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3D microfluidic *ex vivo* culture of organotypic tumor spheroids to model immune checkpoint blockade

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Microfluidic culture has the potential to revolutionize cancer diagnosis and therapy. Indeed, several microdevices are being developed specifically for clinical use to test novel cancer therapeutics. To be effective, these platforms need to replicate the continuous interactions that exist between tumor cells and nontumor cell elements of the tumor microenvironment through direct cell-cell or cell-matrix contact or by the secretion of signaling factors such as cytokines, chemokines and growth factors. Given the challenges of personalized or precision cancer therapy, especially with the advent of novel immunotherapies, a critical need exists for more sophisticated *ex vivo* diagnostic systems that recapitulate patient-specific tumor biology with the potential to predict response to immune-based therapies in real-time. Here, we present details of a method to screen for the response of patient tumors to immune checkpoint blockade therapy, first reported in Jenkins *et al. Cancer Discovery*, 2018, **8**, 196–215, with updated evaluation of murine- and patient-derived organotypic tumor spheroids (MDOTS/PDOTS), including evaluation of the requirement for 3D microfluidic culture in MDOTS, demonstration of immune-checkpoint sensitivity of PDOTS, and expanded evaluation of tumor-immune interactions using RNA-sequencing to infer changes in the tumorimmune microenvironment. We also examine some potential improvements to current systems and discuss the challenges in translating such diagnostic assays to the clinic.

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Introduction

Monoclonal antibodies targeting immune checkpoints (*e.g.* PD-1/PD-L1 axis, CTLA-4) have demonstrated clinical activity in several malignances (Fig. 1).^{1,2} Unlike molecular targeted therapies (*e.g.* BRAF and EGFR inhibitors), which can be employed in patients whose tumors harbor select oncogenic driver mutations (*e.g.* BRAF V600E, EGFR exon19del or

L858R), to date there are no reliable predictors of response for immune checkpoint inhibitors. PD-L1 expression predicts response to anti-PD-1/PD-L1 antibodies in some, but not all patients, and is increasingly recognized as an imperfect marker of activity.³ Intense research efforts are now underway to identify predictive biomarkers of response, toxicity and resistance to immunotherapeutic agents.

There is great interest in studying other immune checkpoints and novel combinations of immune modulatory agents to overcome both innate and acquired resistance to immune checkpoint blockade (ICB).^{4,5} Dual ICB with combination PD-1 and CTLA-4 antibody treatment has recently shown dramatic response rates in patients with metastatic melanoma, however roughly half of patients experienced significant toxicity from the treatment regimen.^{6,7} Several clinical trials of combinations of immunotherapeutic agents with molecular targeted therapies (*e.g.* kinase inhibitors), cytotoxic chemotherapy, and/or radiation are underway, all in the effort to provide long-lasting disease control to more patients.⁵ While the number of these 'rational' combination trials continues

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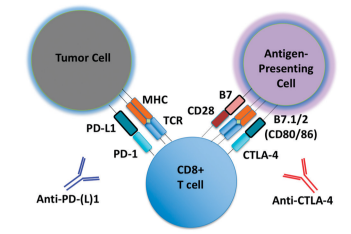


Fig. 1 Schematic of PD-1/CTLA-4 blockade. Schematic detailing basic steps involved in generation of tumor-specific T cells. Shown is a schematic of a tumor cell, CD8 effector T cell, and an antigen-presenting cell (APC), with associated cell-cell interactions *via* PD-1/PD-L1 and CTLA-4/B7. Tumor-associated antigens or neo-antigens are presented by major histocompatibility complex (MHC) on APCs or tumor cells to T cells with appropriate T-cell receptor (TCR). CD28 co-activating receptor on T cells binds B7 on APCs. Anti-PD-(L)1 and anti-CTLA-4 antibodies are shown.

to increase, our understanding of the 'backbone' therapy of PD-1 or CTLA-4 blockade remains far from complete.

Mechanisms of immune escape and resistance to ICB are diverse, owing to insufficient numbers of anti-tumor T cells present, or ineffective immune response due to local (e.g. tumor or stromal derived) factors that impair the cytotoxic immune response.8-11 Approaches to identify cellular, pathologic, immunologic, or molecular features that distinguish responding from non-responding patients have evaluated local features in the tumor microenvironment using immunohistochemical or transcriptomic evaluation of frozen or fixed tissue from biopsies, and circulating levels of select immune cells or cytokines.12,13 As biopsies from patients who have responded to ICB often demonstrate the presence of an inflammatory infiltrate within the tumor, and gene expression profiling studies have confirmed upregulation of pro-inflammatory cytokines and chemokines within tumors following PD-1 blockade, there is increasing interest in understanding the role of the tumor microenvironment in the response to ICB.^{12,13} Development of more sophisticated pre-clinical and ex vivo cancer models that recapitulate human tumor biology may facilitate efforts to predict response to targeted and immune-based therapies.

There is growing interest in capitalizing on the increasing availability of 3D culture systems to tackle important questions in cancer biology, especially with the recent integration of microfluidic systems. 3D-microfluidic culture systems now offer a greater level of precision for evaluating a host of complex biological phenomena in relevant model systems.¹⁴ With the advent of ICB and the expanding number of novel immunotherapies and combination therapies, there is increasing interest in the use of 3D culture systems to interrogate the tumor immune microenvironment.¹⁵

Recently, we described a novel method of profiling response to PD-1 blockade using organotypic tumor spheroids cultured in collagen hydrogels suspended in a 3D microfluidic device.⁵ Patient- or murine-derived organotypic tumor spheroids (MDOTS/PDOTS) retain autologous immune cells and respond to PD-1 blockade ex vivo.^{16,17} Using murine tumors from established PD-1 responsive or resistant tumors, we were able to recapitulate sensitivity and resistance to ICB ex vivo. MDOTS/PDOTS also served as a platform to test novel combination therapies in combination with PD-1 blockade, including inhibitors or TBK1 (ref. 17) and CDK4/6,¹⁷ both of which enhanced response to PD-1 blockade ex vivo and subsequently in vivo efficacy was confirmed. In this paper, we will describe in detail the methods of MDOTS/PDOTS platform ("version 1.0"), and propose some novel applications and future directions.

