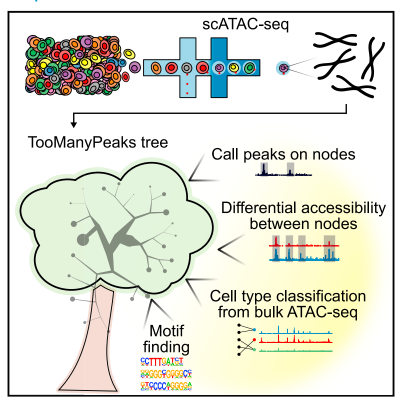
TooManyPeaks identifies drug-resistant-specific regulatory elements from single-cell leukemic epigenomes

Graphical abstract



Authors

Gregory W. Schwartz, Yeqiao Zhou, Jelena Petrovic, Warren S. Pear, Robert B. Faryabi

Correspondence

faryabi@pennmedicine.upenn.edu

In brief

Schwartz et al. present TooManyPeaks, a suite of algorithms to explore and visualize heterogeneity of regulatory elements by using single-cell ATAC-seq data. Using TooManyPeaks's functionalities, they find evidence that heterogeneity of the chromatin accessibility state contributes to the propensity of Notch-mutated T leukemic cells to develop resistance to Notch inhibitors.

Highlights

- TooManyPeaks identifies genomic element heterogeneity from single-cell ATAC-seq
- TooManyPeaks tree shows relationships among cells based on genomic elements
- Genomic element heterogeneity contributes to leukemia drug resistance
- Drug-naive leukemic cells exist with accessibility similar to that of resistant cells







Report

TooManyPeaks identifies drug-resistant-specific regulatory elements from single-cell leukemic epigenomes

Gregory W. Schwartz, 1,2,3 Yeqiao Zhou, 1,2,3 Jelena Petrovic, 1,2,3 Warren S. Pear, 1,3 and Robert B. Faryabi 1,2,3,4,*

- ¹Department of Pathology and Laboratory Medicine, University of Pennsylvania, Philadelphia, PA, USA
- ²Penn Epigenetics Institute, University of Pennsylvania, Philadelphia, PA, USA
- ³Abramson Family Cancer Research Institute, Perelman School of Medicine, University of Pennsylvania, Philadelphia, PA, USA ⁴I ead contact
- *Correspondence: faryabi@pennmedicine.upenn.edu https://doi.org/10.1016/j.celrep.2021.109575

SUMMARY

Emerging single-cell epigenomic assays are used to investigate the heterogeneity of chromatin activity and its function. However, identifying cells with distinct regulatory elements and clearly visualizing their relationships remains challenging. To this end, we introduce TooManyPeaks to address the need for the simultaneous study of chromatin state heterogeneity in both rare and abundant subpopulations. Our analyses of existing data from three widely used single-cell assays for transposase-accessible chromatin using sequencing (scATAC-seq) show the superior performance of TooManyPeaks in delineating and visualizing pure clusters of rare and abundant subpopulations. Furthermore, the application of TooManyPeaks to new scATAC-seq data from drug-naive and drug-resistant leukemic T cells clearly visualizes relationships among these cells and stratifies a rare "resistant-like" drug-naive sub-clone with distinct *cis*-regulatory elements.

INTRODUCTION

Cell-type-specific transcriptional diversity is largely set by the interactions between transcription factors and their cognate cisregulatory elements within accessible chromatin regions. The emergence of single-cell/single-nucleus assays for transposaseaccessible chromatin using sequencing (here, collectively called scATAC-seq) has enabled profiling of accessible cis-regulatory elements (here, interchangeably referred to as the epigenome) for thousands of individual cells. Unique characteristics of scATACseg readouts coupled with the increase in data volume have created a need for efficient computational tools for identifying and visualizing cells with similar chromatin accessibility, including rare populations. Although some scATAC-seq data analysis methods have been proposed, it still remains challenging to simultaneously identify and visualize rare and abundant subpopulations with distinct chromatin structures. To address this need, we introduce TooManyPeaks, which is equipped with several functionalities and provides a standalone end-to-end solution for scA-TAC-seq analysis. We assessed the accuracy and efficiency of TooManyPeaks in identifying and visualizing both rare and abundant populations by using several benchmarks. Given the key role of Notch signals in T cell acute lymphoblastic leukemia (T-ALL), we also used TooManyPeaks to investigate how heterogeneity of cis-regulatory elements influences divergent responses to Notch antagonist gamma-secretase inhibitor (GSI) in T-ALL. TooManyPeaks is open source and available through https:// github.com/faryabib/too-many-cells#too-many-peaks.

RESULTS

TooManyPeaks relates cells with distinct chromatin states

To identify and visualize cell subpopulations with distinct cisregulatory elements from scATAC-seq data, we introduce TooManyPeaks (Figure 1A). TooManyPeaks provides an end-toend solution for scATAC-seq data analysis from chromatin accessibility readouts to multi-scalar renderings of cell group relationships and is integrated into the TooManyCells suite (Schwartz et al., 2020), a platform originally built for single-cell RNA-seq (scRNA-seg) data analysis. To this end, TooManyPeaks implements a number of graph-based algorithms to extract distinct cis-regulatory elements of both rare and abundant subpopulations and creates cell clade relationships from scATAC-seq data (Figure 1A; see STAR Methods). These cell clades are represented by a nested cluster structure in which relationships among the groups are maintained. In contrast to single-resolution clustering algorithms commonly used for scATAC-seq analysis (Li et al., 2020; Pliner et al., 2018; Bravo González-Blas et al., 2019; Cusanovich et al., 2018; Danese et al., 2019; Stuart et al., 2020; Fang et al., 2021), each inner node of the TooManyPeaks output is a cluster at a given resolution and a leaf node is a finer-grain cluster for which any additional partitioning would be as informative as randomly separating the cells (see STAR Methods).

The TooManyPeaks tree-based visualization offers several advantages over "flat" two-dimensional portrayals of data provided by projection-based methods such as t-distributed stochastic





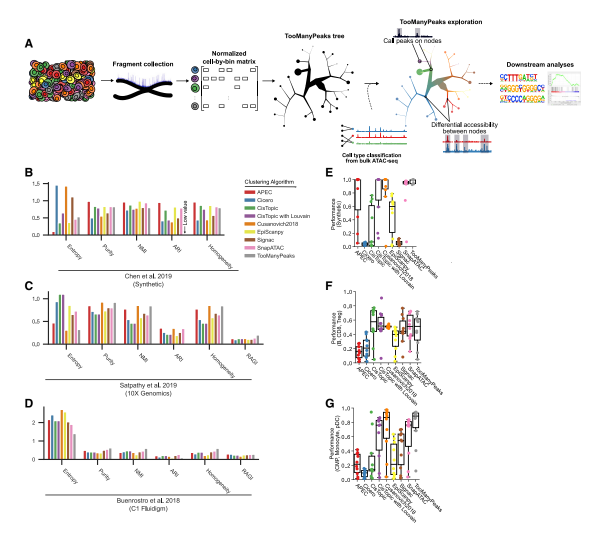


Figure 1. TooManyPeaks overview and performance comparison

(A) Graphical representation of the TooManyPeaks algorithm. Following the arrows from left to right, TooManyPeaks converts scATAC-seq data to a cell-by-bin matrix, binarizes each value (accessible or inaccessible), and identifies and visualizes cell clade relationships by using matrix-free divisive hierarchical spectral clustering (see STAR Methods). TooManyPeaks trees are interpreted by following the cell groups from the root (the largest inner node) to the leaves. A leaf node here is shown as a pie chart of its cell composition. The sizes of a leaf and branches are proportional to the number of cells in the node. TooManyPeaks may then perform several downstream analyses.

