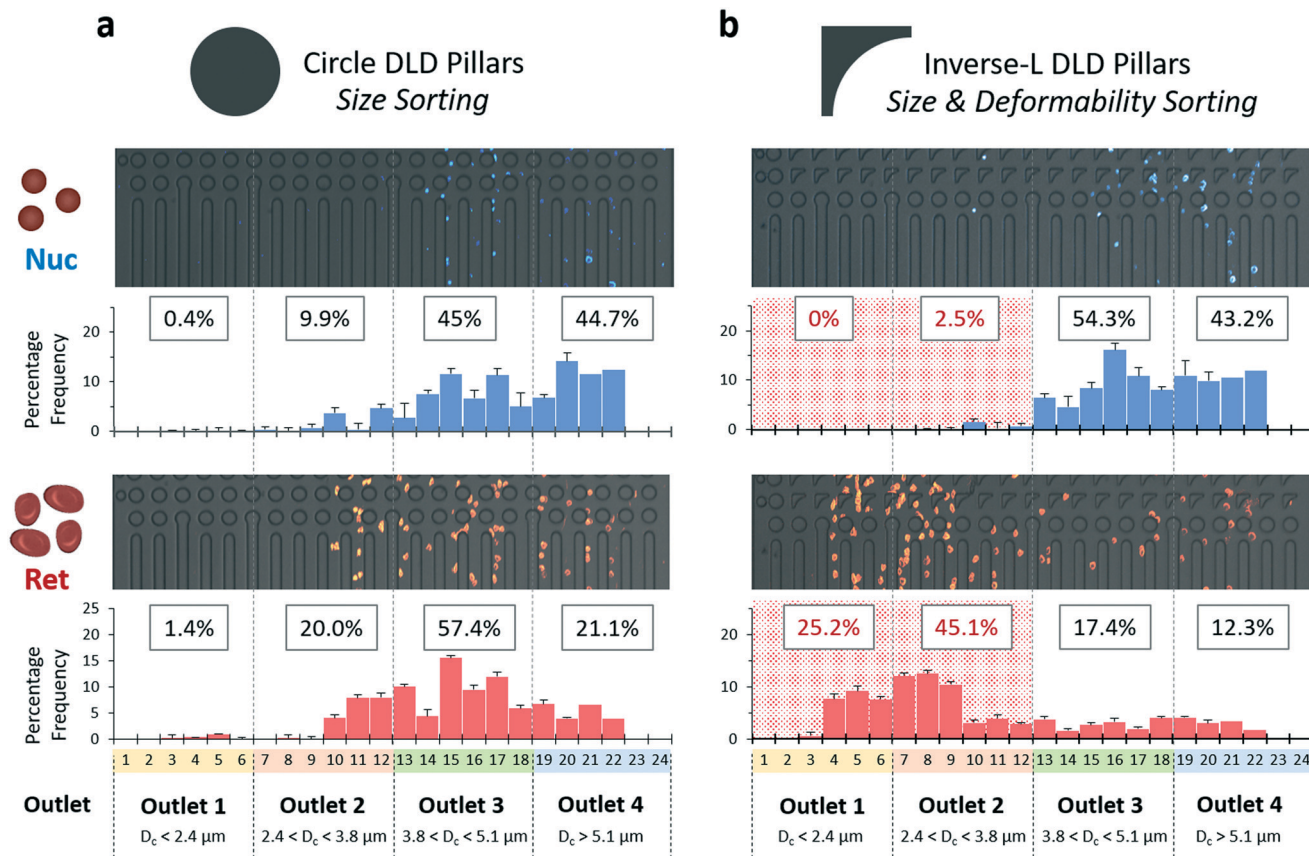


Fig. 5 Sorting characterization of cells by DLD devices. The sorting characteristics of both a) circle DLD pillars and b) inverse-L DLD pillars can be seen respectively. A FACS purified sample of nuclei (Nuc) and reticulocytes (Ret) from the erythroid culture were infused at a particle concentration of 2 million per mL to evaluate the sorting performance of the different pillar structures. The 24 sub-channels correspond to 4 outlets (1, 2, 3 and 4 respectively with increasing D_c size bins). The image for the output region is an image stack from high-speed camera footages with pseudo-coloring. The frequency histogram of the output region was tabulated by particle counting for each sub-channel. Histogram bin and total particle count per histogram range from 150 to 600.

