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Dielectrophoretic enrichment of live chemo-resistant circulating-like pancreatic cancer cells from media of drug-treated adherent cultures of solid tumors†

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Due to low numbers of circulating tumor cells (CTCs) in liquid biopsies, there is much interest in enrichment of alternative circulating-like mesenchymal cancer cell subpopulations from *in vitro* tumor cultures for utilization within molecular profiling and drug screening. Viable cancer cells that are released into the media of drug-treated adherent cancer cell cultures exhibit anoikis resistance or anchorage-independent survival away from their extracellular matrix with nutrient sources and waste sinks, which serves as a pre-requisite for metastasis. The enrichment of these cell subpopulations from tumor cultures can potentially serve as an *in vitro* source of circulating-like cancer cells with greater potential for scale-up in comparison with CTCs. However, these live circulating-like cancer cell subpopulations exhibit size overlaps with necrotic and apoptotic cells in the culture media, which makes it challenging to selectively enrich them, while maintaining them in their suspended state. We present optimization of a flowthrough high frequency (1 MHz) positive dielectrophoresis (pDEP) device with sequential 3D field non-uniformities that enables enrichment of the live chemo-resistant circulating cancer cell subpopulation from an *in vitro* culture of metastatic patient-derived pancreatic tumor cells. Central to this strategy is the utilization of single-cell impedance cytometry with gates set by supervised machine learning, to optimize the frequency for pDEP, so that live circulating cells are selected based on multiple biophysical metrics, including membrane physiology, cytoplasmic conductivity and cell size, which is not possible using deterministic lateral displacement that is solely based on cell size. Using typical drug-treated samples with low levels of live circulating cells (<3%), we present pDEP enrichment of the target subpopulation to ~44% levels within 20 minutes, while rejecting >90% of dead cells. This strategy of utilizing single-cell impedance cytometry to guide the optimization of dielectrophoresis has implications for other complex biological samples.

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Introduction

An overwhelming majority of cancer fatalities are attributed to metastasis,^{1,2} motivating the interest in developing

therapeutics that target this mechanism.³ This need is especially critical for pancreatic cancer arising from pancreatic ductal adenocarcinoma (PDAC), which is the third leading cause of cancer deaths⁴ and has the shortest survival duration, due to its propensity for tumor metastasis.^{5–7} However, due to intra-tumoral heterogeneity, only a subpopulation of the parent cancer cells exhibits the ability for metastasis.⁸ Identifying this subpopulation across each of the multiple steps in the metastatic cascade can be challenging. An important first step in this cascade is the ability of cancer cells from an adherent solid tumor to survive as suspended cells that can enter the circulatory system,⁹ away from their extracellular matrix that provides them nutrient sources and waste sinks. This so-called anoikis resistance or anchorage-independent survival characteristic of chemo-resistant cancer cells after drug treatment serves as a

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pre-requisite for metastasis.¹⁰ Currently, circulating tumor cells (CTCs) serve as the leading model for such subpopulations,¹¹ but the lack of reliable markers for their identification¹² and their occurrence at ultra-low concentrations within liquid biopsies of PDAC patients¹³ has led to much interest in alternative sources of circulating-like cancer cell subpopulations for utilization within drug screening.^{14,15}

Recent studies suggested that live cancer cells released into the suspension of drug-treated adherent cancer cell cultures exhibit chemo-resistance and tumor initiating properties.^{16–18} Furthermore, systematic alterations to their culture conditions, such as shear flow,^{19,20} hypoxia,^{21,22} and nutrients,^{23,24} can be used to generate greater numbers of live suspended cells that exhibit mesenchymal phenotypes. These cultures can serve as an *in vitro* source of circulating-like cancer cells with greater potential for scale-up, in comparison with CTCs. However, selective isolation of low levels of live circulating cells from the drug-treated culture media, while maintaining the cells in their suspended state, is challenging, due to their wide size distribution that overlaps with dead cells in the media.¹⁸ For this purpose, we present utilization of impedance cytometry for optimization of a flowthrough high frequency (1 MHz) dielectrophoresis (DEP) device with sequential 3D field non-uniformities to enrich the live chemo-resistant circulating cancer cell subpopulation from an *in vitro* culture of metastatic patient-derived PDAC cells.

Dielectrophoresis (DEP)²⁵ has been widely utilized for phenotype-selective cell deflection to enable marker-free isolation of CTCs,^{26,27} while maintaining cells in their suspended state,²⁸ including single-cell trapping approaches²⁹ that enable the study of molecular signatures³⁰ and biomechanical characteristics.³¹ However, planar electrode designs limit the throughput for cell deflection due to their limited spatial electric field gradient across the channel depth. While this may be alleviated by using higher voltages, it usually occurs at the cost of inducing irreversible pDEP trapping and cell electroporation. Recent approaches using 3D electrodes to pattern the electric field across the channel depth address this issue.^{32,33} Our prior work created a set of sequential lateral field non-uniformities orthogonal to the sample flow and extending over the device depth,³⁴ to enable DEP deflection at 10–20 $\mu\text{L min}^{-1}$ flow rates, along with an on-chip means for cell exchange into an optimal buffer prior to DEP.³⁵ Since target CTCs have a more folded cell membrane structure and are larger in size on average than the background blood cells,^{36,37} CTCs can be selected over blood cells by positive DEP (pDEP) based on their lower DEP crossover frequency^{38,39} due to their larger size and higher membrane capacitance. However, the live chemo-resistant PDAC cell subpopulation suspended in the culture media exhibits a wide size distribution that overlaps with dead cells suspended in the media, which includes smaller apoptotic cells and larger necrotic cells from the adherent culture. To address the challenge of selecting small live

suspended PDAC cells (<average in the size histogram) with lower DEP trapping force, while rejecting larger dead PDAC cells that exhibit a volumetric enhancement to their DEP trapping force, single-cell impedance cytometry^{40–42} is used to optimize pDEP deflection at high frequencies (1 MHz). This enables biophysical metrics of the cell membrane and cytoplasm to be used along with cell size as bases for pDEP selection of cells. Furthermore, by carrying out cell deflection by pDEP in the MHz range, rather than in the vicinity of its crossover from nDEP (<0.1 MHz),^{27,36} the likelihood of cell viability loss due to transmembrane potential drop that occurs for pDEP near the crossover is reduced.⁴³ Single-cell impedance cytometry data gated by supervised machine learning to identify live *vs.* dead cells, despite their overlapping size distributions, are utilized to identify the DEP frequency range that enables pDEP enrichment of live circulating PDAC cells, as validated by live/dead fluorescence assays. This includes live cells of smaller size (<mean in the histogram) despite the lower volumetric contribution to their DEP trapping force. On the other hand, using size-based microfluidic isolation by deterministic lateral displacement (DLD), while most of the larger sized live cells can be selected into the displaced outlet, substantial numbers of larger-sized dead PDAC cells are also selected into the displaced outlet and smaller-sized live circulating PDAC cells are not selected into the displaced outlet due to their zigzag motion through the posts. In this manner, live chemo-resistant circulating-like pancreatic cancer cells can be enriched by pDEP deflection from the dilute fractions typically generated within *in vitro* cultures (<10% of total cell numbers). This sets the roadmap for future application to enrich CTC-like mesenchymal phenotypes from other *in vitro* cultures for emerging needs within molecular profiling, drug screening and subpopulation classification.

