Label-free enrichment of rare unconventional circulating neoplastic cells using a microfluidic dielectrophoretic sorting device

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Cellular circulating biomarkers from the primary tumor such as circulating tumor cells (CTCs) and circulating hybrid cells (CHCs) have been described to harbor tumor-like phenotype and genotype. CHCs are present in higher numbers than CTCs supporting their translational potential. Methods for isolation of CHCs do not exist and are restricted to low-throughput, time consuming, and biased methodologies. We report the development of a label-free dielectrophoretic microfluidic platform facilitating enrichment of CHCs in a high-throughput and rapid fashion by depleting healthy peripheral blood mononuclear cells (PBMCs). We demonstrated up to 96.5% depletion of PBMCs resulting in 18.6-fold enrichment of cancer cells. In PBMCs from pancreatic adenocarcinoma patients, the platform enriched neoplastic cells identified by their KRAS mutant status using droplet digital PCR with one hour of processing. Enrichment was achieved in 75% of the clinical samples analyzed, establishing this approach as a promising way to non-invasively analyze tumor cells from patients.
Cancer is a major worldwide public health burden and the second leading cause of death in the United States. Advances in medical technologies have led to significant progress in both treatment and diagnosis, yet clinicians rely on invasive and often risky biopsies for cancer detection. To address these limitations, the field of liquid biopsy has rapidly emerged as a non-invasive method to obtain tumor-derived biomarkers for diagnosis, staging, treatment, and prognosis.

Circulating neoplastic cells disseminated from solid tumors are valuable biomarkers reflecting the disease transcriptome, genome, and proteome. Circulating tumor cells (CTCs) are cells disseminated from the tumor, shed into peripheral blood, and capable of recapitulating tumorigenesis. In epithelial-derived tumors, CTCs are phenotypically defined as large cells (30 μm on average) lacking expression of the pan-leukocyte marker, CD45, and expressing cytokeratin (CK). In addition, other markers have been associated with CTCs like epithelial cell adhesion molecule (EpCAM) or E-cadherin (ECAD) for isolation or identification. CTC enrichment and isolation technologies exist, including the FDA-approved CellSearch® technology, to enumerate and phenotype CTCs to guide clinical prognosis. With estimates of one CTC per 10^5–10^8 leukocytes found in peripheral blood of pancreatic ductal adenocarcinoma (PDAC) patients, levels of these rare biomarkers challenge their functional utility, thus improved approaches to capturing these rare cells are needed.

Many groups have focused on using microfluidic sorting, density-based devices, size exclusion filtration, or marker specific isolation methods. These technologies rely on either a single physical property (size, density) or on specific protein expression (EpCAM, ECAD, CK) to differentiate CTCs from peripheral blood mononuclear cells (PBMCs). However, the rarity of CTCs has greatly limited their application in providing multi-omics information that encompasses tumor heterogeneity and has not translated well into widespread clinical use.

In the past 5 years, other promising types of unconventional circulating neoplastic cells have been described such as tumor hybrid cells (THC) and circulating hybrid cells (CHCs). THC are defined as having hybrid phenotypes of a neoplastic cell and a leukocyte with dual expression of CD45 and tumor markers such as EpCAM or CK11–15. In PDAC patients, THC and CHCs have shown better prognostic value than CTCs as their enumeration correlated with stage and survival. While CHCs are more abundant, their physical and phenotypic similarity to PBMCs makes their isolation and analysis a challenge.

To address these challenges, we developed a microfluidic device utilizing the intrinsic dielectrophoretic (DEP) properties of cells to enable their label-free enrichment, in a fashion compatible with both phenotypic and genotypic downstream analyses. Particle movement caused by difference of electrical polarizability between particles and medium in a non-uniform electric field has been thoroughly elaborated by Jones and Poll, and has been applied to separate and isolate biomarkers. When cells are exposed to this electric field, differential polarization is created between the cells and the surrounding media translating into a resultant force. This resultant force can either be positive DEP where cells travel up the electric field (E-field) gradient, or negative DEP where cells travel down the gradient. The overall magnitude of the DEP force is dictated by the E-field gradient and the particle/cell geometry and the directionality of the force is determined by relative permittivity between the cell and the surrounding media and is specifically explained by the Clausius–Mossotti factor.

DEP as a cell differentiation approach can be grouped into two main categories, batch mode or continuous mode processing. Batch mode processing relies on a unique frequency cut-off where a cell’s DEP response changes between positive and negative, for each discrete cell type. Defined frequencies can then be used to separate two cell groups to positive or negative DEP regimes followed by flushing out the negative DEP partition. This approach provides enhanced specificity for discrete cell types expressing narrow dielectrophoretic variance; however, its utility is impeded by low throughput and narrow search space. In contrast, continuous mode processing uses a secondary force (flow, acoustic, or others) to incrementally bias cell trajectory over a period of time.

