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Accessibility
A single-cell and single-nucleus RNA-Seq toolbox for fresh and frozen human tumors

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Single-cell genomics is essential to chart tumor ecosystems. Although single-cell RNA-Seq (scRNA-Seq) profiles RNA from cells dissociated from fresh tumors, single-nucleus RNA-Seq (snRNA-Seq) is needed to profile frozen or hard-to-dissociate tumors. Each requires customization to different tissue and tumor types, posing a barrier to adoption. Here, we have developed a systematic toolbox for profiling fresh and frozen clinical tumor samples using scRNA-Seq and snRNA-Seq, respectively. We analyzed 216,490 cells and nuclei from 40 samples across 23 specimens spanning eight tumor types of varying tissue and sample characteristics. We evaluated protocols by cell and nucleus quality, recovery rate and cellular composition. scRNA-Seq and snRNA-Seq from matched samples recovered the same cell types, but at different proportions. Our work provides guidance for studies in a broad range of tumors, including criteria for testing and selecting methods from the toolbox for other tumors, thus paving the way for charting tumor atlases.
between tissue acquisition and processing teams, posing a challenge in clinical settings. Conversely, single-nucleus RNA-Seq (snRNA-Seq) allows profiling of single nuclei isolated from frozen tissues, decoupling tissue acquisition from immediate sample processing. snRNA-Seq can also handle samples that cannot be successfully dissociated even when fresh, due to size or cell fragility, as well as multiplexed analysis of longitudinal samples from the same individual. However, nuclei have lower amounts of mRNA compared to cells and are more challenging to enrich or deplete for specific cell types of interest. Both scRNA-Seq and snRNA-Seq pose experimental challenges when applied to different tumor types, due to the distinct cellular composition and extracellular matrix (ECM) in different tumors, and thus each assay requires dedicated customizations.

To address these challenges, we developed a systematic toolbox for fresh and frozen tumor processing using scRNA-Seq and snRNA-Seq, respectively (Fig. 1a). The toolbox contains our experimental workflow and methods, computational pipelines and evaluation metrics. To generalize across tumor and sample types, we tested eight tumor types with different tissue characteristics (Fig. 1b), including comparisons of matched fresh and frozen preparations from the same tumor specimen. Our work provides direct recommended protocols for multiple tumor types, decision trees that allow researchers to choose the most suitable protocol for their research goals, and guidelines on how to customize protocols for new tumor and specimen types.

Results
A systematic study to develop an sc/snRNA-Seq toolbox. To develop a toolbox of customized protocols for sc/snRNA-Seq of tumors, we studied eight tumor types with different tissue characteristics (Fig. 1). The tumor types span the following characteristics: different cells of origin (for example, epithelial, neuronal), solid and non-solid, patient ages and transitions (for example, primary, metastatic). We tested varying tissue and sample characteristics including resection, biopsy, ascites and orthotopic patient-derived xenograft (O-PDX). We included samples from non-small cell lung carcinoma (NSCLC), metastatic breast cancer (MBC), ovarian cancer, neuroblastoma, glioblastoma (GBM), pediatric high-grade glioma, chronic lymphocytic leukemia (CLL), pediatric sarcoma and melanoma (Fig. 1b). In total, we analyzed 216,490 cells and nuclei across 23 tumors, from 22 patients spanning 40 sample preparations. We provide a comprehensive analysis summary for each sample tested in a dedicated website (https://tumor-toolbox.broadinstitute.org).

Experimental and computational QCs assess quality and composition. We evaluated and compared protocols based on (1) cell/nucleus quality; (2) number of recovered versus expected cells/nuclei; (3) cellular composition (Fig. 1a). For ‘cell/nucleus quality’, we considered both experimental and computational metrics. Experimentally, we measured cell viability (for scRNA-Seq), the extent of doublets or aggregates in the cell/nucleus suspension and cDNA quality recovered after whole transcriptome amplification. Computationally, we evaluated the percent of reads mapping to the transcriptome, genome and intergenic regions, the number of cells/nuclei exceeding a minimal number of genes and unique transcripts (reflected by unique molecular identifiers, UMIs), the number of reads, transcripts (UMIs) and genes detected per cell/nucleus and the percent of UMIs from mitochondrial genes (see Methods). To compare protocols, when there was a notable difference in sequencing saturation or total reads across samples, we also downsampled reads to equal numbers across samples and then re-estimated and compared their QC metrics. For ‘number of recovered versus expected cells/nuclei’, we evaluated the proportion of droplets scored as likely empty (that is, containing only ambient RNA rather than the RNA from an encapsulated cell), and the proportion of doublets (see Methods). Because the algorithm for scoring empty droplets was developed for cells, we did not use it to evaluate snRNA-Seq. Finally, for ‘cellular composition’, we considered the diversity of cell types captured, the proportion of cells/nuclei recovered from each subset and the copy number aberration (CNA) pattern classes that are recovered in malignant tumors (see Methods). We considered it a virtue if a protocol recovers a larger diversity of cell types, because this facilitates comprehensive studies. However, capture of diverse cell types may not always be a researcher’s desired goal, nor would it always be the most accurate representation of a tumor’s composition (see Discussion). We annotated the malignant cells based on the presence of CNAs (when detectable) and the cell type signature they most closely resembled (see Methods). We conducted initial data analysis using Cumulus, a cloud-based data analysis framework (see Methods and Fig. 1a), and developed a dedicated pipeline for additional quality control, tumor sample characterization and protocol comparison.

Customization of workflows and dissociation protocols for scRNA-Seq of fresh tumors. We customized successful protocols for specimen acquisition and dissociation for scRNA-Seq across five types of fresh tumor (NSCLC, ovarian cancer, MBC, neuroblastoma, GBM and cryopreserved CLL (Fig. 1b)). We constructed workflows that minimize the time interval between removal of the sample from the patient in a clinical setting and its dissociation into cells, to maximize cell viability and preservation of RNA profiles. We determined dissociation conditions for each tumor type and constructed specific steps as a decision tree to adjust for differences between clinical samples (for example, size, presence of red blood cells (RBCs); Fig. 2a and Methods). To choose the best performing dissociation method, when possible, we apportioned large tumor specimens into smaller pieces (~0.5–2 cm), dissociating each piece following a different protocol. When specimen size was limiting (for example, biopsies), optimization spanned multiple samples. We subjected the samples that yielded highly viable single cell suspensions to droplet-based scRNA-Seq (see Methods), to allow sampling of larger number of cells for calculation of QC metrics. We tested protocols several times to confirm similar performance trends.

For dissociation, we selected enzymatic mixtures for processing fresh tissues based on the specific characteristics of each tumor type, such as cell type composition and ECM components, literature review and experience from processing similar human or mouse tissues. For example, to break down collagen fibers in breast cancer (see Methods), we used Liberase TM (see Methods), whereas to break down ECM in GBM, we used papain (cysteine protease). We also included DNase I to digest DNA released from dead cells to decrease viscosity in all dissociation mixtures. In the following, we recommend those methods that broke down the ECM and cell-to-cell adhesions sufficiently, while minimizing processing time, maintaining high viability and supporting cell type diversity in the sample.

Cell-type-specific and cell-composition QCs are important in protocol evaluation. As an example of the optimization process, consider the in-depth analysis of an NSCLC resection sample (sample NSCLC14, Fig. 2b–f and Extended Data Figs. 1–3). We used three processing protocols: (1) collagenase 4 (NSCLC-C4), (2) a mixture of pronase, dispase, elastase and collagenases A and 4 (PDEC) and (3) Liberase TM and elastase (LE), each in combination with DNase I (see Methods). For other tumor types, we show the results of selected protocols out of those tested (Figs. 1b, 2g and 3a–c and Extended Data Fig. 4a,b).

Protocols often performed similarly on standard QC measures (for example, number of cells recovered), but differed markedly in recovered cellular diversity or in the fraction of droplets predicted to contain only ambient RNA (‘empty drops’)—two evaluation criteria that we prioritized. For example, in the NSCLC14 resection sample,
all methods yielded a similar number of cells with high-quality expression profiles (Fig. 2b,e and Extended Data Fig. 1a–c), doublets (Fig. 2c) and CNA patterns in malignant cells (Fig. 2f and Extended Data Fig. 1d). However, only the PDEC and LE protocols recovered fibroblasts and endothelial cells (Fig. 2e and Extended Data Fig. 1c), and NSCLC-C4 had a 100-fold higher fraction of droplets called as ‘empty’ (7% versus 0.08% and 0.04% in PDEC and LE, respectively; Fig. 2d and Extended Data Fig. 1a). The drops
designated ‘empty’ in NSCLC-C4 clustered within macrophages (Fig. 2d and Extended Data Fig. 1c), the most prevalent cell type, suggesting that these cell barcodes either had lower sequencing saturation or that the sample itself had higher ambient RNA content. Although we estimated similarly low levels of ambient RNA across the three protocols (Extended Data Fig. 1e), NSCLC-C4 indeed had lower overall sequencing saturation (Extended Data Fig. 1a).

Comparing QC metrics across protocols can be challenging due to differences in cell type recovery and in sequencing depth between preparations, which we controlled for in the NSCLC14 sample by also evaluating QC metrics within each cell type and downsampling by total reads across protocols (Fig. 2b and Extended Data Fig. 2). For example, when we consider all cells in the NSCLC14 samples, NSCLC-C4 had a significantly higher number of detected genes ($P=1.3 \times 10^{-36}$ versus PDEC; $1.4 \times 10^{-42}$ versus LE, two-sided Mann–Whitney U test), but within B cells, PDEC had a significantly higher number of detected genes ($P=2 \times 10^{-15}$ versus NSCLC-C4; $2 \times 10^{-10}$ versus LE), whereas within epithelial cells, LE had the highest number ($P=5 \times 10^{-8}$ versus NSCLC-C4; $2 \times 10^{-7}$ versus PDEC) (Fig. 2b). Because the number of detected genes (and other metrics) varies between cell types, and cell type composition varies between the protocols (Fig. 2e), it is important to assess cell-type-specific QCs when selecting a protocol. Moreover, the lower sequencing saturation does not directly reflect the performance of the NSCLC-C4 protocol, and downsampling by total reads did not qualitatively change any of our protocol evaluation metrics (Extended Data Fig. 3). Considering all of these features, we selected the PDEC protocol for processing NSCLC tumor samples, as it balances cell type diversity and QCs per cell type.

