Multiplexed enrichment and genomic profiling of peripheral blood cells reveal subset-specific immune signatures

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Specialized immune cell subsets are involved in autoimmune disease, cancer immunity, and infectious disease through a diverse range of functions mediated by overlapping pathways and signals. However, subset-specific responses may not be detectable in analyses of whole blood samples, and no efficient approach for profiling cell subsets at high throughput from small samples is available. We present a low-input microfluidic system for sorting immune cells into subsets and profiling their gene expression. We validate the system’s technical performance against standard subset isolation and library construction protocols and demonstrate the importance of subset-specific profiling through in vitro stimulation experiments. We show the ability of this integrated platform to identify subset-specific disease signatures by profiling four immune cell subsets in blood from patients with systemic lupus erythematosus (SLE) and matched control subjects. The platform has the potential to make multiplexed subset-specific analysis routine in many research laboratories and clinical settings.

INTRODUCTION

Millions of immune cells can be obtained from a small blood draw, yet most methods for immune profiling from clinical samples fail to resolve the biological information contained within these cells. Recently, profiling the immune state of individuals using gene expression analysis of total peripheral blood mononuclear cells (PBMCs) has become instrumental in defining immune signatures and disease states in humans. These studies provide insight into the mechanisms of complex immune responses that occur in infection (1, 2) and autoimmunity (3–5), which are difficult to recapitulate in murine models (6–8). Furthermore, expression signatures can be used to stratify individuals into different disease subtypes (9–13) or to predict individualized clinical prognoses (14–16). However, profiling total PBMCs has a severely limited potential to resolve immune status since PBMCs constitute a complex mixture of specific cell types or cell “subsets” with distinct functions. Furthermore, no effective methods exist to resolve the underlying signatures of immune subsets from a total PBMC dataset. Only recently, gene expression profiles from purified subset samples have been shown to be better discriminants of immune status than total PBMC profiles due to the diversity of leukocyte responses (17–20). In addition, new immune subsets and cellular states, some of which are indicative of impaired immune function, have been discovered through gene expression profiling of PBMCs at the single-cell level (21–24). These observations have stirred interest in probing the gene expression and monitoring the activity of these subsets in particular. As a whole, this developing body of work suggests that molecular profiling of PBMC subsets is poised to become an important tool in basic studies of immune disease as well as a clinical tool useful for predicting and monitoring patient outcomes.

Despite its potential as a tool for immunomonitoring, available methods for subset-specific expression profiling are ill-suited for large studies and clinical translation. First, technologies for cell subset enrichment such as fluorescence-activated cell sorting (FACS) are capital intensive and require substantial attention from highly trained staff. As a result, FACS is challenging to scale for large clinical studies with many samples (e.g., multiple cell subsets across many patients at different time points). In addition, FACS requires a minimum sample input to establish gates for each target subset, which can frustrate its application to low-quantity samples and projects targeting many subsets from each sample. Another popular approach, magnetic affinity cell sorting (MACS), is most often practiced manually or on proprietary instrumentation with a capacity to run batches of 6 to 10 enrichments. Although the potential to automate MACS at higher throughput using pipetting robots exists, this also requires large samples for good results (typically 1 million cells per subset), would be capital intensive, requires extensive custom programming, and is only effective for central processing centers with a large steady supply of samples. Second, the throughput of complex RNA sequencing (RNA-seq) library construction protocols is generally limited by reagent cost and labor. Implementing library construction at high throughput on custom liquid handling systems has been widely demonstrated but suffers the same drawbacks described above in reference to a hypothetical automated MACS approach. Last, single-cell analysis obviates the need for subset purification since each cell is profiled individually. However, many thousands of cells would need to be analyzed from each sample, which is prohibitively expensive today for large studies and particularly inefficient where rare subsets are of interest or immune cells make up a small fraction of the total sample (25).