(B-D) Clustering benchmarks with, from left to right, lower entropy, higher purity, higher normalized mutual information (NMI), higher adjusted Rand index (ARI), higher homogeneity, and higher residual average Gini index (RAGI; not applicable to synthetic data) representing more accurate clustering of simulated bone marrow cells with a moderate noise level of 0.2 (Chen et al., 2019) (B), CD34⁺ hematopoietic progenitor cells profiled using 10x Genomics (n = 7,771 cells) (Satpathy et al., 2019) (C), or Fluidigm C1 (n = 2,954 cells) (Buenrostro et al., 2018) (D).

(E-G) Detection of cells from two "rare" populations mixed with a "common" population was benchmarked. Box-and-whisker plots quantifying the accuracy of rare population detection in controlled admixtures from various datasets (m = 10 admixtures), as follows: n = 1,000 synthetic cells generated by simATAC (Navidi et al., 2021) (E); n = 1,000 B (common), CD8⁺ T ("rare1") and Treg cells ("rare2") (Satpathy et al., 2019) (F); and n = 500 common myeloid progenitors (CMPs) (common), monocytes (rare1), and plasmacytoid dendritic cells (pDC) (rare2) (Buenrostro et al., 2018) (G). Each point represents the average performance of 10 experiments from an admixture (100 admixtures overall). Performance indicates (true rare pairs (cells from the same rare population in the same cluster)/total rare pairs (true rare pairs and cells from different rare populations)). Box-and-whisker plots represent the following: center line, median; box limits, upper (75th) and lower (25th) percentiles; whiskers, 1.5 × interquartile range; points, outliers. See also Figure S1.

neighbor embedding (t-SNE) and uniform manifold approximation and projection (UMAP) (van der Maaten and Hinton, 2008; McInnes et al., 2018). Although frequently used, projection-based methods generally do not report quantitative inter-cluster relationships and lack interpretable visualizations across clustering resolutions (Kobak and Linderman, 2021). To complement these existing single-resolution visualization methods and enable multi-resolution scATAC-seq data exploration, TooManyPeaks provides a fully customizable dendrogram for the visualization of inter-cluster relationships. To facilitate data exploration, we included many features in the TooManyPeaks visualization output including, but not limited to, branch scaling, weighted-average

Report



color blending, and statistically driven tree pruning. TooMany-Peaks can also display outputs of other scATAC-seq clustering algorithms to quantify the relationships among their identified cell populations.

To enable an end-to-end built-in scATAC-seq analysis solution, TooManyPeaks provides several specialized and commonly used functionalities for scATAC-seq data analysis. For example, TooManyPeaks provides an algorithm for celltype annotation based on input reference cis-regulatory elements of fluorescence-activated cell sorting (FACS)-purified cells. TooManyPeaks can also perform cluster- and cell-labelspecific peak calling, as well as differential accessibility analyses across various clustering resolutions. Furthermore, TooMany-Peaks enables several downstream analyses by generating normalized genome browser tracks and incorporating motif analysis methods (Heinz et al., 2010; Bailey et al., 2009) for each population. All TooManyPeaks functionalities can be readily set through its command-line interface (see STAR Methods).

TooManyPeaks accurately segregates and clearly visualizes rare cells

To assess the accuracy of cell clustering, we compared the outputs of TooManyPeaks and seven commonly used scATAC-seq clustering methods, as follows: APEC (Li et al., 2020), Cicero (Pliner et al., 2018), CisTopic (Bravo González-Blas et al., 2019), Cusanovich2018 (Cusanovich et al., 2018), EpiScanpy (Danese et al., 2019), Signac (Stuart et al., 2020), and SnapATAC (Fang et al., 2021). Importantly, these methods use different combinations of features for cell clustering. TooManyPeaks and Cusanovich2018 use latent semantic analysis (LSA) (Deerwester et al., 1990) for producing features in lower dimensional space, whereas CisTopic uses latent Dirichlet allocation (LDA) to identify "topics" as a form of feature definition (Falush et al., 2003). In contrast, Cicero and APEC summarize scATAC-seq signals into gene activity scores and "accessons," respectively. As Cis-Topic recommends density peak clustering but other selected algorithms use Louvain clustering, we also included CisTopic topics as features for Louvain clustering (referred to as CisTopic with Louvain) in our comparative analysis (see STAR Methods).

To assess the performance of each method in identifying homogeneous cell label clusters, we used purity (Manning et al., 2008), entropy (Tan et al., 2019), mutual information (Kvålseth, 2017), adjusted rand index (ARI), homogeneity (Rosenberg and Hirschberg, 2007), and residual average Gini index (RAGI) (Chen et al., 2019). More homogeneous clusters result in higher purity, normalized mutual information (NMI), homogeneity, and RAGI, as well as lower entropy (see STAR Methods). We compared the ability of each algorithm to identify pure cell clusters of synthetic data (Chen et al., 2019; Figure 1B) and phenotypically defined cells within bone marrow and blood samples profiled using 10x Genomics (Satpathy et al., 2019; Figure 1C) or Fluidigm C1 (Buenrostro et al., 2018; Figure 1D) scATACseq platforms. We chose a 5-kb genomic bin size and 50 LSA dimension due to the low variability of performance across parameter choices (Figures S1A and S1B). As expected, TooManyPeaks resulted in low ARI, a measure that is sensitive to true label uncertainty and is biased against multi-resolution clustering

methods (Chen et al., 2019). Nevertheless, TooManyPeaks, Cusanovich2018, and SnapATAC generated the purest clusters in both synthetic and complex real datasets included in this analysis (Figures 1B-1D). Together, these comprehensive analyses indicated the advantage of using TooManyPeaks for clustering individual cells based on their chromatin state, while maintaining their multi-scalar relationships in highly diverse hematopoietic

Although some clustering methods provide resolution parameters to focus on small or large populations, concurrent identification and visualization of rare and abundant cells from scATAC-seq data remain challenging (Lancichinetti and Fortunato, 2011; Fang et al., 2021). Previous clustering benchmarks (Figures 1C and 1D) measured the diversity of cell labels within clusters, yet they did not directly quantify an algorithm's ability to detect rare subpopulations with distinct regulatory elements. To rigorously assess the ability of various scATAC-seq clustering algorithms to simultaneously identify rare and abundant cells, we adapted our previous scRNA-seq rare population benchmark (Schwartz et al., 2020) to scATAC-seq. We used synthetic data (Navidi et al., 2021), 10x Genomics (Satpathy et al., 2019), and Fluidigm C1 (Buenrostro et al., 2018) scATAC-seq datasets and generated several controlled cell admixtures with various ratios of one "common" and two equally abundant "rare" populations. We then assessed how each algorithm separated the two rare populations from each other and from the common population in 10 controlled cell admixtures with various levels of rare populations. We found that TooManyPeaks outperformed all other tested algorithms or tied with SnapATAC in recovering rare populations (Figures 1E-1G).

Feature choice can significantly affect scATAC-seq analysis outputs. Given that several genomic elements could be involved in the regulation of a gene, scATAC-seq data have orders of magnitude more features than scRNA-seq and cannot necessarily be collapsed to the resolution of genes. As such, genomic bins or peaks, defined as equal-sized genomic windows or loci with enriched accessibility in pseudo-bulk ATAC-seq, respectively, are commonly used as scATAC-seq analysis features (Chen et al., 2019). Alternatively, some algorithms use topic and gene activity features. TooManyPeaks can readily compute on all four types of features (Figures S1C-S1H). More importantly, our data revealed that TooManyPeaks, Cusanovich2018, and SnapATAC, which are algorithms that use genomic bin features, show superior performance in detecting rare cells compared to algorithms using peaks or other features (Figures 1F and 1G). Timing benchmark on a set of 2,954 cells (Figure S1I) further showed that TooManyPeaks operates at a comparable rate or faster than other algorithms, even with its multi-resolution output.

