High-content single-cell combinatorial indexing

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Single-cell combinatorial indexing (sci) with transposase-based library construction increases the throughput of single-cell genomics assays but produces sparse coverage in terms of usable reads per cell. We develop symmetrical strand sci (‘s3’), a uracil-based adapter switching approach that improves the rate of conversion of source DNA into viable sequencing library fragments following tagmentation. We apply this chemistry to assay chromatin accessibility (s3-assay for transposase-accessible chromatin, s3-ATAC) in human cortical and mouse whole-brain tissues, with mouse datasets demonstrating a six- to 13-fold improvement in usable reads per cell compared with other available methods. Application of s3 to single-cell whole-genome sequencing (s3-WGS) and to whole-genome plus chromatin conformation (s3-GCC) yields 148- and 14.8-fold improvements, respectively, in usable reads per cell compared with sci-DNA-sequencing and sci-HiC. We show that s3-WGS and s3-GCC resolve subclonal genomic alterations in patient-derived pancreatic cancer cell lines. We expect that the s3 platform will be compatible with other transposase-based techniques, including sci-MET or CUT&Tag.

Single-cell genomics assays have emerged as a dominant platform for interrogating complex biological systems. Methods to capture various properties at the single-cell level typically suffer a tradeoff between cell count and information content, which is defined by the number of unique and usable reads acquired per cell. We and others have described workflows that use single-cell combinatorial indexing (sci)1, leveraging transposase-based library construction2 to assess a variety of genomic properties in high throughput; however, these techniques often produce sparse coverage for the property of interest. Here we describe an adapter-switching strategy, ’s3’, capable of producing one-to-two order-of-magnitude improvements in usable reads obtained per cell for chromatin accessibility (s3-assay for transposase-accessible chromatin, s3-ATAC), whole-genome sequencing (s3-WGS) and whole-genome plus chromatin conformation (s3-GCC), while retaining the same high-throughput capabilities of predecessor ‘sci’ technologies. We apply s3 to produce high-coverge single-cell ATAC with high-throughput sequencing (ATAC-seq) profiles of mouse brain and human cortex tissue; and whole-genome and chromatin contact maps for two low-passage patient-derived cell lines (PDCLs) from a primary pancreatic tumor.

The core component of many sci assays, as well as ATAC-seq, is the use of transposase-based library construction. While the transposition reaction itself (tagmentation) is highly efficient, viable sequencing library molecules are only produced when two different adapters, in the form of forward or reverse primer sequences, are incorporated at each end of the molecule. This occurs only 50% of the time (Fig. 1a). To combat this inefficiency, strategies including the use of larger complements of adapter species3, incorporation of T7 promoters to enable amplification via in vitro transcription4,5, or reverse adapter introduction through targeted6 or random priming7 or ligation8 have been developed; however, these methods are often complex and result in limited efficiency improvements. Here we present a strategy of adapter replacement to produce library molecules tagged with both forward and reverse adapters for top and bottom strands. In addition to overcoming the 50% yield limitation, the efficiency of opposite adapter incorporation is also improved when compared to standard tagmentation. This is due to the use of multiple rounds of extension as opposed to a single extension before PCR. This format permits the use of a DNA index sequence embedded within the transposase adapter complex, enabling single-cell combinatorial indexing (sci) applications, where two rounds of indexing are performed—the first at the transposition stage and second at the PCR stage3,8,9.

Our strategy, symmetrical strand sci (s3; Fig. 1b) uses single-adapter transposition to incorporate the forward primer sequence, the Tn5 mosaic end sequence and a reaction-specific DNA barcode. As with standard tagmentation workflows, extension through the bottom strand is then performed to provide adapter sequences on both ends of each molecule; however, the s3 transposome complexes contain a uracil base immediately following the mosaic end sequence. Use of a uracil-intolerant polymerase therefore prevents extension beyond the mosaic end into the DNA barcode and forward adapter sequence. A second template oligo is then introduced that contains a 3′-blocked (inverted dT) locked nucleic acid (LNA) mosaic end reverse complement sequence with a reverse adapter sequence 5′ overhang. This oligo favorably anneals to the copied mosaic end sequence, due to the higher melting temperature of LNA, and acts as a template for the library molecule to extend through and copy the reverse adapter. This results in all library fragments having both a forward and reverse adapter sequence. The LNA-templated extension is carried out over multiple rounds of thermocycling to ensure maximum efficiency of reverse adapter incorporation, which provides an additional improvement over traditional tagmentation workflows where only a single pre-PCR extension is possible. Furthermore, adapter sequences are designed...
such that standard sequencing recipes can be used instead of the custom workflows and primers that are required for current indexed transposition technologies (Supplementary Tables 1–5)11,12, making use of the TruSeq read 2 and index read 1 sequencing primer and the standard Nextera read 1 and index read 2 primer.

Results

s3-ATAC creates high-content chromatin profiles. We first sought to establish the s3 technique to assess chromatin accessibility. In s3-ATAC, nuclei are isolated and tagmented using our single-ended, indexed transposomes and carried through the adapter-switching s3 workflow (Fig. 1b). To ensure we attain true single-cell libraries without contamination from other nuclei, and minimal barcode collisions, we performed a mixed-species experiment on primary frozen human cortical tissue from the middle frontal gyrus and frozen mouse whole-brain tissue (Fig. 2a). We elected to perform this test on primary tissue samples instead of an idealized cell line set-up, that is, the library complexity. One of our mixed-species plates was sequenced to beyond 50% saturation (duplicate reads/total reads), to represent the sequencing depth obtained where diminishing returns of increased sequence depth become excessive16. For this plate, the mean sequencing saturation per cell was 63.6% and resulted in a median unique read count per cell of 178,069 (mean = 258,859, statistics on all plates can be found in Supplementary Table 6). The human cells reached a mean sequencing saturation of 56.6% with a median unique reads per cell of 99,882 (mean = 146,937) unique reads per cell (Supplementary Table 6). When compared to other single-cell ATAC-seq datasets performed on mouse whole-brain tissue, our mouse s3-ATAC libraries contain substantially greater properly paired, unique, nuclear reads per cell with 13.7-, 6.02- and 6.22-fold improvement compared to single-nucleus ATAC-seq (snATAC), 10x Genomics single-cell ATAC (scATAC) and droplet single-cell ATAC-seq (dscATAC), respectively (Fig. 2d and Supplementary Table 7)13–15. Read-count increases can be indicative of poor ATAC-seq library quality, with increased depth reflecting increased noise and loss of signal at open chromatin regions. To address this, we first assessed read pair insert sizes, revealing the characteristic nucleosome-size banding distribution of ATAC-seq (Fig. 2e)16. We next calculated transcription start site (TSS) enrichment using the approach defined by the ENCODE project (Methods). This produced notable enrichment for both species