Because of these limitations, most clinical gene expression studies are currently limited to whole-blood or total PBMC profiling (9–11, 14, 15), which fail to resolve expression signatures from most cell subsets due to confounding signals from more abundant cell populations. To efficiently identify and monitor important disease
signatures in lower-abundance subsets, we developed a multiplexed microfluidic system that integrates both human PBMC subset enrichment and sequence library construction procedures for subset-specific genome-wide expression measurements by RNA-seq. Our approach has the unique advantage of integrating the subset enrichment and complex multistep RNA-seq library construction protocols to provide an end-to-end solution in a single device that is not provided by any other approach. By integrating the key functionality into a single microfluidic device, we eliminate the need to acquire, program, and monitor complex liquid-handling robots or integrate a constellation of single-purpose proprietary instruments. Here, we show how this integrated workflow can be multiplexed to handle multiple cell subsets and sequence libraries in parallel. Because no comparable integrated instrumentation exists, we benchmark the technical performance of our system against the gold standard approaches for each step that are commonly implemented manually by expert immunology research laboratories.

The microfluidic system carries out multiplexed enrichment of target cell subsets based on affinity for cell surface markers by MACS and high-sensitivity sequence library construction for full-length RNA-seq using Smart-seq chemistry. From an input of 50,000 cells, the device can purify multiple PBMC subsets with high purity and produce highly quantitative gene expression data covering about 10,000 genes in each subset. In testing immune stimulation and challenge in vitro, we highlight the importance of subset-specific profiling by showing differential responses across four selected subsets. Last, we applied the microfluidic device to profile PBMCs of patients with systemic lupus erythematosus (SLE) and identified clear differences in the transcriptomic states of healthy individuals and patients with SLE in multiple immune cell subsets. By integrating multiplexed enrichment and library construction workflows in a single device, our platform has the potential to enable scalable PBMC sample preparation for large clinical studies and allow for both high-resolution and high-throughput profiling of the immune system. We foresee the routine application of this system as a tool to monitor immune responses in clinical studies and help diagnose patients with complex and/or critical immune conditions.

RESULTS

Microfluidic device design
We designed a custom two-layer microfluidic device capable of semiautomated cell isolation, cell disruption, and sequence library construction protocols. This system integrates microfluidic liquid handling with magnetic affinity purification and capability for on-board polymerase chain reaction (PCR). The device is fabricated using established methods for two-layer soft lithography (26), has “large” microliter-scale internal volumes to handle mammalian cell samples, and contains 39 micromechanical valve sets controlled by an external pneumatic valve controller (27). Each operating unit on the device consists of three main chambers, each having different capacities (1, 2, and 4 μl), that are partitioned by microvalves (Fig. 1A). The largest chamber is rectangular in shape and is used for cell isolation (Supplementary Note), while the two smaller “rotary reactor” chambers are used for library construction (28). These reactors are fitted with internal microvalves that are used to formulate sample and reagent combinations and to mix these by perfisalising around the circular channel (Fig. 1B). Bead resuspension is achieved by perfisalising in the rotary reactors and by a moving magnetic field in the large rectangular chamber. A 675-μm-thick silicon wafer was used as the substrate for these microfluidic devices to allow rapid heat transfer during temperature changes called for in the protocol, particularly for PCR (due to its thinness and high thermal conductivity). The substrate thinness also enables small external permanent magnets to closely approach magnetic beads in the device chambers and subject these to strong magnetic forces. The magnets are used to move beads between chambers and hold beads in place during buffer exchange steps. With such device functionality, we are able to automate many steps in the complex protocol for cell sorting, cell disruption, and RNA-seq library construction in a simple microarchitecture (fig. S1). The three-chamber operating unit is modular and constitutes a scalable microarchitecture for devices with variable sample multiplexing capacity. The data presented here were produced using 10-channel devices, although we have fabricated devices with 6 to 30 channels. The devices can also be reused following a simple washing procedure (particular devices were used up to four times in this study).

