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Scalable mesenchymal stem cell enrichment from bone marrow aspirate using deterministic lateral displacement (DLD) microfluidic sorting†

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The growing interest in regenerative medicine has opened new avenues for novel cell therapies using stem cells. Bone marrow aspirate (BMA) is an important source of stromal mesenchymal stem cells (MSCs). Conventional MSC harvesting from BMA relies on archaic centrifugation methods, often leading to poor yield due to osmotic stress, high centrifugation force, convoluted workflow, and long experimental time (~2–3 hours). To address these issues, we have developed a scalable microfluidic technology based on deterministic lateral displacement (DLD) for MSC isolation. This passive, label-free cell sorting method capitalizes on the morphological differences between MSCs and blood cells (platelets and RBCs) for effective separation using an inverted L-shaped pillar array. To improve throughput, we developed a novel multi-chip DLD system that can process 2.5 mL of raw BMA in 20 ± 5 minutes, achieving a 2-fold increase in MSC recovery compared to centrifugation methods. Taken together, we envision that the developed DLD platform will enable fast and efficient isolation of MSCs from BMA for effective downstream cell therapy in clinical settings.

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1. Introduction

The advent of regenerative medicine marks a new paradigm shift in medical treatment through therapeutic cell delivery

by autologous or allogenic means to replace/regenerate damaged cells, thereby creating personalized medicine.^{1,2} The discovery of stromal mesenchymal stem cells (MSCs) has led to the pursuit of pioneering novel therapies due to their tri-lineage differentiation potential into mesodermal cell types *in vitro* (adipocytes, chondrocytes, and osteocytes).³ This unique multilineage differentiation has spearheaded MSC therapies to treat health conditions such as neurological disorders,^{4,5} cardiac ischemia,⁶ diabetes,^{7,8} and cartilage diseases.⁹ Apart from their unique differentiation capabilities, MSCs are also known to exhibit immunomodulatory properties,¹⁰ enabling expanded cell therapeutic applications. These cells can be found in various biological samples such as adipose tissues,¹¹ umbilical cord blood¹² and salivary glands.¹³ A traditional MSC harvesting method is aspiration from human bone marrow (BMA)^{3,14–16} as it has a relatively higher MSC quantity compared to other tissues. Despite the abundance of MSCs in the bone marrow, they only constitute 0.01% and 0.005% of the bone marrow mononucleated cell (BMMNC) population for males and females, respectively.¹⁷ This advocates a critical need for more effective and efficient MSC extraction from biological samples.

Advancements in cell sorting methods exploit cells' biophysical and biochemical properties. Current cell sorting techniques such as fluorescence-activated cell sorting (FACS)

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and magnetic-activated cell sorting (MACS) leverage antibody labeling targeted to cell surface markers.^{18,19} These techniques can reproduce extremely high sorting resolution but are limited by the high cost of antibodies, bulky machinery, and laborious sample preparation. They also require technical expertise and high operating pressure, which may influence MSC consistency and cell integrity.²⁰ Besides, it has been challenging to determine MSC-specific cell surface markers.²¹ Antibody binding on cells may also affect their native biological or signalling pathways,²² thus unsuitable for downstream therapeutic purposes.

The conventional MSC isolation technique from BMA adopts a label-free density-based centrifugation^{23–25} method known as Ficoll density gradient centrifugation (DGC). Through Ficoll DGC, the mononuclear cell (MNC) fraction contains a heterogeneous cell population consisting mainly of monocytes, lymphocytes, hematopoietic stem cells (HSCs), and MSCs. Despite its wide usage, Ficoll DGC has the poorest yield amongst different MSC isolation techniques, including RBC lysis, immunomagnetic depletion, and simple filtration.^{26–28} It was shown that Ficoll DGC typically suffers from significant cell loss amongst human progenitor cells (30% for HSCs and 55% for MSCs).²⁶ MSCs can aggregate to form large clusters, congregating in the polymorphonuclear (PMN) fraction during centrifugation.²⁹ Apart from the modest cell recovery, the workflow for Ficoll DGC (~2 to 3 h) suffers from poor repeatability due to inconsistent preparation protocols, clinical personnel training, and increased exposure to contamination.

Microfluidic technologies have generated substantial advancement in cell sorting technologies over the years.^{20,30} Cell sorting using microfluidics often leverages the intrinsic biophysical properties of cells, rendering the techniques label-free.³¹ They can be classified into two groups: active and passive methods. Active methods exploit externally applied fields or forces such as acoustophoresis,³² electrokinetics,³³ and magnetophoresis³⁴ to enhance separation. In contrast, passive methods rely on inherent hydrodynamic interactions between the channel structures and cells for separation, optimized by manipulating channel geometries.²⁰ Passive methods include inertial focusing,³⁵ deterministic lateral displacement (DLD),³⁶ and hydrodynamic filtration.³⁷ These microfluidic techniques are often easy to operate and integrate and are scalable into clinical sample processing workflows. To our knowledge, no work has been done on directly isolating MSCs from whole undiluted BMA using microfluidic techniques.