**Fast depletion of immune cells for enrichment of malignant and stromal cells.** Because in some tumor specimens the proportion of malignant cells is relatively low and that of immune cells is particularly high, we considered strategies to deplete CD45+ immune cells as a way to both enrich for epithelial cells without specific markers and to maintain any stromal cells. We chose to use MACS MicroBeads with anti-CD45 antibodies rather than sorting by flow cytometry (FACS), because samples are not always available at the designated time for which sorters are booked, and FACS requires MicroBeads with anti-CD45 antibodies rather than sorting by flow cells as a way to both enrich for epithelial cells without specific depletion in NSCLCs (top) and ovarian ascites (bottom) data not shown). For example, we profiled an NSCLC tumor sample (NSCLC17) by scRNA-Seq before and after depletion, finding an increase from 26% to 82% epithelial cells (Fig. 2g and Extended Data Fig. 4a) and virtually no immune cells profiled post depletion. Similarly, in an ovarian ascites sample (HTAPP-727), we recovered 32% epithelial cells by scRNA-Seq post depletion (Fig. 2g and Extended Data Fig. 4b), compared to <1% CD45+ EpCAM+ cells typically found in ovarian ascites by FACS (data not shown). Consistently, FACS shows that CD45+ cell depletion of another ascites sample increased the overall proportion of non-immune (CD45−) cells from 0.75% to 29.4% and increased the proportion of EpCAM+ cells from 0.17% to 4.9% (Extended Data Fig. 4c).

**Successful scRNA-Seq of biopsies and post-treatment samples from diverse tumors.** We successfully applied the scRNA-Seq tool-box to much smaller core biopsy clinical samples from different anatomical sites. For example, in MBC, we applied the LD (Liberase TM and DNase I) protocol to a resection (HTAPP-254) and a biopsy (HTAPP-735) from lymph node metastases from two patients, yielding similarly successful QCs (Fig. 3a–d). The resection and biopsy of the two patients had different cellular compositions (Fig. 3e): the biopsy had a higher proportion of epithelial, endothelial and fibroblast cells and a lower proportion of T cells compared to the resection. We similarly successfully profiled biopsies of MBC liver metastases (HTAPP-285 and HTAPP-963) with the same protocol (Fig. 3a–d), recovering some hepatocytes in addition to a similar range of cell types as was recovered in the lymph node biopsy (Fig. 3e). Thus, this protocol can be used across breast cancer metastases from different anatomical metastatic sites.

The scRNA-Seq toolbox also performs well on samples obtained post-treatment, which can pose challenges as a result of cell death and changes in cell type composition with treatment. For example, both a pre-treatment diagnostic biopsy (HTAPP-312-pre) and a post-treatment resection (HTAPP-312-post) from the same neuroblastoma patient profiled with the NB-C4 protocol yielded high QCs (Fig. 3a–d) and similar CNA patterns in malignant cells (Fig. 3f). More cells, but of fewer cell types, were recovered in the pre-treatment biopsy (4,369 cells: neuroendocrine, T cells and macrophages) than the post-treatment resection (786 cells: neuroendocrine, T cells, macrophages, as well as endothelial cells and fibroblasts) (Fig. 3e), consistent with observed post-treatment fibrosis. We tested an additional dissociation protocol (papain) in a neuroblastoma O-PDX sample (O-PDX1) in a neuroblastoma O-PDX sample (O-PDX1) 1.9×10^6, which is not expected to include non-malignant human cells and indeed resulted in high-quality malignant cell profiles (Fig. 3a–e). The experimental QCs for the papain protocol were superior to those we had observed in other samples with NB-C4, a trend we corroborated in additional neuroblastoma samples that were not shown. Thus, we ultimately selected the papain protocol for neuroblastoma tumors.

**Fig. 2 | Fresh tumor processing and protocol selection for scRNA-Seq.** a, Flow chart recommended for collection and processing of fresh tumor samples. b–f, Comparison of three dissociation protocols applied to one NSCLC sample. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. Bottom: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset. b, Protocol performance varies across cell types. Top and middle: distribution (median and first and third quartiles) of the number of reads per cell, the number of UMIs per cell, the number of genes per cell and fraction of UMIs per cell mapping to mitochondrial genes (Fr. mito. genes) (y axes) in each protocol (x axes) across the entire dataset.

c, The protocols detect similar numbers of doublets. Uniform manifold approximation and projection (UMAP) embedding of single cell profiles (dots) for each protocol, colored by assignment as single cell (gray) or doublet (red). Horizontal bars: bottom: fraction of single (gray) and doublet (red) cells. d, The protocols vary in the number of empty drops. UMAP embedding of single cell profiles (dots) for each protocol, colored by assignment as cell (gray) or empty drop (red). Horizontal bars: bottom: fraction of assigned cells (gray) and empty drops (red). e, The protocols vary in the diversity of cell types captured. UMAP embedding of single cell profiles (dots) from all three protocols, colored by assigned cell subset signature (left) or by protocol (right). Bottom: proportion of cells in each subset in each of the three protocols; k, number of cells passing QC. f, Inferred CNA profiles. Chromosomal amplification (red) and deletion (blue) are inferred in each chromosomal position (columns) across the single cells (rows) using the PDEC protocol. Top: reference cells not expected to contain CNAs in this tumor. Bottom: cells tested for CNAs relative to the reference cells. Color bar: assigned cell type signature for each cell. g, Successful depletion of CD45+ cells. The proportion of cells in each subset without and with CD45+ depletion in NSCLCs (top) and ovarian ascites (bottom) samples is shown; k, number of cells passing QC. n = 1 sample per protocol. The numbers of cells (k) are indicated in e and g. Numbers of epithelial cells from NSCLC-C4, PDEC and LE are k = 1,284, 641 and 260, respectively, and the number of B cells is k = 100, 121 and 78, respectively. ACK, ammonium-chloride-potassium.
In addition to such NSCLC, ovarian cancer ascite, MBC and neuroblastoma samples, we established effective scRNA-Seq protocols for GBM (GBM125), CLL (CLL1) and ovarian cancer tumors (HTAPP-624) (Fig. 3a–e). In particular, in CLL, we successfully recovered the expected cell types from a cryopreserved sample, containing viable cells. This reflects the increased resilience
of immune cells to freezing compared to other cell types, also observed in other settings, and the lack of a dissociation step in CLL scRNA-Seq (see Methods). Cryopreservation, however, can increase the proportion of damaged cells and may not successfully recover all the malignant and other non-malignant cells in the tumor.
Fig. 4 | Frozen tumor processing and protocol selection for snRNA-Seq. a, Flow chart for collection and processing of frozen tumor samples. b–e, Comparison of four nucleus isolation protocols in one neuroblastoma sample. b, Variation in protocol performance: distributions (median and first and third quartiles) of the number of UMIs per nucleus, the number of genes per nucleus and fraction of UMIs per nucleus mapping to mitochondrial genes (y axes) in each protocol (x axis) across all nuclei in the dataset. c, The protocols detect similar numbers of doublets. UMAP embedding of single nucleus profiles (dots) for each protocol is colored by assignment as nucleus (gray) or doublet (red). Horizontal bars (bottom): fraction of single (gray) and doublet (red) nuclei. d, The protocols vary in the diversity of cell types captured. UMAP embedding of single nucleus profiles (dots) from all four protocols is colored by assigned cell subset signature (left) or protocol (right). Bottom: proportion of cells from each subset in each of the protocols. k, number of nuclei passing QC. e, Inferred CNA profiles. Chromosomal amplification (red) and deletion (blue) inferred in each chromosomal position (columns) across the single nuclei (rows) from the TST protocol are shown. Top: reference nuclei not expected to contain CNAs in this tumor. Bottom: nuclei tested for CNAs relative to the reference nuclei. Color bar: assigned cell type signature for each nucleus. n=1 sample per protocol. The numbers of nuclei (k) are indicated in d. OCT, optimal cutting temperature compound.
Fig. 5 | snRNA-Seq protocol comparison across tumor types. a–c. QC metrics: distributions (median and first and third quartiles) of the number of UMIs per nucleus (a), the number of genes per nucleus (b) and the fraction of UMIs per nucleus mapping to mitochondrial genes (c) (y axes) for each sample (x axis). d. Cell type composition, showing the proportion of nuclei assigned to each cell type signature (color) for each sample. n = 1 sample per protocol. The numbers of nuclei (k) are indicated in d.

Four nucleus isolation protocols assessed for snRNA-Seq of frozen tumors. For frozen specimens from solid tumors, we optimized snRNA-Seq, assessing different methods for nucleus isolation (Fig. 4a and Methods) across seven tumor types: neuroblastoma, MBC, ovarian cancer, pediatric sarcoma, melanoma, pediatric high-grade glioma and CLL (Fig. 1b). We initially approximated larger samples or used multiple biopsies to compare four isolation methods: EZPrep, Nonidet P40 with salts and Tris (CST)11, and Tween with salts and Tris (TST)11. The methods differ primarily in the mechanical ples or used multiple biopsies to compare four isolation methods: EZPrep8, Nonidet P40 with salts and Tris (CST)11, and Tween with salts and Tris (TST)11 (modified from ref. 23), CHAPS, with salts and Tris (CST)11, and Tween with salts and Tris (TST)11. The methods differ primarily in the mechanical

TST protocol typically recovers the highest diversity of cell types. Overall, three nucleus isolation methods—TST, CST and NST—had comparable performances based on the assessed nucleus quality (Figs. 4b–d and 5 and Extended Data Figs. 5 and 6), with TST typically yielding the greatest cell type diversity and number of nuclei per cell type, together with the highest expression of mitochondrial genes, and NST typically having the fewest genes per nucleus and lowest diversity of cell types. For example, in neuroblastoma, testing each of the four protocols on a single resection sample (HTAPP-244) yielded a similar number of high-quality nuclei (7,896, 6,157, 7,531 and 7,415 for EZ, CST, NST and TST, respectively) (Fig. 4b–d and Extended Data Fig. 5c) and malignant cells with similarly detectable CNAs (Fig. 4e and Extended Data Fig. 5d; CNAs are less thresholds for transcript (UMI) and gene counts to reflect the lower expected mRNA content in nuclei (see Methods). Experimentally, we added in-process light microscopy QCs to ensure complete nucleus isolation and to estimate doublets, aggregates and debris (Fig. 4a and Methods). As with scRNA-Seq, we tested protocols several times to confirm similar performance trends.
prominent, as expected for this pediatric, low-risk sample). Nuclei prepared with the EZ protocol had lower numbers of UMIs and genes detected (Fig. 4b) compared to the three ST protocols. TST recovered more endothelial cells, fibroblasts, neural crest cells and T cells than the other protocols (Fig. 4d). TST also yielded a higher expression of mitochondrial genes (Fig. 4b), in this and all other tumors tested (Fig. 5c), because the nuclear membrane, endoplasmic reticulum and ribosomes remain attached to the nucleus when using this method. The same trends were preserved in cell-type-specific QCs (Extended Data Fig. 6) and after downsampling by the total number of sequencing reads (Extended Data Fig. 7).