Microfluidic cell sorting and low-input RNA-seq
To validate the performance of our microfluidic device, we independently benchmarked the subset enrichment and RNA-seq workflows (Fig. 1C) against standard protocols for cell sorting and RNA-seq library construction. We tested our workflows using adult PBMC samples from healthy subjects obtained from a commercial supplier (Research Blood Components) at an input level of 50,000 cells per enrichment. We implemented MACS on the microfluidic device and configured a high-resolution eight-color flow cytometry analysis to read out the purity and yield of the resulting enriched cell subset samples (fig. S2). We optimized the conditions for microfluidic cell subset isolation by testing different reagents, incubation times, and washing procedures and compared the results of the optimized microfluidic protocol to conventional benchtop MACS (fig. S3 and Methods).

We tested positive selection of target cells, negative depletion of nontarget cells (where we recover cells that do not bind to the beads), and sequential isolation using both modalities in tandem. The total time required for isolation is about 1 hour. The CD4 and CD8 subsets were isolated using the tandem procedure. First, total T cells were isolated by depleting cells expressing markers for lineages other than T cells. The total T cell population was then positively selected for either CD4 or CD8 to isolate helper and cytotoxic T cell subsets separately. The previous negative selection reduced contamination from nontarget lineages that express CD4 or CD8. B cells and monocytes, on the other hand, were effectively isolated using single positive selection for CD19 and CD14, respectively. The device consistently achieved good purity (80 ± 8%) and excellent yield (76 ± 21%) for multiple targets and modes of isolation (Fig. 2, A and B), leading to 2- to 13-fold enrichment of the selected cell types. While the device workflow does not result in perfect enrichment of the target cell type, we found that the purity of the subsets isolated using the optimized microfluidic protocol was similar to those obtained by benchtop MACS, suggesting that our workflow is on par with current implementations used to process clinical immune cell samples (fig. S3A). In addition, our device achieved better yield with lower inputs than benchtop MACS (fig. S3B). These results show that microfluidic cell sorting with magnetic beads is a viable alternative to conventional sorting approaches and demonstrates the feasibility of subset-specific enrichment with limited quantity samples.
On the basis of the results of our cell isolation testing, we expected to capture thousands of cells in each subset using our microfluidic device. With these relatively low numbers in mind, we implemented a sensitive RNA-seq protocol (Smart-seq2) (29) in the chip with minor modifications. Instead of solid-phase reversible immobilization (SPRI)–based cleanup for RNA extraction, we used custom-prepared poly-dT capture beads (Methods) that captured mRNA molecules in lysate by direct hybridization to enable purification and subsequent solid-phase reverse transcription. Our protocol calls for amplifying the resulting complementary DNA (cDNA) molecules by PCR, purifying the products with SPRI, and subsequently recovering the samples from the device for transposase-based fragmentation and adaptation with subsequent enrichment PCR closely following the standard Smart-seq2 method. The timing for this procedure closely follows the Smart-seq2 protocol (about 2 hours hands-on time plus 5 hours for incubations). The cDNA amplicons from the microfluidic device showed the expected size distribution, and the RNA-seq datasets resulting from these samples show high technical reproducibility (Pearson correlation of 0.88 ± 0.04) and correlate well with libraries produced using the standard Smart-seq2 protocol on the benchtop across four different cell subsets (0.90 ± 0.03) (Fig. 2D, fig. S4, and Table 1). Despite the overall similarity between the gene expression profiles of the four subsets (Fig. 2C), the sequence libraries produced in our workflow can distinguish the subsets based on simple correlation and clustering procedures (Fig. 2E). In addition, the enrichment of polyadenylated RNA in the microfluidic protocol reduced the number of ribosomal RNA reads and improved transcript mapping rates over the standard Smart-seq2 protocol (Table 1 and fig. S5). Combining RNA-seq with cell isolation in an integrated workflow yields libraries of similar quality as standard processing approaches (Table 1 and figs. S6 and S7). These results demonstrate that full-length cDNA synthesis and amplification by PCR can be implemented in a microfluidic device with input from on-device–enriched cell subsets to support RNA-seq and that reduction in the reaction volume and reagent consumption (from 25 to 2 μl) does not negatively affect the quality of libraries obtained.