Previous studies addressing the separation of MSCs have relied mostly on the inertial focusing technique.^{38–40} However, these investigations focus on size-based sorting for separating different MSC subpopulations from a pure culture population of MSCs. A significant constraint posed by this technique is the low sample concentration required for effective sorting to occur, limiting the use in high cell concentration samples such as blood or bone marrow samples. Conversely, DLD has demonstrated proficiency in

processing samples with high concentrations,^{41–43} making it more suitable for clinical applications with the absence of preprocessing steps or reagents. Therefore, this study aims to determine the potential of employing DLD technology for direct BMA processing as an alternative to Ficoll DGC, the preferred method for isolating MSCs. We demonstrate a simple user-friendly workflow for BMA processing using a benchtop microfluidic pressure system. DLD, first described by Huang *et al.*³⁶ in 2004, leverages micropillar arrays to sort cells based on size and deformability with promising recovery rate and purity.⁴⁴ The sorting criteria (critical diameter, D_C) can be calculated by an empirical formula based on the geometrical parameters of the pillar arrays where particles larger than the D_C are displaced laterally, away from their original stream.⁴⁵ Particles smaller than the D_C will remain in their original stream, causing a distinct separation between the larger and the smaller cells. DLD sorting using peripheral blood samples has produced high recovery and purity of nucleated cells.^{41–43} However, the cellular composition between peripheral blood and BMA differs. The latter comprises a higher amount of immature blood cells and nucleated cells with larger variations in cell biophysical properties, such as cell size, as studied by Xavier *et al.*⁴⁶ Thus, the efficiency of direct MSC isolation from undiluted BMA samples remains unknown.

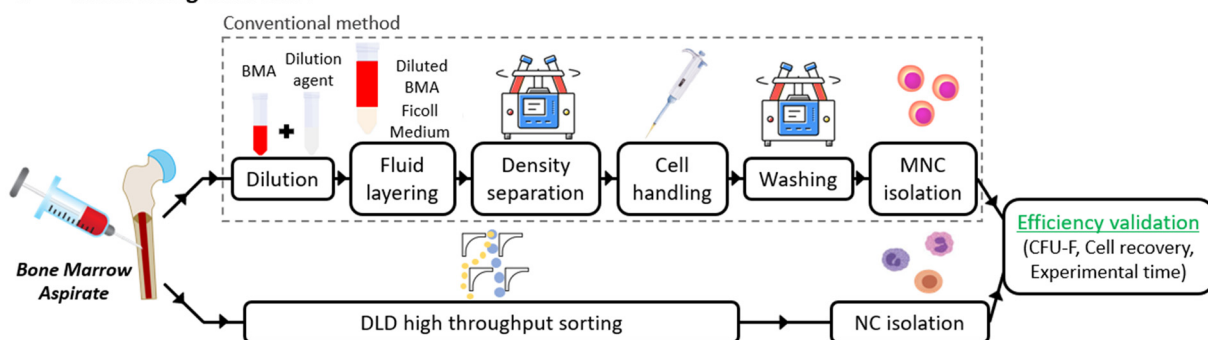
Three parameters, namely recovery rate, total experimental time, and MSC colony forming unit-fibroblast (CFU-F) count validated through FACS analysis, will be used to determine the effectiveness and efficiency of MSC isolation between DLD sorting and Ficoll DGC. To improve throughput scalability, we have also developed a novel multi-chip DLD system that can process 2.5 mL of raw BMA in 20 ± 5 minutes, achieving a 2-fold increase in MSC recovery compared to the Ficoll DGC protocol. Overall, the DLD platform shows excellent potential for fast and efficient isolation of MSCs from BMA in clinical settings.

II. Methods and materials

A. DLD device design

A high-throughput chip consisting of four DLD sorters (Fig. 1b), connected in parallel, was developed to increase the processing speed for BMA. This high-throughput chip involves layering two PDMS substrates where the second PDMS layer comprises channel connectors combining four channels into one, thereby reducing experimental complexity. Each DLD device design is based on a mirrored DLD pillar array and a centric sheath flow. Samples are injected from the sides of the device, and particles larger than the D_C are displaced laterally into the center sheath stream and sorted into the collection outlet (Fig. 2a). Particles smaller than the D_C follow their original stream, and they are not displaced, causing a separation between the smaller and the larger particles. Quantification of cells in BMA from flow cytometry in previous studies^{47–49} provides little information on biophysical properties such as size, shape, and deformability

a MSC sorting from BMA



b Custom Modular DLD Sorting System

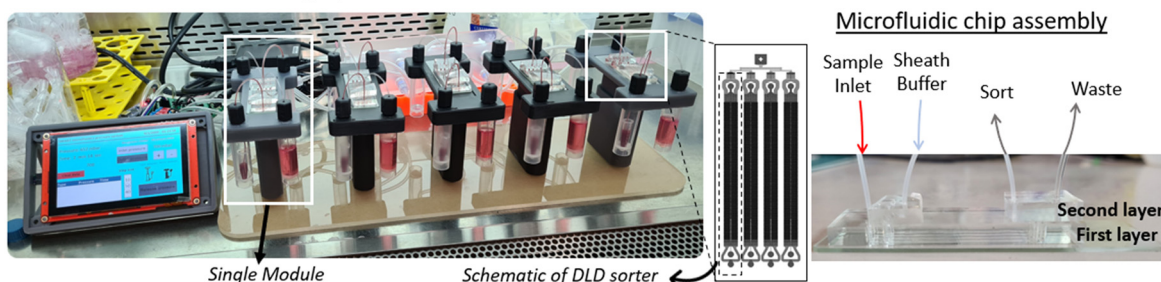


Fig. 1 Schematics of the experimental comparison of DLD *versus* Ficoll DGC for MSC sorting and recovery from BM samples. (a) Experimental overview for bone marrow aspirate using Ficoll DGC and DLD sorting. The product from both methods will be subjected to MSC phenotypic assays to verify the presence of MSCs through its colony-forming capabilities (CFU-F). Cell recovery and experimental time will be used to corroborate the effectiveness and efficiency for both methods. (b) The high-throughput design parallelizes four DLD sorting arrays into a single chip to increase the sorting speed. Throughput is further increased using the custom sorting system which allows multiple chips (up to 5) to operate simultaneously.