Materials and methods

Patient samples

Tumor samples were collected and analyzed according to Dana-Farber/Harvard Cancer Center IRB-approved protocols. Informed consent was obtained from all patients. These studies were conducted according to the Declaration of Helsinki and approved by the MGH and DFCI IRBs.

Syngeneic murine models

All animal experiments were performed in compliance with established ethical regulations and were approved by the Dana-Farber Animal Care and Use Committee. MC38 murine colon adenocarcinoma cells were generously provided by Dr. Gordon Freeman (DFCI) received under an MTA from Dr. Jeffrey Schlom of NCI (Bethesda, MD). CT26 colon carcinoma cells were purchased from ATCC (2015). Mouse tumor implantations were performed as previously described.¹⁷

Device design and fabrication

MDOTS and PDOTS were evaluated using '3-D cell culture chip' (DAX-1, AIM BIOTECH, https://www.aimbiotech.com/), as previously described.^{16,18} Microfluidic device design and fabrication using cyclic olefin polymer (COP) conducted at AIM BIOTECH. Briefly, the single layer slide format (75 mm × 25 mm) device or 'chip', consists of 3 microfluidic chambers each with a central gel channel (width 1.3 mm) flanked by two media channels (width 0.5 mm). The height of the microfluidic chambers is 0.25 mm. Media channels were designed including larger reservoirs to prevent over-aspiration (Fig. 3A–C).

Preparation of MDOTS/PDOTS and *ex vivo* microfluidic 3D culture

MDOTS/PDOTS (S2 fraction; 40–100 μ m) containing tumor, immune, and stromal cells were prepared as described previously.¹⁷ Briefly, fresh tumor specimens (murine and human patients) were received in media (DMEM or RPMI) on ice and

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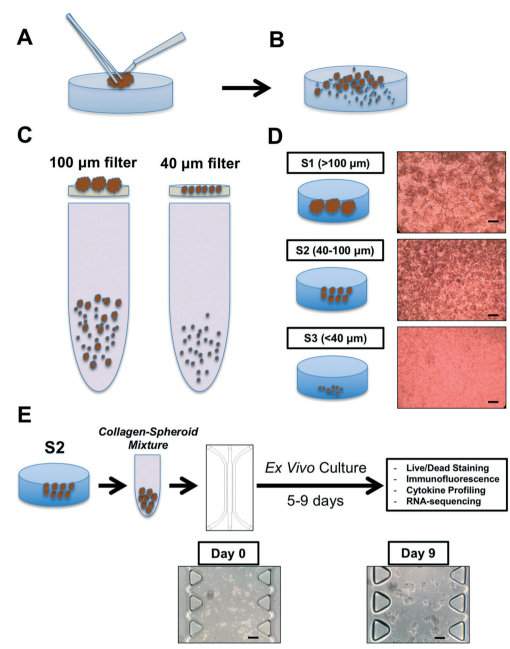


Fig. 2 MDOTS/PDOTS workflow. (A and B), A tumor specimen is received and subjected to physical and enzymatic dissociation (A), yielding dissociated tumor tissue (B) containing spheroids, single cells, and macroscopic tumor. (C and D), This heterogeneous mixture is then sequentially applied to 100 μ m and 40 μ m filters (C) to obtain three separate fractions (D), S1 (>100 μ m), S2 (40–100 μ m), and S3 (<40 μ m). E, The S2 fraction is pelleted and resuspended in collagen to be injected into the microfluidic culture device for subsequent *ex vivo* culture with indicated terminal readouts. Scale bars indicate 100 μ m (D and E).

minced in a standard 10 cm dish using sterile forceps and scalpel. Minced tumor was resuspended in high-glucose DMEM (or RPMI, for CT26) with 100 U mL⁻¹ type IV collagenase, and 15 mM HEPES (Life Technologies, Carlsbad, CA). Samples were incubated at 37 °C and evaluated every 15 minutes to monitor for adequate formation of spheroids with modest release of single cells. Following digestion (range: 15–60 min), equal volumes of media were added to minced tumor specimens and samples were pelleted and resuspended in fresh media and passed over 100 µm and 40 µm filters sequentially to obtain S1 (>100 µm), S2 (40–100 μ m), and S3 (<40 μ m) spheroid fractions, which were subsequently transferred to ultra low-attachment tissue culture plates. An aliquot of the S2 fraction was pelleted and resuspended in type I rat tail collagen and the spheroid–collagen mixture was injected into the center gel region of the 3D microfluidic chamber (10 μ L per each microfluidic chamber). After incubation for 30 minutes at 37 °C in sterile humidity chambers, collagen hydrogels containing PDOTS/MDOTS were hydrated with media with or without indicated therapeutic monoclonal antibodies – MDOTS: isotype control IgG (10 μ g

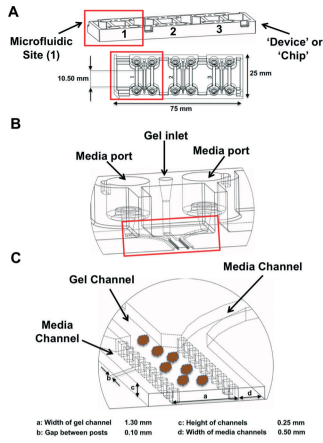


Fig. 3 Microfluidic device. (A), The 3D cell culture chip (AIM Biotech) is shown with three independent microfluidic chambers per chip. Red rectangle identifies a single microfluidic chamber in the 3D cell culture chip. (B and C), Each device contains a center gel region with posts separating the gel region from the anti-parallel side channels. Gel loading port and media ports labeled (B), along with center and side channels (C).

mL⁻¹, clone 2A3) or anti-PD-1 (10 μ g mL⁻¹, clone RMP1-14); PDOTS: anti-PD-1 (pembrolizumab, 250 μ g mL⁻¹), anti-CTLA-4 (ipilimumab, 50 μ g mL⁻¹), or combination (250 μ g mL⁻¹ pembrolizumab + 50 μ g mL⁻¹ ipilimumab), as previously shown.¹⁷