TooManyPeaks classifies and relates cells from mouse bone marrow and spleen

Cell-type classification is one of the major applications of scA-TAC-seq analysis. To this end, we equipped TooManyPeaks with functionality to annotate individual cells based on input reference cis-regulatory element sets, including those from FACS-sorted bulk ATAC-seq data. Briefly, TooManyPeaks implements a fast bipartite-graph algorithm using cosine similarity



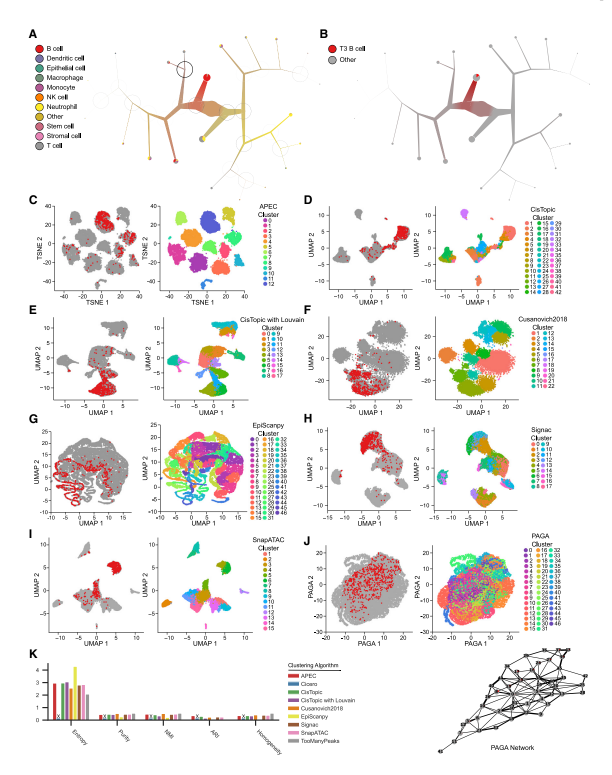


Figure 2. Stratification and annotation of murine bone marrow and spleen cells

(A) The TooManyPeaks algorithm for cell-type annotation based on input reference cis-regulatory elements is used to predict the cell types in mouse bone marrow and spleen (n = 16,749 cells) (Cusanovich et al., 2018). Reference cis-regulatory elements of 92 phenotypically defined progenitor and differentiated hematopoietic cell types are generated from the analyses of bulk ATAC-seq in FACS-sorted cells (Yoshida et al., 2019). A TooManyPeaks tree pruned at median(modularity) + 15 × MAD (modularity) threshold shows major hematopoietic lineages. At each bipartitioning, a darker circle circumference represents higher modularity.

(legend continued on next page)

Report



to assign each cell to one of the reference cell types with a known cis-regulatory element repertoire (see STAR Methods and Figure 1A). To assess the efficacy of the TooManyPeaks cell-type classification, we annotated murine bone marrow and spleen cells (Cusanovich et al., 2018) based on reference cis-regulatory elements defined by bulk ATAC-seq analysis of 92 FACS-purified progenitor and differentiated hematopoietic cells (Yoshida et al., 2019). Visual inspection of the TooManyPeaks tree showed general separation of major phenotypically defined hematopoietic cell types (e.g., B cells segregate into a single branch) without or with modularity-guided pruning (Schwartz et al., 2020; Figures S2 and 2A).

To further inspect the localization of more refined cellular subtypes, we next overlayed the positions of late transitional T3 B cells on the TooManyPeaks tree and projection outputs of all other algorithms included in this analysis including PAGA, which attempts to conserve and display global topology by using a network (Wolf et al., 2019; Figures 2B-2J). Interestingly, Cicero failed to complete the analysis of 16,749 cells. T3 B cells were mostly compartmentalized within a single TooManyPeaks tree branch (Figure 2B), whereas they were spread across the projection plots (Figures 2C-2J, left panels) and separated into multiple clusters (Figures 2C–2J, right panels) with the other algorithms. Notably, T3 B cells were spread out over 13 nodes of the PAGA network (Figure 2J). Furthermore, quantitative assessment of cell-type classification based on reference cis-regulatory elements of hematopoietic cells showed the improved performance of TooManyPeaks compared to all the other algorithms in accurately detecting (Figure 2K) and clearly visualizing (Figure S3) 92 distinct cell types in murine bone marrow and spleen.

Similar to the T3 B cell analysis of single-cell combinatorial indexing ATAC-seq (sciATAC-seq) data (Cusanovich et al., 2018; Figure 2), human hematopoietic stem cells (HSCs) profiled with the Fluidigm C1 platform (Buenrostro et al., 2018) were clearly distinguishable within the TooManyPeaks tree, but not in the projection plots of other algorithms, and the PAGA network (Figure S4).

TooManyPeaks determines the unique chromatin state of GSI-"resistant-like" drug-naive T-ALL cells

Notch mutations are observed in nearly 60% of patients with T-ALL and correlate with poor prognosis (Marks et al., 2009). These observations provide a compelling rationale for focusing on Notch signaling antagonists, such as GSI, as targeted therapies for Notch-mutated T-ALL. Nevertheless, progress toward targeted treatment of Notch-mutated T-ALL has been stymied partly due to a limited understanding of GSI-resistance acquisition. To investigate the underlying mechanisms of GSI resistance, we selected for GSI-resistant T-ALL cells by prolonged treatment of parental NOTCH1-mutated DND-41 cells with a high GSI dose (Schwartz et al., 2020). Given the genetic homogeneity of DND-41 cells and results of earlier studies showing the reversibility of the GSI resistance phenotype (Knoechel et al., 2014), we hypothesized that epigenetic differences contribute to the divergence of parental cells with resistant-like regulatory programs from non-resistant-like parental cells.

To test this hypothesis, we measured the accessibility of chromatin in 7,989 parental and GSI-resistant DND-41 cells. TooMany-Peaks revealed that although parental cells are largely segregated from resistant cells, a rare resistant-like subpopulation of 144 parental cells had a chromatin state similar to that of the GSI-resistant cells (Figures 3A and S5A). Analyses with other selected tools (Figures S5B-S5E and S6A-S6E) showed resistant-like from nonresistant-like parental cells were separated partially. Nevertheless, the flat outputs of these algorithms generally obscured full separation of resistant-like cells from non-resistant-like parental cells. In contrast, the TooManyPeaks tree immediately rendered the relationship between resistant-like parental and resistant cells and clearly placed them within the resistant-cell-dominant subtree (Figure 3A).

To gain insights into transcriptional regulatory programs conferring resistance to GSI, we used TooManyPeaks to directly compare the chromatin accessibility of resistant-like and nonresistant-like parental cells. We identified 28,593 genomic elements with significantly higher accessibility in resistant-like cells (q < 0.05; see STAR Methods), which were collectively enriched with motifs associated with transcription factors with known functions in T cell development, transformation, and malignancies, such as GATA3, RUNX1, and MYC (Figure S7A; Table S1). Integration of scATAC-seq and scRNA-seq data (Table S1) further revealed that MYC had both significantly elevated expression (Figure S7B; Table S1) and higher accessible consensus binding sequences in the resistant-like parental cells (Figure S7C).

Guided by the differential activity of MYC in resistant-like cells, we used TooManyPeaks to map putative MYC regulatory elements in GSI-resistant and non-resistant-like parental cells. Concordant with transcriptional levels, the MYC promoter was active in both non-resistant-like parental and GSI-resistant cells (Figures 3B and 3C; Table S2). Our scATAC-seq data of nonresistant-like parental cells delineated clusters of accessible elements within \sim 2-Mb region 3' of the MYC promoter (Figure 3B). Importantly, we observed marked differences in accessibility of three chromatin regions flanking the MYC promoter when comparing non-resistant-like parental and GSI-resistant cells (Figure 3B). Accessibility of genomic element E1 (~1.42-Mb 3' of the MYC promoter), and E2 (~1.5-Mb 3' of the MYC promoter and proximal to the long non-protein coding gene LINC00977) were significantly (p < 0.05 and q <0.05) reduced in the GSIresistant cells (Figures 3B, 3D, 3E, 3G, and S7D; Table S2; E1: $log_2FC = -1.74$ and E2: $log_2FC = -2.78$). In contrast, genomic element cluster E3 (~1.85-Mb 3' of the MYC promoter) significantly gained accessibility in the GSI-resistant cells (Figures 3B, 3F, 3G, and S7D; Table S2; log₂FC = 0.924). Together, this scATAC-seq analysis revealed significant chromatin restructuring of the MYC locus during GSI resistance development.