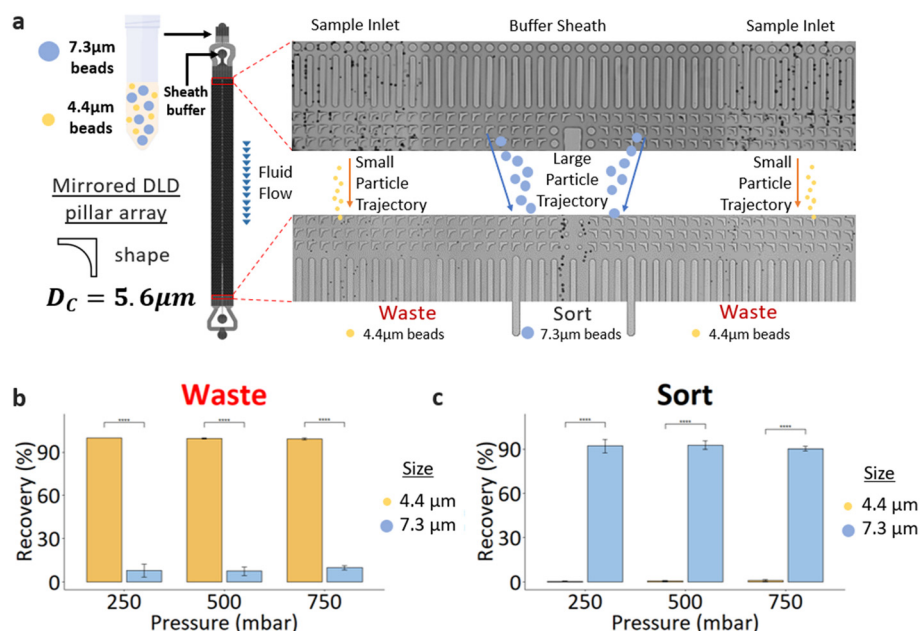


Fig. 2 Characterization of DLD sorting device. (a) Sorting characterization of the DLD device using 4.4 μm and 7.3 μm beads. The mirrored DLD device consists of an input region which has a central sheath buffer flow sandwiched by two sample co-flow streams. The outlet region of the DLD device shows the sorted region with two side waste channels which is characterized by the 4.4 μm and 7.3 μm beads shown in the waste and outlet (sort) region, respectively. (b) Graph showing the bead recovery in both the waste channels. (c) Graph showing the bead recovery in the sort channel. Paired two-tailed Student *t*-test was used to determine the statistical differences between outputs. **** denotes *p* value < 0.0001 for *n* = 4 samples.

of MSCs in BMA. Thus, the effective D_C of the device used in this study is approximately 5.6 μm with an unconventional inverted L-shaped pillar array instead of the original circle pillar array. This value is calculated empirically using an equation developed experimentally by Davis *et al.*⁵⁰ and follows the form:

$$D_C = 1.4G \tan \theta^{0.48} \quad (1)$$

where G refers to the gap size between the pillars and θ is the displacement angle. For a detailed description of the sorting device, please refer to Fig. S1 and S2.† This DLD sorting specification was designed to negatively sort out the smaller and deformable red blood cells in BMA, which have been proven to work best using an inverted L-shaped pillar compared to the conventional circular-shaped pillar.^{43,44}

B. DLD device fabrication and preparation

Fabrication of a silicon master mold follows a standard photolithographic process using a chromed quartz photomask (JD Photodata, UK). An SU-8 mold was made using SU-82020 spun at 4500 RPM, resulting in a final resist thickness of 18 μm . The fabrication of the master mold was made using standard photolithography techniques *via* exposure, post-exposure bake, and development. The master mold underwent a silanization step using trichloro(1H,1H,2H,2H-perfluorooctyl)silane (Sigma-Aldrich, Singapore) to ensure a hydrophobic coating and ease of downstream DLD chip manufacturing. The production of DLD chips follows a soft lithographic process using the master mold. PDMS (Dow Corning, USA) is mixed in a 10:1 ratio with its curing agent and poured over the master mold taped onto a Petri dish. The mixture will be degassed in a desiccator for 30 minutes before baking the degassed PDMS at 80 °C for 2 hours. After curing, a surgical knife is used to cut the PDMS devices from the mold.

Holes of different sizes are punched at the ingress and egress of the desired chip (1 mm for sample inlet, sheath flow, and outlet (sort), 1.5 mm for waste). The resulting PDMS product will be bonded onto a glass slide cleaned with 70% ethanol, using oxygen plasma treatment, and placed onto a hot plate at 140 °C for 10 minutes to ensure complete bonding. PDMS-to-PDMS bonding follows the same protocol as PDMS-to-glass bonding, with the only difference being placing the bonded product into an oven at 80 °C overnight instead of a hot plate.

Prior to experiments using biological samples, DLD chips are primed using 2% (w/v) Pluronic™ F-127 (Sigma, SG) with 20% isopropyl alcohol (IPA) for 30 minutes to prevent nonspecific adsorption and easy bubble removal due to its low surface tension properties from the ingress and 10 minutes from the egress using a pressure pump. Then, the chips will be washed with Dulbecco's phosphate-buffered saline (DPBS) (Thermo Fisher Scientific PTE Ltd, Singapore) or Gibco Dulbecco's modified Eagle's medium (GE

Healthcare, USA) for 5 minutes from the ingress to cultivate a suitable cellular environment as Pluronic™ F-127 is toxic to cells.

C. Single-device characterization

To characterize sorting performance, the DLD chip is placed onto a microfluidic device platform with imaging capabilities powered by an Arduino microcontroller and a positive displacement pump (RS PRO, Singapore). Custom fittings are designed and 3D-printed using low-force stereolithography (Form 3, Formlabs), forming an airtight junction between the sample reservoir and the tubing (Cole-Parmer, USA) using O-rings. The tubing length is 30 cm. Microsphere counting techniques leverage computer vision technology, and cell counting is performed using a hemocytometer.