The CST, NST and TST nucleus isolation methods had similar performance characteristics when tested with MBC, ovarian cancer and pediatric sarcoma samples, with TST providing the most diversity in cell types, especially in non-malignant cells. In MBC, we compared CST and NST in one metastatic brain resection (HTAPP-394) and CST and TST in another metastatic brain resection (HTAPP-589) and in a metastatic liver biopsy (HTAPP-963) (Fig. 5). In all cases, QC statistics (Fig. 5a–c) and CNA patterns (Extended Data Fig. 8a–f) were similar between protocols, and nuclei from epithelial cells were the most prevalent (Fig. 5d). CST and NST captured a very similar distribution of cell types, while TST captured more non-malignant cells, including T cells (Fig. 5d), and a higher fraction of mitochondrial reads (Fig. 5c). In ovarian cancer, CST, NST and TST recovered similar CNA patterns from the same sample (HTAPP-316; Extended Data Fig. 8g–i), but TST captured the greatest cell type diversity (Fig. 5d), whereas NST recovered fewer nuclei, genes per nucleus and UMIs per nucleus (Fig. 5a–c), and had a lower cell type diversity (Fig. 5d), despite having greater overall sequencing depth (73% sequencing saturation versus 57% in CST and 50% in TST). In a rhabdomyosarcoma sample (HTAPP-951), CST and TST captured the same cell types at similar proportions (Fig. 5d) and showed similar CNA patterns (Extended Data Fig. 8j,k).

Overall, we chose the TST protocol for most tumor types and CST for tumors from neuronal tissues, such as pediatric high-grade glioma. With the protocols we selected (Fig. 1b, right column), we profiled additional neuroblastoma tumors (HTAPP-656, O-PDX1) as well as Ewing sarcoma (HTAPP-975), melanoma (MEL112, MEL128), pediatric high-grade glioma (HTAPP-443) and CLL (CLL1) tumor samples—spanning biopsies, resections and treated samples (Figs. 1b and 5). We also tested a pediatric rhabdomyosarcoma sample (HTAPP-951) by two different chemistries for droplet-based snRNA-Seq (V2 versus V3 from 10x Genomics; see Methods), obtaining, overall, similar results in terms of cell types detected, but an improved number of recovered versus expected nuclei and higher complexity per nucleus in V3 (Extended Data Fig. 9).

Different cell composition recovered by scRNA-Seq and snRNA-Seq. We compared scRNA-Seq and snRNA-Seq by testing matching samples from the same specimen each in neuroblastoma (HTAPP-656, Fig. 6a–g), MBC (HTAPP-963, Fig. 6h–n), CLL (CLL1, Extended Data Fig. 10a–g) and O-PDX (O-PDX1, Extended Data Fig. 10h–n). The two approaches typically recovered similar cell types, but sometimes at varying proportions. In both neuroblastoma and MBC, immune cells were much more prevalent in scRNA-Seq, and parenchymal (especially malignant) cells were much more prevalent in snRNA-Seq (Fig. 6a–d,h–k). In all tested tumor types, cells and nuclei readily aligned following batch correction by canonical correlation analysis (CCA, see Methods), grouping by cell type (Fig. 6c,g,l–n and Extended Data Fig. 10e–g,l–n).

Finally, we leveraged the matched data to examine the extent to which expression patterns in cells from dissociated fresh tissue indicate specific stress and compared these to those in nuclei from snap-frozen tissue. To this end, we scored each cell or nucleus in the matched data from recently published ‘dissociation signatures’

![Fig. 6](image_url)
Discussion

Single-cell genomics of clinical tumor specimens obtained as part of routine disease management or through research biopsies should help guide new discoveries and better deployment of therapies, as initial studies have shown in the context of gliomas, melanoma, and other malignancies. However, this requires adaptation of laboratory protocols, initially developed in research settings and requiring very rapid handling of fresh tissue, into the context of clinical sample acquisition and processing. Indeed, these initial studies were applied to small numbers of samples, often from resections, or focused on readily isolated immune cells. Analyzing the larger numbers of samples that would be required in the context of a clinical trial or longitudinal research, involving multiple sites, calls for streamlined and robust protocols. Moreover, systematically characterizing the diverse cells in solid tumors requires robust recovery of cells, many of which, including malignant epithelial cells, are highly sensitive. These challenges are further compounded by the diversity of tissues in which tumors and metastases reside.

Here, we take on these challenges by developing a systematic toolbox for protocols for single-cell and single-nucleus RNA-Seq, with detailed workflows and protocols from sample acquisition to library preparation for processing fresh and frozen clinical tumor samples across eight tumor types, as well as guidelines for testing and selecting protocols for processing future clinical tumor samples. We provide computational pipelines for extensive QCs at https://github.com/klarman-cell-observatory/HTAPP-Pipelines, and all laboratory protocols are provided in detailed form in the open access platform protocols.io.

When selecting a protocol for fresh tissue dissociation, we suggest testing two to three dissociation methods, chosen based on tumor type and tissue composition, and processing according to the fresh sample decision tree (Fig. 2a). We selected the best performing protocol by assessing both experimental and computational QC by Accumax, snRNA-Seq, and CITE-Seq. The best performing protocol for each sample type is presented as a starting point for further optimization. For frozen tissue, researchers should first evaluate tissue dissociation by in-process QCs, especially cell viability and extent of dissociation into single-cell suspension, and only samples that pass those should proceed to scRNA-Seq (for example, in MBC we have previously ruled out the use of Accumax for dissociation (data not shown)). FACS could also be used prior to profiling as an additional QC. It is easier to perform a side-by-side evaluation of nucleus isolation protocols, because they require smaller portions of tissue and can be started at a convenient time. However, in-process experimental QCs for nuclei are less informative and snRNA-Seq is typically required to assess performance.

Because scRNA-Seq and snRNA-Seq vary in their recovered cellular compositions, it is advantageous, when possible, to analyze both fresh and frozen tumor samples. The choice between scRNA-Seq and snRNA-Seq is typically driven by sample availability, logistics and biological questions. scRNA-Seq measures the expression in the whole cell, and the intact cell membrane allows for selection of specific cellular populations and protein profiling by CITE-Seq. In contrast, snRNA-Seq decouples sample procurement from processing, recovers nuclei from hard-to-dissociate samples (for example, bone, adipose, and liver), and allows multiplexing of samples accrued over time, including from banks. This can aid in sample selection and experimental design, reduce batch effects and open the study of rare or unusual samples that may be collected from many sites.

Our toolbox will help researchers systematically profile additional human tumors, leading to a deeper understanding of tumor biology. The toolbox will support charting of high-resolution tumor cell atlases, which will yield insights that inform clinical work and should help improve precision in diagnostics and therapeutics.

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/s41591-020-0844-1.

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Methods

Experimental methods. Human patient samples. External sample cohorts were added to the Broad Institute’s Molecular Classification of Cancer (MCC) protocol (15–370B) and reviewed and approved by the Dana-Farber Cancer Institute’s (DFCI) Institutional Review Board (IRB). No subject recruitment or ascertainment was performed as part of the Broad protocol. Samples added to this protocol also underwent IRB review and approval at the institutions where the samples were originally collected. Specifically, DFCI IRB approved the following protocols: NSCLC (IRB protocol 98-066), MBC (IRB protocol 05-246), neuroblastoma (IRB protocols 11-104, 10-104, 09-160), ovarian cancer (IRB protocol 02-051), melanoma (IRB protocol 11-104), sarcoma (IRB protocol 17-104), GBM (IRB protocol 10-417) and CLL (IRB protocol 99-224), and the St Jude Children’s Research Hospital IRB approved the following protocol: pediatric high-grade glioma (IRB protocol 97BANK).

The XPD 09-234 MAST (Molecular Analysis of Solid Tumor) protocol for creating the neuroblastoma O-PDX sample was reviewed and approved by the St Jude Children’s Research Hospital IRB.

Laboratory animals. For the neuroblastoma O-PDX sample, animal use was restricted to one female nude athymic mouse for para-adenal injection of O-PDX cells. This study was carried out in strict accordance with the recommendations in the Guide to Care and Use of Laboratory Animals of the National Institute of Health. The protocol was approved by the Institutional Animal Care and Use Committee at St Jude Children’s Research Hospital. All efforts were made to minimize suffering. All mice were housed in accordance with approved IACUC protocols. Animals were housed on a 12-12h light cycle (light on 6:00 and off 18:00) and provided food and water ad libitum. All nude female mice were purchased from Charles River Laboratories (strain name SGM3).

Collection of fresh tissue for scRNA-Seq. Collection of fresh solid tumor tissue for NSCLC, ovarian cancer and MBC at Brigham and Women’s Hospital (BWH)/DFCI was performed following protocols established to reduce the time elapsed between removal of the tumor tissue from the body, placement of the specimen in media and processing for scRNA-Seq. To this end, we established procedures between the hospital team (surgeon, clinical research coordinator, clinical pathologist), the coordinating team (project managers/pathology technician) and the processing team (staff scientists/research technicians) before procedure day. This included providing the hospital team with collection containers with appropriate media and predefining allocation priorities to ensure quick handling by the pathology technician of the sample received. On the day of the procedure, timely communication between the teams ensured quick specimen transfer from the hospital team to the research team, timely transport to the Broad Institute for processing, and immediate loading of the single cell suspension into the 10X Genomics Single Cell Chromium Controller (as described in the section ‘Dissection workflow from fresh solid tumor samples’ below). In all cases, the tissue received from the hospital team was examined by the research pathology technician and, following procurement of a specimen for anatomic pathology review, the highest-quality portion (or core) was allocated for scRNA-Seq, placed in liquid medium and transported to the Broad Institute for dissociation following the appropriate protocol (protocols are detailed for each tumor type below). Tissue quality was assessed based on visual examination and rapid pathology interpretation at the time of collection, and determined based on tumor content, necrosis, calcification, fat and hemorrhage.

For ovarian cancer ascites, ~300 ml was usually received from the hospital team within 1 h after being taken out of the body, and contained a vast majority of non-malignant (mainly immune) cells. Hence, all ascites samples were subjected to CD45+ cell depletion (below) to enrich for malignant cells. For CLL, samples were generated from peripheral blood mononuclear cells isolated using density centrifugation (Ficoll-Paque) and stored in freezing medium (FBS + 10% DMSO) in liquid nitrogen until processing.