Results

Device for enriching suspended live PDAC cells

An overview of the device set-up and sample steps for enriching live PDAC cells from the suspension of adherent cultures is shown in Fig. 1. Adhered PDAC cultures (obtained from mouse xenograft models of metastatic tumors derived from patient T608) are subjected to 1 $\mu\text{g mL}^{-1}$ gemcitabine treatment for 48 h (Fig. 1A), which is known to induce apoptosis within the majority of the adhered cells (>50%).^{44,45} The culture supernatant is composed chiefly of suspended dead cells released from the adhered culture, as well as limited numbers of live cells in the suspension (1–10% of total cell numbers depending on culture conditions) that exhibit anoikis resistance (Fig. 1A(ii)). Enrichment of the supernatant sample for live PDAC cells in the suspension is studied in a flowthrough microfluidic device by DEP (Fig. 1B) and DLD methods (Fig. 1C), using sheath flows to focus the cell streamlines in the device. For creating field non-uniformities to initiate DEP, Field's metal is introduced in the liquid state at 65 °C, which solidifies at room



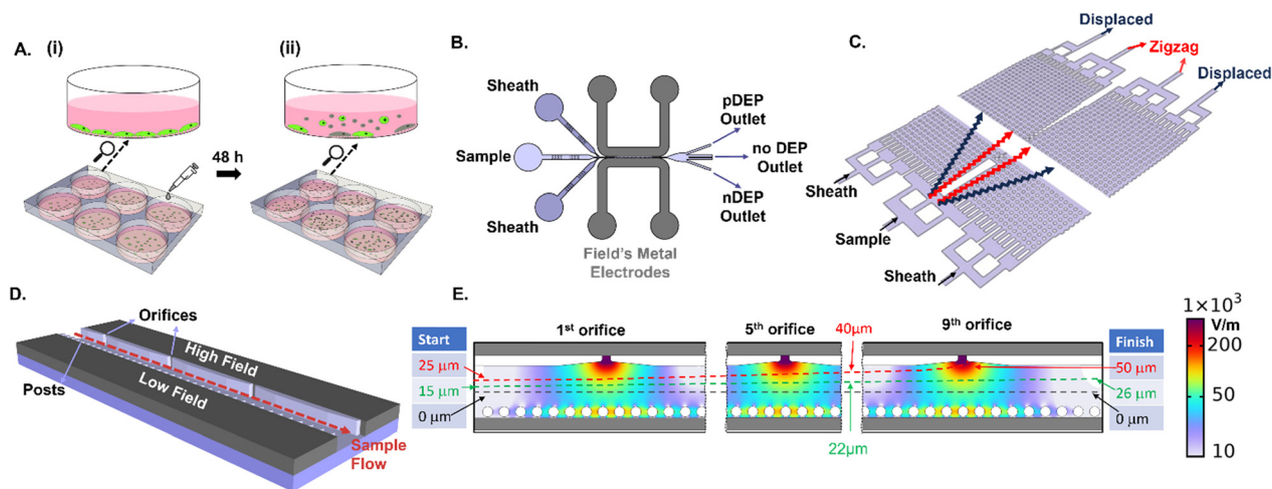


Fig. 1 Overview of the device and sample preparation steps. A. (i) Adherent PDAC cultures are drug treated at $1 \mu\text{g mL}^{-1}$ for 48 h to induce apoptosis and (ii) suspended cells released into the culture media are enriched for the live chemo-resistant population by: B. positive dielectrophoresis (pDEP) and C. deterministic lateral displacement (DLD). D. Schematic 3D view of the DEP device containing sequential field non-uniformities under voltage from fillable Field's metal electrodes in an adjoining channel, using orifices to create high field points and posts that create a continuous metal layer of low field. E. Simulated electric field profiles and flow focused streamline of the cell position at cross-sectional width after pDEP deflection at each orifice from the first orifice (start) up to the ninth orifice (finish). Optimal focusing of the start position (e.g., green streamline) enables successive pDEP deflection along device length without entrapment at the orifices, whereas a start position closer to the orifice (e.g., red streamline) exhibits entrapment due to pDEP at the ninth orifice, and a start position at the center of the channel cross-sectional width (e.g., black streamline) is too far from the high field regions to exhibit significant pDEP deflection.

temperature to fill the electrode channels that adjoin the sample channel. Voltage application initiates spatial 3D field non-uniformities across the sample channel width, due to orifices on one side that create high electrical field points and posts on the other side that create a continuous metal layer of low electric field (Fig. 1D). Focusing of the cell sample streamline with respect to the electric field profiles in the device is a key feature of the design to achieve effective pDEP enrichment without cell viability loss due to pDEP entrapment at electrodes. Using sheath flows, cell streamlines from the sample are focused at the cross-sectional width of $15 \mu\text{m}$ from the channel center towards the high field region (green streamline of Fig. 1E). On the one hand, this position enables continuous pDEP deflection under the high spatial field extent from each of the nine successive orifices, which would not be possible if the cell streamline was focused right at the center

cross-section ($50 \mu\text{m}$ from the channel edge, as per the black streamline of Fig. 1E and S2†). On the other hand, this starting position (green streamline) ensures that after pDEP deflection towards the high field initiated by each of the nine sequential orifices, the net cell streamline leads to minimal entrapment and electroporation at any of the orifice tips, up to the last (or ninth) orifice (for instance, the red streamline of Fig. 1E shows entrapment). This is confirmed by particle tracing simulations and cell imaging experiments.