In total, we generated 2,175 human and 837 mouse single-cell ATAC-seq profiles passing quality filters (Methods) across four PCR indexing plates (Fig. 2a). We then assessed the total unique sequence reads obtained per cell as a function of the total aligned reads, that is, the library complexity. One of our mixed-species plates was sequenced to beyond 50% saturation (duplicate reads/total reads), to represent the sequencing depth obtained where diminishing returns of increased sequence depth become excessive16. For this plate, the mean sequencing saturation per cell was 63.6% and resulted in a median unique read count per cell of 178,069 (mean = 258,859, statistics on all plates can be found in Supplementary Table 6). The human cells reached a mean sequencing saturation of 56.6% with a median unique reads per cell of 99,882 (mean = 146,937) unique reads per cell (Supplementary Table 6). When compared to other single-cell ATAC-seq datasets performed on mouse whole-brain tissue, our mouse s3-ATAC libraries contain substantially greater properly paired, unique, nuclear reads per cell with 13.7-, 6.02- and 6.22-fold improvement compared to single-nucleus ATAC-seq (snATAC), 10x Genomics single-cell ATAC (scATAC) and droplet single-cell ATAC-seq (dscATAC), respectively (Fig. 2d and Supplementary Table 7)13–15. Read-count increases can be indicative of poor ATAC-seq library quality, with increased depth reflecting increased noise and loss of signal at open chromatin regions. To address this, we first assessed read pair insert sizes, revealing the characteristic nucleosome-size banding distribution of ATAC-seq (Fig. 2e)16. We next calculated transcription start site (TSS) enrichment using the approach defined by the ENCODE project (Methods). This produced notable enrichment for both species
at 13.4 for human, well above the ‘ideal’ standard (>7) and 13.5 for mouse, within the acceptable range and just below ideal (>15, Fig. 2f). Similarly, the fraction of reads in pile-up genomic regions (peaks, FRIP) was comparable to other single-cell ATAC technologies at 31.95 and 29.15% as measured using 292,156 and 174,653 peaks for human and mouse cells, respectively. However, FRIP is largely dependent on the number of peaks called, which is influenced heavily by cell number and total sequence depth obtained. When expanding to a human cortex high-depth s3-ATAC-seq peak set, a mean of 48.1% of reads were present in peaks and mean of 78.2% of reads for mouse cells using a high-depth mouse brain ATAC-seq peak set (Supplementary Information and Methods). We further compared peak overlap between the matched datasets to assess any systematic bias in s3-ATAC, with all assays performing comparably with respect to the proportion of overlapping reads for each other assay (Fig. 2g).

**s3-ATAC resolves cell types in the mammalian brain.** With ample signal, we next sought to discern cell types present within the complex tissues. For each species, we used peaks called on aggregate data to construct a count matrix followed by dimensionality reduction using the topic-modeling tool cisTopic, which we then visualized using uniform manifold approximation and projection (UMAP) performed graph-based clustering at the topic level and processed via Signac, Clear separation of cell types was observed using marker gene signal and differential accessibility profiles (Fig. 1h). UMAP projection human cortex cell samples (n=2,175 cells; right). Integration of s3-ATAC mouse data with other datasets. Points colored by cell type (respective of h) or external dataset (gray).

**Fig. 2 | s3-ATAC on human cortex and mouse whole brain.** a, Experimental flow through and plate layout for the mixed-species experiment, including tagmentation and PCR plate conditions per well. b, c, Scatter plots of single-cell libraries with counts of unique reads aligned to mouse or human chromosomes in a chimeric reference genome. Points are colored to reflect species assignment (Methods) in both pretagmentation mixing (b) and posttagmentation mixing (c). d, Comparison of unique read counts per cell, restricted to only properly paired reads for s3-ATAC mouse whole-brain sampled cells to unique read counts produced for previously reported datasets (n=3,034, 4,117, 46,653 and 298 cells, for snATAC, 10x scATAC, dscATAC and ours, respectively). 

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Fig. 3 | s3-ATAC on human cortex inhibitory neurons. a. Subclustering and UMAP projection of human cortical inhibitory neurons (clusters 3 and 4 from h, n = 342). b. Genome coverage track of human inhibitory neurons (n = 342) aggregated over four subclusters for genomic locations overlapping MGE and CGE marker genes LHX6 and ADARB2, respectively. c. Hierarchical clustering of topic weight per cell (top). Hypergeometric test of gene set analysis enrichment for human inhibitory neuron marker genes (bottom; Fisher’s exact test, Methods), with a genome track of SST to delineate MGE cell types (right).

Notably, even with the modest cell count produced by this experiment, the quality improvements allow us to interrogate subclusters of inhibitory neurons previously difficult to distinguish in atlas-level datasets (Fig. 3a)25. With our improved cell depth, we were able to discern caudal and medial ganglionic eminence inhibitory neurons by marker gene coverage plots across 342 GAD1+ cells (CGE and MGE, respectively). From these, we identified 157 GAD1+, ADARB2 + CGE cells and 168 GAD1+, LHX6 + MGE cells (Fig. 3b). We identified 17 cells (subcluster 4) as putative doublets given the coexpression of both LHX6 and ADARB2 and excluded them from subsequent analyses. Aggregated genomic signal over our topic-based dimensionality reduction was used to support our marker gene cell subtype discrimination and describe differentially accessible loci in human cortical inhibitory neurons (Supplementary Information). We grouped topics based on cell embeddings (Fig. 3c, top) through hierarchical clustering, and observed topic-based enrichment of sites overlapping cell type specific marker genes previously defined through transcriptomics (Fig. 3c, bottom)25. This analysis did not identify a specific topic (or topics) associated with MGE/SST+ cells; however, accessibility at SST was elevated specifically in cluster 0 (Fig. 3c, right).