The combination of continuous flow and DEP can result in multiple cell sorting outcomes in high cell density environments translating into a versatile high-throughput methodology for mixed cell populations with variant dielectrophoretic responses. The majority of strategies currently employed for rare cell enrichment are reliant on clearly defined physical properties and minimal phenotypic variation of the target cell population. However, this approach fails for heterogeneous cell populations with high phenotypic and DEP variance making it unlikely for a single condition to be successful. To address these limitations, we designed and optimized a continuous mode DEP based microfluidic device to allow for enrichment of highly heterogenous rare cell populations like CHCs. The workflow was validated using beads, optimized on cell lines and healthy PBMCs, then tested on cancer patient samples.

In this work, PBMCs from a small amount of blood (2 mL) were sorted in a high-density environment where non-target immune cells were depleted from the sample enriching for neoplastic cells in a label-free manner. As a proof of concept, enriched CHCs were subjected to phenotypic and genotypic downstream analysis demonstrating clinically relevant KRAS mutation status in PDAC tumors.

Results and discussion
Design and validation of microfluidic DEP sorting device

**Design.** The concept of enriching a target cell population by depleting non-relevant cells is shown schematically in Fig. 1. First, the sample is infused into the microfluidic DEP sorting device, composed of a microfluidic channel and interdigitated electrode arrays. The microfluidic device was manufactured by standard photolithographic and lift-off techniques where a silicon substrate was patterned with an interdigitated electrode layer and a polydimethylsiloxane (PDMS) layer was bonded on top to form a fluidic channel. To allow cells throughout the channel’s height to be acted upon by DEP forces, the chamber height was designed to be 25 μm or approximately twice the average size of PBMCs. Under continuous flow the sample is centrally confined by two sheath flows providing a small variation of initial speed and position. The interdigitated electrodes were designed to have a width and spacing of 35 μm, approximately 2–3 times the size of PBMCs, to minimize cells being captured on the electrodes by DEP forces (Supplementary Fig. 1). The V-shape electrode geometry was designed to deflect PBMCs in a high cell density environment by utilizing DEP forces to deflect cells along the electrodes while flow forces direct cells along the length of the device. Cells experiencing strong DEP forces are deflected to the side outlets, while less responsive cells are enriched in the center channel (Fig. 1a). In order to maximize the DEP response and minimize cells trapped on electrodes, a conductive buffer at 145 mM (approximately 0.1x PBS) was used during cell enrichment processing for both the sample and sheath inlets. The buffer was chosen to balance cell viability, DEP reactivity, and electrode corrosion. For ease of use and rapid troubleshooting the complete setup was placed under a microscope for continuous viewing where the fluidic channel is visible under a standard 5x objective lens. This setup allows for sample
compatibility with genotypic or phenotypic downstream analysis (Fig. 1a) which can be helpful for understanding cellular clinical significance.37.

Validation. To demonstrate the concept of enriching by depleting a target population we used a sample mix composed of 2 µm (blue) and 10 µm (green) fluorescent polystyrene beads of identical chemistry (Fig. 1b, c). The sample mix was introduced to the inlet of the microfluidic DEP sorting device with AC voltage applied to the interdigitated electrodes, and particle movement was monitored by live imaging. We defined the optimal condition as the parameters where one type of particle was enriched by depletion of the other particle. To find this condition, the frequency was swept from 0.1 MHz to 10 MHz at 0.1 MHz increments at constant voltage. Particles were collected at the outlets and analyzed by flow cytometry.

Under the optimal frequency, both bead populations were closely aligned with the center of the channel while near the sample inlet (Fig. 1b, part I), however only the 10 µm beads were depleted further downstream by DEP forces revealing particle trajectory bias over time and space (Fig. 1b, part II). Under this condition, 95% of the 10 µm beads were depleted from the center due to the application of strong positive DEP forces, while less than 1% of the 2 µm beads were depleted to the sides (Fig. 1c). Figure 1c also shows an example at one of the sub-optimal frequencies that only 19.5% of 10 µm were depleted, suggesting that the magnitude of the DEP force was smaller than the dominant flow forces. To confirm that DEP forces were the primary driver of bead depletion, the device was run without applied DEP and resulted in 99% of both bead populations remaining in the center unaffected by drift. This workflow demonstrated that depletion of one population by DEP can result in the collection of targets with high purity under proper working parameters. To understand how this applied to a complex biological sample mix, we next tested this process on PBMCs and cell lines.