Live/Dead and immunofluorescence staining of MDOTS/ PDOTS

Live/Dead fluorescence staining was performed using AO/PI staining solution, as previously described.¹⁷ Following incubation with AO/PI or Hoechst/PI (20 min, room temp, protected from light) or Hoechst/PI (45 min, 37 °C, 5% CO₂), images were obtained. Image capture and analysis were performed using a Nikon Eclipse 80i fluorescence microscope equipped with Z-stack (Prior), motorized stage (ProScan) and ZYLA5.5 sCMOS camera (Andor) and NIS-Elements AR software package. Live and dead cell quantitation was performed by measuring total cell area of each dye. For direct immuno-fluorescence staining coupled with Live/Dead analysis, unfixed live MDOTS or PDOTS were washed with PBS and

blocked with Fc receptor (FcR) blocking reagent (PDOTS, Miltenyi; MDOTS, BioLegend) for 30 minutes at room temperature. Directly conjugated antibodies for PDOTS were CD326 EpCAM-PE (clone 9C4), CD45-AlexaFluor-488 (HI30), CD8a-AlexaFluor488 (RPA-T8); for MDOTS, CD45-AlexaFluor488 or 647 (30-F11), CD8α-PE (53-6.7; BioLegend). Antibodies were diluted 1:50 in 10 $\mu g mL^{-1}$ solution of Hoechst 33342 (in PBS) and loaded into microfluidic chambers for 1 hour incubation at room temperature in the dark. Spheroids were washed twice with PBS with 0.1% Tween20 followed by PBS. For viability assessment, microfluidic chambers were loaded with 1:1000 solution of calcein AM in PBS.

Live/Dead analysis in 2D conditions

S2 fraction of CT26 tumors was split and seeded in 3-D cell culture chips parallel with flat bottom CellBind surface 384well plate (Corning, #3683). 2D experiment was performed as following: spheroids were loaded in wells of 384-well plate in triplicates in 2 densities of ~15 and ~30 spheroids/well in cultural media containing either IgG or α PD-1 at 10 µg mL⁻¹. Plate was incubated in humidity chamber at 37 °C, 5% CO₂ for 5 days. At the end media was gently removed and 20 µL of AOPI (Nexcelom) was loaded into each well. After 20 minute incubation in the dark at RT cells were imaged on inverted Nikon Eclipse Ti microscope equipped with Nikon DS-Qi1Mc camera using NIS-Elements software. Total area of AO-stained live cells versus propidium iodide-stained dead cells was quantified. CT26 tumor spheroids seeded in both high and low densities produced similar results, graph for the high density is shown.

Cytokine profiling

Media collected from MDOTS/PDOTS culture at the indicated time point were removed (by micropipette) and transferred to 1.5 mL Eppendorf tubes. After collection, tubes containing conditioned media were placed on dry ice before storage at -80 °C. For bead-based cytokine profiling, conditioned media were thawed on ice. Four panels of multiplex ELISA assays were performed utilizing a bead-based immunoassay approach, the Thermo Fisher Immuno-Oncology Checkpoint 14plex Human ProcartaPlex[™] panel 1 (Cat No. EPX14A-15803-901), Bio-Rad Bio-Plex Pro Human Cytokine 27-plex panel (Cat No. m500kcaf0y), and Human Inflammation 37-plex panel (Cat No. 171AL001M), and Human Cytokine 40-plex panel (Cat No. 171AK99MR2). MDOTS/PDOTS conditioned media concentration levels (pg mL⁻¹) of each protein were derived from 5-parameter curve fitting models, and fold changes (relative to untreated or IgG control) were calculated and plotted as log2FC. Lower and upper limits of quantitation (LLOO/ULOO) were imputed from standard curves for cytokines above or below detection. Conditioned media from PDOTS were assayed neat. Only detectable cytokines were included in the analysis.

RNA-sequencing (RNA-seq) and CIBERSORT

RNA-seq was performed as previously described.¹ For RNAseq studies, SI-NET PDOTS were cultured in 3D cell culture chips (AIM BIOTECH). In brief, RNA lysates were prepared from SI-NET PDOTS on day 9 using the lysis buffer from Agencourt RNAdvance kit (using 1:20 proteinase K). Conditioned media was removed (as described above), before 200 mL of lysis buffer (with proteinase K) was added to each microfluidic chamber. Devices were incubated for 25 min at 37 °C, lysates were collected from each microfluidic chamber, and were transferred to RNase-free microcentrifuge tubes, and then stored at -80 °C. RNA were extracted using RNAdvance Tissue kit (Beckman Coulter, Cat. No. A32649). RNA quantity and quality were assessed using Quant-iTTM RiboGreen[™] RNA Assay Kit (Thermo Fisher, Cat. No. R11490) and Agilent Bioanalyzer RNA 6000 pico kit (Agilent, Cat. No. 5067-1513). RNA libraries were prepared from 10 ng RNA per sample using Illumina Truseq RNA Access protocol (Illumina, Cat No. RS-301-2001). RNA-seq was performed at the DFCI Molecular Biology Core Facilities (Illumina NextSeq 500). RNA-seq data were aligned and differential expression analysis were performed using VIPER pipeline, as described.¹⁹ CIBERSORT was performed as described²⁰ (https:// cibersort.stanford.edu).

Results & discussion

Sample preparation: basic principles and practical experience

To recapitulate the biology of the tumor immune microenvironment to model PD-1 blockade, one needs viable tumor tissue, autologous tumor-infiltrating immune cells, and an appropriate device and model extracellular matrix to permit 3-dimensional culture. When devising the workflow for PDOTS/MDOTS processing, we aimed for simplicity. As some immune cells lose viability within hours of collection, we felt it was essential to minimize the time required for tumor specimen processing. At the same time, we avoided added growth factors or cytokines to selectively support the viability or growth of lymphoid or myeloid cells as this might artificially alter biological response to ICB. Therefore, we used traditional cell culture media (DMEM or RPMI) supplemented with 10% FBS. In our initial pilot studies, this standard culture medium supported the growth of both MDOTS and PDOTS so further modification was not necessary.