To further assess the impact of increased coverage in our s3-ATAC data, we performed a random read downsampling analysis followed by peak calling, topic modeling and cell type discrimination (Methods and Supplementary Fig. 1b). As expected, the increased reads produced a greater number of called peaks (Supplementary Fig. 1c), although peak calling is a function of the total reads and can be improved by either greater cell number or greater coverage per cell. When examining the downsamled data performance on cell type discrimination, major cell types could be discerned with only 15 or 20% of reads used for human and mouse datasets, respectively (Supplementary Fig. 1d). However, as reads increased, cell types separated more cleanly and produced additional granularity (for example, inhibitory neurons). As a final examination of the advantages provided by improved library complexity, we assessed the impact on sequencing depth required to reach a comparable number of unique, passing reads per cell in our s3-ATAC library when compared to sci-ATAC, which produces libraries with a lower overall complexity (Supplementary Fig. 1e). When targeting a total of 10,000 unique reads per cell, the s3-ATAC library resulted in the removal of only 1.2% of reads as PCR duplicates when compared to 27–38% for the lower-complexity preparations. This directly translates to a reduction in sequencing costs of roughly one-third, enabling studies that want to profile larger cell numbers at a lower depth to do so more efficiently.

s3 for whole-genome and chromatin conformation capture. We then extend the improvements in data quality produced by s3-ATAC to other sci- workflows. This includes our previously described sci-DNA-seq method26 that produces s3-WGS and a strategy to
incorporate the core components of HiC library preparation but without ligation junction enrichment to produce s3-GCC (Fig. 4a).

Both strategies disrupt nucleosomes to acquire sequence reads uniformly across the genome26, which we also improved for the s3 assay by optimizing fixation conditions. All experiments were performed using the same number of indexed tagmentation reactions and the same number of nuclei deposited into each PCR indexing well to achieve a comparable expected doublet rate as with the s3-ATAC technology, which produced a median of 43,367 reads per cell (mean ± s.d.), comparable to other single-cell genome sequencing techniques (Fig. 4e)10,23,24.

We performed s3-WGS and s3-GCC on two cell lines derived from a single primary pancreatic ductal adenocarcinoma (PDAC) tumor (Fig. 4b). PDAC is a highly aggressive cancer that typically presents at an advanced stage, making early detection and study of tumor progression key25. PDAC studies suffer from a low cancer cell fraction in biopsied samples, thus we used PDCLs maintained at fewer than ten passages. This method allows for multiple modalities of characterization and perturbation, while maintaining a large portion of the heterogeneity present in the tumor sample26. We targeted two PDCLs (referred to as PDAC-1 and PDAC-2) derived from the same parent tumor, which had a driver mutation in the oncogene KRAS (p.G12D) and profound genomic instability, as indicated by karyotyping (Fig. 4c). For our s3-WGS preparations, we produced 773 and 256 single-cell libraries with a mean passing read counts of 1,588,926 (combined median 73.25% sequencing saturation), respectively. The s3-GCC libraries contained 57 and 145 cells produced a mean passing read count of 973,397 (mean ± s.d.); however, this is expected given the widespread copy number alterations present in the samples. In addition to the WGS component, the s3-GCC libraries also contained reads that were identified as chimeric ligation junctions that provide HiC-like chromatin conformation signal, with a distal contact distribution comparable to bulk HiC datasets (Supplementary Fig 3a).

Across both samples, we identified a mean of 118,048 reads per cell that capture genomic contacts at least 50 kb apart from one another, a 14.8-fold improvement over the previous high-throughput single-cell combinatorial indexing technique, sci-HiC27 (Fig. 4g and Supplementary Information) and comparable to low-throughput scHiC methods that process cells individually26, with the exception
of Dip-C, which can achieve exceeding $1 \times 10^9$ contact counts\(^1\). Read pairs spanning $\geq 50$ kb accounted for a median of 15.6 and 17.0% of the total reads obtained per cell, which equates to an enrichment of 361- and 402-fold over that of the s3-WGS libraries for PDAC-1 and -2, respectively (Supplementary Table 8).

s3-WGS and s3-GCC resolve subclonal alterations in PDCLs. We first focused our analysis on the s3-WGS and the WGS component of the s3-GCC libraries to examine the copy number alterations present in the lines. To get a sense of the genomic landscape, we first performed copy number calling on whole-exome sequencing (WES) libraries that were generated using primary tumor tissue and on the PDCL line PDAC-1, derived from the tumor (Fig. 5a). This revealed a profile of copy number aberrations at finer resolution, with a more pronounced profile in the PDCL sample, likely due in part to less euploid stromal cell contamination. We then processed all single-cell libraries using SCOPE\(^2\), which revealed a highly altered genomic landscape within each of the two samples. In line with paired karyotyping and bulk exome data, we see a similar pattern per cell of multi-megabase copy number aberrations when performing breakpoint analysis on 500-kb windows, with a median depth per window of 81 reads. Using the inferred copy number profile within genomic windows for the three samples, GM12878 and two PDAC lines, we performed hierarchical and $K$-means clustering on the Jaccard distance between cell breakpoint copy numbers at two different centroid counts. For our optimal centroid value, we found a relatively clean separation between cell lines ($k=3$), for subclonal analysis we used a higher centroid count at local optima ($k=6$). s3-WGS and s3-GCC cells cluster dependent on cell line, reflecting our ability to capture genome-wide copy number data in our s3-GCC libraries (Fig. 5b).

| Cell 1 | Cell 2 | ... | Cell N |
|-------|-------|-----|-------|
| Chr16 | Chr16 | ... | Chr16 |

![Fig. 5 | s3-WGS for copy number calling and s3-GCC for genome conformation changes.](image)