Optimization of PBMC depletion and hybrid cell line enrichment

PBMC DEP response characterization. As DEP conditions can be optimized to separate discrete physiologic objects, we set out to establish the optimal condition where hybrid cells could be enriched from PBMCs. Cells are biologic entities, thus their conductivity, permittivity, and physiological state are the main drivers of differential DEP response.38 Healthy PBMCs are comprised of heterogeneous populations, but tend to be relatively similar in nature across individuals, therefore healthy PBMCs act as an ideal object to be optimized for depletion. PBMCs are mainly comprised of lymphocytes and monocytes. Of these, T-cells represent the majority (70–80%) of the lymphocytes, therefore we hypothesized that of all healthy PBMCs depleting T-cells would be the primary contributor of neoplastic cell enrichment. First, to understand the DEP response of PBMCs as a collective of cells and as individual subpopulations, we conducted the workflow using PBMCs isolated from whole blood (n = 5) from healthy subjects. PBMCs were diluted in DEP buffer (1 × 10^7 cells/mL), and 1 × 10^6 cells were loaded into the central inlet for each parameter, then subjected to device processing for at least 30 min. Cells were not responsive to DEP at low frequencies, thus we focused analyses when cells were visually responsive, between 13 and 17 MHz at a constant voltage of 8 Vpp. We selected this voltage because cells were minimally reactive below 7 Vpp but were trapped on the electrodes at voltages greater than 9 Vpp. For each frequency, cells were collected.
from all three outlets, then stained with antibodies against CD45 (pan-leukocyte), CD3 (T-cell receptor), CD14 (macrophage), and CD19 (pan B-cell) (Fig. 2a). Depletion of each cell type was calculated by flow cytometry analysis of the outlets where the percent of CD3\(^+\), CD14\(^+\), or CD19\(^+\) cells in the side channels was divided by the percent of the cell of interest from the total cells processed (CD45\(^+\)). A similar trend was observed for all populations, where 15 MHz resulted in maximum depletion from the center channel, which was most pronounced for CD3\(^+\) cells (99.3%), followed by CD14\(^+\) cells (92.5%), and CD19\(^+\) cells (88.5%). Ultimately, only 1.5% of the initial number of PBMCs (CD45\(^+\)) were collected in the central outlet (Fig. 2a). Of the subpopulations, we found CD3\(^+\) T-cells were the most responsive and had similar depletion levels between 14, 15, and 16 MHz at 97%, 99.3%, and 98.3%, respectively. In contrast, depletion of CD19\(^+\) and CD14\(^+\) cells significantly changed from 13 MHz to 15 MHz, suggesting that these cell populations were more sensitive to specific DEP parameters. We corroborated the previous findings by Gascoyne et al. that different cell types have different DEP response profiles\(^{39-41}\). Not only did we observe PBMCs depletion, we found that the PBMCs composition changed post-depletion e.g. there was 7.3-fold and 4.6-fold enrichment of CD19\(^+\) and CD14\(^+\) cells in the center channel, respectively. These results suggest that across PBMC phenotypes there is a cell type-dependent DEP response that supports their differential separation and that depletion of CD3\(^+\) cells could drive enrichment of rare cells.

**In vitro CHC DEP response characterization.** Next, to understand the DEP response of malignant cells we selected an established in vitro hybrid cell line which was studied in depth previously by Gast et al. B16M\(\phi\)-RFP, generated through the fusion of B16F10 and bone marrow-derived macrophages as a model of CHCs\(^{12}\). In addition, a breast cancer cell line, MCF7, was used as a surrogate for traditional CTCs. Previously, Gast et al. characterized the phenotypic hybrid nature of in vitro derived CHCs. Conveniently these hybrids co-expressed nuclear RFP and cytoplasmic YFP facilitating their differentiation from the MCF7-GFP cells by flow cytometry\(^{12}\). To optimize conditions for hybrid cell enrichment, cancer cells were diluted in DEP buffer at 1 x 10\(^7\)/mL and only 1 x 10\(^6\) cells were loaded into the microfluidic DEP sorting device for each parameter. To visualize cancer cell DEP response, fluorescent images were taken at three locations (Fig. 2b), near the inlet (part I), the middle after the first batch of electrodes (part II), and close to the outlets (part III). At 15 MHz/8 Vpp, both B16M\(\phi\)-RFP and MCF7-GFP were aligned with the central inlet flow. As the cells flowed through the device, the majority of MCF7 cells depleted to the sides while the majority of B16M\(\phi\) remained in the center. To quantify each sorting parameter, we processed cells in the device for at least 30 min and analyzed all the outlets by flow cytometry. Across all tested frequencies MCF7 cells were more responsive than B16M\(\phi\) cells. MCF7 cells had a minimum response at 13 MHz where 65% of the cells were depleted and a maximum response at 16 MHz where 92% were depleted to the sides (Fig. 2c). In contrast, the minimum DEP response for B16M\(\phi\) was at 19 MHz where only 6% of the cells were depleted and the maximum DEP response was at 15 MHz where only 29.5% of cells were depleted suggesting that hybrid cells have a unique non-responsive DEP profile (Fig. 2c). In parallel, we analyzed the DEP response of healthy PBMCs (CD45\(^+\)) under the same conditions. Again, PBMCs were highly responsive across all frequencies with a minimum response at