**Gene expression signatures of PBMCs stimulated in vitro**

We assessed how accurately and reproducibly our system can profile the dynamic immune responses of different cell subsets and to what extent the responses were stereotypic or subset specific. We cultured healthy PBMCs and applied three distinct treatments known to affect immune cells [lipopolysaccharide (LPS), interferon-α (IFN-α), and dexamethasone (DEX)] in duplicate. Using the microfluidic device to carry out multiplexed subset enrichments and RNA-seq library construction, we profiled the treatment response of three different subsets (CD4+ T cells, B cells, and CD14+ monocytes). The
device-processed libraries again showed strong reproducibility, co-clustering the duplicates based on differential gene expression responses (Fig. 3A and fig. S8) and accurately recording differences in responses to the three treatments (Fig. 3B and fig. S9).

Furthermore, our results highlight the heterogeneity in response between different cell subsets (Fig. 3, C and D), as evidenced by the minimal overlap in differentially expressed genes and enriched pathways among the three subsets profiled. Even in the canonical Jak-STAT (signal transducer and activator of transcription) signaling pathway, which is known to be directly activated by IFN-α, the pattern of downstream responses varied substantially across the three subtypes studied here (Fig. 3E). These results are consistent with previous reports that type I IFNs can have either proliferative or suppressive effects on lymphocytes, depending on the relative timing of receptor coactivation (30). In addition, these responses are greatly affected by cell-to-cell communication and the interplay between innate and adaptive immunity (31–33). After 24 hours of stimulation, IFN-α induces a strong proliferative response in B cells as shown by the up-regulation of cell cycle and metabolism pathways, while for monocytes, this effect is not observed (Fig. 3D). This finding is in line with previous studies (34–40) showing that IFN-α directly stimulates B cells to induce a strong antiviral response and a commitment to an effector phenotype, while in monocytes, the effect of IFN-α is costimulatory and primes monocytes to differentiate into dendritic cells after further stimulation. Together, these results emphasize the importance of subset-specific profiling to resolve differences that would be obscured in total PBMC expression profiling.

Transcriptomic profiling of SLE patients

To demonstrate the utility of the device for disease studies, we profiled the immune state of five patients with SLE and five healthy control individuals by isolating CD4+ T cells, CD8+ T cells, B cells, and CD14+ monocytes (plus a negative control) in duplicate from cryopreserved PBMC samples (for a total of 100 subset isolations and RNA-seq libraries including controls; table S1). For each sample, 0.5 million cells were split into eight channels on devices to isolate four subsets in duplicate and prepare RNA-seq libraries (Supplementary Methods). While we detected only a few differentially expressed genes when performing single-gene analyses (fig. S10), we found that gene sets with targets of IFN are up-regulated in patients with SLE.
ing disease signatures. B cells instead of total PBMCs. Together, these initial findings show that the IFN signature for SLE may be improved by specifically profiling B cells. This suggests that the diagnostic sensitivity and predictive power of this signature can be found across all the subsets we profiled, the gene expression responses in SLE differ across immune cell subsets (48). Our data show that while this signature can be found across all the subsets we profiled, the difference between healthy and SLE signature scores is much more pronounced in B cells (P = 0.05 for B cells, P ≥ 0.2 for other subsets; with Bonferroni correction for testing multiple subsets) (Fig. 4B). This suggests that the diagnostic sensitivity and predictive power of the IFN signature for SLE may be improved by specifically profiling B cells instead of total PBMCs. Together, these initial findings show that gene expression responses in SLE differ across immune cell subsets and highlight the importance of subset-specific profiling in identifying disease signatures.