It is possible that at certain optimal flow velocities, deformability-based sorting of reticulocytes from nuclei in circle pillars will occur. However, based on the current throughputs and processing timescale, the inverse-L shape DLD pillars showed significant differences in sorting of deformable cells *versus* rigid cells. Approximately 70.3% of reticulocytes tested were collected from outlets 1 and 2, while 97.5% of nuclei were deflected into outlets 3 and 4 (Fig. 5b). Hence, the inverse-L DLD device amplified the sorting sensitivity of deformable reticulocytes from rigid and smaller nuclei, which resulted in a collection of very pure reticulocytes.

2.4.2. Enhancement of sorting purity by adjustment of cell input concentrations. As demonstrated by Vernekar *et al.*,⁵⁴ there is a trade-off between throughput and sorting efficacy in DLD separation. Holm *et al.* simulated and experimentally tested the effects of hematocrit on sorting efficiencies and found that hematocrit above 10% would significantly impact effective size sorting in DLD.⁵³ Here, we explore reducing the hematocrit from ~3–4% to ~0.4% and were able to obtain 98.8% reticulocytes purity an improvement of 1% by diluting the cell culture concentration 8 times from 400 million cells per mL to 50 million cells per mL (Fig. S10, ESI[†]). Therefore,

the trade-off between sorting throughput and efficiency is observed. However, unlike FACS, throughput of DLD microfluidics can be easily scaled through parallelized and stacked microfluidic channels.

2.5. Quality assessment of microfluidic purified reticulocytes

Reticulocytes mature into erythrocytes whose deformability is vital for blood microcirculation. Reduction in deformability will result in poor blood circulation or even more serious clogging and clotting issues within capillaries. Thus, we compared the deformability of reticulocytes before and after sorting to evaluate their quality for downstream RBCs production using micropipette assay. It was found that all sorting methods (FACS, DFF and DLD) did not change reticulocyte cell deformability significantly (Fig. 6a). The mean shear modulus for all categories was seen to fluctuate around 6–10 pN μm^{-1} .

Finally, we verified the viability and functional quality of *in vitro* generated and purified reticulocytes using malaria parasites. The infection of reticulocytes by malaria parasites is an important indicator of their robustness and biological