**Fig. 5 | s3-WGS for copy number calling and s3-GCC for genome conformation changes.** a. WES of primary tumor biopsy and PDAC-1 cell line. Scatterplot of reads per bin with a shading of called copy number variation. b. Single-cell whole-genome copy number calling on 500-kb bins genome-wide. Cells (rows) are hierarchically clustered and annotated by assay, sample and assigned clade (left). Representative single-cell contact maps (raw counts) at 1-Mb resolution for chromosome 16 and ensemble contact map profile at 500-kb resolution with compartment calling. Compartment eigenvectors are plotted above and compared to GM12878 high-depth bulk HiC data from Rao et al.\(^3\). d, UMAP projection of scHiC topic-modeling dimensionality reduction and clustering of single-cell distal contact profiles. e, Putative subclonal translocation on chr12 specific to PDAC-1 (top left) when compared to PDAC-2 (bottom right).
amplification methods, since only reads capturing the breakpoint would provide supportive evidence. To address both of these limitations, we used the HiC-like component of our s3-GCC libraries. Using read pairs spanning ≥50 kb, we produced chromatin contact maps that produced clear chromatin compartmentalization signal (Fig. 3c)27. Single cells were separated by their distal contact information via scHiC topic modeling and observed distinct clusters by PDCLs4. Notably, even at this relatively low sequencing depth, we were able to reliably tell PDCL line sparse contact profiles apart (Fig. 5d and Supplementary Fig. 3b,c). Differences between the aggregated contact maps between clusters were then used to assess unique translocation and inversion events across the sampled cells. This identified a putative intrachromosomal translocation between the 8.5–9.5 and 88.5–91.0-megabase (Mb) regions of chromosome 12 (Fig. 5e), which exhibited contact signal comparable to proximal regions in sequence space (Supplementary Fig. 3d). The putative translocation contained ATP2B1, which is commonly overexpressed in PDAC4 and the tumor suppressor gene DUSP6 (ref. 13), and is only present in PDAC-1.

Discussion

Our s3 workflow provides marked improvements over the predecessor sci platform with respect to passing reads obtained per cell without sacrificing signal enrichment in the case of s3-ATAC, or coverage uniformity for s3-WGS. We also introduce another variant of combinatorial indexing workflows, s3-GCC to obtain both genome sequencing and chromatin conformation, with improved chromatin contacts obtained per cell when compared to sci-HiC. We demonstrate the use of these approaches by assessing two patient-derived tumor cell lines with genomic instability. Our analysis reveals patterns of focal amplification for disease-relevant genes, and uncover wide-scale heterogeneity at a throughput not attainable with standard karyotyping. Additionally, we highlight the joint analysis of our protocols for uncovering the chromatin compartment disrupting effect of copy number aberrations. Furthermore, the s3 workflow has the same inherent throughput potential of standard single-cell combinatorial indexing, with the ability to readily scale into the tens and hundreds of thousands of cells by expanding the set of transposome and PCR indexes. We also expect that this platform will be compatible with other transposase-based techniques, including sci-MET5 or CUT&Tag4. Last, unlike sci workflows, the s3 platform does not require custom sequencing primers or custom sequencing recipes, removing one of the main hurdles that groups may face while implementing these technologies.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41587-021-00962-z.

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Methods

PDCl propagation. Low-passage, PDCLs were propagated from rapidly dissociated PDAC tumors and cultured for continuous propagation in culture medium containing ROCK inhibitor (Y-27632)45. Briefly, approximately 50,000 viable, dissociated tumor cells were plated to a 35 mm diameter, collagen-coated well (Gibco, A11428-02) and passed 1:3 while subconfluent until reaching 85% confluence on a 10-cm diameter dish. From a fraction of these cells, DNA was extracted to validate the presence of KRAS-G12 mutations by droplet digital PCR (BioRad, 1863506) and to validate a short tandem repeat profile that matches normal leukocyte DNA from the same patient (Genetica). PDCLs exhibited morphologies consistent with epithelial tumor cells and abundant KRT expression was detected by immunocytofluorescence using the monoclonal antibodies: AE1/AE3, C-11 and Cam5.2. To ensure that only tumor cells were cultured, G-banded karyotyping was performed by the Knight Diagnostic Research Cytogenetics Laboratory at the Oregon Health and Science University (OHSU). Chromosome spreads from more than 20 cells were observed to ensure that the culture contained 100% tumor cells.

WES and analysis. WES libraries for the patient's blood sample, tumor biopsy and PDCLs were carried out by the Knight Diagnostic Research Cytogenetics Laboratory at OHSU. Libraries were prepared using 500 ng of fragmented genomic DNA using KAPA Hyper-Prep Kit (KAPA Biosystems) with Agilent SureSelect XT Target Enrichment System and Human All Exon V5 capture baits (Agilent Technologies), following the manufacturer's protocols. Sequencing was carried out using the Illumina HiSeq 2500 platform by the OHSU Massively Parallel Sequencing (MPS) Resource. Primer sets were aligned to bwa mem (v.0.7.15-r1140) to GRCh38 (hg38, Genome Reference Consortium Human Reference20 (GCA_000001405.2)21). The data was processed following the best practices workflow for the GATK pipeline (v.4.1.9.0)22. Exome regions annotated as 'protein-coding' were extracted from GenCode (v.35)23 and used as input for s3-WGS.

From the experimentally generated genome-wide single-nuclei data for the PDAC cell line, a formatted stepwise protocol for s3-ATAC is available (https://doi.org/10.17504/protocols.io.beb3jaqn). For s3-ATAC tissue handling, primary samples of C57/B6 mouse whole brain were minced into the chilled NIB-HEPES buffer within a dounce homogenizer. Suspended nuclei were once again held on ice until ready to proceed. A 1-µl aliquot of suspended nuclei was diluted in 90 µl of NIB-HEPES (1:10 dilution) and quantified on either a Hamacostmtm or with a BioRad TC-20 Automated cell counter following the manufacturer's recommended protocol. After an initial cell number was obtained, the stock nuclei suspension was then diluted to a concentration of 1,400 nuclei per µl.