For O-PDX of neuroblastoma samples Foxn1−/− nude mice (Charles River Laboratories) were orthotopically injected via ultrasound-guided para-adenal injection with cells derived from a patient MYCN-amplified neuroblastoma (available as sample SJB8046-X1 through the Childhood Solid Tumor Network)26. A portion of O-PDX tumor was flash-frozen for future scRNA-Seq, while the remainder was subjected to the procedures below.

Preservation of tissue for scRNA-Seq. For those samples that we prospectively collected for scRNA-Seq (neuroblastoma HTAPP-244-SMP-451 and HTAPP-656-SMP-3481), freezing of tumor samples was performed as quickly as possible after sample collection using a standard biobanking technique. The times when samples were frozen were recorded. (Other samples were obtained from tissue banks with a limited record on how they were frozen, which is a typical scenario.) Samples were placed in cryo-tubes without any liquid. Complete removal of liquid from the sample was accomplished by gently wiping it (not patting, as this would damage the tissue) in a Petri dish, then pelleting the sample in a centrifuge. The tubes were then covered in dry ice and transferred to −80°C for long-term storage.

The other frozen samples from scRNA-Seq were obtained from tissue banks as follows: ovarian optimal cutting temperature compound (OCT)-frozen archival samples were obtained from the DFCI Gynecology Oncology Tissue Bank; sarcoma snap-frozen samples were obtained from the Boston Children’s Hospital Tissue Bank; pediatric snap-frozen glioma samples were obtained from the St Jude Children’s Research Hospital Biorepository; and the Boston Children’s Hospital Precision Link Biobank for Health Discovery. MBC OCT-frozen samples were obtained from the Center for Cancer Precision Medicine Bank; snap-frozen melanoma samples were obtained through the laboratory of Dr. C. Yoon at BWH.

Dissociation workflow from fresh solid tumor samples to a single-cell suspension for scRNA-Seq. MBC, NSCLC (protocols PDEC and LE), ovarian cancer solid tumor and neuroblastoma workflows. Fresh tissue dissociation of MBC, NSCLC (protocols PDEC and LE), ovarian cancer solid tumor and neuroblastoma workflows were performed using a similar workflow (Fig. 2a), with different components of the dissociation mixture for each tumor type, as described in the next section.

Samples were transferred from interventional radiology (biopsies) or the operating room (resections) in DMEM (MBC), RPMI (NSCLC) or RPMI with HEPES (ovarian cancer and neuroblastoma) medium. On arrival at the laboratory, the sample was washed in cold PBS and transferred into either a 2 ml Eppendorf tube containing dissociation mixture (for biopsies) or a 5 ml Eppendorf tube containing 3 ml dissociation mixture (for resections). Next, the sample was minced in the Eppendorf tube using spring scissors (Fine Science Tools, cat. no. 15514-12) into fragments under ~0.4 mm, and incubated at 37 °C, while rotating horizontally at ~14 r.p.m., for 10 min. After 10 min, the sample was pipetted 20 times with a 1 ml pipette tip at room temperature and placed back into incubation with rotation for an additional 10 min. The sample was then pipetted again, 20 times with a 1 ml pipette tip, transferred to a 1.7 ml Eppendorf tube and centrifuged at 300–500 g for 4–7 min at 4 °C. The supernatant was removed and the pellet was resuspended in 200–500 µl of ACK (ammonium–chloride–potassium) RBC lysis buffer (Thermo Fisher Scientific, A1094201). The ACK volume added depended on the size of the pellet; while pellet size is hard to quantify, we suggest adding ~100 µl ACK lysis buffer per 100,000 cells, with a minimum volume of 200 µl. The sample was incubated in ACK RBC lysis buffer for 1 min on ice, followed by the addition of cold PBS at twice the volume of the ACK. The cells were pelleted by a short centrifugation at 8 × 4 °C using the short spin setting with centrifugal force ramping up to, but not exceeding, 1500 r.p.m. The supernatant was removed. The pellet color was assessed; if RBCs remained (pellet color pink or red), the ACK step was repeated up to two additional times. To remove cell clumps in the MBC protocol (or sample), the pellet was resuspended in 100 µl of TrypLE (Life Technologies, cat. no. 12604013) and incubated while constantly pipetting at room temperature for 1 min with a 200 µl pipette tip. TrypLE was inactivated by adding 200 µl of cold RPMI 1640 (10% FBS) and the cells were pelleted using short centrifugation as described above. The pellet was resuspended in 50 µl of 0.4% BSA (Ambion, cat. no. AM2661) in PBS. To assess the single-cell suspension, viability and cell count, 5 µl of Trypan blue (Thermo Fisher Scientific, cat. no. T10282) was mixed with 5 µl of the sample and loaded onto an INCYTO C-Chip Disposable Hemocytometer (Neubauer International, VWR, cat. no. 82030-468). The cell concentration was calculated as necessary to a range of 200–2,000 cells per µl. A total of 8,000 cells were loaded into each channel of the 10X Genomics Single Cell Chromium Controller. Due to differences between clinical samples, some steps may need to be repeated or adjusted; for a general overview of guidelines see Fig. 2a.

NSCLC-C4 protocol workflow. A similar workflow was used for protocol NSCLC-C4 with the following modifications. Following mechanical chopping as above, sample was dissociated for 15 min in a 15 ml falcon tube, with a gentle vortex every 5 min, followed by filtration through a 70 µm filter, and washed with 20 ml of ice-cold PBS and centrifuged at 580g for 5 min. RBC lysis was performed similarly to the above workflow by resuspending the pellet in 1 ml ACK lysis buffer with incubation on ice for 1 min. A 20 ml volume of ice-cold PBS was added to quench the ACK lysis buffer, followed by filtration through a 70 µm filter, and centrifugation at 580g for 5 min. Sample NSCLC-C4 was further cleaned using a Viahance dead-cell removal kit (BioPAL, cat. no. CP-50VQ02) according to the manufacturer’s instructions. Cells were then resuspended in PBS and loaded on the 10X Genomics Single Cell Chromium Controller as described above.

GBM workflow. All steps were completed on ice. Each sample was miniced thoroughly in a Petri dish, 4 ml HBSS was added (Life Technologies, cat. no. 14175095), then the sample was transferred to 15 ml tubes and centrifuged at 1,000 r.p.m. for 2 min. After sample collection using a standard biobanking technique, samples were placed in cryo-tubes without any liquid. Complete removal of liquid from the sample was accomplished by gently wiping it (not patting, as this would damage the tissue) in a Petri dish, then pelleting the sample in a centrifuge. The sample was then pipetted up–down 20 times, incubated at 37 °C for an additional 15 min, and pipetted again. After dissociation, the sample was filtered through a 100 µm cell strainer (Fisher Scientific, cat. no. 22–363–549) into a 50 ml conical tube. The tissue fragments left in the cryotube were then frozen in OCT and stored at −80°C for long-term storage.

The other samples from NSCLC-C4 were obtained from tissue banks as follows: ovarian optimal cutting temperature compound (OCT)-frozen archival
(Cedarlane, cat. no. CL5015) or RBC Lysis Solution (10x) (Miltenyi Biotec, cat. no. 130-094-183). The pellet was washed with 10 ml of cold PBS/1% BSA, transferred to a 15 ml tube and centrifuged at 1,200 r.p.m. for 3 min. Supernatant was removed and the pellet was resuspended in 0.4% BSA in PBS. The single-cell suspension was visualized, counted and loaded on the 10x Genomics Single-Cell Chromium Controller as described above.

Dissociation mixtures for different tumor types. Dissociation mixtures were prepared 5–10 min before sample processing from frozen aliquoted stocks, as follows.

MBC LD protocol. A 950 µl volume of RPMI 1640 (Thermo Fisher Scientific, cat. no. I1875093) was used with 10 µl of 10 mg ml⁻¹ DNase I (Sigma Aldrich, cat. no. 11284932001) to a final concentration of 100 µg ml⁻¹, and 40 µl of 2.5 mg ml⁻¹ Liberase TM (Sigma Aldrich, cat. no. 5401127001). Ovarian cancer resection MHTD kit. The dissociation mixture was based on the Miltenyi Human Tumor Dissociation Kit (Miltenyi Biotec, cat. no. 130-095-929). Before starting, enzymes H, R and A were resuspended according to the manufacturer's instructions. Dissociation mix containing 2.2 ml RPMI, 100 µl enzyme H, 50 µl enzyme R and 12.5 µl enzyme A was prepared immediately before use.

Neuroblastoma NB-C4 protocol. Medium 199 with Hanks balanced salts buffer (Thermo Fisher Scientific, cat. no. 14117121, 187.5 µg ml⁻¹ of 2-mg ml⁻¹ pronase (Sigma Aldrich, cat. no. 1065921001) to a final concentration of 1.250 µg ml⁻¹, 27.6 µg ml⁻¹ of 1 mg ml⁻¹ elastase (Thermo Fisher Scientific, cat. no. NC9301601) to a final concentration of 9.2 µg ml⁻¹, 30 µl of 10 mg ml⁻¹ DNase I (Sigma Aldrich, cat. no. 11284932001) to a final concentration of 100 µg ml⁻¹, 30 µl of 100 µg ml⁻¹ Dispase (Sigma Aldrich, cat. no. 4942078001) to a final concentration of 100 µg ml⁻¹, 30 µl of 100 µg ml⁻¹ collagenase A (Sigma Aldrich, cat. no. 10303578001) to a final concentration of 1,500 µg ml⁻¹ and 3 µl of 100 mg ml⁻¹ collagenase IV (Thermo Fisher Scientific, cat. no. NC9386705) to a final concentration of 100 µg ml⁻¹.

NSCLC PDEC protocol. We used 2692 µl HBBS (Thermo Fisher Scientific, cat. no. 141017012, 187.5 µl of 20 mg ml⁻¹ of 1% CHAPS (Millipore Sigma, cat. no. 5401119001) to a final concentration of 100 µg ml⁻¹, 50 µl of 10 mg ml⁻¹ DNase I (Sigma Aldrich, cat. no. 11284932001) to a final concentration of 100 µg ml⁻¹, 46 µl of 1 mg ml⁻¹ elastase (Thermo Fisher Scientific, cat. no. NC9301601) to a final concentration of 9.2 µg ml⁻¹.

NSCLC LE protocol. We used 4.7 ml RPMI 1640 (Thermo Fisher Scientific, cat. no. I1875093, 200 µl of 2.5 mg ml⁻¹ Liberase TM (Millipore Sigma, cat. no. 5401119001) to a final concentration of 100 µg ml⁻¹, 50 µl of 10 mg ml⁻¹ DNase I (Sigma Aldrich, cat. no. 11284932001) to a final concentration of 100 µg ml⁻¹ and 46 µl of 1 mg ml⁻¹ elastase (Thermo Fisher Scientific, cat. no. NC9301601) to a final concentration of 9.2 µg ml⁻¹.