Metrics for analyzing live cells in suspension

The phenotypes of live PDAC cells suspended in the media were compared to those of dead cells in the media, as well as to those of live adherent cells trypsinized from the culture. The metrics of cell size (Fig. 2A(i)), impedance phase (ϕZ in Fig. 2A(ii)), EpCAM (Fig. 2B(i)) and vimentin expression (Fig. 2B(ii)) and vimentin expression

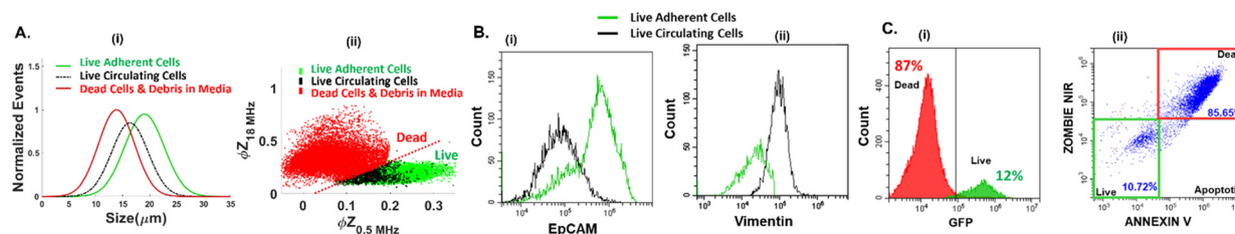


Fig. 2 A. (i) Overlap in electrical cell size distribution of respective cell types; (ii) live cells can be gated from dead cells using ϕZ metrics at multiple frequencies (based on $\sim 10\,000$ events in impedance cytometry – see ESI† Fig. S1 for other metrics and Fig. S8† for gate optimization by supervised learning). B. (i) EpCAM and (ii) vimentin expression (15 000 events in flow cytometry). C. Comparison of the same heterogeneous cell sample for: (i) GFP expression and (ii) live/dead staining, after collection from the pDEP device under E-field OFF conditions (refer to Fig. S3† for collection under E-field ON conditions).



(Fig. 2B(ii)) were used for this purpose. Additional markers, including fibroblast activation protein, have been identified for isolation of the subpopulation of metastatic circulating cells.^{46,47} Using impedance cytometry, four frequencies (0.5 MHz, 2 MHz, 18 MHz, and 30 MHz) are simultaneously applied to compute impedance magnitude ($|Z|$) and phase (ϕZ) at each frequency for each cell. The electrical size of single-cell events can be quantified by normalization of $|Z_{0.5\text{MHz}}|$ to that of polystyrene beads of known size.⁴² This is consistent with forward scattering flow cytometry (ESI,† Fig. S7). The ϕZ value at low vs. high frequency (0.5 vs. 18 MHz in the scatter plot of Fig. 2A(ii)) can be used to gate live suspended cells.⁴⁴ Using supervised learning based on the support vector machine (SVM) model to train impedance metrics with known samples of dead and live PDAC cells, the specific metrics that distinguish live vs. dead cells were identified to arise from comparison of the impedance phase (ϕZ) at low frequency (0.5 MHz) to that at high frequency (18 MHz or 30 MHz). This enabled computation of the hyperplane for label-free gating of live vs. dead cells (ESI,† section B1, Fig. S8 & Methods section). As per the sloped line in Fig. 2A(ii), live cells are roughly in the region of higher $\phi Z_{0.5\text{MHz}}$ levels due to effective electric field screening by their intact plasma membrane, and lower $\phi Z_{18\text{MHz}}$ levels due to lack of alteration in cytoplasmic conductivity by culture media penetration, as would be observed for dead cells with disrupted plasma membranes. In this manner, cell sizes in the heterogeneous sample can be attributed to live vs. dead circulating cells. Based on this, live circulating cells exhibit wide size distributions that overlap with dead cells in the suspension and trypsinized live adherent cells, with their average size falling between those of the dead cells and live adherent cells (Fig. 2A(i)). The remaining live PDAC cells after drug treatment in the adherent culture and in suspension are associated with chemo-resistant subpopulations.⁴⁵ Interestingly, live circulating PDAC cells exhibit a high degree of phenotypic similarity to the chemo-resistant live adherent PDAC cell subpopulation across every measured impedance metric (ESI,† section A, Fig. S1). However, live circulating PDAC cells exhibit lower levels of EpCAM expression and higher levels of vimentin expression vs. live adherent PDAC cells (Fig. 2B(i) and (ii)), indicating the onset of mesenchymal characteristics for this chemo-resistant subpopulation, resembling those of CTCs, thereby highlighting the interest in enriching these circulating-like cancer cells. Due to the low proportion of live circulating cells, GFP expressing PDAC cells are used to optimize conditions for their microfluidic enrichment, since live PDAC cells retain their GFP signal level, while dead cells lose their GFP signal. Using GFP-expressing cancer cells, the GFP expression level (Fig. 2C(i)) corresponds to that from standard live/dead staining (Fig. 2C(ii)), using flow cytometry of a typical heterogeneous sample (12% live cell proportions based on the GFP level vs. 10.72% based on

the live/dead assay under E-field OFF conditions). Results from live/dead staining of the pDEP enriched fraction under E-field ON conditions are shown in ESI,† Fig. S3, to validate pDEP enrichment based on the live/dead assay. While live/dead staining is the accurate live-cell quantification method, it requires extensive sample preparation and staining of the collected fraction at each outlet, which can lead to considerable cell loss. On the other hand, live-cell quantification by monitoring of GFP expression and impedance metrics requires no sample preparation. Hence, we utilize it in subsequent sections to optimize pDEP enrichment.

Optimizing field conditions for live PDAC cell enrichment

The low numbers and proportions of live circulating cancer cells in typical drug-treated samples (1–10% depending on culture conditions) vs. dead cells in the media, make it challenging to optimize pDEP conditions for enriching live circulating cells. Hence, the sample of suspended cells in the drug-treated media was mixed with the trypsinized adherent cell sample in equal parts (2 wells of suspended cells with 2 wells of lifted adherent cells from 6-well plates) for the results reported in Fig. 3 (input sample of ~45% viability based on the GFP expression level) and in majority parts (3 wells of suspended cells with 1 well of lifted adherent cells from 6-well plates) for the results reported in Fig. 4 (input sample of ~25% viability based on GFP expression) and finally, using only suspended cells for the results in Fig. 5 (input sample of ~3% viability based on GFP expression). For quantifying pDEP collection from dilute input samples, the GFP expression level and impedance metrics were used to

