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High-throughput chromatin accessibility profiling at single-cell resolution

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Here we develop a high-throughput single-cell ATAC-seq (assay for transposition of accessible chromatin) method to measure physical access to DNA in whole cells. Our approach integrates fluorescence imaging and addressable reagent deposition across a massively parallel (5184) nano-well array, yielding a nearly 20-fold improvement in throughput (up to ~1800 cells/chip, 4–5 h on-chip processing time) and library preparation cost (~81¢ per cell) compared to prior microfluidic implementations. We apply this method to measure regulatory variation in peripheral blood mononuclear cells (PBMCs) and show robust, de novo clustering of single cells by hematopoietic cell type.

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A central challenge of systems biology is to determine the epigenome of phenotypically distinct cellular states within complex primary tissue. Toward this goal, single-cell chromatin accessibility measurements provide an important epigenetic view of the regulatory landscape within individual cells by capturing the physical accessibility of putative functional elements across the genome^{1–6}. Methods for measuring chromatin accessibility at single-cell resolution, however, are low throughput, depth limited, or require complex molecular processing to generate cellular indexing reagents^{2–5,7}. For ultra-high throughput accessibility profiling applications, combinatorial indexing approaches^{2,7} offer significant promise, yet these methods capture fewer accessible fragments per cell than single-cell isolation technologies^{1,3} and are not amenable to integration with single-cell microscopy or other multi-omic assays that require whole,

live cells. In this report, we describe a high-throughput implementation of single-cell ATAC-seq⁸ (scATAC-seq) that directly integrates fluorescence imaging and provides an extensible foundation for multi-omic epigenetic profiling in single cells.

Results

Implementation of scATAC-seq on nanoliter-scale wells. We have implemented scATAC-seq in small volumes (μ ATAC-seq) using a recently developed nanoliter-scale liquid deposition system (ICELL8 Single Cell System, Takara Bio USA). This approach reduces reagent costs and achieves equal or higher per-cell fragment counts than prior state-of-the-art implementations^{2,3,7}. The workflow—illustrated in Figure 1a—is comprising of the following steps: (1) isolated single cells are stained with Hoechst and

