Live single cell functional phenotyping in droplet nano-liter reactors

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While single cell heterogeneity is present in all biological systems, most studies cannot address it due to technical limitations. Here we describe a nano-liter droplet microfluidic-based approach for stimulation and monitoring of surface and secreted markers of live single immune dendritic cells (DCs) as well as monitoring the live T cell/DC interaction. This nano-liter in vivo simulating microenvironment allows delivering various stimuli reagents to each cell and appropriate gas exchanges which are necessary to ensure functionality and viability of encapsulated cells. Labeling bioassay and microsphere sensors were integrated into nano-liter reaction volume of the droplet to monitor live single cell surface markers and secretion analysis in the time-dependent fashion. Thus live cell stimulation, secretion and surface monitoring can be obtained simultaneously in distinct microenvironment, which previously was possible using complicated and multi-step in vitro and in vivo live-cell microscopy, together with immunological studies of the outcome secretion of cellular function.

The cellular phenotype is a conglomeration of multiple cellular processes, representing varying expression levels of genes and proteins that determine the cell’s particular function in activities such as cellular communication, adhesion, or metabolism. Average measurements of such molecular markers in cell populations obscure inherent cell-cell heterogeneity and restrict the ability to distinguish between the responses of individual cells within a sample. While flow cytometry has traditionally been used to determine single cell phenotypes, it cannot provide continuous measurements of proteins in the same individual cells over time. Recently, the development of microfluidic technology enabled analysis of cell phenomena on the single cell level. The reported platforms, mostly based on micro well technologies, do not provide a controllable micro-environment for cell-cell interaction, however, as no reagent mixing is provided. Moreover, technical manipulation of these platforms prohibits their current use beyond the wall of the developer laboratory. In this paper, we describe a novel approach that allows stimulation and dynamic characterization of live single cells for surface and secreted protein expressions. This approach should have a broad impact on diverse biological systems for the study of cell surface and secretion proteins as potential biomarkers and targets for diagnostics and therapeutics as well as cell-cell interactions imaging. In particular, one clinical application of the described approach for monitoring live cell surface markers and secretion as well as cell-cell interactions can be applied to a DC vaccine evaluation. Evaluation of the efficacy of various therapeutic agents such (i.e. small molecules, antibodies, nucleic acid polymers) could significantly benefit from this sensitive and rapid detection methodology. The correlation between surface protein expression and secretion obtained from cells before and after treatment can be obtained in a dynamic fashion on a single cell level.

Defining the frequencies of specific cell types and states is a primary goal of ex vivo immune system diagnostics. Mature dendritic cells (DCs) play a central role in the onset of immunity or tolerance by presenting antigens to T cells. The functional phenotype of mature DCs, can be characterized by their patterns of cytokine secretion and expression of differentiation/activation cell surface markers, such as co-stimulatory molecules. Population-wide studies of functional phenotype mask the dynamic behavior of individual DCs and are thus often insufficient for characterizing the immuno-regulatory potential of DCs. To address this problem, we aimed to develop a droplet microfluidic platform for single live cell phenotyping. This platform enables activation and monitoring of live single cells for secretion and cell surface markers simultaneously. In particular, we applied this technology to simultaneously stimulate DC maturation with an immunological adjuvant, lipopolysaccharide (LPS), and to monitor two phenotypic markers of DCs. The first of these is cluster of differentiation 86 (CD86), an important cell surface molecule that provides co-stimulatory signals necessary for T cell activa-
We also followed the maturation phenotype of the DC by measuring secreted interleukin (IL)-6, a cytokine that plays an important role in lymphocyte activation. Additional critical step in the generation of antigen specific immune responses that activate naïve T cells in lymph nodes is immunological synapse (IS) formation between mature DC and T cell. Though attempts were made to analyze the dynamic aspects of these cell-cell interactions in microscopic imaging, a robust technique to analyze the formation and cell-cell interaction has lagged. Here we were able monitoring live single cell surface markers during DC/T cell IS formation in a distinct microenvironment, a feat not possible using conventional techniques.