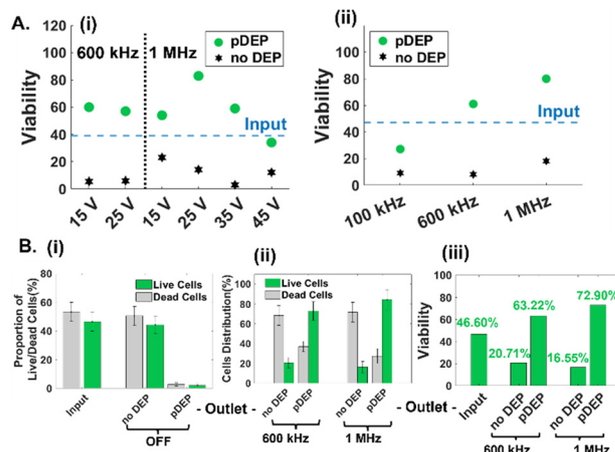


Fig. 3 A. Optimization of applied voltage (i) and frequency (at 25 V_{pp}) (ii), for maximizing enrichment of viable cells (GFP+) in the pDEP fraction using a sample input with ~45% live PDAC cells. B. (i) Proportion of live vs. dead cells (GFP-) in the input at “no DEP” and “pDEP” outlets under field OFF conditions. (ii) Distribution of cells to the “no DEP” and “pDEP” outlets under field ON conditions (25 V_{pp}) at 600 kHz and 1 MHz pDEP trapping frequencies shows the collection efficiency of ~85% for pDEP at 1 MHz. (iii) Selection purity determined based on viable cells % in each outlet at each DEP frequency.



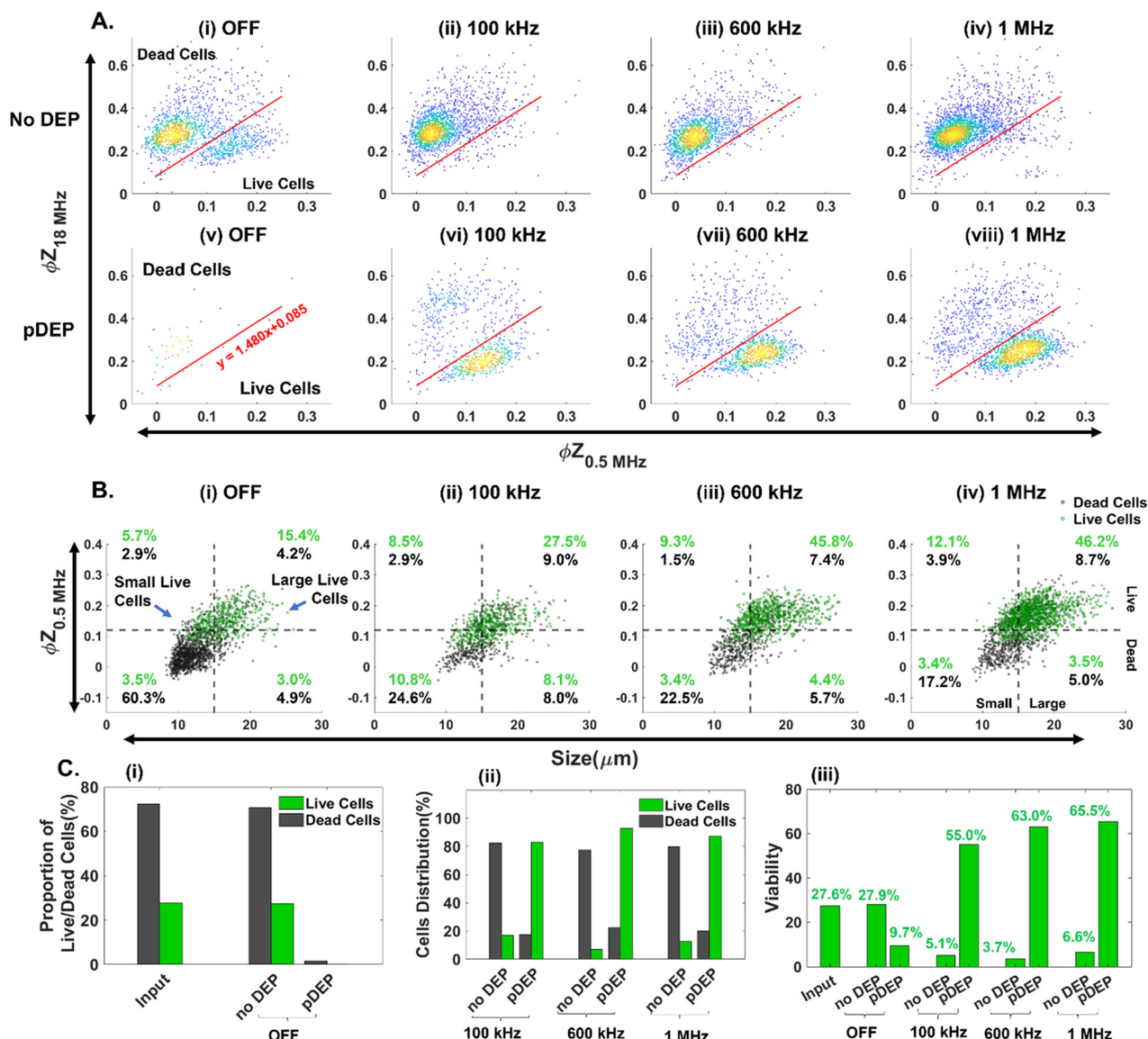


Fig. 4 Using impedance cytometry to optimize the frequency for pDEP enrichment (at 25 V_{pp}) of live cells over the entire size range of an input sample ($\sim 27\%$ live PDAC cells). Single-cell impedance scatter plots of: A) $\phi Z_{18\text{ MHz}}$ vs. $\phi Z_{0.5\text{ MHz}}$ for live cell gating of the sample from “pDEP” and “no DEP” outlets; and B) $\phi Z_{0.5\text{ MHz}}$ vs. electrical size for size-stratified live vs. dead cells in the input and pDEP outlets post-DEP. C) Quantifying collection at the “pDEP” and “no DEP” outlets (25 V_{pp} at 0.1–1 MHz) based on: (i) proportion of live and dead cells; (ii) cell distribution in each outlet at different DEP trapping frequencies and (iii) viability of collected cells. All live vs. dead cell gates (lines in 2D plots) are obtained based on the optimized hyperplane from the SVM supervised learning model.