Generation of MDOTS/PDOTS requires fresh tumor specimens from immune competent mouse models or patients. Several syngeneic murine cancer models have been profiled, including MC38 colon, CT26 colon, B16 melanoma, Lewis lung carcinoma, and GL261 glioblastoma. Diverse solid tumor types have been profiled to date, including primary lesions, lymphadenectomy specimens, pleural effusions, ascites fluid, and resected metastases. Baseline viability and sample size is essential for immune cell profiling and *ex vivo* culture. Surgical (excisional) specimens are ideal for PDOTS preparation, and are similar to the explanted murine tumors used for MDOTS preparation (Fig. 2A). While bacterial contamination was rare, all experiments were performed with antibiotics (1% penicillin–streptomycin). For primary colorectal cancer specimens (exposed to the gastrointestinal lumen and commensal bacteria), addition of metronidazole and gentamicin was required for Gram negative and anaerobic bacteria. For oropharyngeal head and neck squamous cell carcinomas, anti-fungal coverage (*e.g.* amphotericin B) was required to prevent contamination with *Candida*. Bacterial or fungal contamination was not encountered for cutaneous metastases, as the skin was sterilized prior to obtaining the excisional biopsy.

There is no minimum size cutoff, but rather sample quality, cellular viability, and immune cell composition are more important characteristics. For example, fine needle aspirates (FNAs) and core needle biopsies often yield far fewer cells than surgical (excisional) biopsies, and in our hands viability is frequently poorer from needle biopsies. On the rare occasion in which core needle biopsy yields viable cells and spheroids, there is often an insufficient number of PDOTS for extensive *ex vivo* profiling, thereby limiting the number of experimental conditions and replicates that are possible. On the other hand, pleural effusions and ascitic (peritoneal) fluid usually in cancer patients often generate at least 0.5–1 liter of fluid, and for most adenocarcinomas (*e.g.* lung, thyroid, breast) tumor cells are already aggregated with immune cells in multicellular spheroids.

With the exception of pleural effusions and peritoneal fluid (ascites) specimens, the initial step in processing involves physical and enzymatic dissociation using a limited collagenase digestion (see Materials and methods). Type IV collagenase is used in this step as it has low tryptic activity, and it was used at a dilute concentration to avoid complete dissociation of MDOTS or PDOTS to single cells. The amount of time required to digest minced tumor tissue with type IV collagenase (in media) varies from sample to sample, and tumor to tumor. In general more fibrous tumors (*e.g.* mesothelioma) require longer incubation times (up to 30–60 min), compared to most samples (20–30 min). Some murine tumor explants require only very brief collagenase treatment, as longer incubations have yielded an excess of single cells relative to spheroids.

Following physical and enzymatic dissociation, the minced and digested tumor specimen contains a mixture of macroscopic undigested tissue, spheroids, and single cells (Fig. 2B). Spheroids are isolated following passage of the dissociated specimen over a series of filters (100 μ m and 40 μ m) (Fig. 2C and D).

The second fraction (S2; 40–100 μ m) is subsequently used for MDOTS/PDOTS profiling and *ex vivo* culture (Fig. 2E). S1 and S3 fractions can be frozen or banked, or used for immune cell profiling.

3D microfluidic culture

The majority of our studies to date utilized the commercially available DAX-1 3-D cell culture chip (AIM Biotech). Three

self-contained microfluidic chambers are present on a single '3D cell culture chip' (Fig. 3A). Following preparation of the **S**2 fraction, MDOTS/PDOTS are pelleted briefly and resuspended in a neutral pH collagen solution (see Materials and methods). The collagen solution is prepared fresh for each experiment and is always prepared in excess. Pelleted MDOTS or PDOTS are initially re-suspended in a smaller volume of collagen (e.g. 100-200 µL) based on the size of the pellet, and then 10 µL is loaded into a single microfluidic chamber to confirm spheroid density and/or used for automated cell counting. Methods to accurately quantify the number of cells per spheroid (or cells per microfluidic chamber) are limited, but we will adjust the volume of collagen with a given number of spheroids to obtain approximately 10-20 000 cells per microfluidic chamber (based on automated cell counting technologies).

The collagen-spheroid mixture is kept on ice and injected into the upper and lower gel loading ports, such that the collagen-spheroid mixture extends almost half way to the opposite gel loading port, before the pipet tip is transferred to the opposite (lower) gel loading port to inject the remaining spheroids, such that the two pools of spheroids 'meet' in the middle of the gel region (Fig. 3A-C). After all microfluidic chambers are loaded in a similar fashion, they are transferred from the tissue culture hood into sterile humidity chambers and placed in a sterile incubator (37 °C) for 30 min. During this incubation, the culture media is prepared with the various antibody or drug treatments. After the 30 min incubation is complete, each microfluidic chamber is hydrated with 250-300 µL of media (depending on the downstream application and frequency of media changes) containing the desired final concentration of the drug or antibody of interest. The chambers are then returned to the incubator in the aforementioned humidity chamber and can be cultured from 5-9 days (5-6 days routinely, especially for MDOTS which grow quickly).

MDOTS/PDOTS readouts

In the initial publication describing MDOTS/PDOTS methodology (Jenkins *et al.*),¹⁷ we featured five different methods of analysis of MDOTS/PDOTS: light/phase contrast microscopy, time-lapse (live) imaging, immunofluorescence microscopy, Live/Dead imaging, and secreted cytokine profiling. Much of the routine analysis of *ex vivo* response to PD-1 blockade focused on dynamic secreted cytokine changes using beadbased cytokine profiling and fluorescence imaging using viability dyes for quantitative live/dead assessment.