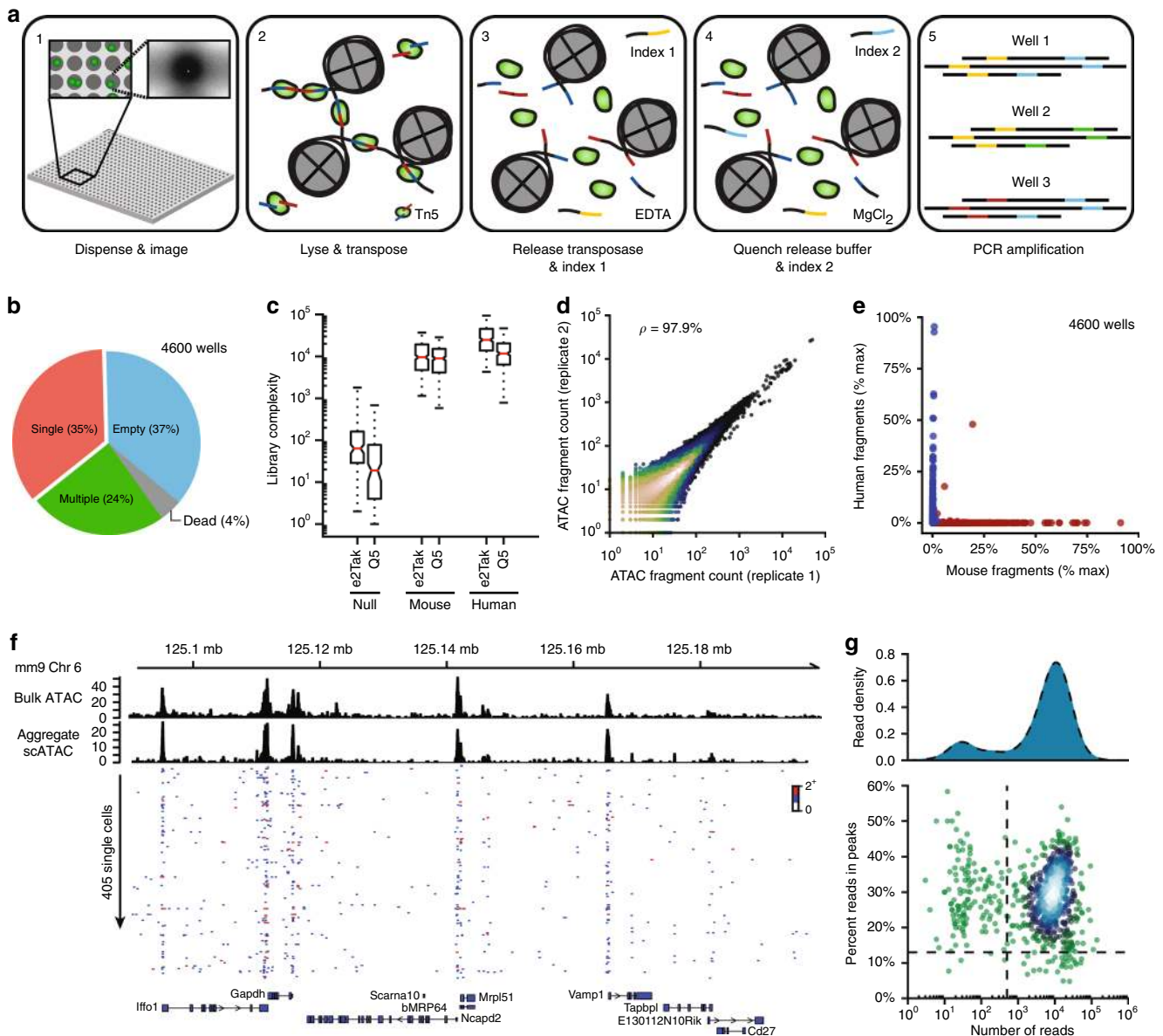


Fig. 1 μ ATAC-seq: a nano-well scATAC-seq implementation on the ICCELL8 platform. **a** μ ATAC-seq workflow. **b** Distribution of cell counts per well measured by fluorescence microscopy (Hoechst). **c** μ ATAC-seq library complexity for null, mouse, and human targeted wells using two separate polymerases (e2Tak and Q5) for well barcoding and amplification ($n = 5000$ wells). For each sample, the box denotes the interquartile range centered at the median (red line), while the whiskers span the 5th and 95th percentile range. **d** Correlation between nano-well chips processed with either a e2Tak (replicate 1) or Q5 polymerase (replicate 2) across all accessible loci. **e** Inter-well mixing of mouse and human μ ATAC-seq fragments. **f** Representative population²² and single-cell ATAC-seq genome tracks for the *Gapdh* locus. **g** Signal-to-background (percent reads in peaks) as a function of read depth ($n = 792$). Only cells lying in the upper right quadrant (marked by dashed lines) are retained for downstream analysis

propidium iodide and stochastically loaded under Poisson statistics (~1 cell per well on average) across 5184 wells under active humidity and temperature control; all wells are then imaged via multi-color microscopy to identify those containing a single-live cell; (2) transposition reagents are added to a selected set of wells (e.g., those containing a single live cell) and incubated at 37 °C for 30 min; (3) the transposition reaction is quenched by incubation with EDTA; (4) MgCl₂ is added in equimolar concentration to quench the chelating capacity of EDTA in preparation for subsequent PCR amplification; (5) PCR reagents are added and μ ATAC-seq fragment libraries are amplified using barcoded primers provided in the prior two steps (see Supplementary Table 1 for reagent loading chart). Following on-chip library construction, indexed μ ATAC-seq libraries are extracted from all nano-wells by centrifugation, purified, and then further amplified as necessary for sequencing (Methods section).