**Results**

Nano-liter droplet microfluidic-based technology for stimulation and monitoring of surface and secreted markers of live single immune DCs. Figure 1a schematically illustrates the droplet microfluidic platform employed by us to activate and to monitor live single cells for secretion and cell surface protein markers simultaneously. The poly(dimethylsiloxane) (PDMS) microfluidic system was fabricated using standard soft lithography methods, combining functionalities of droplet generation and an incubation chamber array of 1000 droplets (Fig. 1a, b). The generation of monodisperse droplets in a micro-channel through shearing flow at a flow-focusing zone is illustrated in Fig. 1b. In this setup, three perpendicular inlet channels form a nozzle. The individual syringe pumps were used to control flow rates of the central stream of the oil phase (1) and the left stream of DCs suspension (2) and the right stream of bioassay reagents (bead sensors and antibodies) (3) (Fig. 1a). Each generated droplet in the array contained fluorescently-labeled detection antibodies (anti-CD86-FITC Abs) that create a localized fluorescent signal on the bead surface in the droplet (Fig. 1c). The advantages of this droplets-based array technique include the physical and chemical isolation of cells in droplets eliminating the risk of cross-contamination and the fast and efficient mixing of the reagents that occurs inside droplets facilitating fast reaction rate. More important, this nano-liter microenvironment allows delivering various stimuli and assay reagents to each cell and gas exchange to ensure viability.

To measure the time dependent CD86 and IL-6 expression of single DCs, the fluorescent signal generated on cell and bead surfaces was recorded each 20 min in the confined volume of droplets and analyzed by the ImageJ software (Fig. 2a). The positive signal was recorded and quantified for each cell in the droplets, and compared to the background empty droplet signal (Fig. 2b). The image sequence was analyzed and the increase in intensity from each cell was plotted over time showing expression kinetics (Fig. 3a, b). After 40 minutes of incubation in LPS nano-liter droplets, the encapsulated cells started to upregulate the surface maturation marker, CD86, and reached saturation at 2 hours and 20 min (Fig. 3a). Fig. 3c describes the fluorescent signal intensity from thirty cells after 2 hours and 20 min in separate droplets, confirming the previously-described inherent CD86 marker heterogeneity. Thus, this method allows us to distinguish the responses of individual cells to LPS stimulation in the confined volume of droplets. In addition to CD86, we were able to monitor LPS induced IL-6 secretion from the same encapsulated single DCs (Fig. 3b). The detection of IL-6 on microsphere surfaces is shown in Figure 2b for single DCs. The signal was detected in droplets after 10 min in the array (Figure 3c). Droplets with LPS stimulation contained IL-6 levels that were significantly higher (three standard deviations) than droplets with no LPS, which showed negligible IL-6 levels.

Monitoring single cell immunological synapse (IS) formation between mature DC and T cell. Immunological synapse (IS) formation between mature DC and T cell is one of critical steps in the generation of antigen specific immune responses that activate naïve T cells in lymph nodes. Here we were able monitoring live single cell surface during DC/T cell IS formation in a distinct microenvironment, a feat not possible using conventional techniques. Previously described droplet microfluidic-based platform was applied to co-encapsulate pairs of cells of different types in one micro-reaction droplet (Fig. 1a). In this setup, the center stream contains oil (1), the left steam contains suspension of T cells (2) and the right steam contains DCs (3). To demonstrate the IS formation, we co encapsulated naïve T cell and LPS matured DC in nano-liter droplets of tubulin marker assay which provides green-fluorescent staining of polymerized tubulin in live cells. Fig. 2c demonstrates the physical interaction and the polymerization...
of tubulin in live mature DCs that encountered naive T cell and establishing signaling zone on live cells. We were able to monitor remodeling of the cytoskeleton and microtubule polymerization in DCs at the IS formed between co-encapsulated pairs of live DCs and CD4+ T cells in droplets.

**Discussion**

Altogether, our work provides a platform technology to stimulate specific cellular processes, such as DC maturation and cell-cell interactions, and to measure the effects of these cellular processes, such as CD-86 and IL-6 expression, in a time dependent manner for single

![Figure 2](image1.png)

**Figure 2** | (a) DCs and bead-based sensors and reagents co-encapsulated in nano-liter reaction droplet containing anti-CD86 fluorescently tagged Abs for cell surface analysis. 1,2. Fluorescence images of droplets captured on a Zeiss 200 Axiovert microscope (a1 × 20 and a2 × 40) after 2 hours and 20 min of incubation. 3. The intensity of the fluorescence recorded for both CD86 protein and IL-6 detection measured using ImageJ software (the fluorescence intensity scale is present on the right). (b, c, d) Physical interaction and the polymerization of tubulin in live DCs that encountered a T cell and establishing signaling zone.