**DISCUSSION**

Through our multiplexed microfluidic workflow, we demonstrate the utility of subset-specific profiling of immune cells and its advantages over conventional total PBMC or total blood transcriptomics. Subset-specific analysis allows ready detection of biological signals from minority subsets by reducing confounding effects from abundant cell populations such as the monocytes that dominated our test samples. Our method is complementary to the application of single-cell transcriptomics approaches. For example, single-cell studies could reveal pathogenic subsets that can be enriched using the microfluidic device for large-scale research studies or clinical diagnostics, even for rare subsets. With this framework, single-cell RNA-seq (scRNA-seq) can be initially applied to a small cohort at a single time point to identify clinically relevant subsets, after which, the integrated subset-specific microfluidic workflow can be used to scale up to a larger cohort with multiple time points, increasing the study’s resolution and statistical power while lowering its cost. Another example would be the application of cell subset enrichment to target cells of interest ahead of scRNA-seq. This type of workflow could markedly improve the efficiency of scRNA-seq studies that target rare cell subsets by reducing the number of nontarget cells that need to be processed and sequenced to gain access to information about rare cells of interest.

**METHODS**

**Study samples**

Human blood samples were obtained either from Research Blood Components (MA, USA) for technical validation experiments or from collections at the Brigham and Women’s Hospital (MA USA) (table S1). Research on the samples was approved by Institutional Review Boards at the participating hospitals.

**Table 1. RNA-seq library statistics for the samples generated in this study.** Two technical replicates are performed for each sample and isolated subset; n represents the total number of RNA-seq libraries generated for each column. Values are shown as means ± SD. rRNA, ribosomal RNA.

|                | Benchtop Lysates (n = 12) | Benchtop PBMCs (n = 10) | Microfluidic PBMCs (clinical study) (n = 24) | Microfluidic PBMCs (clinical study) (n = 32) | Healthy PBMCs (clinical study) (n = 34) |
|----------------|--------------------------|-------------------------|---------------------------------------------|---------------------------------------------|-----------------------------------------|
| Estimated library size (millions) | 16.1 ± 1.7              | 12.4 ± 5.5              | 7.7 ± 5.47                                  | 13.0 ± 8.7                                  | 6.3 ± 4.6                               | 8.9 ± 3.6 |
| Genome map rate (%)         | 91.6 ± 8.7              | 88.9 ± 1.7              | 86.8 ± 2.5                                  | 77.0 ± 16.3                                 | 79.8 ± 6.3                              | 82.5 ± 5.7 |
| Transcript map rate (%)     | 48.0 ± 7.0              | 63.0 ± 10.0             | 45.0 ± 15.0                                 | 65.0 ± 13.0                                 | 61.9 ± 10.5                             | 63.1 ± 8.7 |
| Gene count                  | 13,592 ± 512            | 10,850 ± 481            | 10,336 ± 848                                | 8899 ± 1379                                 | 9091 ± 1956                             | 9817 ± 1757 |
| rRNA (%)                    | 8.67 ± 1.76             | 0.22 ± 0.06             | 2.48 ± 1.17                                 | 0.56 ± 0.37                                 | 1.29 ± 1.07                             | 1.34 ± 1.08 |

Compared to matched healthy control subjects based on gene set enrichment analyses (41), in line with our expectation that gene set enrichment analysis would have higher statistical power than analysis of single gene expression (Fig. 4A). This finding is in agreement with other studies that show the role of type I IFNs in the pathogenesis of SLE (42, 43) and validates the robustness of our method in identifying disease-relevant signatures. We found that IFN targets are enriched across all four subsets, which is expected based on previous profiling studies of cells from SLE patients (20, 44–46). Gene targets of the fusion protein NUP98-HOXA9, a potent driver of myeloid leukemia, were also enriched in all subsets. This supports previously published evidence that dysregulated lymphocyte proliferation is associated with both cancer and autoimmune disease and could explain the increased risk of malignancy in patients with SLE (47).
Fig. 3. Genomic characterization of PBMCs treated in vitro. (A) Unsupervised clustering of untreated (NT) and treated (DEX, IFN, and LPS) PBMC subsets based on differentially expressed genes (false discovery rate < 0.05). Venn diagrams showing numbers of (B) treatment-specific differentially expressed genes for each subset and (C) subset-specific differentially expressed genes for each treatment. (D) Gene set enrichment analysis (Reactome sets, FDR < 0.01) of IFN-treated B cells and monocytes. Red nodes indicate up-regulation, while blue nodes indicate down-regulation. Node sizes are proportional to the number of genes in the gene set, while edge lengths are inversely proportional to the number of overlapping genes between the sets. (E) Normalized fold change (FC) in expression of Jak-STAT pathway genes in IFN-treated samples over untreated controls. Each node shows the fold-change expression of each gene in the three subsets profiled (left block, monocytes; middle block, B cells; right block, CD4 T cells).
Review Boards at the Broad Institute of Massachusetts Institute of Technology (MIT) and Harvard (MA, USA) and Brigham and Women's Hospital (MA, USA). Blood samples from patients with SLE and healthy control donors were drawn with EDTA Vacutainer tubes (BD Biosciences) and processed within 3 hours of collection.