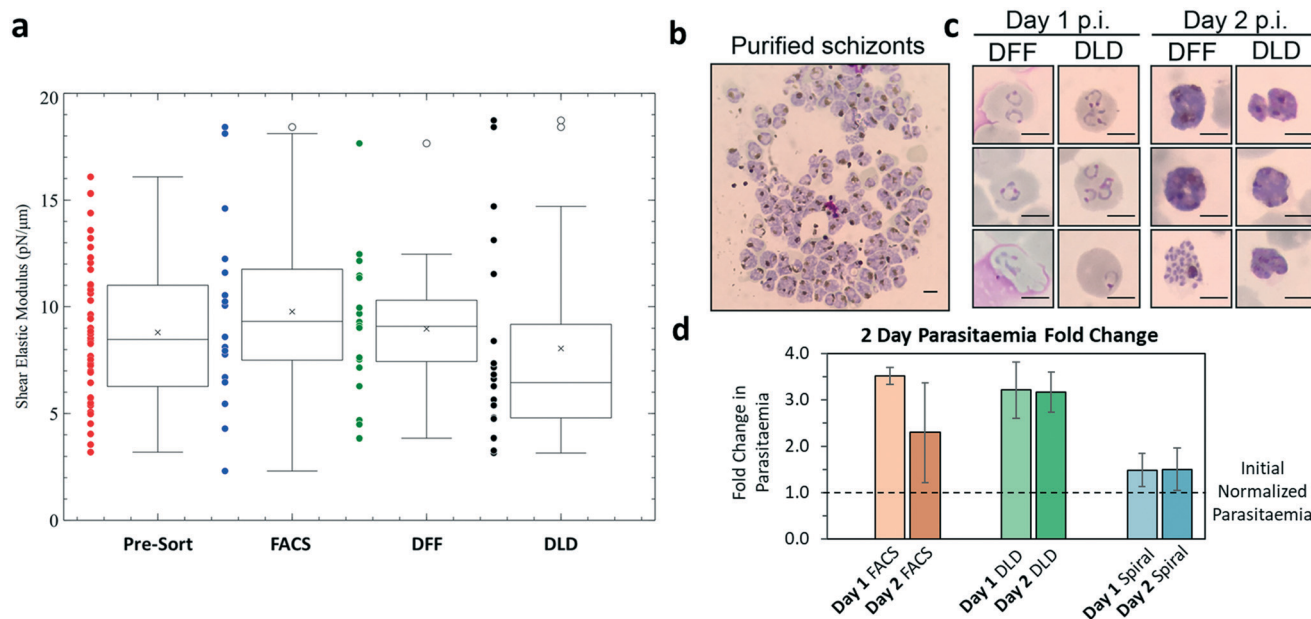


Fig. 6 Validation of biological functionality of microfluidic purified reticulocytes *via* cell deformability measurement and infection of malaria parasites, *Plasmodium falciparum* 3D7 strain. **a**) The shear elastic modulus was analysed by micropipette aspiration of reticulocytes from day 23 crude erythroid culture (red, pre-sort), from purified reticulocytes by FACS (blue), DFF (green) and DLD (red). The mean shear modulus for all categories can be seen to fluctuate around 6–10 pN μm^{-1} ($n = 30$, $\pm\text{SD}$). **b**) Giemsa stain of purified Pf3D7 schizonts (>99% purity) that were inoculated to test the biological functionality of purified reticulocytes. Schizonts were purified using magnetic column. **c**) 5 million purified reticulocytes were inoculated with 0.5% initial parasitemia. Shown are images of infected reticulocytes with Pf3D7 rings on day 1 post infection (p.i.) and schizonts by day 2 p.i. Some cells are infected by more than one parasite, which is due to preferential invasion of reticulocytes by *P. falciparum*. Day 1 and day 2 cultures showed changes in the *P. falciparum* infected cells with day 1 showing ring stage and day 2 showing schizont stage. **d**) Parasitemia fold change quantified by microscopic examination of Giemsa-stained culture smear. The change in parasitemia over two days is compared with initial seeded parasitemia density of a normalized value of 1. Results represent mean values ($\pm\text{SD}$) from two independent experiments. Parasitemia initial concentration of infected cells was loaded with reticulocytes. The graph shows successful invasion and development of parasites for the increase in parasitemia for FACS, DLD and DFF sorted cells.

function, as the parasite development require membrane integrity for efficient nutrient uptake from the medium as well as metabolites available directly from the infected reticulocytes. The microfluidic-sorted reticulocytes were infected by *Plasmodium falciparum* 3D7 strain (Fig. 6b). Fig. 6c shows the ring-stage formation on day 1 and mature schizont development on day 2 within the sorted cells from both DFF and DLD after inoculation with 0.5% parasitemia. The specific invasion of parasites into sorted reticulocytes was further confirmed using CFSE labeling of host cells prior to infection (Fig. S13, ESI[†]). Both DLD and FACS sorted samples showed a 3 fold increase in parasitemia (Fig. 6d) while DFF sorted cells showed a smaller increase of 50%. The compromised infection efficiency of DFF-purified culture as compared to the FACS and DLD-purified cultures was likely due to the residual nuclei, which might inhibit parasite invasion (Fig. S2, ESI[†]). Together, these assays confirmed that biophysical marker-based microfluidic purification of reticulocytes from *in vitro* erythroid culture resulted in viable reticulocytes with normal biophysical property and bio-functionality.

3. Outlook and conclusions

Currently, FACS purification of fluorescently labeled cells is the common method of choice for most biologists, however

the technique has many issues such as cell yield, speed, and cost. These issues are even more problematic when one considers large-scale processing, as in cell therapy for example. As an alternative to utilizing cell surface markers, biophysical markers are relatively new and are gaining momentum as unique biomarker for cell identification. Masaeli *et al.*, Nawaz *et al.* and Nitta *et al.* have developed label-free biophysical cytometers using image analysis and microfluidic structures to measure cell deformation and size albeit with limited sorting throughput of 100 cells per s.^{23,55,56} DFF devices are scalable and have been shown to process samples at mL per minute, while DLD have been shown to sort cell at high cell concentrations. In this study, we further developed the DFF and DLD technology for the label-free biophysical marker sorting of reticulocytes from erythroid culture and compared the outcome with the current gold standard of FACS sorting. A trapezoidal cross-section DFF chip was applied to simultaneously perform reticulocyte purification and media exchange in a continuous perfusion culture, while an inverse-L shape DLD device showed label-free biophysical isolation of reticulocytes with comparable yield and purity to FACS.