Tagmentation plates were prepared by the combination of 420 µl of 1,400 nuclei per µl of solution with 540 µl 2x TD Buffer (Nextera XT Kit, Illumina Inc. FC-131-1024, diluted with ultrapure water) and held in a sample chamber kept at 4 °C. Fluorescent nuclei were then flow sorted gating by size, internal complexity and DAPI fluorescence profile for single nuclei following the same gating strategy as previously described45. Immediately following sorting completion, the plate was sealed and spun down for 5 min at 500 r.c.f. and 4 °C to ensure nuclei were within the buffer.

Nucleosomes and remaining transpositionss were then denatured with the addition of 0.5% (v/v) SDS (roughly 0.01% final concentration) per well of 1 µl of NPM (Nextera XT Kit, Illumina Inc.) per well was subsequently added to perform gap-fill on tagged gDNA, with an incubation at 72 °C for 10 min. Next, 1-µl 1 µM A14-LNA-ME oligo was then added to supply the template for adapter switching (Supplementary Table 3). The polymerase-based adapter switching was then performed with the following conditions: initial denaturation at 98 °C for 30 s, ten cycles of 98 °C for 10 s, 59 °C for 30 s, and 72 °C for 30 s. The plate was then held at 10 °C. After adapter switching 1% (v/v) Triton X-100 in ultrapure H2O (Sigma 93426) was added to quench persisting SDS. At this point, some plates were stored at −20 °C for several weeks while others were immediately processed.

The following was then combined per well for PCR: 16.5 µl of sample, 2.5 µl of indexed primer at 10 µM, 2.5 µl of indexed primer at 150 nM, 0.1 µl of NPM (Nextera XT Kit, Illumina Inc.), 1 µl of 0.5X SYBR Green 1 (Thermo Scientific S7563) for a µl of reaction per well (Supplementary Tables 4 and 5). A real-time PCR was performed on a BioRad CFX with the following conditions, measuring SYBR fluorescence every cycle: 98 °C for 30 s, 16–18 cycles of 98 °C for 10 s, 55 °C for 20 s, 72 °C for 30 s, fluorescent reading, 72 °C for 10 s. After fluorescence passes an exponential growth and begins to inflect, the samples were held at 72 °C for another 30 min then stored at 4 °C. Amplified libraries were then cleaned by pooling 25 µl per well into a 15–ml conical tube and cleaned via a Qiaquick PCR purification column following the manufacturer's protocol (Qiagen 28106). The pooled sample was eluted in 50 µl 10 mM Tris-HCL, pH 8.0. Library molecules then went through a size selection via SPRI selection beads (Mag-Bind TotalPur NGS Omega Biobad 13878-01). Next, 50 µl of vortexed and fully suspended room temperature SPRI beads were combined with the 50-µl library (1x clean up) and incubated at room temperature for 5 min. The reaction was then was placed on a magnetic rack and once cleared, suspended pelleting was performed. The supernatant was sent to the BioRad Sonic Detergent with 80% ethanol. After ethanol was pipetted out, the tube was spun down and placed back on the magnetic rack to remove any lingering ethanol. Next, 31 µl of 10 µM Tris-HCL, pH 8.0 were then used to resuspend the beads off the magnetic rack and allowed to incubate for 5 min at room temperature. The tube was again placed on the magnetic rack and once cleared, the full volume of supernatant was moved to a clean tube. DNA was then quantified by Qubit double-stranded DNA High-sensitivity assay following the manufacturer's instructions (Thermo Fisher Q32851). Libraries were then diluted to 2 ng/µl and run on an Agilent Tapestation 4150 D5000 tape (Agilent 5067-5392). Library molecule concentration within the range of 100–1,000 base pairs (bp) was then used for final dilution of the library. Libraries were then sequenced on the NextSeq 500 system following the manufacturer's recommendations (Illumina Inc. 20029407, 20024904). For greater sequencing effort, select libraries were also sequenced on a NovaSeq S2 flowcell, again following the manufacturer's recommendations (Illumina Inc. 20028315). For both machines, libraries were sequenced as paired-end libraries with ten cycle index reads and 85 cycles for reads 1 and 2.

s3-WGS library generation. A formatted stepwise protocol for s3-WGS is available for review at https://doi.org/10.17504/protocols.io.beb3jaqn. Before processing, the following buffers were prepared: 50 ml of NIB-HEPES buffer as described above, as well as 50 ml of a Tris-based NIB (NIB Tris) variant with the following concentrations: 10 mM Tris-HCL pH 8.0, 1 µM of MgCl2, 0.1% (v/v) IGEPAI CA-630, 0.1% (v/v) Tween and diluted in PCR-grade Ultrapure Water. After dilution, two tablets of Pierce Protease Inhibitor Mini Tablets, EDTA-free (Thermo Fisher, A32955) were dissolved and suspended to prevent protease degradation during nuclei isolation.