Benchmarking analysis of μ ATAC. As an initial test of μ ATAC-seq, we loaded samples into 5000 wells across two nano-well ICELL8 chips. On each chip, 200 wells were loaded with PBS (designated null wells); 1150 wells were loaded with mouse embryonic stem cells (mESCs, ~1 cell per well); and 1150 wells loaded with human lymphoblastoid GM12878 cells (~1 cell per well). This yielded a total of 4600 wells targeted with either human or mouse cells across both chips. Imaging of Hoechst and propidium iodide fluorescence revealed the anticipated fraction of wells containing live single cells (35%, 1616 single cells), consistent with near optimal loading that maximizes the number of single-cell containing wells (Fig. 1b). Barcoded sequencing of each of the 5000 targeted wells revealed 14.3×10^3 (8.1×10^3) median fragments per single human (mouse) cell containing wells ($n = 1616$)—reflecting a two orders of magnitude enrichment over null wells (Fig. 1c and Supplementary Figure 1a,b). These library complexities compare favorably with microfluidic cell capture (5.8×10^3 fragments per GM12878 cell³) as well as combinatorial indexing (2.5×10^3 fragments per GM12878 cell⁷) approaches. The μ ATAC-seq libraries capture both sub-nucleosome as well as nucleosome length fragments, yet, the median fragment length is shorter than that observed using the Fluidigm C1 platform. Consistent with prior bulk and single-cell ATAC-seq libraries, we observe a more than tenfold enrichment for fragments proximal to transcription start sites (TSS) relative to distal regions, reflecting a high fraction of fragments captured within open rather than closed chromatin (Supplementary Figure 2a). Furthermore, we find a high degree of concordance (97.9%) between nano-well chips even when μ ATAC-seq fragments are amplified with different polymerases (Fig. 1d). We further tested the deposition fidelity of the ICELL8 platform, observing both human and mouse cells in fewer than 0.2% of wells (Fig. 1e).

Aggregate single-cell profiles recapitulate population measurements broadly across the accessible genome (Supplementary Figure 2b) as well as specifically at individual genomic loci (Fig. 1f). At single-cell resolution, accessibility profiles are enriched for open chromatin (Fig. 1f, g) in both mESCs (29% reads in peaks, Fig. 1g) and GM12878 cells (22% reads in peaks, Supplementary Figure 2c). Collectively, these data establish the proposed nano-well implementation as a high-throughput framework for scATAC-seq library construction.

Epigenetic signature distinguishes PBMC types. We next asked whether μ ATAC-seq epigenetic profiles are sufficient to distinguish cell types within complex primary tissue. For this purpose, we performed μ ATAC-seq on human peripheral blood mononuclear cells (PBMCs) as well as B, T, CD4⁺ T, CD8⁺ T, and

monocyte cells isolated directly from whole blood (Fig. 2a), yielding 2333 single cells passing all quality control criteria (Methods section). Using ChromVar, a bioinformatic approach described previously⁹, we calculated the relative accessibility of transcription factor (TF) binding motifs in individual cells and found that isolated B, T, and monocyte cells robustly cluster by cell type (Supplementary Figure 3a). By aggregating fragments within single cells that are proximal to a TF motif, this epigenetic signature captures the variation in putative TF binding site accessibility across a population of cells⁹. A relatively small fraction of cells are incorrectly assigned to clusters; however, the frequency of these events as well as the random distribution of these cells within apposing clusters both suggest that isolation impurity upstream of the μ ATAC-seq assay is the primary source of these errors (Supplementary Table 2). PBMC subpopulations co-cluster precisely with the isolated cell types (Fig. 2b, c), showing highly concordant cell type-specific accessibility patterns within appropriate tSNE¹⁰ (t-Distributed Stochastic Neighbor Embedding) clusters (Fig. 2c) as well as k-means clustering across highly variable TF binding motif accessibility patterns (Fig. 2b). Consistent with published gene expression data, we find that the PU.1 binding motif is differentially accessible in monocytes and B cells relative to T cells (Fig. 2c, upper right panel)^{11,12}, the C/EBP α motif is exclusively accessible in monocytes (Fig. 2c, lower left panel)^{13,14}, and RUNX1 motif accessibility is appropriately enriched in T cells—reflecting the broad regulatory role of the RUNX protein family in T lymphocytes (Fig. 2c, lower right panel)¹⁵. These results are highly robust to biological (three human blood donors) and technical variation (Supplementary Figure 3b). To further establish the robustness of clustering by cell type, we independently purified CD4⁺ and CD8⁺ T cells and found that these subtypes co-cluster with independently isolated T cells (Fig. 2c, upper left panel). Collectively, these data suggest that μ ATAC-seq signatures are sufficient for de novo clustering of PBMCs by hematopoietic cell type.