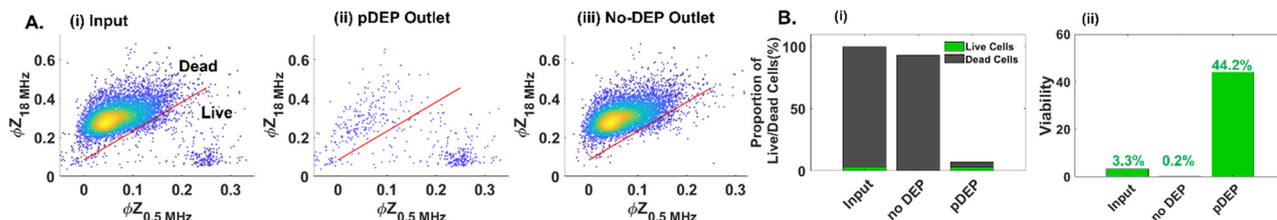


Fig. 5 DEP enrichment of samples with low levels ($\sim 2\text{--}3\%$) of only live PDAC floating cells obtained from the cell culture supernatant after 48 h gemcitabine treatment (*i.e.*, 100% floating cells). A. Impedance cytometry-based live cell gating of input and collected cells in the “pDEP” and “no DEP” outlets. B. Using impedance cytometry to quantify live cell proportions (i) and viability levels (ii) for cells collected in the “pDEP” and “no DEP” outlets. All live vs. dead cell gates (line in 2D plots) are obtained based on the SVM model (Fig. S8†).



minimize sample preparation steps that could lead to cell loss. Following optimization of the cell streamline by flow focusing prior to flowthrough pDEP deflection (Fig. 1E), while ensuring that the deflection does not cause cell entrapment at any of the high field points up to the last orifice, we focused on optimization of field conditions of applied peak-to-peak AC voltage ($15\text{--}45 V_{pp}$) over $100\ \mu\text{m}$ spacing and frequency ($0.1\text{--}1\ \text{MHz}$). As per Fig. 3A(i), while the proportion of viable cells in the pDEP fraction increases from 15 to 25 V_{pp} levels due to greater DEP trapping force, it drops off at higher V_{pp} levels. This is likely due to cell deflection towards the vicinity of high field regions that lead to cell viability loss due to their entrapment and electroporation. Using $25 V_{pp}$, pDEP deflection at $1\ \text{MHz}$ led to higher proportions of viable cells in the pDEP fraction *vs.* that at lower frequencies (Fig. 3A(i) and (ii)). We did not explore frequencies beyond $1\ \text{MHz}$, due to decay in the frequency response of commercial amplifiers.⁴⁸ Based on three independent runs under these optimized conditions ($25 V_{pp}$ at $1\ \text{MHz}$) using an input PDAC sample of $\sim 45\%$ live cells (Fig. 3B(i)), it is apparent that the entire sample (live and dead cells) passes undeflected into the “no DEP” outlet under field OFF conditions. This is apparent in a representative run (Fig. S4[†]), wherein $>95\%$ of the input sample (based on 650 live cell events collected in 15 minutes) is collected into the “no DEP” outlet. Live cells from the input sample are predominantly collected into the pDEP outlet at an $\sim 75\%$ level at $600\ \text{kHz}$ and $\sim 85\%$ level at $1\ \text{MHz}$ pDEP trapping frequency (Fig. 3B(ii)), with minimal loss into the “no DEP” outlet (18% at $600\ \text{kHz}$ and 10% at $1\ \text{MHz}$). The selection purity, as measured by the viability of the collected sample, shows that the input sample of $\sim 45\%$ live cells is enriched to $\sim 63\%$ live cells at $0.6\ \text{MHz}$ and to $\sim 73\%$ live cells under $1\ \text{MHz}$ conditions (Fig. 3B(iii)). For the representative run (ESI[†] Fig. S4), under field ON conditions at $25 V_{pp}$ and $1\ \text{MHz}$, the entire live cell fraction (GFP+) gets deflected to the “pDEP” outlet ($\sim 90\%$), with only limited live cell proportions passing through undeflected to the “no DEP” outlet ($\sim 10\%$). These optimized field conditions ($25 V_{pp}$ at $1\ \text{MHz}$) are subsequently explored at successively lower live cell proportions and using impedance cytometry to optimize DEP enrichment of dilute samples of live floating PDAC cells.

Impedance cytometry to optimize live PDAC cell enrichment

In comparison with GFP expression for flow cytometry-based quantification of live *vs.* dead PDAC cells, impedance cytometry data allow for multiparametric label-free cell quantification, based on their electrical size ($d = \sqrt[3]{|Z|_{0.5\text{MHz}}}$), apoptotic state⁴⁴ (*i.e.*, live *vs.* early apoptotic, late apoptotic and necrotic states based on $\phi Z_{0.5\text{MHz}}$ *vs.* $\phi Z_{18\text{MHz}}$), cell membrane ($|Z|$ and ϕZ in the $1\text{--}10\ \text{MHz}$ range) and cell cytoplasmic properties ($|Z|$ and ϕZ in the $>10\ \text{MHz}$ range). This allows for optimization of pDEP conditions by quantification of field-induced damage to the DEP enriched fractions, without needing to use GFP expressing PDAC cells. Live cell gating can be established based on the SVM model

(optimized gate in Fig. 4A), with live cells roughly showing higher $\phi Z_{0.5\text{MHz}}$ levels (>0.125 at mean cell size) due to effective electric field screening by the intact plasma membrane, and lower $\phi Z_{18\text{MHz}}$ levels (<0.2 at mean cell size) due to the absence of any alteration to their cytoplasmic conductivity by penetration of the culture media that is observed for dead cells with disrupted plasma membranes. Using an input sample of $\sim 27\%$ live PDAC cells, it is apparent that under field OFF conditions, cells pass undeflected into the “no DEP” outlet, with very few cell events in the pDEP outlet (Fig. 4A(i), (v) and C(i)). At successively higher frequencies of $0.1\ \text{MHz}$ (Fig. 4A(ii)), $0.6\ \text{MHz}$ (Fig. 4A(iii)) and $1\ \text{MHz}$ (Fig. 4A(iv)), the number of events in the live cell gates successively increases. Next, we use the impedance scatter plot (Fig. 4B) of $\phi Z_{0.5\text{MHz}}$ *vs.* electrical size of cells to quantify the live cell #s (gated based on the SVM model) that are enriched in the pDEP outlet for each size range of interest. Using the intersection point of $15\ \mu\text{m}$ for the histograms of live circulating cells *vs.* dead cells within the suspension of the culture media after gemcitabine treatment (Fig. 2A(i)), cell events of live circulating cells $< 15\ \mu\text{m}$ are called “small live cells” since they overlap with dead cells in the suspension, while those $> 15\ \mu\text{m}$ are called “large live cells”. Based on this, it is apparent that live cell events increase successively in the “large live cell” gate for pDEP collected fractions at $0.1\ \text{MHz}$ and $0.6\ \text{MHz}$, while remaining substantially unchanged at $1\ \text{MHz}$. On the other hand, pDEP at $1\ \text{MHz}$ is crucial for increasing the proportion of events in the “small live cell” gate of the pDEP collected fractions. The summary plot (Fig. 4C(i) and (ii)) confirms $\sim 80\%$ collection of live cells into the pDEP outlet at $100\ \text{kHz}$ and $\sim 90\%$ collection at $600\ \text{kHz}$ and $1\ \text{MHz}$ DEP trapping frequencies, with $1\ \text{MHz}$ enabling the enrichment of “small live PDAC cells”. Selection purity plots (Fig. 4C(iii)) show that the input PDAC sample of $\sim 27\%$ live cells is enriched in the pDEP outlet to 55% live cells at $0.1\ \text{MHz}$, to 63% at $0.6\ \text{MHz}$ and to 65.5% at $1\ \text{MHz}$.