Ovarian cancer ascites. Ascites samples without spheres were selected and processed as follows. The ovarian ascites sample, CD45⁻ cells were thawed in 10 ml RPMI, pelleted and washed with an additional 10 ml of 1% CHAPS (Millipore Sigma, cat. no. M1028) in ultrapure water. Nuclei isolation from frozen samples for snRNA-Seq. For CST, 1 ml of 2x ST buffer, 980 µl of 1% CHAPS (Millipore, cat. no. 220201), 10 µl of 2% BSA (New England BioLabs, cat. no. R90005) and 10 µl of nuclelease-free water; for TST, 1 ml of 2x ST buffer, 60 µl of 1% Tween-20 (Sigma Aldrich, cat. no. P-7949), 10 µl of 2% BSA (New England BioLabs, cat. no. R90005) and 990 µl of 1x ST buffer; for NST, 1 ml of 2x ST buffer, 100 µl of 0.1 M HEPES (Sigma Aldrich, cat. no. R8545) and 950 µl of 1x ST buffer. 1x ST buffer was prepared by dilution 2x ST with ultrapure water (Thermo Fisher Scientific cat. no. 10977023) in a ratio of 1:1.

Nucleus isolation from frozen samples for snRNA-Seq. On dry ice, tissue was split and subjected to one of three ST-based nucleus isolation protocols8 and the EZ nucleus isolation buffer, as detailed in the following.

Nucleus isolation workflow for ST-based buffers. On ice, a piece of frozen tumor tissue was placed into a well of a 6-well plate (STEM Cell Technologies, cat. no. 3001; with 1 ml of CST, TST or NST buffer. For samples frozen in OCT, an additional step of removing the surrounding OCT and washing any residual OCT from the sample with PBS was performed in a 10 cm Petri dish. Tissue was then chopped using Noyes Spring Scissors (Science Tools, cat. no. 15514-12) for 10 min on ice. For cell pellets, such as for CLL frozen cells, sample was pipetted in the buffer on ice, instead of chopping. The homogenized solution was then filtered through a 40 µm Falcon cell strainer (Thermo Fisher Scientific, cat. no. 08-771-1). An additional 1 ml of the detergent buffer solution was used to wash the well and filter. The volume was brought up to 5 ml with 3 µl of 1x ST buffer. The sample was then transferred to a 15 ml conical tube and centrifuged at 4 °C for 5 min at 500 g in a swinging bucket centrifuge. The pellet was resuspended in 1x ST buffer. Resuspended cell pellet was dependent on the size of the pellet, usually within the range of 100–200 µl. The nucleus solution was then filtered through a 35 µm Falcon cell strainer (Corning, cat. no. 352325). Nuclei were counted using a C-chip disposable hemocytometer.
Nucleus isolation workflow using EZY lysis buffer. Nucleus isolation was done as previously described. Briefly, tissue samples were cut into pieces <0.5 cm and homogenized using a glass Dounce tissue grinder (Sigma, cat. no. D8938). The tissue was homogenized 25 times with pestle A and 25 times with pestle B in 2 mL of ice-cold nucleic EZ lysis buffer. The sample was then incubated on ice for 5 min, with an additional 3 mL of cold EZ lysis buffer. Nuclei were centrifugated at 500g for 5 min at 4°C, washed with 5 mL ice-cold EZ lysis buffer and incubated on ice for 5 min. After centrifugation, the nucleus pellet was washed with 5 mL nucleic acid suspension buffer (NSB; consisting of 1X PBS, 0.01% BSA and 0.1% RNase inhibitor, Clontech Cat. no. 630131A). Isolated nuclei were resuspended in 2 mL NSB, filtered through a 35 μm cell strainer (Corning-Falcon, cat. no. 352235) and counted. A final concentration of 1,000,000 nuclei per mL was used for loading on a 10x channel.

Droplet-based sc/snRNA-Seq. For V2 10x technology, either 8,000 single cells or 10,000 single nuclei were loaded into each channel of a Chromium single-cell 3′ Chip. For V3 10x technology, 8,000 single cells and 8,000 single nuclei were loaded. Single cells/nuclei were partitioned into droplets with gel beads in the Chromium Controller. After emulsions were formed, barcoded reverse transcription of RNA took place. This was followed by CDNA amplification, fragmentation and adapter and sample index attachment, all according to the manufacturer’s recommendations. Libraries from four 10x channels were pooled together and sequenced on one lane of an Illumina HiSeq X, or on one flow cell of a NextSeq, with paired end reads as follows: read 1, 26 nt; read 2, 55 nt; index 1, 8 nt; index 2, 0 nt.

Computational methods. scRNA-Seq data processing. We used Cell Ranger mkFastq (v2.0 and v3.0) (10x Genomics) to generate demultiplexed FASTQ files from the raw sequencing reads. We aligned these reads to the human GRCh38 genome and quantified gene counts as UMI s using Cell Ranger count (v2.0 and v3.0) (10x Genomics). For snRNA-Seq reads, we counted reads mapping to introns as well as exons, as this results in a greater number of genes detected per nucleus, more massively parallel processing and better cell identification power as described. To count introns during read mapping, we followed the approach described at https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/advanced/references. Briefly, we built a ‘pre-mRNA human GRCH38 reference using Cell Ranger mkfkk (v3.0) (10x Genomics) and a modified gene transfer format (GTF) file, where, for each transcript, the feature type had been changed from transcript to exon. The starting GTF files came from refdata-cellranger-GRCh38-1.2.0.targz or refdata-cellranger-GRCh38-3.0.tar.gz, and are available for download at https://support.10xgenomics.com/single-cell-gene-expression/software/downloads/3.0.

To downsample sequencing reads or gene counts (UMIs) when comparing protocols, we used scRNAReads and downsampleMatrix, respectively, from the R package DropletUtils (v1.0.3 or higher). Reads were downsampled to match the protocol with the lowest number of total reads. After downsampling by total reads, we used writeXCounts from DropletUtils and a custom Python script to generate an HDF5 file for input into our analysis pipelines, as described in the sections that follow and in the ‘Code availability’ section.

QC of scRNA-Seq data. To maintain explicit control over all gene and cell quality control filters, in all our downstream analyses we used the raw feature-barcode matrix, rather than the filtered feature-barcode matrix generated by Cell Ranger. We removed low-quality cells by requiring each cell to have a minimum number of UMIs and genes detected. We used different thresholds depending on the experimental modality (single cell or single nucleus) and on the 10x kit (V2 or V3 chemistry). For single nucleus data, we retained nuclei with at least 200 genes and 400 UMIs detected by V2 chemistry and with at least 500 genes and 1,000 UMIs detected by V3 chemistry. For single-cell data, we retained cells with at least 500 genes and 1,000 UMIs detected by either V2 or V3 chemistry. For the V2–V3 comparison in HTAPP-951-SMP-4652 (Extended Data Fig. 9), we used the same thresholds for both chemistries: at least 200 genes and 400 UMIs detected. For both data types, we filtered out those cells or nuclei where ≥20% of UMIs came from mitochondrial genes. Finally, we normalized the total UMIs per cell or nucleus to 100,000 (CPI00K) and log-transformed these values to report gene expression as \( E = \log(CPI00K + 1) \).

We reported the following QC metrics: number of total reads per library sample, sequencing saturation (fraction of reads originating from an already-observed UMI as reported by Cell Ranger count), total recovered cells or nuclei, number of reads per cell or nucleus, number of UMIs per cell or nucleus, number of genes detected per cell or nucleus, fraction of UMIs in a cell or nucleus aligned to mitochondrial genes, fraction of droplets estimated to contain only ambient RNA (‘empty drops’), fraction of cell or nucleus doublets, the number of detected cell types and the pattern of CNAs for malignant cells. For a subset of samples, we also calculated the number of cells or nuclei per detected cell type and the estimated level of ambient RNA in droplets containing cells.

We predicted droplets containing only ambient RNA and no cells using EmptyDrops (part of DropletUtils, v0.3 or higher), with the retain parameter set by the knee of the curve in the barcode rank plot (cell barcodes ranked by their total UMIs). We predicted potential doublets using Scrublet (v0.2) with expected_doublet_rate = 0.06 (ref. 3). We estimated the levels of ambient RNA using SouPx (v0.3.1)10 and a set of cell-type-specific marker genes (Supplementary Table 1). Importantly, we flagged the doublets and empty drops and retained them in our analysis, instead of immediately filtering them out. Droplets that appear to contain doublets or empty drops can arise from many different effects, such as cellular differentiation or insufficient sequencing, and by carrying them through the analysis, potential doublets or empty drops can be more clearly interpreted in the context of the full dataset.

Dimensionality reduction, clustering and visualization. For each tumor sample, we analyzed the filtered expression matrix to identify cell subsets, as previously described. We chose highly variable genes with a z-score cutoff of 0.5 (ref. 4), centered and scaled the expression of each gene to have a mean of zero and standard deviation of one, and performed dimensionality reduction on the variable genes using principal component analysis. We used the top 50 principal components (PCs) as input to Louvain graph-based clustering, with the resolution parameter set to 1.3. For each cluster of cells, we identified cluster-specific differentially expressed genes using the following tests: an AUC classifier, Welch's t-test and Fisher’s exact test. For sets that returned a p value, we controlled the false discovery rate at 5% with the Benjamini–Hochberg procedure. We visualized gene expression and clustering results by embedding cells or nuclei profiles in a Uniform Manifold Approximation and Projection (UMAP) of the top 50 PCs, with min_dist = 0.5, spread = 1.0, number of neighbors = 15 and the Euclidean distance metric.

Annorting cell subsets. For each cell subset identified by clustering, we assigned a cell type from the malignant, parenchymal, stromal and immune compartments of the tumor microenvironment using a combination of differentially expressed genes, known gene signatures (Supplementary Table 1) and SingleR (v0.2.2)11, an automated annotation package. When running SingleR, only cell types assigned to 30 or more cells were considered. When scoring cells for the expression of known gene signatures, we used the AddModuleScore function in Seurat (v2.3.4)12. We note that overlapping expression programs between T cells and NK cells make these cell types sometimes more difficult to identify accurately. We did not distinguish macrophages, monocytes and dendritic cells, and annotated all of these as scoring for ‘macrophages’ signatures.