Discussion

In this report, we have described μ ATAC-seq—a high-throughput, single-cell chromatin accessibility assay that dramatically reduces per-cell costs, requires only commercially available reagents, provides state-of-the-art data quality, and increases throughput nearly 20-fold over existing single-cell capture technologies. Single-cell chromatin measurements present a unique experimental challenge since only two DNA templates are present in a diploid cell. Technical sampling noise as well as biological heterogeneity further confound this problem, resulting in a ~10% observation efficiency of accessible regions in single cells³. Consequently, a few hundred cells are typically required to reliably determine the accessibility landscape of each subpopulation within a mixture of cells. Our approach in this work has been to develop an experimental framework for processing more than a thousand of cells in parallel to determine the accessibility of multiple cell types within a complex tissue. In general, nano-well single-cell sequencing approaches such as μ ATAC-seq are highly extensible, well-suited for multi-omic analysis, and define an important direction for single-cell epigenetic methods development.

Methods

Cell culture. All cell lines were grown at 37 °C with 5% CO₂. GM12878 cells were obtained from the laboratory of Michael Snyder (Stanford University) and were cultured in RPMI 1640 media supplemented with L-glutamine (Thermo Fisher Scientific, MA, USA, Cat. #11875-085) and 10% FBS (Thermo Fisher Scientific, Cat. #10082147); mESC cells (129S1X Castaneous, gift from Howard Chang) were cultured in 15% FBS (HyClone GE Healthcare Life Sciences, SH30070.03E) supplemented with non-essential amino acids, L-glutamine and Leukemia Inhibitory