(B-J) TooManyPeaks tree (B) and UMAP outputs (C-J) colored by T3 B cells (red, left) or cluster label (right) generated by the noted algorithms. (K) Clustering benchmarks with, from left to right, lower entropy, higher purity, higher NMI, higher ARI, and higher homogeneity showing more accurate clustering of phenotypically defined progenitor and differentiated hematopoietic cell types in mouse bone marrow and spleen by TooManyPeaks. An "X" marks algorithms that failed to complete. See also Figures S2, S3, and S4.



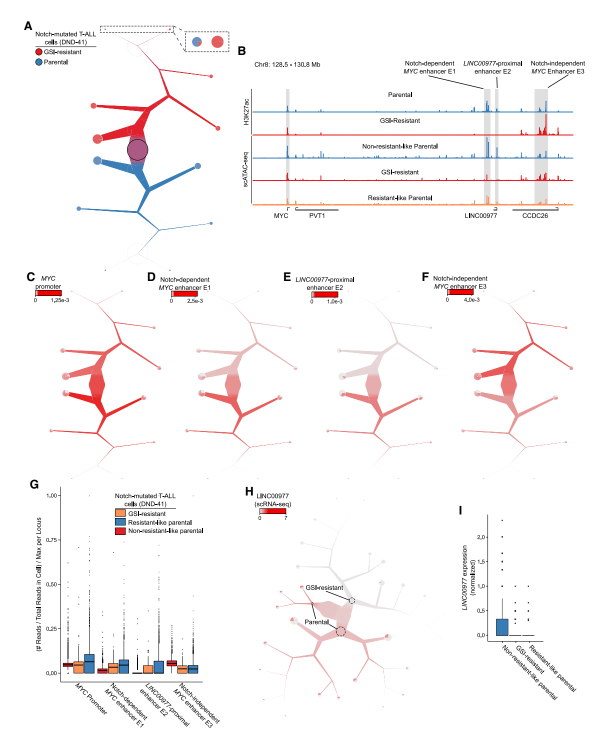


Figure 3. TooManyPeaks identifies genomic elements specific to resistant-like parental T-ALL cells

(A) TooManyPeaks tree of parental (n = 3,831 cells) and GSI-resistant (n = 4,158 cells) DND-41 T-ALL cells showing a resistant-like parental subpopulation of n = 144 cells.

(B) Genome tracks highlight key genomic elements at the MYC locus from 5' to 3', as follows: MYC promoter, Notch-dependent MYC enhancer E1, LINC00977proximal enhancer E2, and Notch-independent MYC enhancer E3. The top two and bottom two tracks show H3K27ac and aggregated scATAC-seq of DND-41 populations in (A), respectively.

(C-F) TooManyPeaks tree as in (A) showing the accessibility of the MYC promotor (C) and enhancers E1 (D), E2 (E), and E3 (F).

(legend continued on next page)

Report



To further elucidate the function of differentially accessible elements at the MYC locus, we complemented our single-cell measurements with chromatin immunoprecipitation sequencing (ChIP-seq) analysis of enhancer histone mark H3K27ac. In concordance with the scATAC-seq data (Figures 3B and 3G), we recapitulated the loss of activity at E1 and E2 and gain of activity in E3 in GSI-resistant cells (Figure 3B). Interestingly, earlier studies showed that although genomic element E1 binds the Notch transcription complex and functions as a Notch-dependent MYC enhancer, E2 does not bind Notch and functions as a Notch-independent MYC enhancer (Yashiro-Ohtani et al., 2014; Herranz et al., 2014; Shi et al., 2013). Together, our bulk ChIP-seq analysis confirmed our scATAC-seq results and further showed differential activity of Notch-dependent and Notch-independent MYC distal enhancers E1 and E3, as well as uncharacterized LINC00977-proximal putative MYC enhancer E2, in GSI-sensitive and GSI-resistant DND-41 cells.

To more directly test whether chromatin accessibility differences in drug-naive cells contribute to the GSI-resistant phenotype, we next benefited from our scATAC-seg data to identify potential chromatin changes underpinning differential MYC expression in the resistant-like compared to non-resistant-like parental cells (Figure S7B). Notch-independent MYC enhancer E3 was similarly accessible in the resistant-like and non-resistant-like parental cells (Figures 3B and 3G; Table S3). Similarly, the accessibility of Notch-dependent MYC enhancer E1 was comparable in these two subpopulations of parental cells (Figures 3B and 3G; Table S3; $\log_2 FC = -0.316$, $q = 2.65 \times$ 10⁻³). In contrast, enhancer E2 accessibility was markedly different between these two parental subpopulations (Figures 3B and 3G). Similar to GSI-resistant cells, enhancer E2 was significantly less accessible in resistant-like than in non-resistant-like subpopulation of parental cells (Figures 3B and 3G; Table S3; $\log_2 FC = -0.802$, q = 0.0286). To assess if the loss of enhancer E2 accessibility may further affect LINC00977 expression, we used TooManyCells (Schwartz et al., 2020) to quantify LINC00977 transcript levels in 7,371 parental and resistant cells (Figures 3H and 3I). This scRNA-seq analysis revealed that LINC00977 expression was markedly lower in the GSI-resistant cells than in non-resistant-like parental cells (Figures 3H and 3I; Table S4; $\log_2 FC = -1.84$, $q < 2.22 \times 10^{-16}$). Notably, in concordance with enhancer E2 accessibility loss (Figures 3B and 3G), we also observed reduced LINC00977 expression in resistant-like compared to non-resistant-like parental cells (Figures 3H and 3I, and Table S1; $\log_2 FC = -0.568$, p = 0.117).

To further elucidate the underlying mechanisms of differential LINC00977-proximal enhancer E2 activity in the two parental sub-populations, we used motif search to explore transcription factors that potentially bound enhancer E2 in non-resistant-like but not resistant-like parental cells (Figure S7E; Table S5). These data revealed the presence of consensus binding motifs of TCF high-mobility group (HMG) family of proteins in the sequences of enhancer E2. Notably, scRNA-seq data showed significant downregulation of TCF-7, the gene encoding for T cell-lineage determinant factor TCF-1 (Johnson et al., 2018), in both GSIresistant and resistant-like parental compared to non-resistantlike parental cells (Figures S7E and S7F; $\log_2 FC = -2.63$, q =9.30×10⁻⁵). Together, these data suggest that in addition to MYC, differential activity of TCF-1 and its cognate regulatory elements such as LINC00977-proximal enhancer E2 may play a role in setting disparate epigenetic transcriptional regulatory programs in resistant-like and non-resistant-like parental subpopulations.

DISCUSSION

We developed TooManyPeaks, which provides complementary algorithms for clustering and visualizing scATAC-seq data. Too-ManyPeaks visualization and clustering are fundamentally different from projection-based visualization and single-resolution clustering. In addition to various visualization features, Too-ManyPeaks provides other capabilities including, but not limited to, flexible genomic feature options and cell type classification based on reference cis-regulatory elements. To enhance usability, TooManyPeaks is extensively documented (https://github. com/faryabib/too-many-cells#too-many-peaks) and is available as an easy-to-install standalone program through Nix or Docker.