DEP enrichment from rare samples of live PDAC floating cells

These optimized field parameters ($25 V_{pp}$ at $1\ \text{MHz}$) and live-cell gating conditions (SVM model for ϕZ) are used to quantify pDEP enrichment of low levels of live PDAC circulating cell samples ($2\text{--}3\%$ live cells in a well plate after $48\ \text{h}$ gemcitabine treatment, as per numbers and sizes in ESI[†] Fig. S7). Based on the ϕZ gate from the SVM model (Fig. 5A), enrichment of low levels of live cells from the input sample (Fig. 5A(i)) into the “pDEP” outlet is apparent, with deflection of only minimal levels of dead cells (Fig. 5A(ii)), while the “no DEP” outlet is composed predominantly of dead cells (Fig. 5A(iii)). This is also apparent in the plots based on the ϕZ gate (Fig. 5B(i)) and GFP expression level (ESI[†] Fig. S5A). The proportion of viable cells increases from $\sim 3\%$ in the input sample to $\sim 44\%$ collected in the pDEP outlet within 20 minutes (Fig. 5B(ii)), based on the ϕZ gate



and based on the GFP expression level in ESI† Fig. S5B. This demonstrates pDEP enrichment from dilute samples ($\sim 3\%$) to $\sim 44\%$, at a collection efficiency of $\sim 60\%$, while rejecting $>90\%$ of dead cells.

DEP vs. DLD for live PDAC cell enrichment

Next, we highlight the importance of utilizing multiple cellular biophysical metrics (size, plasma membrane and cytoplasmic properties) for enriching live circulating PDAC cells with wide and overlapping size distributions with other cell subpopulations in suspension. Impedance cytometry data (Fig. 4B(iv)) suggest that pDEP separation at 1 MHz selects live cells based on their membrane and cytoplasmic properties, with size included in the volumetric contribution to the trapping force. In contrast, microfluidic DLD only selects cells based on their size. Hence, we compare the separation metrics using DLD and pDEP for an input sample composed of $\sim 27\%$ live PDAC cells (combining 3 wells of floating cells and 1 well of trypsinized adhered cells from a 6-well plate). As per Fig. 6A(i), while the input sample is composed predominantly of live cell events that are of greater cell size than dead cell events (top right quadrant vs. bottom left quadrant), there are substantial numbers of live cell events with cell size $<15\ \mu\text{m}$ (*i.e.*, small live cells in the top left quadrant). Previously, we established that pDEP deflection at 1 MHz is needed to enrich small live cells $<15\ \mu\text{m}$ (Fig. 4B(iv)), whereas pDEP deflection at lower frequencies increases only the live cell populations of size

$>15\ \mu\text{m}$. It is also noteworthy that this input sample has a substantial number of events of size $>15\ \mu\text{m}$ in the gate of dead cells (right bottom quadrant of Fig. 6A(i)), which likely arises from necrotic cells released from the adherent culture under drug treatment. Using a DLD array designed for a cut-off size (D_C) of $15\ \mu\text{m}$,⁴⁹ we would expect that most of the live cells would be collected in the displaced outlet and most of the dead cells would be collected in the zigzag outlet. While this is indeed the case based on Fig. 6A(ii), wherein dead cells are collected in the zigzag outlet (left bottom quadrant) and live cells are collected in the displaced outlet (right top quadrant), the zigzag outlet also includes substantial live cell events for small cells ($<15\ \mu\text{m}$ in the left top quadrant) and the displaced outlet also includes substantial dead cell events arising from large cells ($>15\ \mu\text{m}$ in the bottom right quadrant). On the other hand, using pDEP deflection at 1 MHz, the subpopulation of small live cells ($<15\ \mu\text{m}$) is collected in the pDEP outlet along with other live cells (top left and top right quadrants of Fig. 6A(iii)), while the large dead cells ($>15\ \mu\text{m}$) remain undeflected and are collected in the “no DEP” outlet along with other dead cells (bottom left and bottom right quadrants of Fig. 6A(iii)). The summary plots of collected cell events at each outlet classified in Fig. 6B(i) for the proportion of live vs. dead cell events and in Fig. 6B(ii) for % viable cells show the poor separation purity of DLD vs. the high purity obtained with pDEP. Also refer to ESI† Fig. S6 for collected cell #s to compare live cell enrichment after pDEP vs. DLD. To illustrate the biophysical basis that allows pDEP to select small live cells despite the