Compared with conventional cell purification by FACS, the DFF and DLD microfluidic purification methods were label-free and high-throughput (Table 1). 1 million cells can be



Table 1 Comparison of different methods for reticulocyte purification

	FACS	DLD	DFF re-circulation
Recovery (cell no.)	30.6% (20.8 million)	27.6% (18.7 million)	72.3% (49.1 million)
Purity	99.7%	97.4%	80.0%
Throughput	~8000 cells per s	~20 000 cells per s	~16 000 cells per s
Processing time (total cell no.)	4 hours (10^8 cells)	1.5 hours (10^8 cells)	2 hours (10^8 cells)
Cell density	1 million per mL	400 million per mL	4 million per mL
System cost	>\$500 000 (FACS)	\$1500–\$15 000 (syringe pump + Mold)	\$1500–\$15 000 (peristaltic pump + Mold)
Cost per sort	~\$35 (nuclear staining)	~\$10 PDMS Chip	~\$10 PDMS Chip
Sorting method	Cell nucleus staining	Cell size and deformability	Cell size

processed within approximately 1 minute with a single DFF microfluidic chip, in contrast to at least 10–15 minutes by FACS. Note that a single pass through DFF will completely purify 100 million cells within 30 min at a sorting concentration of 4 million per mL, although in this study a circulated sorting approach was applied to progressively purify reticulocytes in a continuous perfusion culture setup. DFF only achieved 80.0% reticulocyte purity, as compared to greater than 97% with DLD and FACS. This is mainly due to its lower efficiency in separating free nuclei from reticulocytes. Narrower DFF chip designs hold a potential to achieve better separation resolution, however, the sorting speed and reticulocyte recovery would be inevitably compromised.

The performance of DLD and FACS are comparable with close to 97.4% purity and 27.6% recovery. Unlike FACS where cells experience high shear stress resulting in potential cell death, DLD is advantageous as the sorting is relatively gentle on the cells. Reticulocytes tethered to expelled nuclei during the enucleation process are observed to remain intact during the sorting process (Fig. S12 and Movie S4, ESI†). During the infusion process, the settling of cells in the syringe and tubing results in loss during infusion. Loss of cells during the insertion of tubing into the DLD chip is inevitable to ensure a bubble-free tubing connection. Notwithstanding, DLD sorting using inverse-L pillars shows engineering precision and control over the selection of biological cells using biophysical properties of size and deformability.

In this study, we have demonstrated the robustness of both DFF and DLD techniques for processing more than 100 million cells in short-term operations (few hours). The lateral position of cells in DFF systems are a function of flow rate, thus fluctuations in flow rates would inevitably compromise the robustness of cell separation. The DLD microfluidic method is less sensitive to changing flow rates as the sorting is performed within Reynolds number <1 where effects of inertial forces is negligible. By employing a circulating DFF sorting in the bioreactor setup, we minimise the effects of fluid flow fluctuation raised from peristaltic pump pulsation (Fig. 3). The use of circulated sorting in long-term culture impose an extremely high requirement on the microfluidic chip's resistance to clogging. The large cross-sectional area and high sorting speed of DFF guarantee very low chance of clogging. In contrast, the same operational modality may not

be ideal for DLD, as cell aggregates and debris may gradually clog up the dense pillars. Although DLD has a higher chance of clogging than DFF in general, the clogging in DLD usually takes place in a gradual manner, and would not impair the sorting until certain threshold of the pillar structures are clogged; while when clogging happens in DFF especially at the outlet bifurcation, the cell separation is likely to be affected immediately.

Combined, these results demonstrate that microfluidic biophysical sorting methods, if properly optimized, are able to provide comparable level of sorting purity, but with significantly higher efficiency and lower cost and technical difficulty. The potential applications of these biophysical cell-sorting technologies using label-free biophysical markers extend beyond reticulocyte purification into realms of disease diagnostics, bacterial pathogenicity and stem cell therapy. With the emerging large-scale applications such as cell therapy production, these are important advantages over more convention techniques such as FACS and MACS. The different features of DFF and DLD also make them suitable for different applications respectively. In scale-up manufacturing processes, DFF should be firstly considered for its' high volumetric throughput. In scenarios where cell purity is pivotal, DLD's high precision feature holds great advantage.