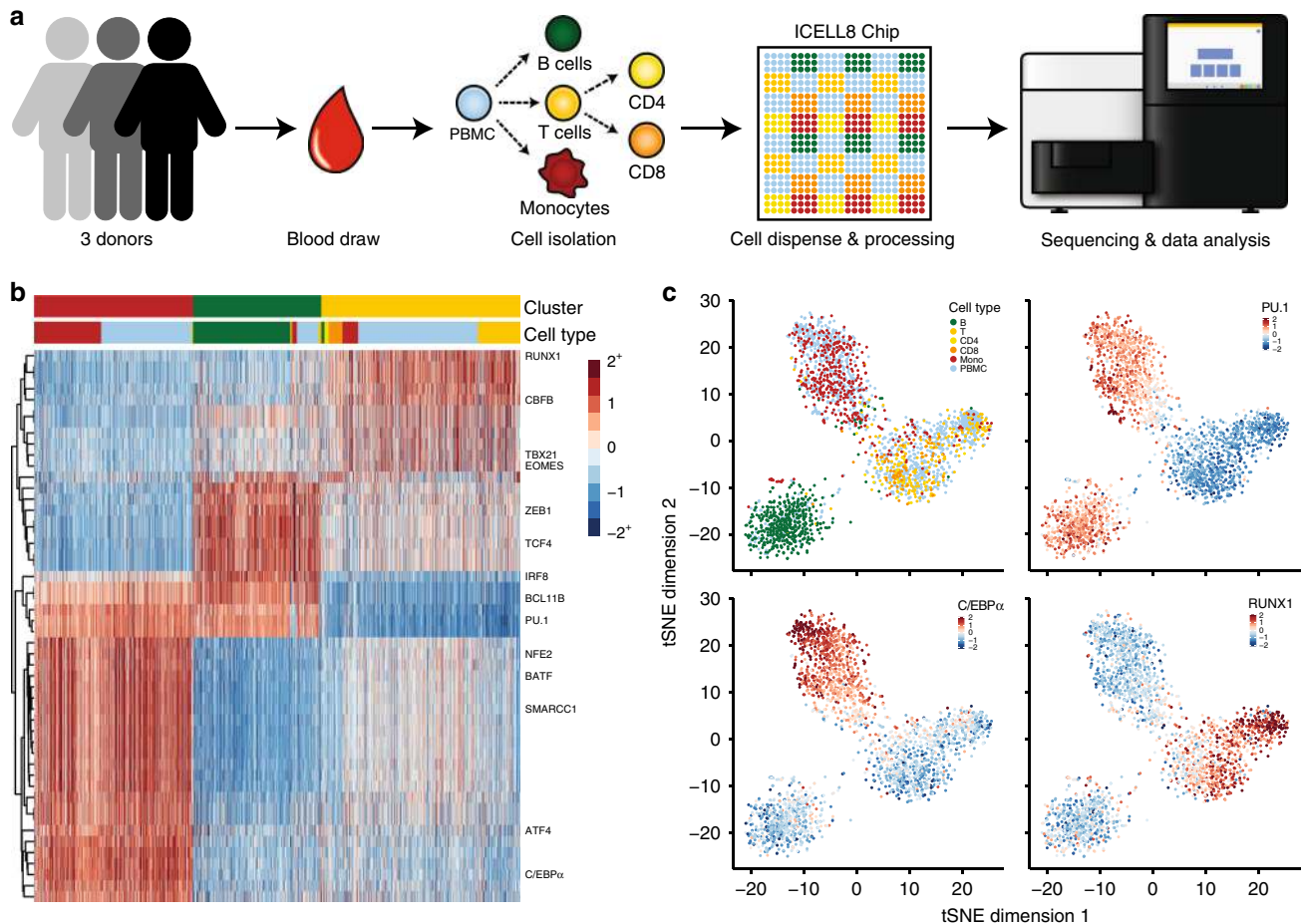


Fig. 2 De novo identification of hematopoietic cell types by μ ATAC-seq. **a** Human PBMC isolation and μ ATAC-seq workflow. **b** Hierarchical (TF motifs) and k-means (cells) clustering of accessibility deviation z-scores across 2333 single cells (columns) of the 50 most variable TF motifs (rows). Colors correspond to cell types defined in **a**. **c** tSNE visualization of accessibility deviations at TF motifs for cells in **b**. Cells are either colored by cell type (upper left panel) or by the accessibility deviation z-score for the specified TF motif

Factor (LIF, Invitrogen, Cat. #A35935). Adherent mESCs were washed twice in 1X PBS and detached using trypsin (Sigma, MO, USA) for 5 min. Cells were diluted in their respective media, collected by centrifugation at 400 g for 5 min, and then resuspended in media.

Immune cell isolation from whole blood. Monocytes, T cells, CD4⁺ T cell, CD8⁺ T cells, and B cells were isolated from whole blood (AllCells, CA, USA) using EasySep Direct Human cell isolation kits (STEMCELL Technologies, MA, USA) according to the manufacturer's protocol. Isolated PBMCs (AllCells, CA, USA) were thawed in RPMI and washed once in media before staining the cells as described below. All human cells were obtained from AllCells with explicit consent to publish data for broad genomic release.

ICELL8 workflow. Cells were stained with Hoechst and propidium iodide using the ReadyProbes Cell Viability Imaging Kit (Thermo Fisher Scientific) for 20 min in media at 37 °C, then washed twice in cold 0.5X PBS. Cells were counted and dispensed into nano-wells using the SMARTerTM ICCELL8[®] Single-Cell System (Takara Bio USA, CA, USA, Cat. #640000) at 25 cells/ μ l in 0.5X PBS, 1X Second Diluent (Takara Bio USA, Cat. # 640196) and 0.4 U/ μ l RNase Inhibitor (New England Biolabs [NEB], MA, USA) into a SMARTer ICCELL8 250v chip (Takara Bio USA, Cat. #640183). Control wells containing 1X PBS (25 μ l) and fiducial mix (25 μ l) (Takara Bio USA, Cat. #640196) were included in the source loading plate (see source plate loading chart in Supplementary Table 1). The on-chip deposition volume was 40 nl for all reagent delivery steps. The chips were maintained at 16 °C or lower between all reagent loading steps. Following cell deposition, chips were sealed with SMARTer ICCELL8 imaging film (Takara Bio USA, Cat. #640014) and centrifuged at 400 g for 5 min at 4 °C and imaged with a 4 \times objective using Hoechst and propidium iodide fluorescence. Images were analyzed using automated microscopy image analysis software (CellSelect, Takara Bio USA). Immediately