Evaluation of immune-mediated tumor cell killing by effector CD8 T cells is central to the evaluation of *ex vivo* profiling of PD-1 blockade. In our initial publication,¹⁶ and companion study featuring PDOTS/MDOTS,¹⁷ dual labeling fluorescence dyes for both qualitative and quantitative Live/ Dead imaging was performed with acridine orange (AO) and propidium iodide (PI). AO/PI labeling for Live/Stain fluorescence Live/Dead staining has been in use for several decades, and are commercially available, inexpensive, and easy-to-use. PI is a cell impermeable DNA dye that only stains cells following membrane damage (e.g. necrosis or late apoptosis) after which it binds nuclear DNA. AO is a membrane-permeable nucleic acid-binding dye. Therefore, live nucleated cells emit green fluorescent signal (AO permeable, PI excluded), whereas dead cells become permeable to PI, and pure red fluorescence is evident in dead cells due to quenching of green fluorescence by PI due to Forster (fluorescence) resonance energy transfer (FRET). Shown are MC38 MDOTS treated with isotype control IgG antibody or anti-PD-1 antibody (Fig. 4A) where treatment with the anti-PD-1 antibody results in CD8 T-cell-mediated tumor cell killing, as previously demonstrated.¹⁷ Modifications to this protocol have been used with Hoechst 33342 (Ho) as a viability dye, instead of AO (Fig. 4B). Ho/PI staining can be a useful alternative for viability staining of cells with large cell size, high cytoplasm: nucleus ratio, and/or multiple cellular projections (e.g. CT26 murine colon carcinoma cells), to make threshold-setting more feasible. Quantification of Ho/PI and AO/PI fluorescence, as shown for CT26 MDOTS, demonstrates that sensitivity to PD-1 blockade is maintained in 3D microfluidic culture, but cannot be detected using MDOTS in 2D culture using 384-well plates (Fig. 4C and D).

In addition to use of fluorescent dyes, direct immunofluorescence labeling can also readily be performed using MDOTS and PDOTS.^{16,17} Standard immunofluorescence labeling protocols, coupled with viability dye evaluation using calcein AM (Fig. 5A-C), can also be employed to evaluate viability of different cell types. Calcein AM is cleaved to its fluorescent form by intracellular non-specific esterases, and in combination with Hoechst can be used to quantify cellular viability. Direct immunofluorescence using cell surface proteins to delineate tumor cells and immune cells is of great utility, especially in PDOTS specimens that demonstrate great inter-sample variability (Fig. 5A-C). Shown here are PDOTS from high-grade serous carcinoma (HGSC) of the ovary and non-small cell lung carcinoma (NSCLC), where tumor cells are readily detected using EpCAM antibodies. Staining for CD8 readily identifies effector CD8 + T cells present in PDOTS, which are necessary effectors following treatment with anti-PD-(L)1 antibodies.¹² To date, we have focused on using direct immunofluorescence (IF) coupled with viability dyes in live, intact cells although multiplexed IF is under investigation. For example, PDOTS from patients with melanoma do not express EpCAM, and most reliable melanoma markers are intracellular proteins, which cannot be readily visualized in live, unpermeabilized cells. More sophisticated imaging, including evaluation of intracellular markers requires fixation and permeabilization, which precludes use of non-fixable viability dyes. Ideal panels would permit detection of tumor cells and immune cells (lymphoid and myeloid), expression of PD-1 and PD-L1, as well as viability in distinct cellular subsets.

Secreted cytokine profiling using multiplexed bead-based kits permits analysis of a large number of inflammatory

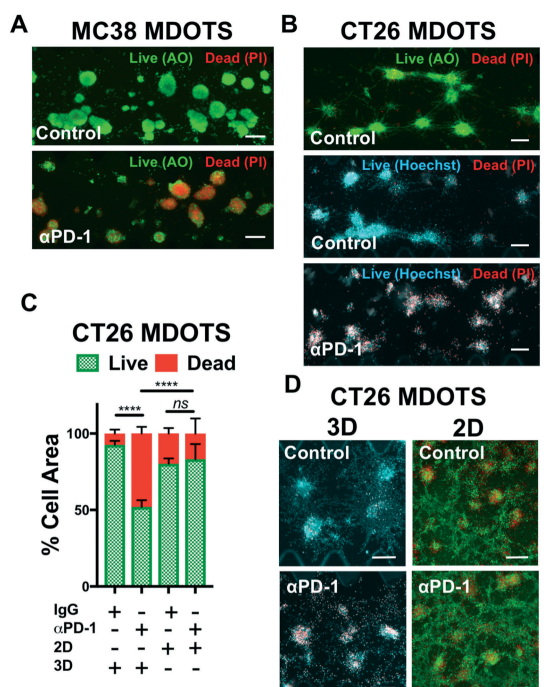


Fig. 4 Live/Dead imaging and analysis of murine-derived organotypic tumor spheroids. (A), Acridine orange (AO) and propidium iodide (PI) staining of MC38 MDOTS on day 6 of *ex vivo* culture, comparing control (isotype control IgG, 10 μ g mL⁻¹) with anti-PD-1 (10 μ g mL⁻¹). B, AO/PI and Hoechst/PI staining of CT26 MDOTS on day 5 of *ex vivo* culture, comparing control (isotype control IgG, 10 μ g mL⁻¹) with anti-PD-1 (10 μ g mL⁻¹). C and D, Live/Dead analysis (C) and fluorescence images (D) of CT26 MDOTS treated with IgG or anti-PD-1 (10 μ g mL⁻¹) for 5 days in 3D micro-fluidic culture ("3D") compared to 384-well plates ("2D") (3D – Ho/PI; 2D – AO/PI) (****p < 0.0001, ns = not significant; Kruskal–Wallis with multiple comparisons; $n \ge 3$). Scale bars indicate 200 μ m (A, B and D).

cytokines and chemokines from a small volume of conditioned media, serum, or plasma. In profiling PDOTS or MDOTS, conditioned media is collected at indicated time points, or upon termination of *ex vivo* culture, and frozen (at -80 °C), banked, and later batched for pooled analysis using either murine or human commercially available cytokine kits. For a given sample, routine Live/Dead analysis (Fig. 6A) can be formed after media collection and banking for bead-based cytokine profiling (Fig. 6B). In the PDOTS sample shown from a patient with a small intestinal neuro-endocrine tumor (SI-NET), dual immune checkpoint blockade with α PD-1 + α CTLA-4 enhanced immune-mediated