Using the unique capabilities of TooManyPeaks, we identified a rare resistant-like population of Notch-mutated T-ALL DND-41 cells with chromatin accessibility more similar to GSI-resistant cells than non-resistant-like parental cells. Our new scATACseq data also suggested regulatory element markers of cells with a propensity for developing GSI resistance and signify potential transcription factor drivers of the resistance phenotype.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
 - Lead contact
 - Materials availability
 - Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - GSI-resistant T-ALL cell culture
- METHOD DETAILS
 - GSI-resistant T-ALL single-cell ATAC-sequencing
 - H3K27ac ChIP-seq
- QUANTIFICATION AND STATISTICAL ANALYSIS
 - TooManyPeaks analysis of scATAC-seq data
 - Accessibility matrix
 - Tree of single-cell clades
 - Peak calling and downstream analyses
 - Classification based on reference elements

⁽G) Box-and-whisker plot showing normalized accessibility at each locus in (B) for each population from (A).

⁽H) TooManyCells tree of gene expression showing elevated LINC00977 levels in the parental population (n = 7,371 cells).

⁽I) Box-and-whisker plot quantifying upper-quartile-normalized LINC00977 expression in each population from (H). See also Figures S5, S6, and S7 and Tables S1, S2, S3, S4, and S5.





- Clustering benchmarks
- Rare population benchmarks
- Timing benchmark
- T-ALL scATAC-seq statistical analyses
- T-ALL scATAC-seq motif analyses
- Statistical parameter definitions

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j. celrep.2021.109575.

ACKNOWLEDGMENTS

We thank Golnaz Vahedi and her laboratory for their support. This work was supported by R01-CA230800 and Susan G. Komen CCR185472448 (to R.B.F.), R01-CA215518 (to W.S.P.), and T32-CA009140 (to G.W.S.).

AUTHORS CONTRIBUTIONS

Conceptualization: R.B.F. and G.W.S.; methodology: G.W.S. and R.B.F.; software: G.W.S.; investigation: G.W.S., R.B.F., and Y.Z.; formal analysis: G.W.S., R.B.F., J.P., and Y.Z.; resources and reagents: R.B.F. and W.S.P.; writing-review & editing: G.W.S., W.S.P., R.B.F., Y.Z., and J.P.; writing-original draft: G.W.S. and R.B.F.; supervision: R.B.F.; funding acquisition: R.B.F.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: August 31, 2020 Revised: March 30, 2021 Accepted: July 29, 2021 Published: August 24, 2021

REFERENCES

Amemiya, H.M., Kundaje, A., and Boyle, A.P. (2019). The ENCODE Blacklist: Identification of Problematic Regions of the Genome. Sci. Rep. 9, 9354.

Bailey, T.L., Boden, M., Buske, F.A., Frith, M., Grant, C.E., Clementi, L., Ren, J., Li, W.W., and Noble, W.S. (2009). MEME SUITE: tools for motif discovery and searching. Nucleic Acids Res. 37, W202-W208.

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: A practical and powerful approach to multiple testing. J. R. Stat. Soc. Series B Stat. Methodol. 57, 289-300.

Bravo González-Blas, C., Minnoye, L., Papasokrati, D., Aibar, S., Hulselmans, G., Christiaens, V., Davie, K., Wouters, J., and Aerts, S. (2019). cisTopic: cisregulatory topic modeling on single-cell ATAC-seq data. Nat. Methods 16,

Buenrostro, J.D., Corces, M.R., Lareau, C.A., Wu, B., Schep, A.N., Aryee, M.J., Majeti, R., Chang, H.Y., and Greenleaf, W.J. (2018). Integrated Single-Cell Analysis Maps the Continuous Regulatory Landscape of Human Hematopoietic Differentiation. Cell 173, 1535-1548,e16.

Butler, A., Hoffman, P., Smibert, P., Papalexi, E., and Satija, R. (2018). Integrating single-cell transcriptomic data across different conditions, technologies, and species. Nat. Biotechnol. 36, 411-420.

Chen, H., Lareau, C., Andreani, T., Vinyard, M.E., Garcia, S.P., Clement, K., Andrade-Navarro, M.A., Buenrostro, J.D., and Pinello, L. (2019). Assessment of computational methods for the analysis of single-cell ATAC-seq data. Genome Biol. 20, 241.

Cusanovich, D.A., Hill, A.J., Aghamirzaie, D., Daza, R.M., Pliner, H.A., Berletch, J.B., Filippova, G.N., Huang, X., Christiansen, L., DeWitt, W.S., et al. (2018). A Single-Cell Atlas of In Vivo Mammalian Chromatin Accessibility. Cell 174, 1309-1324.e18.

Danese, A., Richter, M.L., Fischer, D.S., Theis, F.J., and Colomé-Tatché, M. (2019). EpiScanpy: Integrated single-cell epigenomic analysis. bioRxiv. https://doi.org/10.1101/648097.

Deerwester, S., Dumais, S.T., Furnas, G.W., Landauer, T.K., and Harshman, R. (1990). Indexing by latent semantic analysis. J. Am. Soc. Inf. Sci. 41, 391–407.

Falush, D., Stephens, M., and Pritchard, J.K. (2003). Inference of population structure using multilocus genotype data: linked loci and correlated allele frequencies. Genetics 164, 1567-1587.

Fang, R., Preissl, S., Li, Y., Hou, X., Lucero, J., Wang, X., Motamedi, A., Shiau, A.K., Zhou, X., Xie, F., et al. (2021). Comprehensive analysis of single cell ATAC-seq data with SnapATAC. Nat. Commun. 12, 1337.

Gini, C. (1997). Concentration and dependency ratios. Riv. Polit. Econ. 87, 769-792.

Heinz, S., Benner, C., Spann, N., Bertolino, E., Lin, Y.C., Laslo, P., Cheng, J.X., Murre, C., Singh, H., and Glass, C.K. (2010). Simple combinations of lineagedetermining transcription factors prime cis-regulatory elements required for macrophage and B cell identities. Mol. Cell 38, 576-589.

Herranz, D., Ambesi-Impiombato, A., Palomero, T., Schnell, S.A., Belver, L., Wendorff, A.A., Xu, L., Castillo-Martin, M., Llobet-Navás, D., Cordon-Cardo, C., et al. (2014). A NOTCH1-driven MYC enhancer promotes T cell development, transformation and acute lymphoblastic leukemia. Nat. Med. 20, 1130-1137.

Hubert, L., and Arabie, P. (1985). Comparing partitions. J. Classif. 2, 193-218.

Johnson, J.L., Georgakilas, G., Petrovic, J., Kurachi, M., Cai, S., Harly, C., Pear, W.S., Bhandoola, A., Wherry, E.J., and Vahedi, G. (2018). Lineage-Determining Transcription Factor TCF-1 Initiates the Epigenetic Identity of T Cells. Immunity 48, 243–257.e10.

Kent, W.J., Zweig, A.S., Barber, G., Hinrichs, A.S., and Karolchik, D. (2010). BigWig and BigBed: enabling browsing of large distributed datasets. Bioinformatics 26, 2204-2207.

Knoechel, B., Roderick, J.E., Williamson, K.E., Zhu, J., Lohr, J.G., Cotton, M.J., Gillespie, S.M., Fernandez, D., Ku, M., Wang, H., et al. (2014). An epigenetic mechanism of resistance to targeted therapy in T cell acute lymphoblastic leukemia. Nat. Genet. 46, 364-370.

Kobak, D., and Linderman, G.C. (2021). Initialization is critical for preserving global data structure in both t-SNE and UMAP. Nat. Biotechnol. 39, 156-157.

Kvålseth, T.O. (2017). On normalized mutual information: Measure derivations and properties. Entropy (Basel) 19, 1-14.

 $Lancichinetti, A., and \ Fortunato, S. \ (2011). \ Limits of modularity \ maximization \ in$ community detection. Phys. Rev. E Stat. Nonlin. Soft Matter Phys. 84, 066122.

Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25, 1754-1760.

Li, B., Li, Y., Li, K., Zhu, L., Yu, Q., Cai, P., Fang, J., Zhang, W., Du, P., Jiang, C., et al. (2020). APEC: an accesson-based method for single-cell chromatin accessibility analysis. Genome Biol. 21, 116.

Manning, C.D., Raghavan, P., and Schütze, H. (2008). Introduction to Information Retrieval (Cambridge University Press).