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**Fig. 2 DEP response characterization of PBMCs from healthy donors and cancer cell lines.** a. PBMC subpopulation DEP response as presented by depletion of CD3\(^+\) (blue line), CD14\(^+\) (red line), CD19\(^+\) (green line), and CD45\(^+\) (purple line) cells across frequencies, 13 MHz to 18 MHz at a constant 8 Vpp by flow cytometry. b. Fluorescent images near the inlet (Part I), at the center between batches of electrodes (Part II), and close to the outlet (Part III) of the microfluidic channel revealed depletion of MCF7 (green) and less reactive B16M\(\phi\) (red) at 15 MHz and 8 Vpp. Scale bar is 500 \(\mu\)m. c. Percentage depletion of B16M\(\phi\), MCF7, and PBMCs from the center outlet across frequencies, 13 MHz to 19 MHz, at a constant 8 Vpp by flow cytometry. d. Center enrichment heatmap with voltage (V) in the y axis and frequency (MHz) in the x axes (n = 1–5, independent experiments). e. Percentage side loss heatmap with voltage (V) in the y axis and frequency (MHz) in the x axes (n = 1–5, independent experiments).
19 MHz with 77% of the cells depleted and a maximum response at 15 MHz with 96.5% of the cells depleted from the center channel (Fig. 2c).

Overall, we found a larger differential response between B16Mϕ–RFP and PBMCs as compared to MCF7-GFP cells and PBMCs despite their close proximity in size to PBMCs suggesting that label-free size-independent enrichment was possible within our workflow. This also suggests that B16Mϕ–RFP cells inherently have different dielectric properties likely resulting from their hybrid nature. Although numerous studies focus on CTC enrichment and isolation, these strategies rely on either the inherent large size of CTCs or specific membrane protein expression. CHCs are a promising alternative as they are more abundant than CTCs, however there is a lack of technologies focused on their enrichment, a critical factor in developing CHCs as a cancer biomarker. This data highlights the potential of using DEP to enrich for CHCs and potentially other rare cellular biomarkers that are difficult to differentiate phenotypically.

Hybrid cell line spiked into healthy peripheral blood samples. A number of technologies focused on rare cell analysis rely on the isolation of single cells, which is challenging. Recently, the development of high-throughput single-cell technologies allows for the analysis of thousands of cells and demonstrates the need for rare cell enrichment rather than pure isolation. For most enrichment strategies there is a tradeoff between total enrichment and overall cellular biomass loss during processing due to a limit of detection (LOD) threshold. To find the optimal frequency and voltage parameters to enrich for CHCs, we evaluated the enrichment of B16Mϕ–RFP cells from PBMCs. PBMCs isolated from healthy subjects were spiked with 50,000 B16Mϕ cells (50,000 per 1 × 106 PBMCs), then subjected to DEP for at least 30 min, and analyzed for cells types isolated from all outlets by flow cytometry. Enrichment of hybrids in the center channel and depletion in the side channels was calculated using flow cytometric data for each outlet (see Eq. 1 and Eq. 2). Unlike other strategies for rare cell isolation focused on the absolute number of recovered rare cells, our strategy utilized the ratio of rare cells to total PBMCs as the rating of enrichment with the goal of enriching samples above the LOD for downstream analytics. This process was repeated for six frequencies, from 13–18 MHz and three voltages for each frequency, 7–9 Vpp. A heatmap was generated to assess center channel hybrid enrichment (fold change) and side channel hybrid loss (percentage loss) taking into account both the frequency and voltage applied to the device (Fig. 2d, e). The enrichment varied from 1.4-fold and 2.8-fold at 13 MHz/7 Vpp and 19 MHz/9 Vpp, respectively, to 18.6-fold and 12.7-fold at 15 MHz/8 Vpp and 15 MHz/9 Vpp, respectively. The highest B16Mϕ enrichment identified was 18.6-fold (n = 5) at 15 MHz/8 Vpp (Fig. 2d). The voltage was a strong driver of the magnitude of DEP response with greater loss of B16Mϕ cells to the sides at higher voltages (Fig. 2e). For example, at 15 MHz, there was 6.5% loss to the sides at 7 Vpp, 18.6% at 8 Vpp, and 22.7% at 9 Vpp. Overall, we found that 15 MHz and 8 Vpp provided optimal DEP parameters for B16Mϕ hybrid enrichment and potentially for patient-derived CHCs. These trends observed follow previously describe bell-shaped DEP response curves. In these models as well as in the data presented here, a narrow range of maximum DEP response can be appreciated. Overall, our unique strategy to deplete healthy PBMCs from the sample drives enrichment of the target tumor cell population.

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\text{Center Enrichment} = \frac{\text{# of B16Mϕ in center channel after sorting}}{\text{# of B16Mϕ in total}} \quad (1)
\]

\[
\text{Side Loss} = \frac{\text{# of B16Mϕ in side channels after sorting}}{\text{# of B16Mϕ in total}} \quad (2)
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According to current literature there are approximately 30 CHCs per 500,000 PBMCs in PDAC patients representing only 0.006% of total cells in blood. This overall rarity highlights a clear need for CHC enrichment for downstream analyses such as mutational profiling using ddPCR or genomic surveying using single-cell sequencing, both of which have a limit of detection (LOD) of 0.1%. With 18.6-fold enrichment using the described workflow, the microfluidic DEP sorting device described could enrich above 0.1% LOD suggesting there is utility targeting the depletion of healthy PBMCs to facilitate a more comprehensive understanding of poorly understood circulating neoplastic cells.