We identified the malignant cells by inferring chromosomal CNAs from the gene-expression data using inferCnv (v1.1.0)13. On a sample-by-sample basis, we used the immune and endothelial cells as a healthy reference to estimate CNAs in the malignant cells. We created the count matrix file and annotation file for inferCnv by randomly subsampling the counts data to sample at most 2,000 cells or nuclei. We created a gene ordering file from the human GRCh38 assembly, which contains the chromosomal start and end positions for each gene. To run inferCnv, we used a cutoff of 0.1 for the minimum average read counts per gene among reference cells or nuclei, clustered according to the annotated cell types, downsampled the output, ran a hidden Markov model (HMM) to predict cell CNA level, implemented inferCnv’s 16 HMM model, and requested eight threads for parallel steps.

Comparing sc- and snRNA-Seq data. To compare profiles between sc- and snRNA-Seq data collected from the same sample, we used a batch correction approach.

We performed batch correction using CCA as implemented in Seurat (v2.3.4)14. We selected 1,500 genes that were variable across both the cell and nucleus data, used those genes as input to RunCCCA to compute the first 20 canonical components, and aligned the first 12 canonical components with AlgHiSubspace. The aligned canonical components represent a co-embodiment of the cell and nucleus data, and we carried out clustering in this dimensionality-reduced space using FindClusters. Following batch correction by CCA, we scored the dissociation signature from ref. 15 (from their Supplementary Table 5) on our matched cell/nuclei samples using the AddModuleScore function in Seurat (v2.3.4)14.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this Article.

Data availability
All main and extended data figures have associated raw data. Raw data will be available in the controlled access repository dbGaP (https://www.ncbi.nlm.nih.gov/dbgap) under dbGaP Study Accession phs001983.v1.p1. Raw data will also be available in the controlled access repository DUOS (https://duos.broadinstitute.org/), under DUOS Dataset IDs DUOS-000111, DUOS-000112, DUOS-000113 and DUOS-000114. The counts matrices and metadata for each sample will be
publicly available in the Gene Expression Omnibus [https://www.ncbi.nlm.nih.gov/geo/] under data repository accession no. GSE140819. Finally, we provide a website that displays a comprehensive analysis summary for each sample tested [https://tumor-toolbox.broadinstitute.org].

Code availability
We implemented all initial analysis steps, from FASTQ files to clustering cell subsets, in Cumulus,28 which may be executed in both a cloud-based environment and locally. Pipelines were written in the Workflow Description Language (WDL) and run on Cromwell in the Terra Cloud platform [https://app.terra.bio/], and data were stored in Google Cloud Platform storage buckets. Cumulus workflows are publicly available at https://github.com/klarman-cell-observatory/Cumulus. We built and ran an additional pipeline for subsequent QC steps, including detecting empty drops and doublings, annotating cell subsets, evaluating cell type specific QC s and inferring CNAs. This pipeline was implemented in R (v3.5 or higher) by converting the single-cell AnnData objects from Cumulus into Seurat objects, and was used to compare and evaluate processing protocols. Our analysis pipeline, including complete example analysis for one scRNA-Seq sample and one one mRNA-Seq sample, will be made publicly available at https://github.com/klarman-cell-observatory/HTAPP-Pipelines.

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Author contributions
M.S., C.R.M.P., O.A., O.R.-R. and A. Regev conceived and led the study. M.S. designed protocols and carried out experiments together with J. Waldman, I.W., S.V., J. Wu, M.-J.S., S.N., D.D., M.N., A.G.P., S.G. and L.N. C.B.M.P. and O.A. designed and performed computational analysis, with input and assistance from A.M.T., L.J.-A., O.C., J.K., M.H., C.S., G.S.-R. and T.L.T. G.S.-R. contributed CNA inference analyses. I.W. and S.V. performed FACS analysis and scRNA-Seq of ovarian ascites and NSCLC CD45 depletion. P.J.T. carried out NSCLC specimen collection and protocol optimization. E.D. and C.S. provided protocols for scRNA-Seq. A.W. provided input for the scRNA-Seq PDEC protocol. A.G.P. provided input and performed the neuroblastoma papan protocol. S.H.G. provided input for CLL. S.G. provided and carried out the GBM protocol. J.-V. managed the study and tissue acquisition for all samples except neuroblastoma O-PDX, which A.K. managed. B.L., Y.R. and Y.G. contributed Cumulus expertise. A. Rotem, C.J.W., B.I., R.H., F.S.H., C.H.Y., A.N.H., S.B., M.L.S., R.B., E.H.S., M.R.C., M.A.D., N.R.C., U.A.M., N.W. and B.E.J. supervised research, provided tissues and shared clinical insights. M.S., C.B.M.P., O.A., O.R.-R. and A. Regev wrote the manuscript, with input from all authors.

Competing interests
A. Regev is a founder and equity holder in Celsius Therapeutics, an equity holder in Imunitas, and a SBAM member of Syros Pharmaceuticals, Thermo Fisher Scientific, Asimov and NeoGene Therapeutics. M.S., O.A., E.D., O.R.-R. and A. Regev are co-inventors on patent applications filed by the Broad Institute for inventions relating to work in this manuscript, such as in PCT/US2018/068066 and US provisional application no. 62/745,259. C.J.W. is a founder and member of the scientific advisory board of Neon Therapeutics and receives research funding from Pharmacyclics. A. Rotem is a consultant and equity holder in Celsius Therapeutics. F.S.H. reports grants, personal fees from Bristol-Myers Squibb and Novartis and personal fees from Merck, EMD Serono, Takeda, Surface, Genentech/Roche, Compass Therapeutics, ApiCure, Bayer, Aduro, Sanofi, Pfizer, Pionyr, Verastem, Torque and Rheos. In addition, F.S.H. has a patent ’Methods for Treating MICA-Related Disorders’ (#20100111973) with royalties paid, a patent ’Tumor Antigens and Uses Therof’ (#7250291) issued, a patent ‘Anti-epitope-2 Biomarkers Predictive of Anti-immune Checkpoint Response’ (#20170248603) pending, a patent ’Compositions and Methods for Identification, Assessment, Prevention and Treatment of Melanoma using PD-L1 Isomorphs’ (#20160340407) pending, a patent ’Therapeutic Peptides’ (#2016044716) pending, a patent ’Therapeutic Peptides’ (#2014004112) pending, a patent ’Therapeutic Peptides’ (#20170008962) pending, a patent ’Therapeutic Peptides’ (patent no. 9402905) issued and a patent Methods of using ‘Pembrolizumab and Trebananib’ pending. R.H. has received research support from Novartis and Bristol-Myers and is a consultant for Tango Therapeutics. R.B. has received research support from Roche, Genentech, Merck, Siemens, Verastem, Grifostine, Epizyme, Medgenome and HTG and has equity in Navigation Sciences. A.N.H. has received research support from Novartis, Amgen, Pfizer, Roche/Genentech and Relay Therapeutics. B.I. is a paid consultant for Merck. U.A.M. received consulting fees from Novartis and Merck.

Additional information
Extended data is available for this paper at https://doi.org/10.1038/s41591-020-0844-1. Supplementary information is available for this paper at https://doi.org/10.1038/s41591-020-0844-1.

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Peer review information Joao Monteiro was the primary editor on this article and managed its editorial process and peer review in collaboration with the rest of the editorial team.