**Isolation of PBMCs from whole blood**

Cells were isolated from whole blood samples using density gradient centrifugation. Whole blood was diluted 1:1 with 1× phosphate-buffered saline, layered on top of Ficoll-Paque PLUS (GE Healthcare), and centrifuged at 1200g for 20 min. The PBMC layer was retrieved, resuspended in 10 ml of RPMI-1640 (Gibco), and centrifuged at 300g for 10 min. The cells were counted using a manual hemocytometer, resuspended in fetal bovine serum (FBS) (Gibco) with 10% dimethyl sulfoxide (Sigma), and aliquoted in 1-ml cryopreservation tubes at a concentration of 5 million cells/ml. The tubes were kept at −80°C overnight and then transferred to liquid nitrogen for long-term storage. Before processing, cells were thawed at 37°C for 3 min, resuspended in 10 ml of RPMI-1640 supplemented with 10% FBS (Gibco), and centrifuged at 300g for 5 min. The cells were then resuspended in the desired concentration or buffer, depending on the experiment.

**Microfluidic device design and fabrication**

The microfluidic device was fabricated using a previously published protocol (27) with minor modifications. Flow layer molds were patterned in three steps: (i) rectangular (75 μm), (ii) rectangular (200 μm), and (iii) rounded (60 μm). All silicon wafers were precoated with hexamethyldisilazane (Sigma) before spin coating. Rectangular features were prepared by spin-coating SU-8 2075 (MicroChem) on a silicon wafer. The coated wafers were patterned by ultraviolet (UV) exposure (OAI 206 mask aligner) through a mask printed at 20,000 dots per inch (Fineline Imaging, design files are included in the Supplementary Materials). The features were then developed using an SU-8...
Magnetic affinity cell isolation and microfluidic implementation

Magnetic affinity cell sorting was done using commercially available EasySep kits (CD14, CD19, CD4, and CD8 positive isolation II and T cell negative isolation) from STEMCELL Technologies. To implement the isolation protocols on the device, the buffers were modified and volumes were scaled accordingly. EasySep buffer (STEMCELL Technologies) was supplemented with 10% FBS (Gibco) and 0.2% Pluronic-F127 (Sigma), to reduce nonspecific cell adhesion in the polydimethylsiloxane channels. The microchannels were also preincubated with 1% Pluronic-F127 (Sigma) before cell isolation. Neodymium magnets (Grainger) with a 43-lb pull were used for all magnetic capture steps.

Flow cytometry and FACS

For assessment of isolation purities, flow cytometry was performed using the Cytoflex system (Beckman Coulter). For RNA-seq library validation experiments and benchmark comparisons, PBMCs were sorted using the MoFlo Astrios (Beckman Coulter). Lysate pools were generated by sorting 5000 cells into 20 μl of TCL buffer (Qiagen) and 1 μl protease K (20 mg/ml; Qiagen) and stored at −80°C to maintain RNA integrity. The following panel was used for both purity assessment and sorting: DAPI, CD45 BV605, CD3 AF700, CD4 FITC, CD8 PE, CD14 APC, CD19 PE-Cy7, and CD56 BV650 (all IgG1, BioLegend). Flow cytometry data were analyzed using FlowJo v10.1.