In conclusion we have generated reticulocytes from *in vitro* erythroid culture derived from HSPCs, where the end-stage erythroid cultures contain a complex mixture of progenitor cells, expelled nuclei and reticulocytes. To effectively purify the reticulocytes, we developed two label-free and scalable microfluidic techniques based on cell size and deformability. The DFF system, employing a trapezoidal cross-section design to enhance separation resolution and throughput, retained a high 72.3% recovery of reticulocytes while achieving 80.0% purity in circulated perfusion cultures. The DLD system on the other hand achieved 97.4% purity with a lower 27.6% recovery by adopting an inverse-L shape pillar structure, which selectively deflect rigid particles into waste outlets and retains the softer and deformable reticulocytes. Compared to the current gold standard FACS sorting, these two microfluidic sorting technologies showed great potential for large scale processing and purification of erythroid cultures. This study sheds light on large-scale high-purity label-free reticulocyte production for therapeutic and research purposes.



4. Experimental section

Ethics statement

All the experimental methods were carried out in accordance with the approved guidelines. The blood donation from healthy individuals was approved by Institutional Review Board (IRB) of National University of Singapore (NUS-IRB Ref Code: 10-285). Written approval consents were obtained prior to blood collection.

In vitro erythroid culture

A total of 1 million adult peripheral blood CD34⁺ HSPCs (Fred Hutchinson Cancer Center), mobilized by granulocyte-colony stimulating factor were thawed gradually. Cells were then placed in expansion medium (day 0–5), differentiation phase 1 (day 5–9) and 2 (day 9–12) media, as reagents and cell density are described previously.¹³ Aliquots of 50 million erythroid cells were made on day 12 and cell pellets (centrifuged at 600 g, 3 minutes) were gently resuspended in 50 μ L of FBS. 550 μ L of freezing solution, 15% DMSO and 75% FBS, were added gently and samples were frozen gradually at -80 $^{\circ}$ C, then transferred to liquid nitrogen. When reticulocytes were desired, cryo-preserved erythroblasts were gradually thawed as for HSPCs. Thawed cells were then placed in differentiation phase 3 (day 12–16), 4 (day 16–20) and 5 (day 20–23) media.¹³ At the end of differentiation stage 5, each batch of aliquoted 50 million *in vitro* erythroid culture resulted on average in 735 million erythroid cells. The final expansion for the entire erythroid culture is approximately 14 700 fold, which could yield close to 15 billion erythroid cells from 1 million HSPCs.

Flow cytometry based sorting of reticulocytes

On day 23 of erythroid culture, cells were stained in a batch of 50 million cells and maximum total of 200 million cells at once for sorting by BD ARIA II (BD Biosciences). Erythroid cells were centrifuged at 600 g for 3 minutes, washed in 1 mL of Hank's balanced salt solution (Thermo Fisher Scientific, 14025076), then stained in 1 mL of HBSS with 18 μ L of SYTOX red (Thermo Fisher Scientific, S34859) and 22.5 μ L of 1:100 SYTO16 (Thermo Fisher Scientific, S7578) for 10 minutes at room temperature in the dark.^{41,42} Cells were washed once in 1 mL of HBSS, resuspended in 1 mL HBSS, then filtered through 70 μ m pre-separation filters (Miltenyi Biotec, 130-095-823) into 5 mL Polystyrene round bottom tube (Corning, 352058). Filters were rinsed with additional 200 μ L of HBSS and tubes were capped and stored on ice until the acquisition. Reticulocytes were sorted with 100 μ m nozzle, 2- or 4-way purity precision mode and 6000 to 10 000 events per second. Reticulocytes were collected into 15 mL Falcon tubes coated with 3 mL of FBS to minimize the hemolysis. Sorted cells were centrifuged at 600 g for 5 minutes and resuspended in plain IMDM (Thermo Fisher Scientific, 12440) and stored at 4 $^{\circ}$ C until downstream experiments. For microscopic categorization of Heilmeyer

stages, 1 μ L of packed reticulocytes were mixed with 1 μ L of new methylene blue dye (Streck, 340235-6) for 15 minutes at room temperature. Thin-blood smears were prepared on a glass slide, dried immediately and quantified on a microscope (ESI† Fig. S11).

Design and fabrication of spiral DFF microfluidic chip

The mold for producing the DFF microfluidic chip was designed with SolidWorks software (Dassault Systèmes, France), and fabricated from a piece of 50 \times 50 \times 8 mm aluminum plate using micro-milling technology (Whits Technologies, Singapore). The chip possessed a DFF microchannel with one inlet and two outlets (Fig. 1). The two outlets were named as inner and outer outlets respectively, with the inner outlet being closer to the center of spiral. The spiral DFF microchannel had 8 loops with diameter decreasing from 26.2 mm to 10.6 mm (from inlet to outlets). The cross-section of the microchannel had a trapezoidal shape with 500 μ m width, 70 μ m inner height, and 100 μ m outer height. The ratio of inner to outer outlet width was approximately 1:1 (250 μ m each).