Days old.
s3-WGS library preparation was performed on cell lines as follows. For patient-derived PDCL cell lines, cell lines were plated at a density of 1×10⁶ on a T25 flask the day before processing. At collection, cells were washed twice with ice-cold 1× PBS (VWR 75800-986) and then trypsinized with 5 ml of 1× TrypLE (Thermo Fisher 12604039) for 15 min at 37°C. Suspended cells were then collected and pelleted at 300 g, r. c.f. at 4°C for 5 min. For suspension-growth cell lines (GM12878), cells were pipetted from growth medium and pelleted at 300 g, r. c.f. at 4°C for 5 min. Following the initial pellet, cells were washed with ice-cold 1 ml of NIB-HESPES twice. After the second wash, pellets were then resuspended in 300 µl of NIB-HESPES. Nuclei were aliquoted and quantified as described above, then aliquots of 1 million nuclei were generated based on the quantification. The aliquots were pelleted by a 300 g, r. c.f. centrifugation at 4°C for 5 min and resuspended in 5 ml of NIB-HESPES. Next, 246 µl of 16% (w/v) formaldehyde (Thermo Fisher 28906) were then added to nuclear suspensions (final concentration of 0.75% formaldehyde) to light-crosslink nuclei. Nuclei were quenched via incubation in formaldehyde solution for 10 min on an orbital shaker set to 50 rpm. Suspensions were then pelleted at 500 g, r. c.f. for 4 min at 4°C and supernatant was aspirated. Pellet was then resuspended in 1 ml of NIB Tris Buffer to quench remaining formaldehyde. Nuclei were again pelleted at 500 g, r. c.f. for 4 min at 4°C and supernatant was aspirated. The pellet was washed once with 500 µl of 1× NEBuffer 2.1 (NEB B72025) and then resuspended with 760 µl of 1× NEBuffer 2.1. Then 40 µl 1% SDS (v/v) was added and sample was incubated on a ThermoMixer at 300 g, r. c.f. set to 37°C for 20 min. Nucleosome depleted nuclei were then pelleted at 500 g, r. c.f. at 4°C for 5 min and then resuspended in 50 µl of NIB Tris. A 3 µl aliquot of nuclei was taken and digested 1:10 in NIB Tris then quantified as described above. Nuclei were diluted to 500 nucle per µl with addition of NIB Tris, based on the quantification. Depending on experimental setup, the 420 µl of nuclei at 500 nucle per µl were then combined with 540 µl of 2× TD buffer. Following this, nuclei were tagged, stained, and flow sorted, gDNA was gap-filled and adapter switching was performed as described for the s3-ATAC protocol. Library amplification was performed by PCR as described above with fewer cycles (13–15) likely due to more initial capture events per library. Libraries were then cleaned, size selected, quantified and sequenced as described previously.

s3-GCC library generation. A formatted stepwise protocol for s3-WGS is available for review at: https://doi.org/10.17504/protocols.io.beb4jaqw. The same cultured cell line samples were collected as described for s3-WGS library generation, and processed from the same pool of fixed, nucleosome depleted nuclei. Following quantification of nuclei, the full remaining nuclear suspensions (roughly 2–3 million nuclei per sample) were pooled respective samples of 3 µl aliquot of nuclei was taken and digested 1:10 in NIB Tris then quantified as described above. Nuclei were diluted to 500 nucle per µl with addition of NIB Tris, based on the quantification. The initial processing of all library types, we focused our analysis on the mouse brain libraries. We plotted our PCR plate sequenced to 36.4±1.4% unique reads/total reads for comparison to three other single-cell ATAC-seq methods that have been applied to postnatal mouse whole brain (n=3,034, 5,336 and 46,653 for smATAC, 10x Genomics scATAC and dscATAC, respectively)⁵⁰–⁵⁵. Our data were filtered to just unique reads that were properly paired via the samtools view ‘-f 2’ argument to allow for proper read-count comparison across datasets that report read pairs or fragments. Data passing self-reported filters were used for comparison and plotted with the ggplot geom_boxplot function (Fig. 2d). Welch's two-sample t-tests comparisons between unique read cell were calculated with the t.test function in base R for a one-sided alternative hypothesis. For peak overlap comparison, we added an additional sci-ATAC low sequencing effort dataset performed on adult mouse flash frozen brain tissue. We then counted unique peak overlaps between datasets and plotted as stacked bar plots via ggplot geom_bar function (Fig. 2e).

To assess tagmentation insert size, samtools (Supplementary Fig. 6). Those not passing this filter (2.7%, 19/687 cells for pretagmentation barnyard) were then split from the original FastQ files to be aligned to the proper hg38 or mm10 genomes with bwa mem as described above. Human and mouse assigned cellIDs were then processed in parallel for the rest of the analysis. After alignment, reads were again deduplicated to obtain proper estimates of library complexity (Supplementary Fig. 6).

Tagmentation insert quantification. To assess tagmentation insert size, samtools s3-ATAC analysis. Barnyard analysis. With single-cell libraries distinguished, we next quantified contamination between nuclei during library generation. We calculated the read count of unique reads per cellID aligning to either human reference or mouse reference chromosomes (Fig. 1c). CellIDs with ≥90% of reads aligning to a single reference genome were considered bona fide single cells. Those not passing this filter (2.7%, 19/687 cells for pretagmentation barnyard) were considered collisions. Collision rate was estimated to account for cryptic collisions (mouse cell–mouse cell or human cell–human cell) by multiplying by two (final collision rate of 5.5%, Fig. 2b). Bona fide single-cell cellIDs were then split from the original FastQ files to be aligned to the proper hg38 or mm10 genomes with bwa mem as described above. Human and mouse assigned cellIDs were then processed in parallel for the rest of the analysis. After alignment, reads were again deduplicated to obtain proper estimates of library complexity (Supplementary Fig. 6).
Cis-acessibility networks were generated through the Signac wrapper of Cicero (v.1.3.4.10)\(^1\). Genome track plots with cis-accessibility network linkages were generated through Signac function CoveragePlot for marker genes previously described in our Supplemental Fig. 1\(^1\). We inspected accessibility between clusters in one by one, and one by rest comparisons were generated using Signac function FindMarkers using options: test.use = `LR` and only.pos = `T`, with latent. 

**Subsampling.** For subsampling analysis, the processed, deduplicated bam files were split by cellID into single-cell bam files. Each bam file was then subsampled randomly using samtools view -s a argument for 0.5, 1, 2, 5, 10, 15, 20, 40, 50, 60 and 80%, respectively. Following this single-cell subsampled bam were collated respective of downsampling percentage and processed through peak calling, dimensionality reduction and projection as described above with the following exception. Topic model generation was limited to ten and 20–30 topics. Numbers of peak callable per downsampled dataset were plotted via geom_bar. The final projections were plotted via geom_point with the color of the cell type assignment in the full dataset (Supplementary Tables 6 and 7 and Supplementary Information).

**Cross-platform integration.** Data used in library complexity and peak overlap comparisons were integrated using the Signac package as follows\(^1\). Counts matrices were generated using the platform-defined peak regions and formatted as Seurat objects. For each integration, our s3-ATAC mouse data, we generated a new counts matrix was generated from the platform-defined peaks. Following this, counts matrices were merged and latent semantic indexing (LSI) was used for reduction. Harmony was used to integrate, and a UMAP projection was generated using dimensions 2–30 for sci-ATAC, scATAC and snATAC datasets and 2–40 for dscATAC\(^1\). Integrated plots were generated using our defined cell types (Fig. 2i).