Marks, D.I., Paietta, E.M., Moorman, A.V., Richards, S.M., Buck, G., DeWald, G., Ferrando, A., Fielding, A.K., Goldstone, A.H., Ketterling, R.P., et al. (2009). T-cell acute lymphoblastic leukemia in adults: clinical features, immunophenotype, cytogenetics, and outcome from the large randomized prospective trial (UKALL XII/ECOG 2993). Blood 114, 5136-5145.

McInnes, L., Healy, J., and Melville, J. (2018). Umap: Uniform manifold approximation and projection for dimension reduction. arXiv, 1802.03426. https:// arxiv.org/abs/1802.03426.

Navidi, Z., Zhang, L., and Wang, B. (2021). simatac: a single-cell atac-seq simulation framework. bioRxiv. https://doi.org/10.1101/2020.08.14.251488.

Newman, M.E.J., and Girvan, M. (2004). Finding and evaluating community structure in networks. Phys. Rev. E Stat. Nonlin. Soft Matter Phys. 69, 026113.

Petrovic, J., Zhou, Y., Fasolino, M., Goldman, N., Schwartz, G.W., Mumbach, M.R., Nguyen, S.C., Rome, K.S., Sela, Y., Zapataro, Z., et al. (2019).

Report



Oncogenic Notch Promotes Long-Range Regulatory Interactions within Hyperconnected 3D Cliques. Mol. Cell 73, 1174-1190.e12.

Pliner, H.A., Packer, J.S., McFaline-Figueroa, J.L., Cusanovich, D.A., Daza, R.M., Aghamirzaie, D., Srivatsan, S., Qiu, X., Jackson, D., Minkina, A., et al. (2018). Cicero Predicts cis-Regulatory DNA Interactions from Single-Cell Chromatin Accessibility Data. Mol. Cell 71, 858-871.e8.

Quinlan, A.R., and Hall, I.M. (2010). BEDTools: a flexible suite of utilities for comparing genomic features. Bioinformatics 26, 841-842.

Robinson, M.D., McCarthy, D.J., and Smyth, G.K. (2010). edgeR: a Bioconductor package for differential expression analysis of digital gene expression data. Bioinformatics 26, 139-140.

Rosenberg, A., and Hirschberg, J. (2007). V-measure: A conditional entropybased external cluster evaluation measure. In Proceedings of the 2007 Joint Conference on Empirical Methods in Natural Language Processing and Computational Natural Language Learning (EMNLP-CoNLL), Association for Computational Linguistics, pp. 410-420.

Sandelin, A., Alkema, W., Engström, P., Wasserman, W.W., and Lenhard, B. (2004). JASPAR: an open-access database for eukaryotic transcription factor binding profiles. Nucleic Acids Res. 32, D91-D94.

Satpathy, A.T., Granja, J.M., Yost, K.E., Qi, Y., Meschi, F., McDermott, G.P., Olsen, B.N., Mumbach, M.R., Pierce, S.E., Corces, M.R., et al. (2019). Massively parallel single-cell chromatin landscapes of human immune cell development and intratumoral T cell exhaustion. Nat. Biotechnol. 37, 925-936.

Schwartz, G.W., Zhou, Y., Petrovic, J., Fasolino, M., Xu, L., Shaffer, S.M., Pear, W.S., Vahedi, G., and Faryabi, R.B. (2020). TooManyCells identifies and visualizes relationships of single-cell clades. Nat. Methods 17, 405-413.

Shannon, C.E. (1948). A Mathematical Theory of Communication. Bell Syst. Tech. J. 27, 623-656.

Shi, J., Whyte, W.A., Zepeda-Mendoza, C.J., Milazzo, J.P., Shen, C., Roe, J.-S., Minder, J.L., Mercan, F., Wang, E., Eckersley-Maslin, M.A., et al. (2013).

Role of swi/snf in acute leukemia maintenance and enhancer-mediated myc regulation, Genes Dev. 27, 2648-2662.

Stuart, T., Srivastava, A., Lareau, C., and Satija, R. (2020). Multimodal singlecell chromatin analysis with signac. bioRxiv. https://doi.org/10.1101/2020.11. 09.373613

Tan, P.-N., Steinbach, M., Karpatne, A., and Kumar, V. (2019). Introduction to Data Mining, second edition (Pearson).

van der Maaten, L., and Hinton, G. (2008). Visualizing data using t-sne. J. Mach. Learn. Res. 9, 2579-2605.

Wolf, F.A., Hamey, F.K., Plass, M., Solana, J., Dahlin, J.S., Göttgens, B., Rajewsky, N., Simon, L., and Theis, F.J. (2019). PAGA: graph abstraction reconciles clustering with trajectory inference through a topology preserving map of single cells. Genome Biol. 20, 59.

Yashiro-Ohtani, Y., Wang, H., Zang, C., Arnett, K.L., Bailis, W., Ho, Y., Knoechel, B., Lanauze, C., Louis, L., Forsyth, K.S., et al. (2014). Long-range enhancer activity determines Myc sensitivity to Notch inhibitors in T cell leukemia. Proc. Natl. Acad. Sci. USA 111, E4946-E4953.

Yoshida, H., Lareau, C.A., Ramirez, R.N., Rose, S.A., Maier, B., Wroblewska, A., Desland, F., Chudnovskiy, A., Mortha, A., Dominguez, C., et al.; Immunological Genome Project (2019). The cis-Regulatory Atlas of the Mouse Immune System. Cell 176, 897-912.e20.

Zhang, Y., Liu, T., Meyer, C.A., Eeckhoute, J., Johnson, D.S., Bernstein, B.E., Nusbaum, C., Myers, R.M., Brown, M., Li, W., and Liu, X.S. (2008). Modelbased analysis of ChIP-Seq (MACS). Genome Biol. 9, R137.

Zhou, Y., Zhou, B., Pache, L., Chang, M., Khodabakhshi, A.H., Tanaseichuk, O., Benner, C., and Chanda, S.K. (2019). Metascape provides a biologist-oriented resource for the analysis of systems-level datasets. Nat. Commun. 10,





STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit polyclonal anti-H3 acetyl-K27	Active Motif	Cat# 39133; RRID:AB_2561016
Chemicals, peptides, and recombinant proteir	ns	
Recombinant Protein G Agarose	Invitrogen	Cat# 15920-010
Proteinase K	Invitrogen	Cat# 25530-049
RNase A	Roche	Cat# 10109169001
γ-Secretase Inhibitor XXI (compound E)	Calbiochem	Cat# 565790
RPMI 1640	Corning	Cat# 10-040-CM
HyClone Fetal bovine serum	Thermo Fisher Scientific	Cat# SH30070.03
L-glutamine	Corning	Cat# 25-005-CI
Penicillin-Streptomycin	Corning	Cat# 30-002-CI
MEM Non-Essential Amino Acids	GIBCO	Cat# 11140-050
Sodium Pyruvate	GIBCO	Cat# 11360-070
Glycine	Invitrogen	Cat# 15527-013
Pierce 16% Formaldehyde	Thermo Fisher Scientific	Cat# 28908
Trizma Hydrochloride Solution, pH 7.4	Sigma-Aldrich	Cat# T2194-100ml
Sodium Chloride Solution, 5M	Sigma-Aldrich	Cat# 59222C-500ml
Magnesium Chloride Solution, 1M	Sigma-Aldrich	Cat# M1028-100ml
Nonidet P40 Substitute	Sigma-Aldrich	Cat# 74385-5I
MACS BSA Stock Solution	Miltenyi Biotec	Cat# 130-091-376
Flowmi Cell Strainer, 40 μm	Bel-Art	Cat# H13680-0040
Digitonin	Thermo Fisher Scientific	Cat# BN2006
Dulbecco's Phosphate-Buffered Salt Solution 1X	Corning	Cat# 21031CV
Critical commercial assays		
KAPA Library Quant Kit	Roche	Cat# KK4824
D1000 ScreenTape	Agilent	Cat# 5067-5582
D1000 Reagents	Agilent	Cat# 5067-5583
High Sensitivity D1000 ScreenTape	Agilent	Cat# 5067-5584
High Sensitivity D1000 Reagents	Agilent	Cat# 5067-5585
QIAquick PCR Purification Kit	QIAGEN	Cat# 28106
NEBNext Ultra II DNA Library Prep Kit	NEB	Cat# E7645S
Chromium Single Cell ATAC Library & Gel Bead Kit, 4 rxns	10X GENOMICS	Cat# PN-1000111
Chromium i7 Multiplex Kit N, Set A	10X GENOMICS	Cat# PN-1000084
Chromium Chip E Single Cell ATAC Kit, 48 rxns	10X GENOMICS	Cat# PN-1000082
NextSeq® 500/550 High Output Kit v2 (75 cycles)	Illumina	Cat# FC-404-2005
NextSeq® 500/550 High Output Kit v2 (150 cycles)	Illumina	Cat# FC-404-2002
Deposited data		
Raw and analyzed scATAC-seq data	This paper	GEO: GSE155916
Raw and analyzed ChIP-seq data	This paper	GEO: GSE171098
Bulk ATAC-seq of purified progenitor and differentiated hematopoietic cells	Yoshida et al., 2019; https://doi.org/10. 1016/j.cell.2018.12.036	GEO: GSE100738