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Extended Data Fig. 1  | scRNA-Seq protocol comparison for a single NSCLC sample. (a) Sample processing and QC overview. For each protocol, shown are the number of cells passing QC, and the number of sequencing reads and sequencing saturation across all cells. The remaining metrics are reported for cells passing QC: the median number of reads per cell, median number of UMIs per cell, median number of genes per cell, median fraction of UMIs mapping to mitochondrial genes, fraction of cell barcodes called as empty droplets and fraction of cell barcodes called as doublets. (b) Read mapping QCs. The percent of bases in the sequencing reads (y axis) mapping to the genome, transcriptome and intergenic regions (x axis) across the three protocols (colored bars). (c) Cell type assignment. UMAP embedding of single cell profiles from each protocol colored by assigned cell type signature. (d) Inferred CNA profiles. Chromosomal amplification (red) and deletion (blue) inferred in each chromosomal position (columns) across the single cells (rows) from the NSCLC-C4 (left) and LE (right) protocols. Top: reference cells not expected to contain CNA in this cancer type. Bottom: cells tested for CNA relative to the reference cells. Color bar: assigned cell type signature for each cell. (e) Ambient RNA estimates. Estimates of the fraction of RNA in each cell type derived from ambient RNA contamination (y axis), with cell types ordered by their mean number of UMIs/cell (x axis). Red line: global average of contamination fraction; Green line: LOWESS (locally weighted scatterplot smoothing) smoothed estimate of the contamination fraction within each cell type, along with the associated binomial 95% confidence interval (Clopper-Pearson interval).
Extended Data Fig. 2 | Cell type specific QC metrics for scRNA-Seq protocol comparison in a single NSCLC sample. Cell type specific QC metrics for NSCLC14. Distribution (median and first and third quartiles) of the number of reads per cell, number of UMIs per cell, number of genes per cell and fraction of UMIs mapping to mitochondrial genes in each cell (y axes) in each of the three protocols (x axis), for cells passing QC from each cell type (rows). n = 1 sample per protocol. Number of B cells (k) from NSCLC-C4, PDEC and LE, respectively, is: 100, 121, 78; endothelial cells: 0, 920, 1,078; epithelial cells: 1,284, 641, 260; fibroblasts: 0, 1,476, 1,403; macrophages: 3,306, 911, 727; mast cells: 0, 119, 77; T cells: 449, 723, 722.
Extended Data Fig. 3 | scRNA-Seq protocol comparison for NSCLC following read down-sampling. Shown are analyses for NSCLC14 (as in Extended Data Figs. 1 and 2), but after the total number of sequencing reads within each sample was down-sampled to match the protocol with the fewest total sequencing reads. (a) Sample processing and QC overview. For each protocol, shown are the number of cells passing QC. The remaining metrics are reported for those cells passing QC: median number of UMIs per cell, median number of genes per cell, median fraction of UMIs mapping to mitochondrial genes in each cell, fraction of cell barcodes called as empty droplets and fraction of cell barcodes called as doublets. (b, c) Overall and cell types specific QCs. Distribution (median and first and third quartiles) of the number of UMIs per cell, number of genes per cell and fraction of gene expression per cell from mitochondrial genes (y axes) in each of the three protocols (x axis), for all cells passing QC (b) and for cells from each cell type (c, rows). (d,e) Relation of empty droplets and doublets to cell types. UMAP embedding and fraction (horizontal bar) of single cell (gray), empty droplet (red, d) and doublet (red, e) profiles for each protocol. (f) Cell type assignment. UMAP embedding of single cell profiles from each protocol colored by assigned cell type signature. n = 1 sample per protocol and number of cells (k) is indicated in (a). Number of B cells (k) from NSCLC-C4, PDEC and LE, respectively, is: 114, 157, 78; endothelial cells: 0, 879, 1,078; epithelial cells: 1,283, 644, 260; fibroblasts: 0, 1,439, 1,403; macrophages: 3,278, 853, 727; mast cells: 0, 106, 77; T cells: 432, 663, 722.
Extended Data Fig. 4 | CD45+ depletion protocol enriches for non-immune cells in freshly processed NSCLC and ovarian ascites. (a, b) QCs. Distribution (median and first and third quartiles) of the number of reads per cell, number of UMIs per cell, number of genes per cell and fraction of gene expression per cell from mitochondrial genes (y axes) for all cells passing QC from NSCLC (a) before and after CD45+ cell depletion, and for ovarian ascites (k = 2,998 and 10,716 cells, respectively) or (b) after CD45+ cell depletion (2,359 cells) (x axis). n = 1 sample per protocol. (c) CD45+ cell depletion estimates in ovarian cancer ascites by FACS. Flow-cytometry comparison of single cells isolated without (top) or with (bottom) depletion of CD45+ cells. Cells were gated by FSC and SSC (first column), doublets removed using FSC-A and FSC-H (second column), 7-AAD gating of dead cells to identify live cells (third column), the distribution of immune and non-immune cells quantified using a CD45 antibody (fourth column) and the distribution of EPCAM+ cells quantified using an EPCAM antibody (fifth column). (Efficient removal of CD45+ cells from ovarian cancer ascites was also demonstrated with an independent sample from a different patient (data not shown).) Number of cells without and with depletion, respectively are: 10,000, 10,000 (1st column), 3,468, 2,256 (2nd column), 3,467, 2,251 (3rd column), 2,936, 2,174 (4th and 5th columns).
Extended Data Fig. 5 | See next page for caption.
Extended Data Fig. 5 | snRNA-Seq protocol comparison in a single neuroblastoma sample. (a) Sample processing and QC overview. For each protocol, shown are the number of nuclei passing QC, number of sequencing reads and sequencing saturation across all nuclei. The remaining metrics are reported for those nuclei passing QC: median number of reads per nucleus, median number of UMIs per nucleus, median number of genes per nucleus, median fraction of UMIs mapping to mitochondrial genes in each nucleus and fraction of nucleus barcodes called as doublets. (b) Read mapping QCs. The percent of bases in the sequencing reads (y-axis) mapping to the genome, transcriptome and intergenic regions (x-axis) across the four protocols (colored bars). (c) Cell type assignment. UMAP embedding of single nucleus profiles from each protocol colored by assigned cell type signature. (d) Inferred CNA profiles. Chromosomal amplification (red) and deletion (blue) inferred in each chromosomal position (columns) across the single nuclei (rows). Top: reference nuclei not expected to contain CNA in this cancer type. Bottom: nuclei tested for CNA relative to the reference nuclei. Color bar: assigned cell type signature for each nucleus. n=1 sample per protocol and number of nuclei (k) is indicated in (a).
Extended Data Fig. 6 | Cell type specific QC metrics for snRNA-Seq protocol comparison in a single neuroblastoma sample. Cell type specific QCs for HTAPP-244-SMP-451. Distribution (median and first and third quartiles) of the number of reads per nucleus, number of UMIs per nucleus, number of genes per nucleus and fraction of UMIs mapping to mitochondrial genes in each nucleus (y axes) in each of the four protocols (x axis), for nuclei passing QC from each cell type (rows). n = 1 sample per protocol. Number of endothelial nuclei (k) from EZ, CST, NST and TST, respectively, is: 69, 32, 91, 95; erythrocyte: 0, 18, 0, 15; T cell: 157, 171, 229, 337; neuroendocrine: 7,379, 5,728, 6,790, 6,477; neural crest: 18, 27, 50, 67; macrophage: 119, 107, 189, 230; fibroblast: 138, 74, 182, 194; zona glomerulosa: 16, 0, 0, 0.
Extended Data Fig. 7 | snRNA-Seq protocol comparison in a single neuroblastoma sample following read down-sampling. Shown are analyses for NB HTAPP-244-SMP-451 (as in Extended Data Figs. 5 and 6), but after the total number of sequencing reads within each sample was down-sampled to match the protocol with the fewest total sequencing reads. (a) Sample processing and QC overview. For each protocol, shown are the number of nuclei passing QC. The remaining metrics are reported for those nuclei passing QC: median number of UMIs per nucleus, median number of genes per nucleus, median fraction of UMIs mapping to mitochondrial genes in each nucleus and fraction of nucleus barcodes called as doublets. (b,c) Overall and cell types specific QCs. Distribution (median and first and third quartiles) of the number of UMIs per nucleus, number of genes per nucleus and fraction of UMIs mapping to mitochondrial genes in each nucleus (y axes) in each of the four protocols (x axis), for all nuclei passing QC (b) and for nuclei from each cell type (c, rows). (d) Relation of doublets to cell types. UMAP embedding and fraction (horizontal bar) of single nucleus (gray) and doublet (red) profiles for each protocol. (e) Cell type assignment. UMAP embedding of single nucleus profiles from each protocol colored by assigned cell type signature. n=1 sample per protocol and number of nuclei (k) is indicated in (a). Number of endothelial nuclei (k) from EZ, CST, NST, TST is: 53, 31, 95, 98; erythrocyte: 0, 0, 0, 23; T cell: 146, 177, 230, 345; neuroendocrine: 7,407, 5,726, 6,776, 6,454; neural crest: 14, 27, 50, 69; macrophage: 123, 104, 196, 240; fibroblast: 111, 81, 177, 186; zona glomerulosa: 21, 0, 0, 0.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | Inferred CNA profiles from snRNA-Seq in diverse tumors. Chromosomal amplification (red) and deletion (blue) inferred in each chromosomal position (columns) across the single nuclei (rows) from three MBC samples (a-f), one ovarian cancer sample (g-i) and one sarcoma sample (j, k). Top: reference nuclei not expected to contain CNA in this cancer type. Bottom: nuclei tested for CNA relative to the reference nuclei. Color bar: assigned cell type signature for each nucleus. n = 1 sample per protocol and number of nuclei (k) per sample: MBC HTAPP-963-SMP-4741—9,857 (CST), 7,260 (TST); MBC HTAPP-394-SMP-1561—6,948 (CST), 8,058 (NST); MBC HTAPP-589-SMP-2851—7,858 (CST), 8,373 (TST); ovarian HTAPP-316-SMP-991—9,026 (CST), 5,970 (NST), 10,493 (TST); sarcoma HTAPP-951-SMP-4652—7,858 (CST), 4,458 (TST).
Extended Data Fig. 9 | snRNA-Seq protocol comparison of V2 and V3 chemistry from 10x Genomics on a resection of sarcoma. (a) Sample processing and QC overview. For each protocol, shown are the number of nuclei passing QC, after the total number of sequencing reads from the V3 protocol data was down-sampled to match the number of reads in the V2 data. The remaining metrics are reported for those nuclei passing QC: median number of UMI s per nucleus, median number of genes per nucleus, median fraction of UMIs mapping to mitochondrial genes in each nucleus and fraction of nucleus barcodes called as doublets. (b) Overall QCs. Distribution (median and first and third quartiles) of number of UMIs per nucleus, number of genes per nucleus and fraction of UMIs mapping to mitochondrial genes in each nucleus (y axes) for all nuclei passing QC. $n = 1$ sample per chemistry type and number of nuclei ($k$) is indicated in (a).
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Comparison of scRNA-Seq and snRNA-Seq from the same tumor sample. (a–g) CLL. UMAP embedding of scRNA-Seq and snRNA-Seq profiles of the same CLL sample combined by CCA\textsuperscript{24} (Methods) showing profiles (dots) from both (a), scRNA-Seq (b) and snRNA-Seq (c), colored by assigned cell type signatures. (d) Proportion of cells from each subset in the two protocols. k: number of cells or nuclei passing QC. (e–f) Same UMAP embedding as in (a), colored by cells or nuclei (e) or unsupervised clustering (f). (g) Fraction of cells and nuclei in each cluster. n = 1 sample per protocol and number of cells and nuclei is indicated in (d). (h–n) O-PDX neuroblastoma. As in (a–g) for an O-PDX neuroblastoma sample. n = 1 sample per protocol and number of cells and nuclei is indicated in (k). (o–r) Dissociation signatures are more prominent in cells than in nuclei from the same tumors. Left and middle: UMAP embedding of scRNA-Seq (left) and snRNA-Seq (middle) profiles (dots) of the same tumor combined by CCA\textsuperscript{24} and colored by the score of a dissociation signature (color bar). Right: Distribution of dissociation signature score (y axis; median and first and third quartiles) in cells (green) and nuclei (orange). (o) neuroblastoma (3,449 cells, 7,810 nuclei), (p) MBC (5,163 cells, 7,260 nuclei), (q) CLL (2,562 cells, 2,339 nuclei), (r) O-PDX neuroblastoma (3,495 cells, 4,946 nuclei).
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For all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.

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- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection We did not use any software for data collection.

Data analysis

1) Cell Ranger mkfastq (v2.0 and v3.0) (10x Genomics) to generate demultiplexed FASTQ files from the raw sequencing reads
2) Cell Ranger count (v2.0 and v3.0) (10x Genomics) to align reads and quantify gene counts as UMIs
3) Cell Ranger mkref (v3.0) (10x Genomics) to build a custom reference
4) R (v3.5 or higher) for gene expression analyses
5) RStudio (v1.2.1335) for running R analyses
6) Python (v3.7) for gene expression analyses
7) DropletUtils (v1.0.3 or higher, R package, http://bioconductor.org/packages/release/bioc/html/DropletUtils.html) to estimate droplets containing only ambient RNA
8) Scrublet (v0.2, Python package, https://github.com/AllonKleinLab/scrublet) to estimate droplets contain doublets
9) SoupX (v0.3.1, R package, https://github.com/constantAmateur/SoupX) to estimate ambient RNA in droplets that also contain cells
10) SingleR (v0.2.2, R package, https://github.com/dviraran/SingleR) for automated draft annotation
11) Seurat (v2.3.4, R package, https://satijalab.org/seurat/install.html) as a framework for additional quality control steps, cell-subset annotation, and cell/nuclei batch correction
12) inferCNV (v1.1.0, https://github.com/broadinstitute/infercnv) for inferring chromosomal copy number aberrations (CNAs) from the gene-expression data
13) Cumulus (https://github.com/klarman-cell-observatory/Cumulus), developed by Bo Li and his colleagues, is used to perform all major single-cell and single-nucleus RNA-Seq data analysis
14) BD FACSDiva Software (v8.0.1) for flow cytometry analysis.
15) FlowJo (v10.5.3) for flow cytometry plotting

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.
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Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

All main and Extended Data figures have associated raw data. Raw data will be available in the controlled access repository dbGaP (https://www.ncbi.nlm.nih.gov/gap/), under the dbGaP Study Accession phs001983.v1.p1; raw data will also be available in the controlled access repository DUOS (https://duos.broadinstitute.org), under the following DUOS Dataset IDs: DUOS-000111, DUOS-000112, DUOS-000113, and DUOS-000114. The counts matrices and metadata for each sample will be publicly available in Gene Expression Omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/ under data repository accession no. GSE140819. Finally, we provide a website that displays a comprehensive analysis summary for each sample tested (https://tumor-toolbox.broadinstitute.org).