D. Modular high-throughput sorting

For high-throughput sorting targeted for clinical applications, a new benchtop multi-chip microfluidic platform was developed, which allows multiple chips to operate concurrently. We have established a clear, easy-to-follow experimental workflow to complement our system. Tubing, fittings, and the prototype are sterilized in a biological safety cabinet (BSC) under UV light for 30 minutes before use. Experiments using BMA are also performed in the BSC to ensure sterility.

A pressure sensor (AMS 5915, AMSYS, Germany) and pump setup was assembled to control the desired operating pressure. To offset the pressure drop at the tubing and connections, we integrated a simple pressure cutoff control in our PCB design that monitors the pressure continuously and activates the pump whenever the sensor detects a drop below the 10% threshold. Pneumatic solenoid valves (Parker Hannifin, USA) were connected to a manifold inside the pressure chamber to enable proper distribution of pressure to the correct ports. The pressure chamber with manifold, modular unit, and human-machine interface (HMI) stand is fabricated using 3D printing. The system components can be found in Fig. S3.†

E. Peripheral blood samples

Peripheral whole blood was obtained following informed consent under the approval of the ethics committee according to a protocol permitted by the SingHealth Centralised Institutional Review Board (CIRB Ref: 2022/2322). 100 μL of whole peripheral blood was processed from a 3 mL venous blood draw into a 3 mL EDTA tube. The blood sample was processed within 4 hours of the blood draw to prevent inherent aggregation and degradation of samples. The blood samples were stored at room temperature to ensure minimal biophysical changes to the nucleated cells.

F. Bone marrow aspirate samples

Approximately 50 mL of bone marrow was harvested from the iliac crest of three healthy donors aged 26–52 under local anesthesia, following informed consent and strict adherence to ethical guidelines reviewed by the National Healthcare Group, Singapore (NHG DSRB 2019/00284). The aspiration was performed using a Jamshidi needle; the aspirate was collected into heparinized syringes and transferred to sterile containers. The bone marrow aspirate was transported to the National University of Singapore Tissue Engineering Programme (NUSTEP) for further processing. From the 50 mL aspirate, 5 mL was used for Ficoll DGC and DLD sorting at the Bioprocessing Technology Institute, A*STAR, Singapore. The remaining aspirate was not used in this study.

G. BMA processing using Ficoll DGC

Heparinized bone marrow aspirate was first diluted with an equal amount of D10 culture medium. This culture medium consisted of Dulbecco's modified Eagle's medium (DMEM), 1000 mg mL⁻¹ glucose supplemented with 10% fetal bovine serum and 2 mM L-glutamine. The diluted aspirate is carefully layered on top of 1.078 g mL⁻¹ Ficoll-Paque PREMIUM (Cytiva, USA) at a ratio of 2.5 mL diluted aspirate to 2 mL Ficoll-Paque. The solution was centrifuged at 400g for 30 minutes at ambient temperature without brake mode. Next, the interface layer containing the mononuclear cells was collected and washed twice by centrifugation at 200g for 15 minutes at maximum brake mode using complete DMEM.

H. Mononuclear cell culture

Following the user guide, cells obtained from the Ficoll DGC method and DLD sorting were counted using a NucleoCounter NC-250 analyzer (Chemometec, Denmark). Cells (3.5×10^6) were seeded into a 25 cm³ tissue culture flask (Thermo Fisher Scientific, Singapore) with 5 mL of D10 culture medium. DNase I (Roche, Germany) was added into the culture medium at 20 U mL⁻¹ concentration. Culture medium change was done at three- to four-day intervals. The spent culture medium was centrifuged at 380 RCF for 5 minutes at each culture medium change. The supernatant was discarded, and the pellet was resuspended in 5 mL fresh culture medium. DNase I at 20 U mL⁻¹ was added. The fresh medium with resuspended cells was seeded into the cell culture. Cells were cultured until 80% confluency was reached. These cells are termed passage 0 mesenchymal stem cells (P0 MSCs).

I. Mesenchymal stem cell culture

Confluent P0 MSCs cultured in 25 cm³ tissue culture flasks were passaged using 0.25% trypsin-EDTA dissociation solution (Gibco, Singapore). The passaged cells were counted using NucleoCounter NC-250 and seeded into a 75 cm³ tissue culture flask (Thermo Fisher Scientific, Singapore) with 15 mL

of D10 culture medium at a seeding density of 5000 cells per cm³. Cells were cultured until 80% confluency was reached. These cells are called passage 1 mesenchymal stem cells (P1 MSCs) at this stage. These P1 MSCs were then further passaged and seeded into (i) a 175 cm³ tissue culture flask (Thermo Fisher Scientific, Singapore) with 35 mL D10 culture medium at a seeding density of 5000 cells per cm³ and cultured until 80% confluent, and (ii) a 6-well tissue culture plate (Thermo Fisher Scientific, Singapore) with 2 mL D10 culture medium at a seeding density of 65 cells per well and cultured for 14 days with culture medium change on day 7 for CFU-F analysis. These cells were termed passage 2 mesenchymal stem cells (P2 MSCs).

J. MSC quality analysis: colony-forming unit-fibroblast (CFU-F)

P2 MSCs cultured on a 6-well tissue culture plate at a seeding density of 65 cells per well for 14 days with culture medium change on day 7 were used for CFU-F analysis.⁵¹ The cells were washed with sterile phosphate-buffered saline (PBS) (Gibco, Singapore) and stained with 2 mL of CFU-F stain for 30 minutes at room temperature. The CFU-F stain was prepared by dissolving a 0.5% weight-to-volume ratio of crystal violet (Sigma-Aldrich, Singapore) in a 25% volume-to-volume ratio of methanol (Sigma-Aldrich, Singapore) in deionized water. The cells were then rinsed with PBS thrice to remove excess stain. The plate containing the stained cells was dried. Once the plate had dried, the number of colonies formed was counted. A colony is defined as a region with a minimum of 50 cells. Any areas with less than 50 cells were not considered a colony and thus were not counted.