following imaging, the Tn5 transposition mix (2X TD buffer [20% dimethylformamide, 20 mM Tris-HCl, pH 7.6, 10 mM MgCl₂], 100 μ l Tn5 transposase [Nextera DNA Library Prep Kit, Illumina, CA, USA] per ml Tn5 transposition mix, 0.2% Tween 20, 0.2% NP40, and 0.02% Digitonin [Promega, WI, USA]) was dispensed. Chips were then sealed with imaging film, centrifuged at \sim 3000 g for 5 min at 4 °C and incubated for 30 min at 37 °C. To index the whole chip, 72 i5 and 72 i7 previously published, custom indices (Supplementary Table 3)³ were dispensed at 6.25 μ M concentration with EDTA and MgCl₂, respectively. To release the bound Tn5 transposase, 60 mM EDTA was dispensed together with the i5 indexes. After sealing the chip, it was centrifuged at 3000 g for 3 min and incubated for 30 min at 50 °C. Prior to performing PCR on-chip, the chelating capacity of EDTA was suppressed by dispensing 60 mM MgCl₂ together with the i7 indices. Chips were then sealed with imaging film, centrifuged, and incubated at room temperature for 5 min. Finally, a PCR mix (5x Q5 [NEB] or e2TAK [Takara Bio USA] reaction buffer, 1 mM dNTPs [Thermo Fisher Scientific], and 100 U/ml Q5 [NEB] or 50 U/ml e2TAK polymerase [Takara Bio USA], respectively) was dispensed and 14 cycles of PCR were performed on-chip after sealing with TE Sealing film (Takara Bio USA, Cat. #640109) and centrifuging at \sim 3000 g (3 min) as follows: 5 min at 72 °C and 30 s at 98 °C followed by 14 cycles of 10 sec at 98 °C and 90 s (Q5 polymerase) or 150 s (e2TAK polymerase) at 72 °C, with a final extension of 2 min at 72 °C. PCR products were extracted by centrifugation at \sim 3000 g for 10 min using the supplied SMARTer ICCELL8 Collection Kit (Takara Bio USA). All dispense and sealing steps were followed by centrifugation at \sim 3000 g for 3 min. All on-chip thermal cycling was performed using a SMARTer ICCELL8 Thermal Cycler (Takara Bio USA).

Off-chip purification and additional amplification. The collected PCR product was purified using MinElute PCR purification columns (Qiagen, Germany) following the manufacturer's instructions. Due to the large sample volume, the PCR product was split across four MinElute columns, eluted in 10 μ l volumes, and

subsequently pooled. To remove free PCR primers, which would induce index-swapping during additional rounds of off-chip amplification, we performed two rounds of bead clean-up using Ampure XP beads (Beckman Coulter, CA, USA) in a 1:1.2 ratio. The beads were incubated for 8 min with the PCR product, washed twice in 70% ethanol, and eluted in 20 μ l ultrapure water (Thermo Fisher Scientific). Further amplification was required only for the mouse and human mixing experiment. PBMCs libraries generated on-chip were directly sequenced following column and bead purifications.

The number of required off-chip amplification cycles was determined by running a 20 μ l qPCR reaction (2 μ l PCR product, 0.5 μ M oligo C [Illumina P5], 0.5 μ M oligo D [Illumina P7], 0.6X SYBR Green I [Thermo Fisher Scientific], and 1X NEBNext High-Fidelity 2X PCR Master Mix [NEB]): 30 s at 98 °C, followed by 20 cycles of 10 s at 98 °C and 30 s at 63 °C and 1 min at 72 °C. The remaining 18 μ l PCR product was amplified the number of PCR cycles corresponding to 1/3 of the maximum fluorescence intensity. The amplified PCR product was then purified and concentrated using a Qiagen MinElute column.

DNA sequencing. All libraries were sequenced on a NextSeq 500 (Illumina) using the high output v2 kit (Illumina) in 76 \times 8 \times 8 \times 76 cycle mode, although 38 bp \times 8 \times 8 \times 38 bp sequencing is sufficient. On average, ~50 K reads were sequenced per cell. Due to the nature of the sequencing libraries 30–40% phiX control v3 (Illumina) was spiked in and 1.5 pM were loaded onto the flow cell.

Per cell cost estimate. The per cell library preparation cost is conservatively estimated (assuming only 1200 single cells captured per chip) at 81¢/cell: (1) Takara Bio ICELL8 chip (52¢/cell), (2) Illumina Tn5 (24¢/cell), (3) e2Tak polymerase (4¢/cell), (4) other reagents contribute <1% additionally. The additional per cell sequencing cost at the depth used for this report (assuming a 75 cycles NextSeq 500/550 High Output v2 Kit) is approximately 17¢/cell.