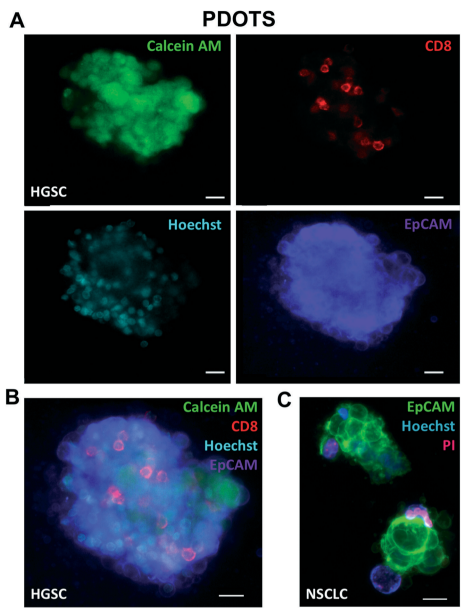


Fig. 5 Fluorescence imaging of patient-derived organotypic tumor spheroids. (A and B), Baseline IF staining of HGSC PDOTS demonstrating viable cells (calcein AM; green), CD8 T cells (red), tumor cells (EpCAM; purple), and all nucleated cells (Hoechst; blue). C, Overlay IF image of NSCLC PDOTS demonstrating EpCAM positive tumor cells (green), all nucleated cells (Hoechst; blue), and dead cells (PI; red). Scale bars indicate 20 μm (A–C).

killing compared to single-agent PD-1 blockade and CTLA-4 blockade (Fig. 6A). For the PDOTS specimen featured (Fig. 6B), the evolution of cytokine and growth factor secretion is evident over time (days 1, 3, 6, and 9), with the clearest differences between the treatment groups evident by day 9 (*e.g.* IL-8, VEGF, Il-12 (p70), CCL4). A single PDOTS (or MDOTS) sample yields >200 μ L of media, enough for four separate bead-based profiling runs (each requiring 50 μ L of conditioned media).

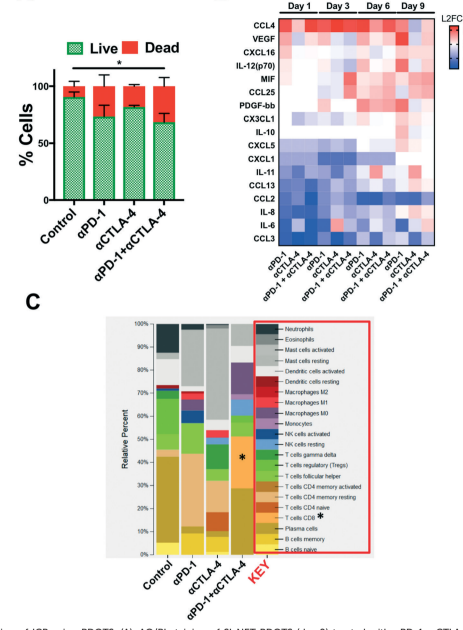
Despite the ease of using bead-based profiling to evaluate secreted cytokines and growth factors from conditioned media, most panels are not exhaustive and running multiple panels on individual samples is impractical and costly. Furthermore, for discovery efforts and pathway analysis, secretion profiling is unlikely to evaluate a sufficient number of effectors, growth factors, and chemoattractants. Lastly, molecular analysis (*e.g.* RNA-seq) provides a large data set that can be interrogated to infer changes in immune cell populations (*e.g.* CIBERSORT²⁰). We performed a pilot study to determine if bulk RNAsequencing (RNA-seq) could be performed on PDOTS and conducted CIBERSORT analysis to determine if *ex vivo* treatment with α PD-1 +/- α CTLA-4 influenced relative number of immune cells (inferred by RNA-seq). RNA was purified using a modified lysis protocol utilizing proteinase K to digest collagen gels and cellular membranes, Α

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Fig. 6 *Ex vivo* profiling of ICB using PDOTS. (A), AO/PI staining of SI-NET PDOTS (day 9) treated with α PD-1, α CTLA-4, and α PD-1 + α CTLA-4 compared to untreated control PDOTS. (B), Heatmap of changes in secreted cytokines from PDOTS (SI-NET); represented as L2FC relative to untreated control at each time point. (C), Inferred changes in PDOTS immune cell populations using CIBERSORT from SI-NET PDOTS RNA-seq.

total RNA from the mixed cell population was isolated (see Materials and methods), and was used as input for library prep for RNA-seq. CIBERSORT was performed (as described²⁰), and demonstrated relative expansion of CD8 T cells and M0 macrophages in PDOTS treated with dual ICB (α PD-1 + α CTLA-4) relative to control or single agent ICB-treated PDOTS. While this is an initial proof of concept experiment, it suggests that RNA-seq from PDOTS in 3D microfluidic culture is feasible, and that CIBERSORT or similar computational methods can be used to infer immune cell changes in response to *ex vivo* ICB.

Discussion

We have previously demonstrated that patient- and murinederived organotypic tumor spheroids retain relevant lymphoid and myeloid subsets of immune cells, grow readily in 3D microfluidic culture in collagen hydrogels, recapitulate sensitivity and resistance to PD-1 blockade *ex vivo* using established immune competent murine tumor models, to test novel combination therapies using a PD-(L)1 backbone, and demonstrated the utility of PDOTS profiling to identify and nominate candidate biomarkers.¹⁷ Here, we review this

Table 1 Summary of the advantages/limitations of MDOTS/PDOTS microfluidic culture models relative to other in vivo and in vitro cancer models. Sev-
eral typical references are provided for each type of culture