K. MSC quality analysis: cell surface marker (FACS)

P2 MSCs cultured in a 175 cm³ tissue culture flask at a seeding density of 5000 cells per cm³ until 80% confluency were used for cell surface marker (FACS) analysis. Firstly, cells were washed with sterile PBS, dissociated from the tissue culture flask using 0.25% trypsin-EDTA, and counted using NucleoCounter NC-250. 1×10^5 cells were used for each cell surface marker analysis. The positive selection markers used for the study were CD73, CD90, CD105, and CD146, and the negative selection markers were CD34 and CD45. Cells were washed in FACS buffer, which was made by dissolving a 1% weight-to-volume ratio of bovine serum albumin (Sigma-Aldrich, Singapore) in PBS. The washed cells were then added to a 96-well V-bottom plate (Greiner Bio-One, Germany) and individually incubated with each fluorophore-conjugated antibody of positive and negative selection markers (Biolegend, USA) on ice for 30 minutes. The cells were washed with FACS buffer to remove excess unbound antibodies and resuspended in FACS buffer. According to the user manual, cells were analyzed using Novocyte (ACEA Biosciences, USA).

III. Results

A. DLD beads sorting characterization and optimization

To calibrate the DLD chip, polystyrene microspheres (Bangs Laboratory, USA) of sizes 4.4 μm and 7.3 μm were used at a stock 1.125% (w/v) concentration. This test illustrates the size-based sorting capabilities as the rigid beads are larger and smaller than the D_C (5.6 μm). As the pressure and corresponding flow rate increased, the sorting of 7.3 μm beads showed a minor decrease in bead recovery ($92.14 \pm 3.85\%$ to $90.25 \pm 1.52\%$). Using the inverse L-shaped pillars could result in possible anisotropic effects as shown by Vernekar *et al.*⁵² By comparing the sorting efficiency of 4.4 μm beads, the DLD device with a D_C of 5.6 μm performs as designed in the operating pressure range of 250 to 750 mbar. The 7.3 μm beads accounted for 7–10% of the total count at the waste outlet (Fig. 2b) and were not deflected *via* DLD (see ESI† Video S1 and S2). This is likely due to the functional design of the device, where the inlet sample flow interacts with the DLD array side walls, resulting in inefficient sorting near the walls.

B. Peripheral blood sorting

Peripheral blood was used to elucidate the cell deformability effects and sorting efficiency of the DLD device. Unlike rigid beads, it is known that cell deformability impacts the DLD sorting efficiencies.^{42,46,53,54} The same DLD device used in

Fig. 2 was scaled linearly up to 4 \times by connecting the devices in parallel with a second layer microfluidic sample distribution channel at the input and output. Undiluted and unfiltered whole blood was processed using the device, as shown in Fig. 3a and ESI† Video S3 and S4.

To facilitate the output cell count, cells are DNA stained with Hoechst 33342 fluorescent stain solution and counted with a hemocytometer using brightfield and DAPI microscopy. With fluorescence field imaging, the cell recovery rate is quantified through manual counting and carried out as follows:

$$\text{Recovery (\%)} = \frac{\text{Nucleated cell count in outlet}}{\text{Nucleated cell count in outlet and waste}} \times 100\%$$

The corresponding results (Fig. 3c) showed that the cell recovery decreases as pressure increases. Theoretically, the DLD device design with $D_C = 5.6 \mu\text{m}$ should result in a 100% efficiency for both sorted (PBMC size $>7.23 \pm 0.3 \mu\text{m}$)⁵⁵ and unsorted products (RBC size $\sim 3 \mu\text{m}$, platelets size $= 3.64 \pm 0.72 \mu\text{m}$).^{56,57} However, in comparison to the bead sorting, peripheral blood results showed different efficiencies across different pressure values, which underscore the need to account for an additional cell deformability factor. Decoupling size and deformability effects in DLD sorting remain challenging, and thus, various groups often use the effective size parameter to quantify the deformed size of a