The top piece of the DFF microfluidic chip was produced from the mold using standard soft lithography. Liquid phase polydimethylsiloxane (PDMS) elastomer (10:1 mixture of base and curing agent. Sylgard 184, Dow Corning, USA) was poured into the mold, and cured at 80 $^{\circ}$ C in an oven. After solidification, three tubing accessing holes with 1.5 mm diameter were punched at inlet and outlets, and a 3 mm-thick PDMS base was irreversibly bonded to the top piece of the chip with air plasma treatment (COVANCE, Femato Science, Korea). The assembled chip was placed at 80 $^{\circ}$ C overnight in an oven for tight bonding.

DFF-based purification of reticulocytes with spiral microfluidic chip

The cultured erythroid cells were suspended in erythroid culture medium at concentration of 1 million cells per mL, and pumped into the chip at 0.8 mL min⁻¹. Purified reticulocytes were collected from the outer outlet of the chip, and unwanted erythroblasts and free nuclei were removed from the inner outlet (Fig. 1).

Re-circulation setup for DFF microfluidic purification of reticulocyte

Fifty milliliters of cell suspensions at 2 million cells per mL were loaded in a 125 mL spinner flask (ProCulture, Corning). A magnetic stirrer (Thermo Fisher Scientific, USA) was used to agitate the cell suspension at 35 rpm to avoid cell settlement during the re-circulated sorting process. The cell suspension was pumped into the DFF chip through silicone tubing (Masterflex L/S 14, Cole-Parmer) with a peristaltic pump (Cole-Parmer) at 0.8 mL min⁻¹ (Fig. 3a). Cells separated to the outer outlet of the chip were circulated back to the spinner flask through silicone tubing (Cole-Parmer). Cells separated to the inner outlet of the chip were collected



using a 50 mL Falcon tube (BD Biosciences, USA) as waste. All the silicon tubing and the spinner flask were autoclaved before use.

DFF microfluidic purification of reticulocyte in continuous perfusion culture

A perfusion culture was setup inside CO₂ cell culture incubator by connecting a 125 mL spinner flask to an automated perfusion culture controller system (WhirlCell, Singapore), which utilizes a DFF microfluidic chip as the cell retention device. For clearer imaging and illustration purpose, the DFF device was unmounted from a removable cartridge of controller machine. Similarly, 2 million cells per mL suspension in the spinner flask was injected into the DFF chip through the pumps at 0.8 mL min⁻¹ controlled by the perfusion system. The DFF chip separated erythroblasts and free nuclei in suspension, and discard them with the old media into a waste bottle. Meanwhile, the purified reticulocytes were circulated back to the spinner flask. With old media being continuously removed, the perfusion system sensed a weight change in the culture vessel, and automatically fed fresh media into the vessel to maintain the working volume at 100 mL.

DLD microfluidics design and fabrication

DLD is a sensitive size-based sorting technique, using a regularly spaced pillar array where the separation can be determined by the established empirical formula:

$$D_c = 1.4g \tan \theta^{0.48} \quad (1)$$

where g is the regular spacing between pillars and θ is the offsetting angle of the pillars.⁵² We designed two DLD chips to compare conventional circle pillar and inverse L-shape pillars. These devices contain three segments of sorting of D_c size 2.4 μm , 3.8 μm and 5.1 μm respectively (Fig. 4 and S6, ESI†). Based on eqn (1), the gap size of the pillar array is fixed at 12 μm while varying the θ would determine the specified D_c . $\theta_1 = 1.0^\circ$, $\theta_2 = 2.6^\circ$ and $\theta_3 = 4.9^\circ$ are selected for the three DLD zones 1, 2 and 3 respectively. As the DLD pillars are arranged in an angle θ , each adjacent pillar is shifted by λ/N where $\sin \theta = 1/N$. A complete DLD segment is defined when a particle is displaced laterally by λ after passing N number of pillars (Fig. S6d, ESI†). Each DLD zone will consist of 6 repeating DLD segments and a complete displacement will shift a particle laterally by 6λ . Hence, each of the four outputs is designed for a shift of 6 sub-channels which correspond to the various sizes such that outlet 1 would contain cells less than a D_c of 2.4 μm , *i.e.* these cells are not displaced laterally within the device as their apparent size is smaller than the designed 2.4 μm ; outlet 2 would contain cells between D_c of 2.4–3.8 μm ; outlet 3 contains cells between 3.2 and 5.1 μm while outlet 4 contain cells larger than 5.1 μm . Outlet 1 is designed to be the collection outlet for reticulocytes as the inverse-L shape is predicted to