(Continued on next page)

Cell Reports Report





Continued			
REAGENT or RESOURCE	SOURCE	IDENTIFIER	
10x Genomics scATAC-seq of CD34+ hematopoietic	Satpathy et al., 2019; https://doi.org/10. 1038/s41587-019-0206-z	GEO: GSE129785	
progenitor cells Fluidigm C1 scATAC-seq of	Buenrostro et al., 2018; https://doi.org/10.	GEO: GSE96769	
CD34+ hematopoietic progenitor cells	1016/j.cell.2018.03.074	GEO. GOESO/103	
sciATAC-seq of murine marrow and spleen cells	Cusanovich et al., 2018; https://doi.org/10. 1016/j.cell.2018.06.052	GEO: GSE111586	
scRNA-seq of GSI-resistant DND-41 cells	Schwartz et al., 2020; https://doi.org/10. 1038/s41592-020-0748-5	GEO: GSE138892	
Experimental models: Cell lines			
DND-41	DSMZ	ACC 525	
Software and algorithms			
APEC v1.2.2	Li et al., 2020; https://doi.org/10.1186/ s13059-020-02034-y	https://github.com/QuKunLab/APEC	
Cicero v1.9.1	Pliner et al., 2018; https://doi.org/10.1016/j. molcel.2018.06.044	https://github.com/cole-trapnell-lab/ cicero-release	
CisTopic v0.3.0	Bravo González-Blas et al., 2019; https://doi.org/10.1038/s41592-019-0367-1	https://github.com/aertslab/cisTopic	
Cusanovich2018	Cusanovich et al., 2018; https://doi.org/10. 1016/j.cell.2018.06.052	This paper https://github.com/faryabib/ CellReports_TooManyPeaks_analysis	
EpiScanpy v0.3.0	Danese et al., 2019; https://doi.org/10. 1101/648097	https://github.com/colomemaria/ epiScanpy	
Seurat v3.2.3	Butler et al., 2018; https://doi.org/10.1038/ nbt.4096	https://github.com/satijalab/seurat	
Signac v1.1.0	Stuart et al., 2020; https://doi.org/10.1101/ 2020.11.09.373613	https://github.com/timoast/signac	
SnapATAC v1.0.0	Fang et al., 2021; https://doi.org/10.1038/ s41467-021-21583-9	https://github.com/r3fang/SnapATAC	
tsne v0.1.3	van der Maaten and Hinton, 2008	https://github.com/jdonaldson/rtsne/	
TooManyPeaks v2.2.0.0	This paper https://doi.org/10.5281/zenodo. 5130671	https://github.com/faryabib/ too-many-cells#too-many-peaks	
TooManyPeaks analysis code	This paper https://doi.org/10.5281/zenodo. 5130655	https://github.com/faryabib/ CellReports_TooManyPeaks_analysis	
R wrapper for TooManyCells v0.1.1.0	Schwartz et al., 2020; https://doi.org/10. 1038/s41592-020-0748-5	https://github.com/GregorySchwartz/tooManyCellsR	
umap-learn v0.4.6	McInnes et al., 2018; https://doi.org/10. 21105/joss.00861	https://github.com/lmcinnes/	
HOMER v4.9	Heinz et al., 2010; https://doi.org/10.1016/j. molcel.2010.05.004	http://homer.ucsd.edu/homer	
bedtools v2.30.0	Quinlan and Hall, 2010; https://doi.org/10. 1093/bioinformatics/btq033	http://bedtools.readthedocs.io/en/stable	
BWA v0.7.13	Li and Durbin, 2009; https://doi.org/10. 1093/bioinformatics/btp324	http://bio-bwa.sourceforge.net	
Cell Ranger ATAC v1.2.0	Satpathy et al., 2019; https://doi.org/10. 1038/s41587-019-0206-z	https://support.10xgenomics.com/ single-cell-atac/software/pipelines/latest/ what-is-cell-ranger-atac	
Picard v2.1.0	Broad Institute	https://github.com/broadinstitute/picard	
Trim Galore v0.4.1	Babraham Bioinformatics	https://www.bioinformatics.babraham.ac.uk/projects/trim_galore	
UCSC tools v404	Kent et al., 2010; https://doi.org/10.1093/bioinformatics/btq351	https://github.com/ucscGenomeBrowser/kent	





RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Robert B. Faryabi (email:faryabi@pennmedicine.upenn.edu).

Materials availability

This study did not generate new unique reagents.

Data and code availability

DND-41 T-ALL scATAC-seg and ChIP-seg data have been deposited at the Gene Expression Omnibus and are publicly available as of the date of publication. Accession numbers are listed in the Key resources table.

In addition, Bulk ATAC-seq of purified progenitor and differentiated hematopoietic cells, 10x Genomics scATAC-seq, Fluidigm C1 scATAC-seq, sciATAC-seq, and GSI-resistant scRNA-seq data are existing, publicly available data. The accession numbers for these datasets are listed in the Key resources table.

All original software code has been deposited at https://github.com/faryabib/too-many-cells#too-many-peaks (source), https://hub. docker.com/repository/docker/gregoryschwartz/too-many-cells/ (Docker), https://cran.r-project.org/web/packages/TooManyCellsR (R wrapper), and https://github.com/faryabib/CellReports_TooManyPeaks_analysis (analysis code) and is publicly available as of the date of publication. DOIs are listed in the Key resources table.

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

GSI-resistant T-ALL cell culture

DND-41 cells (DSMZ, cat# ACC525) were purchased from the Leibniz-Institute DSMZ-German Collection of Microorganisms and Cell Lines. These male cells were cultured in RPMI 1,640 (Corning, cat# 10-040-CM) supplemented with 10% fetal bovine serum (Thermo Fisher Scientific, cat# SH30070.03), 2 mM L-glutamine (Corning, cat# 25-005-Cl), 100 Ug/mL and 100 μg mL⁻¹ penicillin/streptomycin (Corning, cat# 30-002-Cl), 100 mM nonessential amino acids (GIBCO, cat# 11140-050), 1 mM sodium pyruvate (GIBCO, cat# 11360-070) and 0.1 mM of 2-mercaptoethanol (Sigma, cat# M6250). All cells were grown at 37°C and 5% CO2 with media refreshed every 3-4 days. Cells were regularly tested for mycoplasma contamination.