![Figure 3](image2.png)

**Figure 3** | (a) The increase in accumulative CD86 expression over the time in response to LPS stimulation. (b) The increase in IL-6 secretion over the time in response to LPS stimulation. RS stands for relative signal. (c) Inherent CD86 marker heterogeneity.
live cells in population. In the future, this technology could be applied to characterize functional phenotypesp in various heterogeneous populations at a single cell level. Furthermore, we were able to monitor micro-size remodeling of the cytoskeleton and microtubule polymerization in DCs at the IS formed between co-encapsulated pairs of live DCs and CD4+ T cells in droplets. In addition, to demonstrate the heterogeneity in cell-cell interactions of T cell and DC populations we co-encapsulate two naive T cells and four previously LPS matured DCs in single droplets to monitor the cell-cell interaction. Figure 2d show that not all cells are interacting to create IS thus demonstrating that due to the heterogeneity in DC maturation process not all DC are able to interact and activate naive T cells. Furthermore the distinguish differences in the morphology of resting and interacting live cells are clearly can be seen in the nano-liter reaction droplets. Thus the developed approach allows to dynamically monitoring the interaction events that transpire during cell-cell communication in the immune system on a single cell level to avoid ambiguities that arise due to heterogeneous responses in cell-cell communication.

Methods

Materials. Rat FITC-anti-CD86 antibody (GL1) was obtained from Abcam and used at concentration of 1 μg/10^6 cells. Biotin-anti-IL-6 Ab (504601) for bead conjugation was obtained from BioLegend and FITC-anti-IL-6 Ab (11-7061-81) for IL-6 detection was obtained from eBioscience and used at concentration of 1 μg/mL.

Microfluidic device fabrication. Microfluidic flow chambers were fabricated using soft lithography at the MGH/CEN Facility. Negative photo resist SU-8 2100 (MicroChem, Newton, MA) was deposited onto clean silicon wafers to a thickness of 150 μm, and patterned by exposure to UV light through a transparency photomask (CAD/Art Services, Bandon, OR). The Siylgard 184 poly(dimethylsiloxane) (PDMS) (Dow Corning, Midland, MI) was mixed with crosslinker (ratio 10:1), poured onto the photomask patterns, degassed thoroughly and cured for 12 hours at 75°C. Next, we peeled the PDMS devices off the wafer and bonded to glass slides after oxygen-plasma activation of both surfaces. The device is composed of a droplet forming nozzle and a storage array for 10^3 droplets. To improve the wetting of the channels with mineral oil in the presence (1%, v/w) of the surfactant (span80), prior to the experiments the microfluidic channels were treated with Pico-SurTM 2 (Dolomite Microfluidics, UK) by filling the channels with the solution as received and then flushing them with air. Tygon Micro Bore PVC Tubing 100f, 0.010” ID, 0.030” OD, 0.010” Wall (Small Parts Inc, FL, USA) were connected to the channels and to the syringes. We used 1 ml syringes to load the fluids into the devices, while syringe pumps (Harvard Apparatus, MA) controlled the flow rates.

Microsphere sensors preparation. ProActive® Streptavidin Coated Microspheres (10 μm) (Bang Laboratories Inc., USA) were conjugated with the biotinylated- anti-IL-6 (Abcam® MA, USA) according to the manufacturer’s protocol. Unbounded active sites were blocked with StarlingBlockTM (thermo Scientific, USA) for one hour. Next, we washed the beads with the Phosphate Buffer Saline (PBS) (Sigma, USA), diluted to the final concentration 0.5 mg ml^-1, and stored at 4°C.

Cell isolation and stimulation. 6–8 week old female C57BL/6 mice were obtained from the Jackson Laboratories. Bone marrow dendritic cells (BMDCs) were collected from femora and tibiae and plated on non-tissue culture treated plastic dishes in RPMI medium (Gibco, Carlsbad, CA, Invitrogen, Carlsbad, CA), supplemented with 10% FBS, L-glutamine, penicillin/streptomycin, MEM non-essential amino acids, HEPES, myc, and 0.5% (mg/mL) QPS, and RPMI medium (Gibco, Carlsbad, CA). At day 7 floating cells were collected and used as GM-CSF derived dendritic cells. Cells were stimulated with LPS (rough, ultra-pure E. coli K12 strain LPS, 100 ng/mL; Invitrogen, San Diego, CA) for 3 days. Where indicated, experiments were performed in the presence (1%, v/v) of the surfactant (Span80) and air. Tygon Micro Bore PVC Tubing 100f, 0.010” ID, 0.030” OD, 0.010” Wall (Small Parts Inc, FL, USA) were connected to the channels and to the syringes. We used 1 ml syringes to load the fluids into the devices, while syringe pumps (Harvard Apparatus, MA) controlled the flow rates.

Image analysis. Fluorescence images of droplets were captured on a Zeiss 200 Axiovert microscope using an AxioCam MRm digital camera. For the cell studies, fluorescence signals were captured separately using appropriate filter sets for FITC

Image processing and analysis was conducted using ImageJ software.

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Author contributions

T.K. supervised the project, designed the experiment, conducted experiments, analysed data and wrote the manuscript, A.G. conducted experiments, analysed data and drafted the manuscript, M.Y. reviewed the manuscript.

Additional information

Competing financial interests: The authors declare no competing financial interests.