Finally, since a strong electric field can induce cell death, which would disrupt downstream analyses, we assessed cell viability throughout the DEP workflow. Non-viable cells can lead to aggregation and clogging, which reduces device performance and sensitivity for isolating cellular biomarkers. The low conductive DEP buffer was designed to maintain cell viability while allowing for DEP separation. In order to study cell viability, the effect of DEP buffer alone and under applied DEP was tested. A549 cells were placed in DEP buffer and cell viability was assessed over time using propidium iodide staining via flow cytometry. A549 cells had 98% viability after 30 min in DEP buffer and 97% after 120 min (Supplementary Fig. 2). To measure cell viability post-DEP processing at 15 MHz/8 Vpp, PBMCs were stained with calcien AM from the outlets and immediately measured via flow cytometry (Supplementary Fig. 2). The side outlets had 99% viability while the center outlet had 97% viability, suggesting that enrichment via DEP is compatible with analytic techniques that require cell viability, morphology, and nucleic acid integrity to be maintained.

Label-free Enrichment of KRAS mutant cells from PDAC patient samples. To evaluate the ability of the microfluidic DEP sorting device to provide relevant clinical information when assessing peripheral blood from cancer patients, we set out to demonstrate that DEP-enriched CHCs could be used to evaluate KRAS mutational status. Whole blood from four patients with PDAC (stage III-IV) was collected. All patients had clinically confirmed KRASmut primary tumors (Table 1). The majority of PDAC tumors harbor KRAS mutations and its presence in circulation, from circulating cells or cell-free DNA, is clinically

| Patient ID | Gender | Age | Primary location | Intact primary | Stage at collection | Metastasis | Treatment | 1° Tumor KRASmut |
|------------|--------|-----|----------------|----------------|-------------------|-----------|-----------|-----------------|
| PDAC3      | F      | 75  | Pancreas       | yes            | IV                | lung      | yes       | p.G12R          |
| PDAC4      | F      | 75  | Pancreas       | yes            | III               | no        | no        | p.G12R          |
| PDAC1      | F      | 53  | Pancreas       | yes            | IV                | liver     | yes       | p.G12D          |
| PDAC2      | M      | 76  | Pancreas       | yes            | III               | no        | no        | p.G12D          |

Primary (1°) tumor biopsies were clinically tested for KRAS.

Table 1 Patient demographics.
relevant for cancer detection\textsuperscript{37}. Moreover, our group has previously found that a subpopulation of FACS isolated CHCs have KRAS mutations as detected by ddPCR\textsuperscript{12,13}. To confirm the presence of CHCs in peripheral blood specimen prior to subjecting to DEP, PBMCs were isolated from whole blood and analyzed as we have previously reported\textsuperscript{12,13}. Briefly, PBMCs were adhered to glass slides then stained with antibodies against CD45 and CK, and visualized by fluorescence microscopy (Fig. 3a). The size of CHCs (\( n = 25 \)) from all patients was 9.05 ± 0.89 \( \mu \)m, and was comparable to the size of PBMCs (\( n = 25 \)) 8.57 ± 0.89 \( \mu \)m, confirming the non-significant difference in size between these two populations (Fig. 3b). As indicated, this negates the possibility of size-based isolation as a valid isolation method for CHCs further supporting the utility of label-free DEP enrichment\textsuperscript{12}. In addition, these samples had non-detectable levels of CTCs (CD45- /CK\textsuperscript{+}) by IHC, further supporting the promising nature of CHCs as an important cancer biomarker due to their detectability and potential for harboring clinically relevant information.

To demonstrate DEP-enriched CHCs harbored mutant KRAS, a smaller aliquot of patient PBMCs were subjected to DEP. A pre-sort aliquot was retained as a control ("Pre-sort", Fig. 3d). Specimens were loaded onto the device (1 × 10\(^6\) cells) and all samples were processed for at least 1 h at 15 MHz and 8 Vpp, the optimal conditions for PBMC depletion. Cells were collected from side and center outlets and each channel was subjected to DNA extraction for downstream evaluation of KRAS mutations and total DNA was quantified using Qubit\textsuperscript{TM} kits (Fig. 3c). Based on DNA concentration we found that an average of 92.6% ± 2.07 (\( n = 5 \), independent experiments) measured by ddPCR. Typically KRAS positive cells are heterozygous therefore 0.26% of the cells were positive, corresponding to one KRAS mutant cell for every 385 wild-type cells post device processing\textsuperscript{51–53}. Previously Dietz et al. found an average of 30 CHCs (\( n = 5 \)) for every 500,000 PBMCs in PDAC patients and found that only 9.1% of CHCs were KRAS mutant, therefore the enrichment presented here could be as high as 476-fold. Considering that no CTCs were found by IHC, we believe that KRAS\textsuperscript{mut} cells are likely CHCs or other important circulating tumor-derived cells. The level of enrichment presented here is sufficiently above the LOD for current single-cell technologies which could allow for a deeper understanding of circulating cancer cells and associated biology.