Data analysis. Illumina sequencing reads in BCL format were demultiplexed by single-cell barcode to fastq files using bcl2fastq (Illumina) according to the manufacturer's manual. Reads were trimmed using Cutadapt¹⁶ (parameters: -a Trans2_rc = CTGTCTCTTATACACATCTCCGAGCCACGAGACA, Trans1_rc = CTGTCTCTTATACACATCTGACG CTGCCGACGA) and aligned to either the human (hg19) or mouse (mm9) genomes using Bowtie2¹⁷. Mitochondrial reads were removed prior to downstream analysis. PCR duplicates were identified and removed if either the start or end position was shared with another sequencing read. Library complexity estimates were obtained using the Picard Tools MarkDuplicates utility (<https://broad-institute.github.io/picard/>), except for empty well where too few reads were present for a robust estimate; in the latter case, the library complexity was estimated as the number of unique reads observed. Accessible chromatin regions (peaks) were determined using MACS2¹⁸ (parameters: --format BAMPE --nomodel --call-summits --nolambda --keep-dup all) for mouse embryonic stem cells (mESCs) and human lymphoblastoid (GM12878) cells. A previously published accessible peak set for hematopoiesis was used for PBMC, T- and B-cell analysis¹. Single cells were selected based on imaging using the supplied ICELL8 CellSelect software (Takara Bio USA). Primary PBMCs with fewer than 500 unique (non-mitochondrial) reads or with <20% (10–15% for mESCs and GM12878 cells) of mappable reads lying within peaks were eliminated from subsequent analysis. Bias-corrected deviations in accessibility near transcription factor motifs were calculated using ChromVar⁹. Bias-corrected deviations were linearly transformed to truncated z-scores with minimum and maximum values of -2 and 2, respectively. K-means clustering ($k = 3$) was performed on the 50 most variable transcription factor motifs to assign each single cell to a specific cluster. Transcription factors (rows) were then hierarchically clustered using the ward.D2 agglomeration method^{19,20} within the R pheatmap package²¹, while single cells (columns) were ordered by assigned cluster and cell type (Fig. 2b). Visualizations of clustering and tSNE¹⁰ analyses were constructed using R scripts.

Data availability

The sequencing data that support the findings of this study are available in Figshare under the following DOIs: Metadata: doi: [10.6084/m9.figshare.7006154.v1](https://doi.org/10.6084/m9.figshare.7006154.v1); Human monocyte cells: doi: [10.6084/m9.figshare.7005707.v1](https://doi.org/10.6084/m9.figshare.7005707.v1); Human lymphoblast cells (GM12878): doi: [10.6084/m9.figshare.7005713.v1](https://doi.org/10.6084/m9.figshare.7005713.v1); Human peripheral blood mononuclear cells (PBMCs): doi: [10.6084/m9.figshare.7005752.v1](https://doi.org/10.6084/m9.figshare.7005752.v1); Mouse embryonic stem cells (mESCs): doi: [10.6084/m9.figshare.7005710.v1](https://doi.org/10.6084/m9.figshare.7005710.v1); Human CD8⁺ T cells: doi: [10.6084/m9.figshare.7005701.v1](https://doi.org/10.6084/m9.figshare.7005701.v1); Human CD4⁺ T cells: doi: [10.6084/m9.figshare.7005698.v1](https://doi.org/10.6084/m9.figshare.7005698.v1); Human T cells: doi: [10.6084/m9.figshare.7005683.v1](https://doi.org/10.6084/m9.figshare.7005683.v1); Human B cells: doi: [10.6084/m9.figshare.7005539.v1](https://doi.org/10.6084/m9.figshare.7005539.v1). All other data are available from the authors upon reasonable request.

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Author contributions

A. Mezger, S.K., S.L., and W.G. conceived the study and designed all experiments; A. Mezger and S.K. analyzed the data and performed all experiments with the help of I.M.; K.B. assisted with imaging analysis; A. Mir, M.B., A.F., P.F., and W.G. supervised the study; A. Mezger, S.K., and W.G. wrote the manuscript with input from all authors.

Additional information

Supplementary Information accompanies this paper at <https://doi.org/10.1038/s41467-018-05887-x>.

Competing interests: I.M., A. Mir, M.B., and A.F. are employees at Takara Bio USA, Inc. S.K. presented the described work on behalf of Takara Bio USA, Inc. at the Advances in Genome Biology and Technology General Meeting (2018), but has no financial interest in this work. W.G. is a scientific co-founder of Epinomics and a consultant for 10X genomics. Stanford University has filed a provisional patent application (US20160060691A1; Status: Components are pending) on the ATAC-seq methods described and W.G. is named as an inventor. A. Mezger, K.B., P.F., and S.L. declare no competing interests.

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