selectively deflect rigid particles along the gradient while allowing the deformable reticulocytes to slip. As reticulocytes are very similar to RBCs, 2.4 μm cut-off size would not deflect these cells and enable the separation of rigid nuclei and deformable RBCs despite a smaller diameter of the condensed nuclei. The DLD PDMS device is made from an SU-8 mold which is fabricated using standard photolithography techniques. A chrome mask is used to develop these fine circle and inverse-L shape features. Briefly, SU-8205 is spin-coated on a 4 inch silicon wafer to a height of 18 μm and a soft bake step is performed. The SU-8 is exposed under a UV-mask aligner and the DLD features are developed after a post-exposure bake. The final SU-8 mold is treated with trichloro(1*H*,1*H*,2*H*,2*H*-perfluorooctyl)silane (Sigma, Singapore) using chemical vapour deposition for an hour. This process renders the surface of the silicon wafer hydrophobic due to the carbon-fluorine groups in the molecule which prevents PDMS from sticking and the final device is easily peeled off from this SU-8 mold.

DLD device preparation and setup

DLD microfluidic chips were UV-sterilized and mounted on an inverted microscope (IX71, Olympus, Japan) equipped with a high-speed CCD camera (Phantom v9, Vision Research, USA) for real-time visualization of cell focusing and separation inside the channel. The recorded videos were processed using ImageJ software (NIH, USA) for images. Buffers or cell suspension were loaded in syringes (Terumo Medical, Japan), and pumped into the chip through UV-sterilized polyethylene tubing (Scientific Commodities, BB31695-PE/2) using syringe pumps (PHD 2000, Harvard Apparatus, USA). The chips were first primed with 1% poloxamer 188 (Sigma-Aldrich, P5556) in Milli-Q water to avoid sticking of cells to the channel walls during sorting, followed by 1× PBS to wash off the residual water inside the microchannel.

DLD sorting parameters for erythroid culture

The 100 million erythroid culture cells were first filtered using a 40 μm filter to remove large clumps of cells. The cells were washed in DMEM medium and added ~2000 units of DNase for 15 min (Sigma-Aldrich, 10104159001) to remove extracellular DNA, which would trap cells within the device. The cells were directly concentrated to 50 million per mL for characterisation tests and 400 million per mL (final volume of 250 μL) for high-throughput sorting. All sorting flows were performed at 3 μL per minute syringe infusion rate. To prevent cell settlement within the syringe, the Harvard pump was oriented vertically such that the syringe plunger is perpendicular to the ground. 4 outlets holes of 1.0 mm in diameter were punch into the PDMS device. 1.09 mm outer diameter polyethylene tubing (Scientific Commodities, BB31695-PE/2) of approximately 7 cm long was inserted into each outlet. The tubes were inserted into a custom orifice of



2 mm in the cap of the 1.5 mL Eppendorf tube. All components were sterilized using UV exposure for 10 min.

Microscopic and flow cytometric monitoring of erythroid cells

Cell number and viability were monitored after each phases of erythroid culture using trypan blue and counting chamber. A total of 5 million erythroid cells were used for Wright-Giemsa stain (Sigma-Aldrich, WG16). Erythroid cells were washed in 1× PBS buffer then thin-blood smears were prepared on a glass slide. Samples were immediately air dried, covered with the dye for 30 seconds then rinsed thoroughly with water. Images were captured using a light microscope (Olympus, CX31), iPhone (Apple, 6S) and iDu LabCam iPhone 6 Microscope adapter (iDu Optics). This image capturing methods were also used for Heilmeyer and Giemsa stained slides in this study. Erythroid cells (approximately 100 000 cells) after each phases of culture were stained with following anti-human antibodies: CD49d-PE (Biolegend, 9F10), CD36-FITC (Biolegend, 5-271), CD71-PeCy7 (Biolegend, CY1G4), CD235 α -PE (Biolegend, HI264) and CD34-APC (Biolegend, 561). All cells were stained in FACS buffer (0.2% BSA, 0.05% sodium azide in PBS) for 10 minutes on ice, washed twice in FACS buffer and resuspended in FACS buffer containing 1:1000 DAPI (Thermo Fisher Scientific, 62248) for the viability analysis. Stained live cells were acquired using BD LSR II (BD Biosciences) and analyzed by FlowJo software (Tree Star).