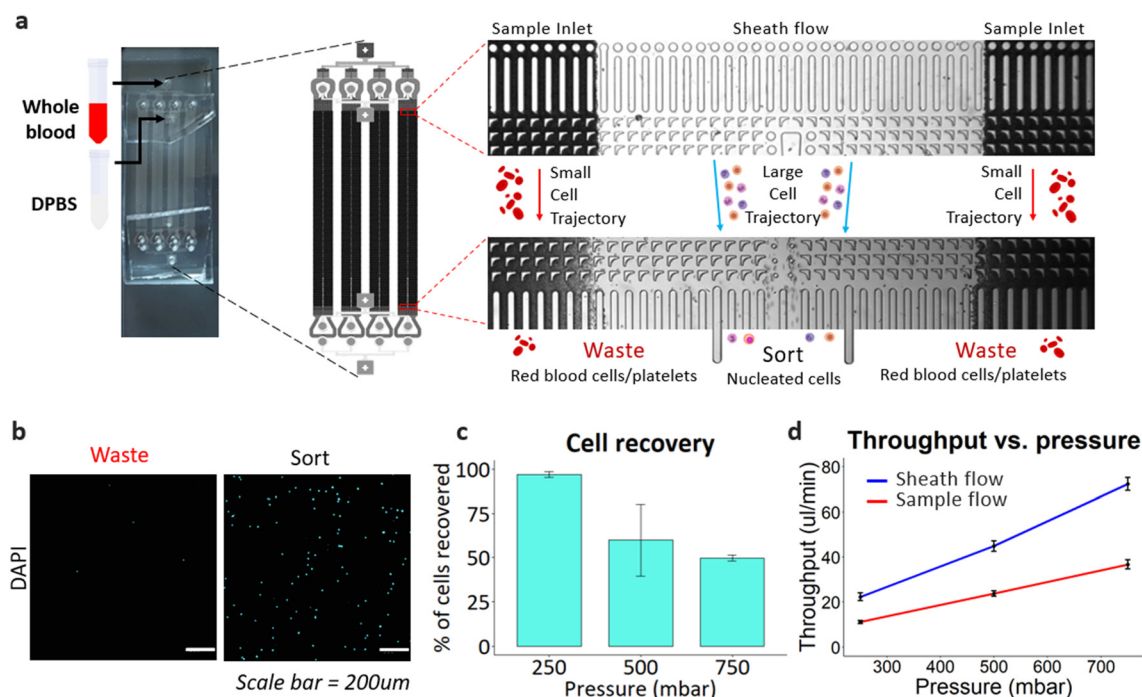


Fig. 3 Characterizing whole blood nucleated cell sorting in parallel DLD devices. (a) PDMS DLD chip shown with 4 \times parallel devices to increase sorting throughput. The input and output region of the DLD device can be seen with the sample stream co-flow with a sheath buffer stream. Nucleated cells were collected in the 'sort' stream while the waste stream consists of RBCs and platelets. (b) Fluorescence images showing the nucleated cells stained with Hoechst 33342 for counting of nucleated cells in the respective sorted outlets at 250 mbar. (c) Cell recovery from the DLD chip was measured with respect to varying flow rates. (d) Graph showcasing the corresponding flow rates ($\mu\text{L min}^{-1}$) and pressures used (mbar) for a fixed volume (100 μL). All experiments were performed in triplicates ($n = 3$).

biological cell.^{53,54,58} This observation for stem cell sorting is made following studies done by Xavier *et al.*, where they verified the deformability effects in DLD sorting using MG-63 and HL-60 cells⁴⁶ with varying flow rates. Therefore, at higher flow rates, the increased forces acting upon the cell induce a larger change in size as compared to that in the literature, which explains the decrease in efficiency (Fig. 3c). However, we have decided to use the highest pressure for our BMA experiments for two reasons. The trade-off between high throughput and cell yield must be considered based on different applications. For this study, achieving high throughput is critical for manufacturing-scale cell separation and favours higher flow and volume processing rates. Secondly, peripheral blood is also known to express pseudoplastic properties.⁵⁹ Since viscosity is inversely proportional to flow rate, we would want blood/BMA to exhibit less viscosity at a higher flow rate, which is beneficial to achieve maximum volume processing rates. In this case, we assume that the properties of BMA and peripheral blood are similar due to the abundance of RBCs. The throughput follows a linear trend with pressure (Fig. 3d) and these corresponding values can be found in Table S1.†

C. MSC isolation from human bone marrow aspirate using DLD

To validate our proof of concept that DLD sorting works better than Ficoll DGC, we performed two comparative sorting experiments: Ficoll centrifugation and DLD sorting. The Ficoll DGC procedure has been explained in section IIG. Microfluidic research on MSC cell sorting is commonly performed using cell lines, cultured cells which are spiked samples and not physiologically and clinically relevant.^{26,27,46,60,61} Therefore, it is critical to process BMA samples directly for clinical impact. The primary goal of this study is to develop a proof-of-concept sorting of BMA and evaluate the quality of sorted cells in comparison to Ficoll DGC. As part of efforts to increase throughput, five high-throughput chips were used, resulting in twenty parallel DLD devices operating concurrently. The workflow is described in Discussion S1.†

Our undiluted BMA sample had an initial cell concentration of $8.59 \pm 3 \times 10^6$ cells per mL and was subjected to a 20 μm filter before sorting to remove unwanted coagulation (see Fig. S5†). After sorting, we combined the outlet (sort) vials from each module into a 15 mL vial before data analysis. This was done likewise for the waste vials. The measured cell recoveries between peripheral blood and BMA at 750 mbar differ slightly ($49.8 \pm 1\%$ and $43.9 \pm 6.3\%$, respectively). Since the design of our DLD array

is based on the static cell size, this illustrates the difference between nucleated cell sizes in BMA from that in the literature (Table 1). One of the reasons for the size disparity could be due to interactions with the DLD pillars causing cells to deform and change in size. Despite the lower cell recovery rate, the yield is better than that of Ficoll DGC by a factor of two (Fig. 4a).

In contrast, Ficoll DGC has a fixed, established protocol, and the time taken is independent of sample volume. The Ficoll DGC method can take 2–3 hours (volume processed can range from 2 to 35 mL with dilution), and a series of centrifugation steps are needed to achieve better results. The time plot in Fig. 4a includes both processing and handling time. Processing time refers to the total sorting/isolation duration, and handling time refers to the time taken to handle the cells before/during the sorting, which includes washing and dilution. Studies have also shown that the Ficoll medium is toxic to cells,⁶² and multiple centrifugation might increase cell death due to prolonged exposure to the Ficoll medium. The results showed that DLD sorting reproduced a comparable number of colonies per million cells to that of Ficoll DGC at P0 (Fig. 4a). However, since the cell recovery is two times higher than that of Ficoll DGC, the total number of colonies and MSC recovery is significantly higher for DLD sorting. To verify that these colonies are MSCs, we performed a phenotypic assay after the second sub-cultivation (passage 2) to examine for MSC surface marker expression (CD73, CD90, CD105, CD146) and clear of the hematopoietic cell lineage (CD34[−], CD45[−]) using FACS as iterated under section IIK. The findings (Fig. 4b) indicate that the MSC population is similar in the Ficoll DGC colonies. However, the total number of MSCs is still higher in DLD sorting due to higher cell yield. The quality and purity of the cells were also assessed using the same markers, showcasing that the DLD sorting method is able to effectively isolate the MSC population with minimal changes to the quality of MSCs (Fig. 4b and S6 and Table S2†). We also performed a CFU assay in passage 2 showing minimal differences in quality (Table S3†). These outcomes further substantiate the DLD method for successful isolation and enrichment of MSCs.