Low-input RNA-seq, microfluidic implementation, and sequencing

RNA-seq was performed using Smart-Seq2 (29) with minor modifications. Cells were sorted into 19 μl of TCL buffer and 1 μl protease K (20 mg/ml; Qiagen), and their RNA was purified by a 2.2× SPRI cleanup with RNAClean XP magnetic beads (Agenecut) before reverse transcription. For the microfluidic implementation of the protocol, Tween 20 (Teknova) was added to all reactions at a final concentration of 0.5%. For mRNA capture, a biotinylated oligo (/5BiosG! - AAGCAGTGGTATCAACGCAGAGTAC-30T-VN) (Integrated DNA Technologies) was attached to streptavidin magnetic beads (New England Biolabs) following the manufacturer’s protocol. The beads were then used to capture mRNA from the lysates and were washed with 10 mM tris-HCl (pH 7.5), 0.15 M LiCl, 1 mM EDTA, and 0.5% Tween 20. The beads were then resuspended in the reverse transcription mix, following the same steps as the published protocol. cDNA processed on the benchtop and microfluidic device were amplified for 18 and 22 cycles, respectively. After amplification and cleanup, libraries were quantified using a Qubit fluorometer (Invitrogen), and their size distributions were determined using the Agilent Bioanalyzer 2100. After normalizing the amplicon concentrations to 0.1 to 0.2 ng/μl, sequencing libraries were constructed using the Nextera XT DNA Library Prep Kit (Illumina), following the manufacturer’s protocol. All RNA-seq libraries were sequenced with 38 × 37 paired-end reads using a MiSeq or NextSeq (Illumina).

In vitro stimulation of PBMCs

Healthy PBMCs were resuspended in RPMI-1640 supplemented with 10% FBS and 1× penicillin-streptomycin (Gibco). Cells were cultured at a density of 1 M/ml and stimulated with LPS (5 μg/ml) (eBioscience), DEX (100 nM) (Millipore), IFN-α (250 U/ml) (Abcam), or no treatment. The cells were cultured for 24 hours at 37°C in a 5% CO2 environment before processing through the microfluidic device.

RNA-seq data analysis

RNA-seq libraries were sequenced to a depth of 5 to 15 million reads per sample. All technical validation libraries were subsampled to 10 million reads to remove potential confounding effects of sequencing depth. Sequencing reads were aligned to the UCSC hg19 transcriptome using STAR (49) and used as input to generate QC statistics with RNA-SeQC (50). RSEM (51) was used to generate an expression matrix for all samples. Both raw count and transcripts per million data were analyzed using edgeR and custom R scripts. Lowly expressed genes with log2 (counts per million) less than 5 were filtered out before analysis. Gene set analyses were performed using the Kolmogorov-Smirnov test implementation in GAGE (52). Cyto-scape and the enrichMap (53) module extension were used to visualize pathway-specific differential expression data.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/5/1/eaau9223/DC1

Fig. S1. Detailed schematic of integrated workflow implemented on microfluidic chip.
Fig. S2. Flow cytometry gating strategy.
Fig. S3. Chip sorting optimization and comparison with benchtop magnetic isolation.
Fig. S4. Representative Bioanalyzer traces for conventional benchtop and microfluidic device.
Fig. S5. Differential gene expression analysis between microfluidic and benchtop RNA-seq libraries.
Fig. S6. Chip enrichment validation by gene expression profiles.
Fig. S7. Comparison between conventional and device-processed RNA-seq libraries (full workflow).
Fig. S8. RNA-seq correlation plots for in vitro–treated PBMCs.
Fig. S9. Gene set enrichment analysis (Reactome sets) comparing treated PBMCs versus control.
Fig. S10. Volcano plots showing differentially expressed genes between patients with SLE and matched healthy controls for four different subsets.
Fig. S11. Heat map showing relative IFN-signature scores across different cell types of 10 patients.

Supplementary Note

Supplementary Methods

Table S1. Ex vivo treatment differential expression (DE) results.
Table S2. Patient information for clinical study.
Table S3. SLE DE and gene set enrichment analysis results.
Data file S1. Device design.
Data file S2. Gene expression matrices.

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B cell, T cell and myeloid cell transcriptomes display unique profiles and each subset

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