P. falciparum culture

P. falciparum 3D7 strain was maintained in RPMI-based malaria culture medium, MCM, (RPMI 1640 supplemented with HEPES, hypoxanthine, sodium bicarbonate and Albumax II) at 2.5% hematocrit. Schizonts were purified by LS magnetic column⁵⁷ (Miltenyi Biotec, 130-042-401) and inoculated to 5 million host cells in 200 μ L of MCM in 96-well plate. MCM was replaced every other day. Parasitemia was monitored daily by microscopy with Giemsa-stained thin blood smears.

Immunofluorescence detection of *P. falciparum* parasites

To confirm the successful invasion, csRET (20 million) was stained in 0.5 μ M of CellTrace CFSE Proliferation Kit (Thermo Fisher Scientific, C34554) following the manufacturer's protocol. Stained cells were either resuspended in pre-warmed culture medium for *P. falciparum*. As a control, mature RBCs (20 million) was also stained in 2.5 μ M of CellTrace Far Red Proliferation Kit (Thermo Fisher Scientific, C34564). The efficiency of cell labeling was verified by acquiring the samples with MACSQuant Analyzer 10 (Miltenyi Biotec). A total of 5 million labeled cells were inoculated with purified schizonts for the parasite culture.

P. falciparum samples were settled on poly-L-lysine coverslips (VWR, BD354085) for 15 minutes, supernatant

were removed, air-dried immediately and fixed with ice-cold methanol for 5 minutes. Samples were stained at 37 °C, by first blocking with 3% BSA for 30 minutes, probed with 1:800 dilution of PfEXP2 antibody (Genscript, generated using the peptide KNIESGKYEFDVD) for 45 minutes and 1:1000 dilution of goat anti-rabbit Alexa Fluor 568 (Thermo Fisher Scientific, A-11010) for 45 minutes. Samples were also stained for DNA using 1:1000 dilution of Hoechst 33342 (Thermo Fisher Scientific, 62249) for 10 minutes at room temperature, and then mounted on a glass slide using a drop of ProLong Diamond Antifade Mountant (Thermo Fisher Scientific, P36970). Images were taken using confocal microscope (Zeiss, LSM700).

Micropipette data acquisition

Shear elastic moduli of reticulocytes were measured using the micropipette aspiration technique. The set-up comprises a micromanipulator (TransferMan NK2, Eppendorf) mounted on a microscope (IX71, Olympus). First, borosilicate glass capillaries (1.0 mm outer diameter, 0.75 mm inner diameter, Sutter Instruments Co.) were prepared using a micropipette puller (P-97, Sutter Instruments Co.). The tips of the pulled glass capillaries were then cut using a microforge (MF-900, Narishige Group) to inner diameters ranging from 1.5 to 1.6 μ m. After mounting the micropipette, each reticulocyte was aspirated at a constant rate of 10 mL per hour using a syringe pump (NE-1000, New Era Pump Systems Inc.). Images of the reticulocyte entering the micropipette were acquired at 1 frame per s using a camera (GP-660 V, Kunshan Gaopin Precision Instrument Co. Ltd.) connected to the microscope.

Micropipette data analysis

The raw images captured were analyzed using imageJ software (NIH, USA). More specifically, the length of the reticulocyte in the micropipette for each frame and the inner diameter of the micropipette were measured. Subsequently, these values were fitted into the model by Chien *et al.*⁵⁸ Two-tailed student's *t*-test with unequal variance was then employed to compare the different treatments with the control.

Fluid streamline simulation

The simulation of fluid streamline was performed using Comsol Multiphysics 5.3a. The cad drawing was importing into the simulation and a single-phase laminar flow physics module was used to simulate the flows. The shallow channel option was checked with a device height of 18 μ m. The results was exported into streamlines and flow 2D flow profile.

Statistical analysis

Statistics were assessed using GraphPad Prism 5 (GraphPad Software). Statistical significance was calculated using a



Mann–Whitney test and 2 way ANOVA, Sidak's multiple comparison test. *P* value of <0.05 was considered significant.

Conflicts of interest

There are no conflicts to declare.

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