IV. Discussion

Stem cell therapy is the new disruptive innovation in regenerative medicine. Multiple biological tissues have been reported to contain MSCs, and identifying the most effective MSC isolation technique from a suitable source is critical for sorting functional MSCs. Despite having a low MSC yield, Ficoll DGC has long been the primary method for MSC isolation from BMA (Table 2).

Table 1 Bone marrow nucleated cell size from the literature^{46,48,49}

	Myeloid cells	Monocytes	Blast cells	Lymphocytes	BM stem cells
Size (μm)	15 ± 5	16 ± 4	13 ± 5	6–15	9.1 ± 1.6

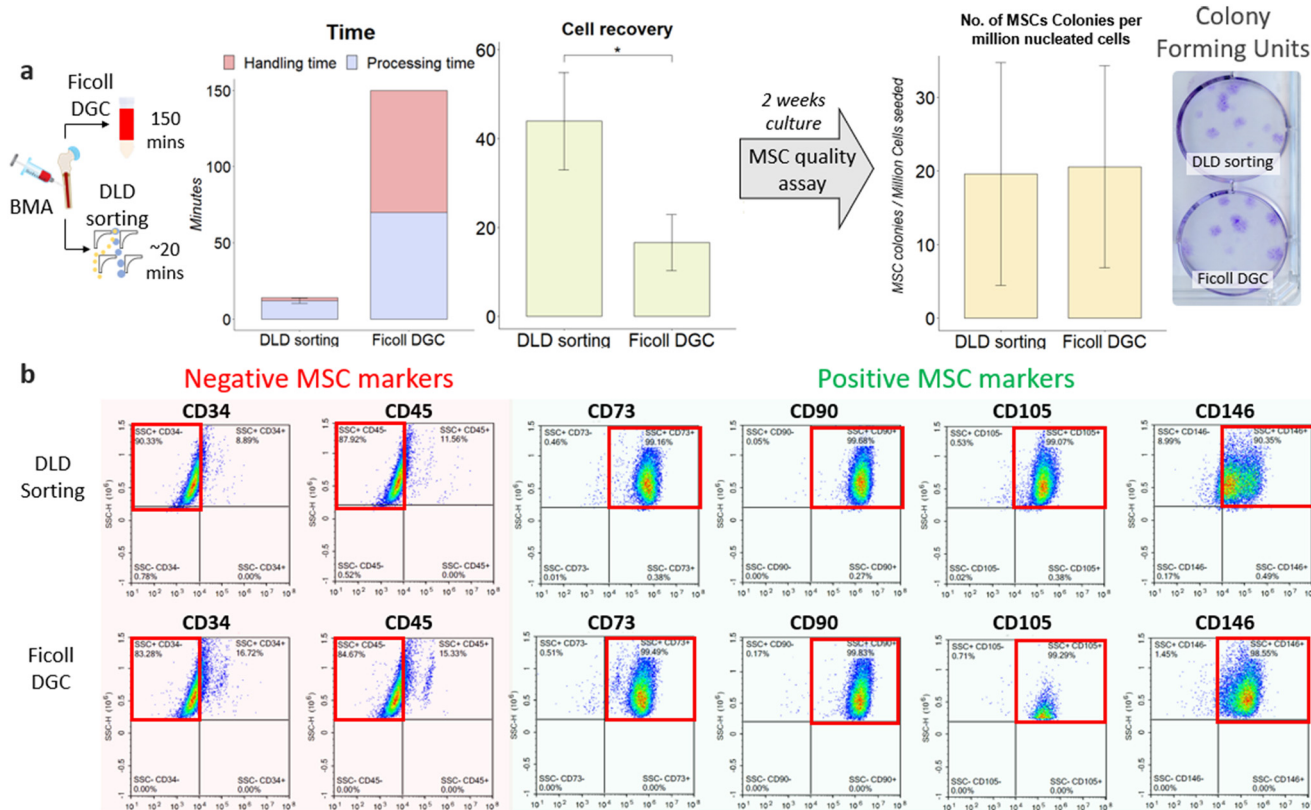


Fig. 4 Sorting efficiency and recovery of MSCs from BM samples using the DLD chip and Ficoll DGC. (a) BMA processing using both DLD sorting and Ficoll DGC was performed to validate the effectiveness of MSC enrichment through CFU count, cell recovery and time taken ($n = 3$). (b) Flow cytometry analysis of MSC phenotypic cell surface markers on sorted cells using DLD and Ficoll DGC.

Comparing the recovery across all MSC harvesting methods, our current DLD design offers one of the best alternatives with no sample dilution or manual washing steps (handling time). “Time taken” in Table 2 refers to the reported processing time of these methods and does not include sample handling time such as pipetting, centrifuge

braking or sample filtration which could plausibly increase the experimental timing significantly. However, what sets the DLD sorting technology apart from the rest of these methods is the absence of sample preparation and pre- and post-sort washing steps and potential scalability. The precision and continuous flow single-step sorting also eliminates variability