**Subclustering.** After gross cell type assignemnt of mouse and human cell lines, human inhibitory neurons (GAD1\(^\text{total read depth}\) and the function normalize_codex2_ns_noK, such that normalization steps. To this end, we used GM12878 lymphoblastoid cell line as our normal diploid samples and used an priori estimate of 2.6N based on averaged PDCL karyotyping results (Fig. 4c). We then used the SCOPe function normalize_scope_foreach with the following options: K = 5, T = 1.6 to normalize read number per cell. The following were included as genomic regions of interest: chromosome and inferred copy number per breakpoint by cell segment_\text{CBS} allowing for a simple nested structure of copy number changes \((\text{max.m.n} = 1)\). To plot inferred copy number per cell, we used the R library ComplexHeatmap (v.2.5.3) by function Heatmap\(^1\). Pairwise distance between cells was generated by Jaccard distance through the R library phyloteny (v.0.4.0) on dimension categorized as neutral (2N), amplified (\(>2N\)) or deleted (\(<2N\)). Cells then underwent hierarchical clustering by the ward.2 dist argument in the function hclust. The resultant dendrogram was then cut into both three and six clades based on the two independent \(k\) value searches using the find_k function in the R library dendextend (v.1.14.0) given a range of two to five and five to ten clusters, respectively (Fig. 3b). Categorical variables were then normalized into pseudobulk clades for higher resolution copy number calling. After combining counts data across 50-kb bins (and filtered as described above), we had six clades with 154, 250, 360, 100, 268 and 133 cells, with mean reads per bin of 1,207, 2,442, 4,662, 2,071, 2,700 and 9,416, respectively. These pseudobulk sampled were then normalized as described above with clade 6, containing 83.45% GM12878 cells (111/133 cells). The genome per sample was then segmented as described above and normalized reads per bin as well as segmentation calls were plotted with ggplot geom_point and geom_rect functions (Supplementary Fig. 2a). Select genomic locations of recurrently mutated genes were visualized and plotted using IGV with five bins (250kb) up- and downstream from the TSSs (Supplementary Fig. 2b).

**Generation of GCC contact profiles.** s3-GCC contact profile raw counts were generated for cellIDs passing the read count and SCOPe filters (215 cells) as follows. For initial plotting of single-cell profiles, paired-end read bam files were filtered for an insert length of 250 kb via pysam\(^1\) and output as upper-triangle triple-sparse format at 1-Mb bin sizes. Raw contact matrices were then plotted with R and ComplexHeatmap (Fig. 5c, left). Merged ensemble plots were generated by summing single-cell contact matrices generated as described above for 500-kb bins. Followin this, we performed dimensionality reduction and clustering analyses using a topic-modeling approach. We treated the GCC portion of single-cell sequencing fragments (read pairs separated by a genomic distance higher than 20kb) as traditional distal interactions. We analyzed these cells using our previously established topic model for analysis and characterization of single-cell HiC data\(^1\). In the topic-modeling framework, each cell is treated as a mixture of topics where each topic corresponds to a set of distal interactions. The model is trained in an unsupervised manner to find the optimum number of topics that best describes the data and associates each distal interaction with a probabilistic mixture of topics.

We trained a topic model using the GCC data with the default parameters in Kim et al. However, we altered one parameter, which is the range of distal interactions that are input into the model. Due to high coverage of s3-GCC assays, we opted for distal interactions that are separated by a genomic distance of 20Mb or less, as opposed to original parameter where we used interactions that are separated by distances lower than 10Mb. After training, we found that the number of topics of topics that best describe the data is 15. We visualized cells using UMAP and formed the closest clusters from two to five clusters. The results validate the HiC-like characteristics of GCC data and further show that we can capture the subtle differences in chromatin organization of the two lines.

**Compartment calling.** We called compartments from pseudobulk HiC contact matrices and calculated contact probabilities as described in the original HiC paper by Lieberman-Aiden et al.\(^1\). We briefly describe these methods here. To obtain compartment calls, we first normalized using iterative correction and eigenvector decomposition (ICE) and removed the distance effect in HiC matrices for each chromosome at 50-kb resolution. For each normalized HiC contact matrix, we calculated the Spearman Correlation Coefficient matrix from the normalized HiC matrix, which yields a correlation matrix that we can represent as a two-dimensional array. The correlation matrix is reduced to one dimensional representation by performing eigendecomposition; the first eigenvector typically yields the compartment calls, which closely tracks the two-dimensional array pattern in one dimension. For comparison we used compartment calls from Rao et al.\(^1\) on the GM12878 cell line (Fig. 5c, right).

**Calculation of contact probabilities.** To calculate intrachromosomal contact probabilities for each genomic distance, we first calculated the mean HiC signal at a given genomic distance for a given resolution. For a HiC matrix binned at 500-kb resolution, the mean HiC signal for distances less than 500k is the mean of the diagonal, the mean HiC signal for distances between 500 and 1,000 kb is the mean of the outermost band along the diagonal, and so forth. After calculating the mean HiC signal vector for the every intrachromosomal matrix, we obtain a mean HiC signal vector at 500-kb resolution that contains 500 elements, ranging from 0 to around 250 Mb, since the largest chromosome (Chr1) is approximately 250 Mb long. To convert this vector into probabilities, we simply divide the vector by the sum. When plotting contact probabilities, we typically omit the visualizing the contact probabilities.
for distances larger than 200 Mb, as the mean HiC signal at such long distances is both sparse and noisy. External bulk HiC datasets have been downloaded from the ENCODE consortium’s data portal, https://www.encodeproject.org/ via accession codes ENCSR194SRI, ENCSR346DCU, ENCSR444WCZ and ENCSR079VIJ (Supplementary Fig. 3a).