**Fig. 3 Downstream analysis of clinical PDAC samples processed with the DEP device.** a In situ immunofluorescence microscopy analysis of PBMCs before device processing, arrows indicate presence of CHCs (Hoescht\textsuperscript{+}/CD45\textsuperscript{−}/CK\textsuperscript{+}). Scale bar is 20 \( \mu \)m. b Average diameter (\( \mu \)m) of CHCs and PBMCs from in situ immunofluorescence images. c Average clinical sample DNA extracted from cells collected from all outlets (\( n = 4 \), independent experiments). d Percent KRAS\textsuperscript{mut} copies for each patient (\( n = 4 \), independent experiments) from all outlets compared to average KRAS\textsuperscript{mut} copies for screen-negative controls (\( n = 5 \), independent experiments) measured by ddPCR.
Conclusion

In this study, we present a unique approach addressing the challenge of targeting heterogeneous circulating biomarkers by focusing on healthy PBMCs with defined DEP response to reveal disease-specific characteristics. A DEP-driven microfluidic device fabricated with a V-shape electrode design is ideal for such a strategy. Thus, allowing for continuous mode processing of high cell density samples in a short amount of time. The utility of the workflow was optimized for the depletion of PBMCs permitted maximal enrichment for a CHC model cell line. Using this approach, the device was used to process PBMCs from four PDAC patients and was coupled with genotypic downstream analysis using ddPCR. This workflow allowed for the identification of isolated cells with KRAS mutations above the limit of detection in three out of four patients from only 2 mL of peripheral blood with 1 h of sample processing. We demonstrated DEP-isolated CHCs harbored KRAS mutations as a proof of concept, but the overall strategy is applicable to other cancers or diseases with associated cellular biomarkers to provide clinically relevant readouts.

Future work will focus on coupling this technology with droplet-based single-cell technologies where both phenotypic and genotypic information can be studied simultaneously. With further biological insights into these potential cellular biomarkers we aim to develop clinical assays beyond KRAS mutation status. With the short sample processing time, low input volume, device reproducibility, and compatibility with downstream analytics, our method of cell enrichment by DEP shows promise as a clinical technology with the potential to improve current liquid biopsy strategies of cancer diagnosis and treatment monitoring.

Materials and methods

Fabrication and operation of microfluidic DEP sorting device. The microfluidic DEP sorting device described here consists of two layers: a silicon chip with an interdigitated patterned electrode layer and a microfluidic PDMS chamber bonded through plasma treatment and heat curing. The bottom layer consists of three parts, a silicon substrate with 500 nm of wet-growth silicon dioxide (SiOx) on both sides, and 90 nm thick of e-beam deposited platinum electrodes as the conductive components for DEP actuation, and silicon dioxide passivated the edge of the electrodes. The electrodes were interdigitated-separated into two batches to form multiple independent electrodes pairs to actuate the sample in the device. The edge of the electrode is designed be 150 µm from the wall of flow channel, the silicon dioxide passivation avoided cells being trapped at the region where DEP forces are much stronger than the flow stress. The passivation ensures enough shear force to carry sorted cells along the direction of flow.

The device has two DEP actuation regions that are connected to two independent pairs of electrode pads. Each electrode pad has two identical circular areas, one was wired bonded by fast drying silver paint (Ted Pella Inc.) and connected to power amplifier (Model 9260, Tabor Electronics Inc) in series with a waveform generator (33622 A, Keysight), the other electrode pad was used to monitor voltage and frequency on the device by connected with an oscilloscope (DSOX2024A, Keysight) at the beginning of each experiment and every time parameters were adjusted. The wired device was placed on a water-cooled aluminum heat sink to keep device at or below room temperature prior to the experiment. DEP buffer was placed under vacuum for at least 30 min to degas it. 10 µm FluoSpheres™ Polystyrene Microspheres and 2.0 µm FluoSpheres™ Carboxylate-Modified Microspheres were diluted to a total of 1 × 10⁶ beads. The bead mixture was then centrifuged and suspended on 100 µl of degassed cold DEP buffer. This solution was then introduced on the microfluidic device and different conditions as described in Fig. 1c were applied. For each condition, the solution was processed for at least 30 min, outlets collected, measured using BD FACs Symphony and analyzed with FlowJo.

Cell culture. MCF7 (breast adenocarcinoma) were purchased from American Type Culture Collection (ATCC, Manassas, VA) and maintained in Dulbecco’s modified Eagle’s Media (DMEM) supplemented with 10% FBS and 1% Penicillin Streptomycin Glutamine (PSG). B16F10-macrophage hybrid (B16Mφ) were previously generated and maintained in Dulbecco’s modified Eagle’s Media (DMEM) supplemented with 10% FBS and 1% Penicillin Streptomycin Glutamine (PSG) [12].