Table 2 Comparison of MSC sorting techniques from BMA

Separation techniques	MSC harvesting methods/medium	Bone marrow aspirate processing	MNC recovery	Time taken	Label free	Ref.
Density gradient centrifugation	Ficoll-Paque™	Centrifuged at 200g for 10 minutes	13.5%	70 minutes	Yes	63
	Percoll	1 : 2 PBS dilution	15.7%	30 minutes	Yes	64
	Ficoll-Paque® PREMIUM		30.9 ± 9.3%	20 minutes	Yes	
Cell lysis	BD Vacutainer® Cell Preparation Tube™ (CPT)	Undiluted	22.9 ± 9.3%	10 minutes	Yes	27
	Red blood lysis		Reported as 10 times better than Ficoll centrifugation	10 minutes	Yes	
Immuno-depletion	MACS (removal of CD15+ PMNs)	1 : 10 PBS/3% FCS + 5 mM EDTA (RBC lysed)	55%	20 minutes	No	26
	PluriBeads (removal of CD15+ PMNs)		58%	40 minutes	No	
Filtration	Porous biological material	1 : 2 alpha-MEM containing 10% [v/v] fetal bovine serum (RBC lysed)	41.3 ± 15.5% (carrier material dependent)	Not reported	Yes	65
Microfluidics	DLD sorting	Undiluted	43.9 ± 6.3%	13 minutes	Yes	NIL

associated with the cell handling steps. For the multistep Ficoll DGC process, the same personnel performing the technique will exhibit some form of variability due to human error, which is almost impossible to eliminate. DLD sorting also inherently integrates a washing step where sorted cells are migrated into the buffer stream, thus removing the need for additional washing steps.

The current sorting efficiency for MSCs is $43.9 \pm 6.3\%$, and improvements on the nucleated cell recovery and processing time can be achieved by further optimizing microfluidic DLD specifications by reducing the critical size specification to sort out smaller ones or more deformable cells at the same or higher flow rate. It can be hypothesized that sorting smaller nucleated (mononuclear and polynuclear) cells will obtain a higher yield of MSCs than sorting only mononuclear cells *via* Ficoll DGC. Early evidence from this paper has shown that mononuclear cell sorting loses a large fraction of MSCs. This was also validated by Horn *et al.*,²⁷ who employed a whole blood lysis protocol to extract all nucleated cells, showing that there exists a MSC population that is missed by mononuclear cell sorting methods. Given the similar cell recovery values between peripheral blood and BMA at 750 mbar, we are confident that our DLD device can achieve a higher cell recovery ($\sim 80\text{--}90\%$) if deformability effects are negated (running at 250 mbar) at the expense of throughput. Also, expanding these cells to the desired cell quantity is often necessary to meet the requirements for MSC-based therapies. However, increased passaging is known to impair MSC plasticity and contribute to genomic instability, telomere shortening, and heterogeneity,⁶⁶ which limits the therapeutic effects of MSCs, thereby reinforcing the need for an effective sorting method. While the CFU potential is comparable between the two approaches, DLD offers a higher cell recovery, translating to a higher MSC recovery. Given the rare occurrence of MSCs in BMA, a higher yield during sorting can lead to exponential cell numbers in downstream expansion.

While microfluidic chip stacking has been explored by various groups^{67,68} to enable scaling of sample processing, this paper investigates the possibility of scaling the sorting process using a modular sort unit that can be seamlessly integrated into a central pressure distribution setup. The microfluidic setup and processing workflow can also be used for leukapheresis with very high yield at lower flow rates. Future development of this prototype can focus on integrating portability, streamlined functionality, and user-friendly design to enable operation by non-trained users/clinicians. Without the need for multiple washing steps (Table 2), the major benefit of using the DLD sorting technique is the integration with bioreactors in a closed-loop processing of cell separation and manufacturing with the potential to reduce contamination and workforce cost and to improve consistency.

Isolating cells by label-free parameters such as size and deformability enables the enrichment of specific cell functions or phenotypes. The next questions to be answered

following the sorting are: Do the sorted cells have differing growth potentials? What is the differentiation potential of MSCs to be of a particular phenotype for regenerative medicine? To quantify the functional qualities of these DLD-sorted MSCs, it is paramount that stemness and immunomodulation are validated through functional assays such as RT-PCR and differentiation assay, which serves as a motivation for future study. A previous study by Yin Lu *et al.* has ascertained that MSCs of sizes $17\text{--}21\text{ }\mu\text{m}$ have the most potential for chondrogenesis using inertial sorting.³⁹ While downstream quantitative assays to test for the mesoderm cell lineage differentiation were not explored in this study, the utility of DLD sorting goes beyond a sample processing tool with the potential for specific phenotypical enrichment for more efficacious cell manufacturing.

V. Conclusion

In this study, we have developed a multi-chip DLD microfluidic platform for MSC sorting from BMA samples directly and demonstrated superior separation performance compared to Ficoll DGC (conventional standard). Improving the chip throughput and automation in our system makes this clinical workflow faster ($\sim 20\text{ min}$), more user-friendly and more robust for MSC isolation. With its label-free approach, this platform technology is low-cost, and we can envision it to be readily adapted to sorting other cell types by controlling DLD pillar shapes and geometries.

Author contributions

N. T. K. Z., K. K. Z., H. W. H., S. O., S. M. C., and J. H. designed aspects of the project study. N. T. K. Z. and K. K. Z. designed and fabricated the chip. N. T. K. Z., K. K. Z., G. C. R., and Y. D. H. optimized and performed the experiments. N. T. K. Z., K. K. Z. and G. C. R. designed, fabricated, and calibrated the electronics and sorting setup. N. T. K. Z. and K. K. Z. analysed the data. N. T. K. Z. coded the algorithms for software and hardware. J. H. H. P. performed the bone marrow aspiration. T. K. L., M. L., J. L., and S. O. performed the patient recruitment and post-sort cell measurements and FACS profiling. N. T. K. Z., K. K. Z., S. M. C., H. W. H. and J. H. wrote the manuscript. All authors reviewed and approved the manuscript prior to submission.

Conflicts of interest

We declare no conflict of interest.

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