Type culture	Cancer models	Characterist	tics/advantages	Limitations	
	3D microfluidic culture of: ^{26–32}	Culture conditions	 Ideal to study immune-tumor interaction in 3D microenvironment Capable of modeling complex tumor microenvironment (TME) and extra-cellular matrix (ECM) Use of patient-derived and mouse specimens (PDOTS, MDOTS) or cell lines (cell line spheroids) 	Culture limitations	 Inability to recapitulate biological <i>in viv</i> interactions within the entire animal (except for body on a chip platforms) Variability in number of spheroids within the device Difficult to maintain long-term culture (months)
			• Dynamic multicellular co-culture		• Difficult to provide correct cell culture medium
			 Reproduces paracrine and contact interactions Accounts for 3-dimensional cancer cell growth Mimics local <i>in vivo</i> organization Medium-term culture (1–2 weeks) 		• Risk of contamination during handling
	Cell line spheroids ^{18,29,30,32}	Material & methods	 Requires low number of cells Ability to modulate cytokine/gradients Reduces reagents 	Technical issues	• Low reproducibility and variability in data (PDOTS)
	MDOTS ^{16,17}		Possibility to include fluid flow stimuli with pumpsMicrofluidic devices are scalable		 Inability to reproduce same experimen (PDOTS) unless after "Bio-banking" of sample and create cell lines from patie Difficult to evaluate/extract results
	PDOTS ^{16,17}	Results & potentiality	 (size, number of cells) Reproducible experiments (cell line, MDOTS) Imaging in real-time 		 Requires cell sorting to collect protein lysate and RNA from each cell population Requires experienced operator(s) and training
			 Capable of evaluating drug toxicity and drug metabolism Live/Dead assays Cytokine profiling High reproducibility with same mouse background (MDOTS) Can be applied for migration studies (immune cells) Ease of bulk protein RNA collections Low cost (cell line spheroids) Potential for personalized medicine 		 Low throughput screening (potential medium to high throughput screening) High cost (MDOTS, PDOTS)
In vitro ex vivo	2D standard cell culture ³¹⁻³⁵	Culture conditions	 Ideal to study single cancer cell autonomous processes Use of patient-derived and commercial cell line 	Culture limitations	 Inability to recapitulate biological <i>in vivo</i> interaction within entire human body Static 2-dimensional culture
			Simple technical cultureReproducible experiments		 Lack of the TME Fails to account for 3-dimensional cancer cell growth
		Material & methods	 Low-, medium- to long-term culture Require cells, cell culture medium and culture dishes 		 Lack ECM Lack immune cells Potential genetic changes of cancer cells over time No multicellular co-culture No possibility to include fluid flow stimuli with pumps Low theoremut corroning
		Results & potentiality	 Potential change of the genetic background of original cancer cells Live/Dead assay Cytokine profiling Imaging in real-time Easy methods to collect protein lysates and RNAs Easy evaluate/extract results capable 	Technical issues	• Low-throughput screening

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Table 1 (continued)

Type culture	Cancer models	Characteristics/advantages		Limitations	
			of evaluating drug toxicity and drug metabolism • Low costs • High-throughput screening (up to 96- or 384 well plates)		
	Standard transwell culture ^{36–40}	Culture conditions	• Ideal to study paracrine signaling, chemotaxis (immune cells) and vascular permeability (drugs)	Culture limitations	• Inability to recapitulate biological <i>in vivo</i> interaction
			 Modulate cytokine/gradients 		• Do not mimic contact interactions in th TME
			 Simple technical culture Dynamic multicellular co-culture 2D coating with ECM Medium, to long term culture 		• Low mimic of <i>in vivo</i> organization
		Material & methods	 Medium- to long-term culture Require cells, cell culture medium, transwell insert (membrane) and culture wells 	Technical issues	• Gravity force (g) may affect results
		Results & potentiality	 Possible apply trans-endothelial flow with custom made/commercial platforms Capable of evaluating drug toxicity and drug metabolism Cytokine profiling Easy collect protein lysate and RNA from each cell population without sorting Easy/reproducible results Low costs High-throughput screening (up to 96- or 384 well plates) 		• Difficult imaging (depends on membrane transparency)
In vitro in vivo ex vivo	Circulating tumor cells (CTCs) ^{24,41–46}	Culture conditions	 Not invasive methods of isolation from blood Multicellular co-culture Medium-term culture (1-2 weeks) Versatile and compatible with multiple platforms and type of culture (3D culture, organoids, <i>in vivo</i> mouse models) 	Culture limitations	 Difficult to provide protocols/medium for culture Lacks native immune and stromal cells Possible different biology between circulating tumor cells and tumor within native microenvironment
		Material & methods	• Use of patient-derived specimens	Technical issues	 Often present only in patients with large disease burden Takes time to propagate sufficient material for drug screening/testing
		Results & potentiality	 Potential for personalized medicine Imaging in real-time Following propagation, CTCs can be used for anti-neoplastic drug testing 		 Difficult evaluate/extract results Medium- to high costs Low to medium throughput screening
In vitro ex vivo	Organoids ^{21,22,47–49}	Culture conditions	 Ideal to recapitulate the pathophysiology of the original tumor Model complex tumor microenvironment TME Single/multicellular co-culture Account for 3-dimensional cancer cell growth Mimic <i>in vivo</i> organization Multiple methods of isolation from peripheral blood Amenable to repeat evaluation ("living biobank") Medium- to long-term culture (up to months) 	Culture limitations	 Lacks native immune and stromal elements Takes time to propagate sufficient material for drug screening/testing Difficult to provide correct protocols/ cell culture medium Risk of contamination for high handling level
		Material & methods	• Use of patient-derived specimens or cell lines	Technical issues	 Difficult to evaluate/extract results Require cell sorting to collect protein lysate and RNA from each cell population (multi-cellular organoids)

Table 1 (continued)