**Data availability**

The data discussed in this publication have been deposited in the National Center for Biotechnology Information’s (NCBI’s) Gene Expression Omnibus ( GEO) and are accessible through GEO Series accession number GSE174226. External single-cell ATAC datasets were downloaded from GEO sample accession number GSM2686124 for snATAC, and external sites for dscATAC [https://github.com/buenroostrolab/dscATAC_analysis_code/blob/master/mousebrain/data/mousebrain-master_dataframe.rds] and 10X Genomics scATAC [https://cf.10xgenomics.com/samples/cell-atac/1.1.0/atac_v1_adult_brain_fresh_3k]. The external single-cell WGS dataset was downloaded from NCBI BioProject PRJNA326989 (https://www.ncbi.nlm.nih.gov/sra/PRJNA326989). Single-cell HiC datasets were downloaded from the 4D Nucleosome project (https://data.fnucleosome.org/publications/048d4558-2ace-41d2-ac6c-f12ac38067c4f/#expets-table). External bulk HiC datasets have been downloaded from the ENCODE consortium’s data portal, https://www.encodeproject.org/ via accession codes ENCSR194SRI, ENCSR346DCU, ENCSR444WCZ and ENCSR079VIJ. Source data are provided with this paper.

**Code availability**

Code and custom scripts used in this study are available at https://github.com/adeylab/scitools and https://mulquer.github.io/.

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**Author contributions**

R.M.M., D.P., F.J.S. and A.C.A. conceived the study. R.M.M. performed all 3s experiments and led all analysis under the supervision of A.C.A. D.P. and F.Z. performed additional experiments under the supervision of F.J.S. B.L.O. and F.J.S. provided support for R.M.M. and advice on analysis. C.A.T. contributed to the analysis of cell types in the s3-ATAC datasets. I.L. generated PDCL cell lines and performed characterization of the lines under supervision of R.C.S. I.L. and R.C.S. contributed to the analysis of PDAC s3-WGS and s3-GCC datasets. The paper was written by R.M.M. and A.C.A. All authors reviewed and contributed to the paper.

**Competing interests**

D.P., F.Z. and F.J.S. are employees of Scale Bio. R.M.M., D.P., F.Z., F.J.S. and A.C.A. are authors on licensed patents that cover components of the technologies described in this paper. This potential conflict of interest for A.C.A. and R.M.M. has been reviewed and managed by OHSU.

**Additional information**

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  - CFX Manager v3.1 for real-time PCR, and NextSeq500 NCS v4.0 or NovaSeq 6000 for sequencing.

- **Data analysis**
  - Bcl2fastq (Illumina Inc, v2.19.0) was used to generate fastq files from bcl files. Bwa mem (v0.7.15-r1140) for sequencing read alignment.
  - Macs2 (v2.1.1.1006309) for chromatin accessible peak calling. Scitoools suite of single-cell analysis tools for data pre-processing (github.com/adaylab/scitools). BEDOPS (v2.4.36) was used for transcription site enrichment calculations. R version 4.0.0 was used for analysis. The following packages were used for analyses cisTopic(v0.3.0), EnsentDb.Musculus (v79.2.99.0), EnsentDb.1hapiens (v86.2.99.0), monacle3 (v0.2.3.0), ggplot2 (v3.3.2), Seurat (v3.2.1) and Signac (v1.1.0).

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | No statistical analyses were performed to pre-determine sample size. The number of single-cell libraries to be generates was selected for comparison with other single-cell chromatin accessibility data sets. These external data sets were also used to infer expected effect sizes, technical limitations and budget considerations. Single-cell whole genome and GEC library sizes were chosen to demonstrate the technology. |
| Data exclusions | Reads were excluded if indexes did not match predetermined barcode sequences. Barcodes (used to identify cellIDs) were filtered as described in the manuscript and required reads to have a Q-score of >= 10. |
| Replication | s3-ATAC was used to generate a total of 3 independent experiment replicates. All attempts at replication were successful. Single-cell libraries were treated as independent measurements of the biological systems used. |
| Randomization | Organism allotment was not random, as data generated was used to demonstrate primarily quality metrics. Cell libraries were inherently randomized during the s3 protocol after a mixing and diluting step. |
| Blinding | Cell libraries were inherently blinded during the PCR phase of the s3 protocol after a mixing and diluting step. |

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

| Materials & experimental systems | Methods |
|---------------------------------|---------|
| n/a                             | n/a     |
| ☒ Antibodies                   | ☒ ChiP-seq |
| ☒ Eukaryotic cell lines        | ☒ Flow cytometry |
| ☒ Palaeontology and archaeology| ☒ MRI-based neuroimaging |
| ☒ Animals and other organisms  |         |
| ☒ Human research participants  |         |
| ☒ Clinical data                |         |
| ☒ Dual use research of concern |         |

Eukaryotic cell lines

Policy information about cell lines

| Cell line source(s) | GM12878, Coriell; Patient-derived cell lines (PDCls) |
|---------------------|------------------------------------------------------|

Authentication

From a fraction of these PDCls cells, DNA was extracted to validate the presence of KRAS-G12 mutations by ddPCR (Bio-Rad, 1863506) and to validate an STR profile that matches normal leukocyte DNA from the same patient (Genetica). PDCls exhibited morphologies consistent with epithelial tumor cells and abundant KRT expression was detected by immunofluorescence using the monoclonal antibodies: AE1/AE3, C-11, and Cam5.2. To ensure that only tumor cells were cultured, G-banded karyotyping was performed by the Knight Diagnostic Research Cytogenetics Lab at OHSU.

Mycoplasma contamination

Lines were not tested for mycoplasma contamination.

Commonly misidentified lines (See iGCLC register)

No commonly misidentified lines were used in this experiment.
### Animals and other organisms

Policy information about *studies involving animals; ARRIVE guidelines* recommended for reporting animal research

| Laboratory animals | The mouse used for whole brain experiments were C57Bl/6J, aged within 8-12 weeks. All mouse cages were kept on a 12 h light/dark cycle at a temperature of 70°F and within a humidity range of 30-70%. |
|---------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Wild animals        | No wild animals were used in this study.                                                                                                                                                           |
| Field-collected samples | No field-collected samples were used in this study.                                                                                                                                           |
| Ethics oversight    | All work overseen and approved by OHSU Institutional Animal Care and Use Committee.                                                                                                             |

Note that full information on the approval of the study protocol must also be provided in the manuscript.