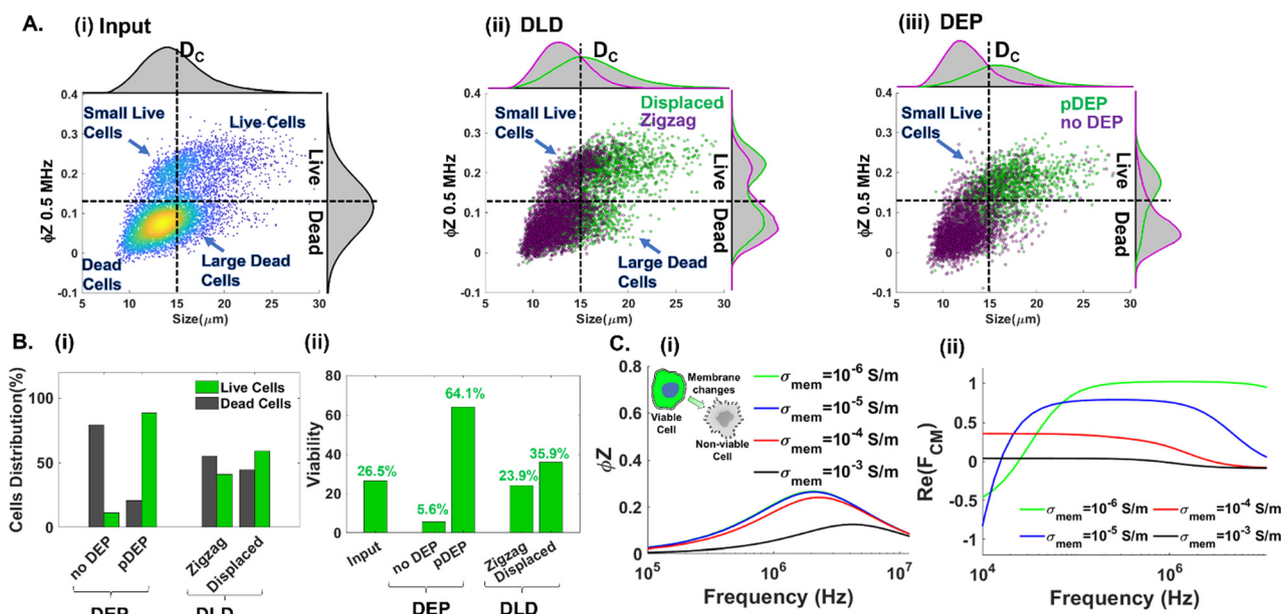


Fig. 6 Enrichment of live PDAC cells by DLD vs. DEP. **A.** Impedance scatter plots with respective histograms for: (i) the input sample (50% suspended cells and 50% trypsinized adhered cells), (ii) post-DLD enrichment and (iii) post-DEP enrichment. **B.** Summary plots of cells within the input and DEP enriched and DLD enriched outlets based on: (i) distribution of live/dead cells and (ii) % viable cells at each outlet. **C.** Frequency response of: (i) the impedance phase for viable (low membrane conductance) and non-viable cells (high membrane conductance) and (ii) $\text{Re}(f_{CM})$ for viable and non-viable cells shows that $\text{Re}(f_{CM})$ drops off to zero at $\sim 1\ \text{MHz}$ for $\sigma_{\text{membrane}} > 10^{-4}\ \text{S m}^{-1}$, causing pDEP force to sharply drop off irrespective of cell size.



lower volumetric contribution to their net trapping force and reject large dead cells despite the higher volumetric contribution to their trapping force, we consider simulations of the frequency dispersion of the impedance phase (ϕ_Z) (Fig. 6C(i)) and the real part of the Clausius Mossotti factor ($\text{Re}(f_{\text{CM}})$) (Fig. 6C(ii)) for live cells with cell membranes of low conductivity ($\sigma_{\text{mem}} \leq 10^{-4} \text{ S m}^{-1}$) and for dead cells with cell membranes of higher conductivity ($\sigma_{\text{mem}} \geq 10^{-3} \text{ S m}^{-1}$). The ϕ_Z frequency dispersion is relatively stable for σ_{mem} levels $\leq 10^{-4} \text{ S m}^{-1}$ (*i.e.*, live cells), but sharply drops off for σ_{mem} levels $\geq 10^{-3} \text{ S m}^{-1}$ (*i.e.*, dead cells), especially in the 1–2 MHz range. As a result, live cells maintain interfacial polarization under DEP at 1 MHz due to sharp differences in cell *vs.* media dielectric properties that cause high positive $\text{Re}(f_{\text{CM}})$ values at 1 MHz, whereas dead cells do not maintain sharp differences in cell *vs.* media dielectric properties, thereby exhibiting sharp drop-offs in $\text{Re}(f_{\text{CM}})$ values in the 1 MHz region. Hence, despite the large volumetric contribution to the DEP trapping force from large dead cells ($>15 \mu\text{m}$), their low $\text{Re}(f_{\text{CM}})$ value leads to minimal pDEP. Similarly, despite the small volumetric contribution to the DEP trapping force from small live cells ($<15 \mu\text{m}$), their high $\text{Re}(f_{\text{CM}})$ value at 1 MHz causes them to be selected under pDEP deflection.

Conclusions

We present a flowthrough positive dielectrophoresis (pDEP) device to enrich live circulating cancer cells at rare levels (down to $\sim 3\%$) with chemo-resistant properties (*i.e.*, high live cell numbers with drug treatment: ESI† Fig. S7A) and overlapping cell size distributions with apoptotic and necrotic cells in the culture suspension, by using impedance cytometry to optimize the pDEP frequency for selection of live cells over the entire size range (10–25 μm). A support vector machine (SVM) model is trained with live and dead cell samples to optimize gating of the impedance metrics of live *vs.* dead cells within the heterogeneous sample of broad cell size distributions, which is validated using cancer cells that lose GFP expression upon viability loss. While the subpopulation of large live circulating PDAC cells ($>15 \mu\text{m}$) can be enriched by pDEP at lower frequencies (0.1 and 0.6 MHz), pDEP at 1 MHz is essential for enriching the subpopulation of small live PDAC cells ($<15 \mu\text{m}$) which exhibits mesenchymal characteristics. This small live cell fraction cannot be enriched using size-based microfluidic DLD, due to size overlaps with apoptotic cells. Based on simulated frequency dispersions of the impedance phase and f_{CM} , we show that live cells can maintain interfacial polarization under DEP conditions at 1 MHz due to sharp differences in cell *vs.* media dielectric properties that cause high positive f_{CM} values. This enables enrichment despite lower volumetric contribution to DEP trapping force for cells of smaller size. On the other hand, dead cells (including smaller sized apoptotic and larger sized necrotic cells) are unable to maintain sharp

differences in cell *vs.* media dielectric properties, thereby exhibiting sharp drop-offs in f_{CM} values in the 1 MHz region causing minimal to no pDEP. As a result, low levels of live circulating PDAC cells with wide size distributions can be enriched from the drug-treated media of adherent cultures. Based on multiparametric information from impedance cytometry (cell size, viability, membrane capacitance, cytoplasmic conductivity, *etc.*) and its minimal sample preparation needs, we envision its broader utilization to optimize microfluidic enrichment using versatile sample types.