Cell line PBMC DEP analysis. Prior to each experiment DEP buffer was placed under vacuum for at least 30 min to degas it. Desellated healthy participant whole blood was obtained from the CEDAR repository following IRB protocol into either a Heparin (BD367874) or ACD (BD364606) vacutainer tubes. Within 15 min of the blood draw, 8-10 mL of whole blood was processed following the SepMate™ (STEMCELL Technologies, cat# 85450) protocol. Cells were counted using the Countess™ II (Applied Biosystems) automated cell counter and diluted to 1 × 10⁶ cells/mL in cold PBS and stored on ice. Cell lines were trypanized, washed, and reconstituted in DEP buffer. Prior to loading into the DEP device, 100 µl of cell solution was diluted to 1 mL in DEP buffer, centrifuged for 5 min at 300g, and reconstituted in 60-80 µl DEP buffer. Reconstituted cells were loaded into the center channel inlet and collected for at least 30 min at each specific frequency and voltage configuration. In the case of the spinning experiment, 50,000 cells were added per 1 × 10⁶ PBMCs prior to loading into the device. Each outlet was collected, transferred to standard 5 mL round bottom polystyrene flow cytometry tubes, stained with AF647-CD45(BioLegend, clone H130), PE-CD3 (BioLegend, clone HI100), FA488-CD19 (BioLegend, clone HIB19) and BV500-CD14 (BD Horizon, clone M5E9) on ice at least 30 min, measured using BD FACS Canto II flow cytometry, and analyzed with FlowJo. Percentage of depletion and fold of enrichment of specific subpopulation of PBMCs were calculated using Eq. 3 and
Viability testing. To measure the viability of cells in DEP buffer alone, A549 cells were plated in either DMEM or DEP buffer on ice and serially sampled every 30 min. Each sample was stained with propidium iodide (PI), measured on BD flow cytometry Symphony, and analyzed with FlowJo. PI+ cells were considered dead or dying. To measure the viability of cells post-DEP sorting through the device, cells were collected at each outlet and stained with Calcin AM (Invitrogen™ C3099). As before, cells were flow cytometry analyzed and Calcin AM positive cells were considered viable.

Clinical sample processing for immunofluorescence. PMBCs separation using Ficoll-Paque was described in56 with some modifications. 5–10 mL of whole blood was 1:1 diluted to DPBS with 2% FBS at room temperature. Diluted blood sample was layer on top of 20 mL of Ficoll-Paque PLUS (GE Healthcare), and centrifuged at 800 rcf for 20 min. PMBCs layer was carefully extracted and resuspended in DPBS with 2% FBS. To perform with concentrated cells, the resuspended sample was centrifuged at 400 rcf for 5 min followed by resuspension with intended volume of DPBS with 2% FBS.

For evaluation of CHCs, samples were first treated with a 5% bovine serum albumin solution, followed by TrueBlack Lipofuscin Autofluorescence Quencher (Biotium), and Image-iT FX Signal Enhancer (Invitrogen), then stained with fluorescent antibodies. The percentage of YAP+ and CHC+ cells in all three outlets was 1:1 diluted to DPBS with 2% FBS at room temperature. Diluted blood sample was centrifuged at 400 rcf for 5 min followed by resuspension with intended volume of DPBS with 2% FBS. To perform with concentrated cells, the resuspended sample was centrifuged at 400 rcf for 5 min followed by resuspension with intended volume of DPBS with 2% FBS.

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25. Becker, F. F. et al. Separation of human breast cancer cells from blood by differential dielectric affinity. *Proc. Natl Acad. Sci. USA* **92**, 860–864 (1995).

26. Yang, J. Cell separation on microfabricated electrodes using dielectrophoretic/gravitational field-flow fractionation. *Anal. Chem.* **71**, 911–918 (1999).

27. Gascoyne, P. R. C., Noshari, J., Anderson, T. J. & Becker, F. F. Isolation of rare cells from cell mixtures by dielectrophoresis. *Electrophoresis* **30**, 1388–1398 (2009).

28. Esramzadeh, H., Beischlag, T. V., Cox, M. E., Parameswaran, A. M. & Park, E. J. Detection and isolation of circulating tumor cells: principles and methods. *Biotecnol. Adv.* **31**, 1063–1084 (2013).

29. Di Trapani, M., Manaresi, N. & Medoro, G. DEPArray™ system: an automatic image-based sorter for isolation of pure circulating tumor cells. *Cytom. Part A* **93**, 1260–1266 (2018).

30. Hu, X. et al. “Marker-specific sorting of rare cells using dielectrophoresis. *Proc. Natl Acad. Sci. USA* **102**, 15757–15761 (2005).

31. Smith, A. J. et al. Rapid cell separation with minimal manipulation for autologous cell therapies. *Sci. Rep.* **7**, 1–15 (2017).

32. Balasubramanian, P. et al. Antibody-independent capture of circulating tumor cells of non-epithelial origin with the ApoStream® system. *PloS ONE* **12**, e0175414 (2017).

33. Nie, X. et al. High-throughput dielectrophoretic cell sorting assisted by cell sliding on scalable electrode tracks made of conducting-PDMS. *Sens. Actuators B: Chem.* **327**, 128873 (2021).