Type culture	Cancer models	Characteristics/advantages		Limitations	
		Results &	 Imaging in real-time Capable of evaluating drug toxicity and drug metabolism Easy – bulk protein lysates and RNA extractions Low to medium costs Potential for personalized medicine 		• Low to medium throughput screening
In vivo	Xenografts mouse models ^{23,31,33,50–54}	Culture conditions	• Ideal to study biological <i>in vivo</i> interaction within the entire animal body <i>in vivo</i> culture system using patient-derived specimens	Culture limitations	• Time and labor-intensive
			• Account for 3-dimensional cancer cell growth		• Challenging imaging in real-time
			• Mimic <i>in vivo</i> organization and TME		• Requires experienced operator(s) and training
			Multicellular co-cultureLong-term culture (over months)		 Genetic differences between species Complex infrastructure and specific technical skills required
			• Incompatible with high-throughput screening		• Lack of immune cells
		Material & methods Results &	 Fluid flow stimuli by <i>in vivo</i> circulation Require mouse and animal facility Patient derived xenograft (PDXs) derived 	Technical issues	 Require long culture to quantify resul Challenging variability in data Collect protein lysate and RNA
					Generic protein syster and RNAafter sacrifice mouseHigh costsLow throughput screening
In vivo	Immune-competent mouse	Culture conditions	• <i>In vivo</i> culture system using patient-derived specimens	Culture limitations	• Time and labor-intensive
	models ^{31,33,54–56}	conditions	• Biological <i>in vivo</i> interaction within the entire animal body	minutions	• Challenging imaging in real-time
			• Include immune interactions		• Requires experienced operator(s) and training
			• Account for 3-dimensional cancer cell growth		• Complex infrastructure and specific technical skills required
			• Mimic <i>in vivo</i> organization and TME		• Limited number of potential drug combinations
			• Long-term culture (over months)		 Drugs and therapeutic antibodies against mouse targets may differ from human targets Only mouse cells study
		Material & methods	• Require mouse and animal facility	Technical issues	Require long culture to quantify results
		Results &	• Capable of modeling heterogeneity		
		potentiality	of <i>in vivo</i> response and resistance • Capable of evaluating drug toxicity		• Challenging variability in data
			and drug metabolismFluid flow stimuli by <i>in vivo</i> circulation		• Collect protein lysate and RNA after
			• Useful for evaluating drugs whose mechanism of action takes time (<i>e.g.</i>		sacrifice mouseHigh costs
			 epigenetic modifying agents) MDOTS derived from immune-competent mouse models can be cultured in 3D microfluidic device or 		• Low throughput screening

previously published method in greater detail with emphasis on the methods of MDOTS/PDOTS preparation and readouts, and discuss future directions with this novel technology (*e.g.* RNA-seq), and provide further evidence 3D microfluidic culture of MDOTS recapitulates response to PD-1 blockade which is lacking using traditional 2D culture methods.

The PDOTS/MDOTS platform offers several advantages over existing methods to evaluate tumor-immune responses

to ICB (see Table 1), providing a window into the tumor immune microenvironment and enabling evaluation of acute and dynamic responses to ICB to using relevant murine models and patient samples. Unlike organoids,^{21,22} patientderived xenografts (PDXs),²³ and circulating-tumor cells (CTCs),²⁴ MDOTS and PDOTS do not require days or weeks of tissue manipulation or propagation, and contain autologous, tumor-infiltrating immune cells. Importantly, MDOTS/PDOTS are not to be confused with organoids, a distinct 3D culture method developed by Hans Clevers and Toshiro Sato, which leverages native tissue stem cells to propagate normal or transformed tissue.²⁵ Organoids, like PDXs and CTCs, can be used as part of personalized medicine efforts, but lack native stromal and immune cells, thus precluding study of tumorimmune interactions. Furthermore, generation of sufficient quantity of organoids takes time, thus limiting rapid drug screening capabilities.

MDOTS/PDOTS offer significant advantages over organoids, PDXs, and CTCs for ex vivo profiling tumor-immune responses to ICB, however there are several important limitations inherent in the current version of the platform. First, the current version of the MDOTS/PDOTS platform is currently only capable of evaluating tumor-immune interactions of immune cells that have already infiltrated the tumor. In other words, PDOTS/MDOTS "v1.0" cannot recapitulate T-cell priming (which occurs primarily in lymph nodes) or recruitment of naïve immune cells to the tumor microenvironment. Second, we have yet to perform comprehensive evaluation of tumor, stromal, and immune changes through the course of ex vivo culture, both in control conditions and in response to ICB. Thirdly, use of core needle biopsies and FNAs is challenging, and excisional specimens are preferred to yield sufficient number & quality of MDOTS/PDOTS for ex vivo profiling at present, lastly, the influence of device dimensions, biophysical parameters, interstitial flow, hypoxia, and metabolic on tumor-immune interactions, especially cytokine production remains poorly understood and will require further investigation.

It should be noted that the size of the gel region and the size of the media channels are among the variables that may differ among different versions and types of microfluidic devices that could conceivably influence absolute and relative cytokine changes. The impact of different device parameters (e.g. device height, gel region width) on cytokine elaboration is under investigation, but remains far from complete. Identifying the physical components & dimensions of the microfluidic device that enables studying ICB will be important, especially when considering building more advanced mediumand high-throughput systems to test multiple drugs and/or antibodies in combination. An important co-variate in this analysis is the composition of the device itself. For example, the AIM microdevice is plastic, whereas other materials (e.g. polydimethylsiloxane, PDMS) used for 'home grown' device fabrication exhibit different properties and characteristics. This is a particularly important consideration when testing ICB in combination with small molecules (commonly prepared in DMSO), as PDMS is known to adsorb small hydrophobic molecules,²⁶ likely influencing drug delivery to MDOTS/PDOTS and ultimately drug efficacy.

While the long-term potential of MDOTS/PDOTS technology as a predictive tool is not yet clear, MDOTS/PDOTS are showing great promise in the evaluation of novel combination therapies and in the discovery of novel biomarkers. Important future directions include extending the lifetime of *ex vivo* cultures (*e.g.* tumor vascular networks), performing clinical validation studies (*e.g.* co-clinical trials or adaptive clinical trials using *ex vivo* responses to guide clinical treatment), and developing medium- and highthroughput versions of the system. With further technology development along with improved data analytics and bioinformatics, we anticipate the MDOTS/PDOTS platform will form the foundation of functional precision immuneoncology efforts.

Conflicts of interest

A. Aref, D. A. Barbie, and R. W. Jenkins have ownership interest in pending U.S. Patent Application No. 15/540346. R. D. Kamm has ownership interest (including patents) in AIM Biotech, D. A. Barbie is a consultant/advisory board member for N of One.

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