Methods

Cell samples

PDAC tumor samples were generated from specimens collected in collaboration with the University of Virginia Biorepository and Tissue Research Facility after approval of the University of Virginia Institutional Review Board for Health Sciences Research and after informed written consent from patients. Tumors were propagated in the pancreas of immunocompromised mice. Xenograft lines were established, transfected with GFP, and selected using puromycin and maintained in RPMI 1640 with 10% FBS and 2 mM glutamine.

Drug treatment

GFP transfected tumor cells were plated at 4×10^5 cells per ml in 6-well plates and allowed to adhere overnight in complete media (RPMI + 10% FBS). The cells were then treated with gemcitabine at a concentration of $1 \mu\text{g ml}^{-1}$ for 48 h. Floating cells were collected from the medium supernatant in the wells (centrifuged at 300g for 10 min). Adherent cells were washed in $1 \times$ PBS (Thermo Fisher) and then subjected to a 0.05% trypsin treatment (Thermo Fisher) for 10 min, and then complete media were added. The adherent cells were then centrifuged at 300g for 10 min. Cell pellets of each sample type were then resuspended in 1 mL of DEP buffer (sucrose and BSA, conductivity $45 \mu\text{S cm}^{-1}$) and counted using a hemocytometer. Floating cell samples were pDEP enriched, mixing in sample wells from adherent cells in some cases (Fig. 3 and 4), to increase live cell events for optimizing the DEP device.

Device design and fabrication

The DEP device as shown in Fig. 1B consists of two sheath inlets and one sample inlet. The active region of the device consists of a microfluidic channel flanked by two electrode channels that adjoins sequential orifices (20 μm) on one side and PDMS posts (25 μm diameter, 40 μm spacing) on the other side, to create spatial field non-uniformities across the channel cross-section orthogonal to the sample flow. The device was fabricated in a single layer to a depth of 50 μm using SU8 photolithography on a 4 inch silicon wafer. The silicon wafers were then silanized to allow for subsequent de-



molding of PDMS. PDMS (Dow Chemicals) was poured onto the silicon master for micromolding at a ratio of 10:1 (base to crosslinker) and allowed to crosslink at 70 °C for 12 hours. Devices were then punched to create inlets and outlets for the electrodes, sample, and sheath channels. Devices were bonded to glass slides using oxygen plasma. Electrode channels were filled with liquefied Field's metal as described previously.^{34,35} Briefly, the device was immersed in a water bath at 65 °C and the liquefied Field's metal (RotoMetals) was introduced through a syringe using positive pressure. After complete filling of the electrode channel, the device was allowed to cool at room temperature resulting in solidification of the metal. DLD devices were fabricated as per prior reports.⁴⁷

Device operation

Prior to introducing sample and sheath flows, the sample inlet and outlets were filled with 3% BSA (in 1× PBS) and left at room temperature for 30 min. The BSA solution was then removed. Sample and sheath fluid flows were introduced in their respective inlets using syringe pumps (Cetoni GmbH). Net flow rates of 2.64 $\mu\text{L min}^{-1}$ were used for pDEP enrichment, based on a cell sample flow rate of 0.24 $\mu\text{L min}^{-1}$, sheath flow near the orifice edge at 0.9 $\mu\text{L min}^{-1}$ and sheath flow near the posts at 1.5 $\mu\text{L min}^{-1}$. The chip was placed on a microscope stage equipped with a CMOS camera (Hamamatsu) for imaging cell streamlines. A function generator and amplifier were connected to the electrodes to deliver the studied voltage and frequency range.

Flow cytometry

After DEP enrichment, samples were stained using APC Annexin V (Biolegend) and Zombie NIR (Biolegend) to quantify viability and apoptotic levels using a Beckman Cytoflex flow cytometer and data were analyzed using Beckman CytExpert software. For analysis of EpCAM levels, samples were stained for 30 min with PE anti EpCAM (CD326) antibody (Biolegend). The samples were then centrifuged and washed twice before analysis. Cells were gated based on forward (FSC) and side scatter (SSC) to exclude debris. To exclude doublets, single cells were then gated based on side scatter – area vs. height. As a measure of viability, the intrinsic GFP signal was also measured and analyzed.

Impedance cytometry

Fractions at each outlet from the DEP device were centrifuged at 300g for 5 minutes and the resulting cell pellet was resuspended in 1× PBS along with 7 μm polystyrene reference beads for single-cell impedance cytometry using an impedance spectroscope (HF2IS, Zurich Instruments).⁵⁰ The measured current was converted to voltage using a current amplifier (HF2TA, Zurich Instruments) at a gain factor of 1000 and a sampling rate of 115 000 samples per s. Lock-in amplification was used to separate the real and imaginary

signal components at each frequency, from which the impedance magnitude and phase were derived. Cells and beads (7 μm polystyrene, Sigma Aldrich) were co-flown through a glass chip with top and bottom gold electrodes to acquire the impedance phase and magnitude at four frequencies (0.5 MHz, 2 MHz, 18 MHz, and 30 MHz) concurrently.

Data analysis

Impedance cytometry data were plotted and analyzed using MATLAB. The impedance signal of the cells was normalized by dividing it by the mean impedance of the reference polystyrene beads that show a frequency independent impedance response. Cells and beads were gated separately using impedance data at 30 MHz. The electrical diameter was calculated using the size of the reference beads and the normalized impedance magnitude at 0.5 MHz ($d = \sqrt[3]{|Z|_{0.5\text{MHz}}}$).

Machine learning

Machine learning (ML) algorithms were implemented using MATLAB. The data set composed of the phase and magnitude at 0.5, 2, 18 and 30 MHz, the electrical diameter and the impedance magnitude opacity was classified as live and dead subpopulations using SVM (support vector machine) models trained with a sample of untreated live cells and heat treated dead cells, using 70% data for testing and 30% for validation. Details are available in ESI† section B1, Fig. S8.

COMSOL simulation

To measure the spatial extent of the electric field from the high field point and its influence on particle deflection, the electric current module, flow module and particle tracing modules of COMSOL were used as described previously.³⁴

Author contributions

A. Rane: conceptualization, investigation, formal analysis, writing; J. Jarmoshti: conceptualization, investigation, formal analysis, writing; A. Siddique & Sara Adair: methodology, investigation; K. Torres-Castro & C. Honrado: methodology, data curation; T. W. Bauer: resources, supervision; N. S. Swami: conceptualization, methodology, formal analysis, resources, writing, supervision, project administration, funding acquisition.

Conflicts of interest

The authors have no financial interests or other conflicts of interest related to this work.

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