34. Wu, Y., Ren, Y., Tao, Y., Hou, L. & Jiang, H. High-throughput separation, trapping, and manipulation of single cells and particles by combined dielectrophoresis at a bipolar electrode array. *Anal. Chem.* **90**, 11461–11469 (2018).

35. Song, H. et al. Continuous-flow sorting of stem cells and differentiation products based on dielectrophoresis. *Lab Chip* **15**, 1320–1328 (2015).

36. Jiang, A. Y. L. et al. High-throughput continuous dielectrophoretic separation of neural stem cells. *Biomicrofluidics* **13**, 064111 (2019).

37. Cheng, J. et al. Nanotechnology-assisted isolation and analysis of circulating tumor cells on microfluidic devices. *Micromachines* **11**, 774 (2020).

38. Crane, J. S. & Pohl, H. A. Theoretical models of cellular dielectrophoresis. *J. Theor. Biol.* **37**, 15–41 (1972).

39. Shim, S., Stemke-Hale, K., Noshari, J., Becker, F. F. & Gascoyne, P. R. C. Dielectrophoresis has broad applicability to marker-free isolation of tumor cells from blood by microfluidic systems. *Biomicrofluidics* **7**, https://doi.org/10.1063/1.4774307 (2013).

40. Gascoyne, P. R. C., Shim, S., Noshari, J., Becker, F. F. & Stemke-Hale, K. Correlations between the dielectric properties and exterior morphology of cells revealed by dielectrophoretic field-flow fractionation. *Electrophoresis* **34**, 1042–1050 (2013).

41. Gascoyne, P. R. C. & Shim, S. Isolation of circulating tumor cells by dielectrophoresis. *Cancers (Basel)* **6**, 545–579 (2014).

42. Jiang, T. et al. Dielectrophoresis response of water-in-oil-in-water double emulsion droplets with singular or dual cores. *Micromachines* **11**, 1121 (2020).

43. Motria, K. et al. Clonal evolution of acute myeloid leukemia revealed by high-throughput single-cell genomics. *Nat. Commun.* **11**, 1–17 (2020).

44. Dong, L., Wang, S., Fu, B. & Wang, J. Evaluation of droplet digital PCR and next generation sequencing for characterizing DNA reference material for KRAS mutation detection. *Sci. Rep.* **8**, 9650 (2018).

45. Performance of the Tapestr™ Platform for Single-Cell Targeted DNA Sequencing [White Paper]. Mission Bio (2019). https://missionbio.com/wp-content/uploads/2019/10/WhitePaper_MissionBio_TapestrPlatform_RevA.pdf.

46. Azzurra, D. et al. Nanofluidic digital PCR for KRAS mutation detection and quantification in gastrointestinal cancer. *Clin. Chem.* **58**, 1332–1341 (2012).

47. Waters, A. M. & Der, C. J. KRAS: The critical driver and therapeutic target for pancreatic cancer. *Cold Spring Harb. Perspect. Med.* **8**, a031435 (2018).

48. Ming, W. et al. Two distinct subtypes revealed in blood transcriptome of breast cancer patients with an unsupervised analysis. *Front. Oncol.* **9**, 985 (2019).

49. Lian, J., Yue, Y., Yu, W. & Zhang, Y. Immunosenesence: a key player in cancer development. *J. Hematol. Oncol.* **13**, 1–18 (2020).

50. Kulemann, B. et al. Pancreatic cancer: circulating tumor cells and primary tumors show heterogeneous KRAS mutations. *Sci. Rep.* **7**, 1–11 (2017).

51. Almoguera, C. et al. Most human carcinomas of the exocrine pancreas contain mutant c-K-ras genes. *Cell* **53**, 549–554 (1988).

52. Soh, J. et al. Oncogene mutations, copy number gains and mutant allele specific imbalance (MASI) frequently occur together in tumor cells. *PLoS ONE* **4**, https://doi.org/10.1371/journal.pone.0007464 (2009).

53. Aguirre, A. J. et al. Activated Kras and Inktta/Arf deficiency cooperate to produce metastatic pancreatic ductal adenocarcinoma. *Genes Dev.* **17**, 3112–3126 (2003).

54. Tapestr™. Tapestr™ Single-cell Custom DNA and Protein Panels Uncover genotypic and phenotypic insights simultaneously from single cells. (2020). Accessed 16 Mar 2021 [Online]. Available: www.missionbio.com.

55. Qian, J. et al. A pan-cancer blueprint of the heterogeneous tumor microenvironment revealed by single-cell profiling. *Cell Res.* **30**, 745–762 (2020).

56. El-Benna, J. & My-Chan Dang, P. Analysis of protein phosphorylation in human neutrophils. *Methods Mol Biol.* **412**, 85–96 (2007).

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Author contributions

J.M.M., A.A.S., and Y.-J.C. designed, performed, and analyzed the experiments. J.M.M., A.A.S., and Y.-J.C. wrote and edited the manuscript. J.M.F., J.B.A., and Y.-J.C. initiated the idea and obtained funding.

Competing interests

The authors declare no competing interests.

Additional information

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