Field-specific reporting

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Life sciences study design

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

Sample size

For each sample, an input of 8,000 single cells or 8,000-10,000 single nuclei were loaded into each channel of the 10x Genomics Single-Cell Chromium Controller. These loading values were chosen to balance the probability of forming doublets with the goal of having maximal cell recovery and sufficient cell/nuclei recovery to reveal the heterogeneous landscape of the tumors.

Data exclusions

We removed low quality cells by requiring each cell to have a minimal number of UMIs and genes detected. We used different thresholds depending on the experimental modality (single cell or single nucleus) and on the 10x kit (V2 or V3 chemistry). For single nucleus data, we retained nuclei with at least 200 genes and 400 UMIs detected by V2 chemistry and with at least 500 genes and 1,000 UMIs detected by V3 chemistry. For single cell data, we retained cells with at least 500 genes and 1,000 UMIs detected by either V2 or V3 chemistry. For the V2-V3 comparison in HTAPP-951-SMP-4652 (Extended Data Fig. 9), we used the same thresholds for both chemistries: at least 200 genes and 400 UMIs detected. For both data types, we filtered out those cells or nuclei where >20% of UMIs came from mitochondrial genes.

Replication

Each biological sample is unique to a patient due to tumor heterogeneity, and furthermore, tissue samples from the same patient tumor may have intra-tumor heterogeneity. All single-cell dissociation protocols and single-nuclei isolation methods that we recommend were tested on more than one patient tumor sample (biological replicate) and protocol performance was consistent across the different samples tested. For under-performing protocols, we generally do not include replicate samples.

Randomization

We do not have experimental groups.

Blinding

We do not have experimental groups.

Reporting for specific materials, systems and methods

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Materials & experimental systems

n/a

Involved in the study

☐ Antibodies
☐ Eukaryotic cell lines
☒ Palaeontology
☐ Animals and other organisms
☐ Human research participants
☒ Clinical data

Methods

n/a

Involved in the study

☒ ChIP-seq
☐ Flow cytometry
☒ MRI-based neuroimaging

Antibodies

Antibodies used

1) FITC anti-human CD45 Antibody, BioLegend #304006, https://www.biolegend.com/en-us/products/ficlt-anti-human-cd45-
antibody-707, Clone HI30, Lot #B226081, used at 1:200 dilution
2) CD45 MicroBeads, human, Miltenyi # 130-045-801, https://www.miltenyibiotec.com/US-en/products/macs-cell-separation/ cell-separation-reagents/microbeads-and-isolation-kits/tumor-cells/cd45-microbeads-human.html
3) CD326 (EpCAM)-PE, human, Miltenyi Biotech #130-113-264, https://www.miltenyibiotec.com/US-en/products/macs-flow-cytometry/antibodies/primary-antibodies/cd326-epcam-antibodies-human-hea-125-1-50.html#pe:for-100-tests, Clone HEA-125, Lot #5190328519, used at 1:50 dilution
4) APC anti-human CD14, BioLegend #367118, https://www.biolegend.com/nl-nl/products/apc-anti-human-cd14-antibody-12901, Clone 63D3, Lot #B262993, used at 1:20 dilution
5) PE-cy7 anti-human CD24, BioLegend #311120, https://www.biolegend.com/nl-nl/products/pe-cy7-anti-human-cd24-antibody-6126, Clone ML5, Lot #B226384, used at 1:20 dilution

**Validation**

All of the antibodies used in this study were validated for use in human specimens by the manufacturers, as indicated below:

FITC anti-human CD45 Antibody, BioLegend #304006:
Application: FC - Quality tested (FC: Flow cytometric analysis of antibody surface-stained cells.)
Recommended Usage: Each lot of this antibody is quality control tested by immunofluorescent staining with flow cytometric analysis.

CD326 (EpCAM)-PE, human, Miltenyi Biotech # 130-113-264:
Peripheral blood leukocytes mixed with cells from a breast cancer cell line (SK-BR-3) were stained with CD326 (EpCAM) antibodies and analyzed by flow cytometry using the MACSQuant® Analyzer.

APC anti-human CD14, BioLegend #367118:
Application: FC - Quality tested (FC: Flow cytometric analysis of antibody surface-stained cells.)
Recommended Usage: Each lot of this antibody is quality control tested by immunofluorescent staining with flow cytometric analysis.

PE-cy7 anti-human CD24, BioLegend #311120:
Application: FC - Quality tested (FC: Flow cytometric analysis of antibody surface-stained cells.)
Recommended Usage: Each lot of this antibody is quality control tested by immunofluorescent staining with flow cytometric analysis.

**Animals and other organisms**

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

**Laboratory animals**
The neuroblastoma O-PDX was propagated in one female nude adult athymic Foxn1-null mouse (Charles River Laboratories, strain code 553) via para-adrenal injection. At the time of injection, the mice are 6-8 weeks in age, and it takes 4-8 weeks for the O-PDX to grow.

**Wild animals**
This study did not involve wild animals.

**Field-collected samples**
This study did not involve field-collected samples.

**Ethics oversight**
Animal use was restricted to 1 female nude athymic mouse for para-adrenal injection of O-PDX cells. This study was carried out in strict accordance with the recommendations in the Guide to Care and Use of Laboratory Animals of the National Institute of Health. The protocol was approved by the Institutional Animal Care and Use Committee at St. Jude Children’s Research Hospital. All efforts were made to minimize suffering. All mice were housed in accordance with approved IACUC protocols. Animals were housed on a 12-12 light cycle (light on 6 am and off 6 pm) and provided food and water ad libitum. Athymic nude female mice were purchased from Charles River Laboratories (strain code 553).

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

**Human research participants**

**Policy information about studies involving human research participants**

**Population characteristics**
This research was not designed as a population study. Only a small number of samples (2-7) are profiled and analyzed per cancer type. Most samples are from adults, with the remaining samples being pediatric (pediatric high-grade glioma and neuroblastoma).

**Recruitment**
Patients were not actively recruited for this secondary-use study. Instead, patients were recruited under the initial IRB protocols approved by our collaborating institutions (see "Ethics oversight" section). External sample cohorts were then added to the Broad’s Molecular Classification of Cancer protocol (15-370B) and reviewed and approved by the Dana Farber Cancer Institute (DFCI) IRB. Patient population compositions are not expected to impact our results as our analyses were done on a per sample basis, rather than on patient populations.

**Ethics oversight**
Ethics oversight for the Molecular Classification of Cancer protocol (15-370B) is performed by the DFCI IRB. Samples added to this protocol also underwent IRB review and approval at the institutions where the samples were originally collected. Specifically, Dana-Farber Cancer Institute IRB approved the following protocols: lung cancer (IRB protocol 02-051), melanoma (IRB protocol 11-104), ovarian cancer (IRB protocol 02-051), melanoma (IRB protocol 11-104), sarcoma (IRB protocol 17-104), GBM (IRB protocol 10-417), and chronic lymphocytic leukemia (IRB protocol 99-224), and the St. Jude Children’s Research Hospital IRB approved the following protocol: pediatric high-grade
The XPD 09-234 MAST (Molecular Analysis of Solid Tumor) protocol for creating the neuroblastoma O-PDX sample was reviewed and approved by the St. Jude Children’s Research Hospital IRB.

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

**Flow Cytometry**

**Plots**

- Confirm that:
  - The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).
  - The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a 'group' is an analysis of identical markers).
  - All plots are contour plots with outliers or pseudocolor plots.
  - A numerical value for number of cells or percentage (with statistics) is provided.

**Methodology**

**Sample preparation**

For flow cytometry analysis of CD45+ depletion in the ovarian cancer ascites sample, cells were resuspended in PBS complemented with 2% fetal bovine serum and stained with FITC anti-human CD45 antibody (BioLegend #304006CD45, 1:200 dilution), PE anti-human EPCAM antibody (Miltenyi Biotech #130-113-264, 1:50 dilution), APC anti-human CD14 (BioLegend #367118, clone 63D3, 1:20 dilution), and PE-cy7 anti-human CD24 (BioLegend #311120, clone ML5, 1:20 dilution) for 20 minutes, and with 7-AAD (Invitrogen #A1310, 1:200 dilution) for 5 minutes. The same cells were also used for single-stain and unstained controls in order to perform compensation and adjust gating.

**Instrument**

BD LSRFortessa Cell Analyzer (Cat. No. 647177)

**Software**

BD FACSDiva Software Version 8.0.1; plots were generated with FlowJo Version 10.5.3

**Cell population abundance**

We used a CD45+ depletion strategy to prepare an ovarian ascites sample for scRNA-Seq. To assess how well our CD45+ depletion strategy worked, we took a sample of these prepared cells, with and without the CD45+ depletion, and performed flow cytometry. CD45- cells were enriched from 0.75% to 29.4% of the population, as determined using the anti-CD45 antibody. EpCAM+ cells were enriched from 0.17% to 4.9%, as determined by the PE anti-human EPCAM antibody.

**Gating strategy**

Cells were gated by FSC and SSC (35% of events retained for no depletion, 23% of events retained for depletion of CD45+ cells), doublets removed using FSC-A and FSC-H (100% singlets for no depletion, 99.8% singlets for depletion of CD45+ cells), live cells identified using 7-AAD (84.7% of cells retained for no depletion are live, 96.6% of cells retained for depletion of CD45+ cells are live), the distribution of immune and non-immune cells quantified using the CD45 antibody (99.3% of cells retained for no depletion are CD45+, 70.5% of cells retained for depletion of CD45+ cells are CD45+), and the distribution of EPCAM+ cells quantified using the EPCAM antibody (0.17% of the cells retained for no depletion are EPCAM+, 4.92% of cells retained for depletion of CD45+ cells are EPCAM+).

Tick this box to confirm that a figure exemplifying the gating strategy is provided in the Supplementary Information.