IC₅₀ values for gamma-secretase inhibitor (GSI) compound E (Calbiochem, cat# 565790) were calculated from dose-response curves using CellTiter Glo Luminescent Cell Viability Assay (Promega, cat# G7571). Briefly, 1,000 treatment-naive DND-41 cells in 5 replicates/condition were plated in 96-well plates with vehicle or increasing concentrations of GSI (0.016, 0.031, 0.062, 0.125, 0.25, 0.5, 1, 2 μM). Luminescence was measured on day 7 with CellTiter Glo Luminescent Cell Viability Assay according to the manufacturer's instructions. DND-41 IC₅₀ of GSI was determined to be 5 nM.

To generate GSI-resistant cells, DND-41 treatment-naive cells were cultured in the presence of 125 nM GSI for at least six weeks. The establishment of GSI-resistance was determined with IC50 assay as described above. GSI-resistant DND-41 cells can tolerate 10 mM GSI with less than 20% cell death. Short-term DMSO treatment was performed on treatment-naive DND-41 cells with 125 nM DMSO for 24 hours.

METHOD DETAILS

GSI-resistant T-ALL single-cell ATAC-sequencing

We performed single-cell ATACseq for parental and GSI-resistant DND-41 cells following manufacture's instructions for Chromium Single Cell ATAC Library & Gel Bead Kit and Chromium Chip E Single Cell ATAC Kit (10x Genomics). Briefly, we loaded cells onto independent channels of a Chromium Controller for targeted recovery of 4,000 cells per condition. We assessed libraries with Agilent TapeStation using High sensitivity D1000 chip and quantified using KAPA Library Quantification Kits for Illumina platform (KAPA Biosystems, Roche, cat# KK4824). We performed paired-end sequencing on NextSeq 550 using 150 cycles High Output kit.

We performed FASTQ file generation and alignment to hg19 using Cell Ranger ATAC v1.2.0 (Satpathy et al., 2019) default arguments. We aggregated these cells using Cell Ranger. We sequenced parental and GSI-resistant cells at 253,594,800 and 252,343,254 read pair depth, respectively. In total, 8,041 cells passed the Cell Ranger QC and showed the typical "knee" plots indicating high quality from DMSO-treated parental (3,887) and GSI-resistant (4,154). We used sequence fragments of parental and resistant cells, with median of 13,238 and 33,523 per cell respectively, either directly as TooManyPeaks and SnapATAC inputs, or indirectly as Cicero, CisTopic, EpiScanpy, Cusanovich2018, APEC, Signac, and PAGA via pseudo-bulk ATAC-seq peak calling.

H3K27ac ChIP-seq

We performed H3K27ac ChIP-seq as previously described (Petrovic et al., 2019). Briefly, we sonicated and cleared chromatin samples prepared from 107 fixed cells with recombinant protein G-conjugated Agarose beads (Invitrogen, cat# 15920-010) and

Cell Reports Report



subsequently immunoprecipitated these cells with antibodies recognizing H3K27ac (Active Motif, cat# 39133). We captured Antibody-chromatin complexes with recombinant protein G-conjugated Agarose beads, washed them with Low Salt Wash Buffer, High Salt Wash Buffer, LiCl Wash Buffer and TE buffer with 50mM NaCl and eluted them. After reversal of cross-linking, we performed RNase and Proteinase K (Invitrogen, cat# 25530-049) treatment and purified DNA with QIAquick PCR Purification Kit (QIAGEN, cat# 28106). We then prepared libraries using the NEBNext Ultra II DNA library Prep Kit for Illumina (NEB, cat# E7645S). We validated indexed libraries for quality and size distribution using a TapeStation 2200 (Agilent). We performed paired-end sequencing (38 bp+38 bp) on a NextSeq 550.

Reads from H3K27ac ChIP-seg experiments were trimmed with Trim Galore (version 0.4.1, https://www.bioinformatics.babraham. ac.uk/projects/trim_galore/) with parameters -q 15-phred33-gzip-stringency 5 -e 0.1-length 20. Trimmed reads were aligned to the Ensembl GRCh37.75 primary assembly including chromosome 1-22, chrX, chrY, chrM and contigs using BWA (version 0.7.13) (Li and Durbin, 2009) with parameters bwa aln -q 5 -l 32 -k 2 -t 6 and paired-end reads were group with bwa sampe -P -o 1000000. Reads mapped to contigs, ENCODE blacklist and marked as duplicates by Picard (version 2.1.0, https://broadinstitute.github.io/picard/) were discarded and the remaining reads were used in downstream analyses and visualization. Bedgraph of reads normalized to reads per million (RPM) from ChIP-seq were generated with bedtools genomecov (Quinlan and Hall, 2010). Genome-wide uploadable bigWig files were generated with UCSC tools (version 329) (Kent et al., 2010) bedGraphToBigWig.

QUANTIFICATION AND STATISTICAL ANALYSIS

TooManyPeaks analysis of scATAC-seq data

TooManyPeaks is a collection of specialized functionalities and entry points for the parent suite TooManyCells, and extends TooManyCells to analyze chromatin accessibility of individual cells. TooManyPeaks provides many additional functionalities such as processing genomic region features (e.g., parsing regions for merging features) for sequence fragment file and / or peak matrix input, binning of regions, filtering out "black list" regions, and dimensionality reduction with LSA. In addition TooManyPeaks provides Too-ManyCells with several new entry points: peaks for peak finding with MACS2 (customizable with any other program), motifs for de novo or known motif search with MEME or HOMER (customizable with any other program), and classify for cell-type assignment from bulk ATAC-seq. While both MACS2 and MEME are included in the Nix derivation, any command line program can be integrated into the TooManyPeaks framework for use with these entry points.

Accessibility matrix

Given a sequence fragment file where each line consists of an initial three BED columns followed by cell barcode and duplicate count columns, TooManyPeaks initially generates an $m \times n$ matrix **M** of m observations (cells) and n genomic loci (equal-size genomic bins or pseudo-bulk ATAC-seq peaks) features, where $\mathbf{M}(i,j)$ is the number of counts for cell i at feature j. When starting from a fragments file, "black list" regions of the genome are known to have high signal (Amemiya et al., 2019). All analyses presented here that originate from a sequence fragment file filter out known "black list" regions with erroneously high signal (Amemiya et al., 2019), unless otherwise specified using-blacklist-regions-file. The width of genomic bin features across all the cells specified using-binwidth, which is set here to 5000 bp unless stated otherwise. By default, TooManyPeaks converts the matrix into a binary matrix to represent accessible or inaccessible sites.

Tree of single-cell clades

Potentially due to the "curse of dimensionality," the large number of features in scATAC-seq data may result in every cell being an outlier, which in turn leads to low modularity in the initial bi-partitioning and stops the tree generation prematurely. To avoid this situation, we use latent semantic analysis for dimensionality reduction (here using 50 dimensions with-lsa) (Deerwester et al., 1990). TooManyPeaks passes this reduced feature space matrix to TooManyCells, which generates a tree of single-cell relationships based on the accessibility of their chromatins. Briefly, we generate a tree of cell clade relationships by recursively bi-partitioning the cells using an efficient matrix-free divisive hierarchical spectral clustering (Schwartz et al., 2020). To simultaneously detect large and small populations and avoid creating arbitrary small clusters, we use Newman-Girvan modularity (Newman and Girvan, 2004) as a stopping criterion for recursive cell bi-partitioning. The TooManyPeaks divisive hierarchical spectral clustering algorithm produces a nested cluster structure where relationships among the groups are maintained.

Peak calling and downstream analyses

Each node in the tree contains a collection of cells. The TooManyPeaks peaks entry point can be used to directly call peaks using MACS2 that is integrated into TooManyPeaks. TooManyPeaks calls peaks at each node specified (or all nodes) for downstream differential peak calculations. TooManyPeaks can be instructed with the-bedgraph option to generate bedGraph and bigWig files to visualize chromatin accessibility of each node on a genome browser. Given labels such as cell type or disease state, TooManyPeaks can also make tracks filtered for a label for a set of nodes. Transcription factor binding sequence motif search programs MEME and HOMER are integrated into TooManyPeaks and can be used to identify de novo motifs and search for known motifs for each node. TooManyPeaks motif analysis options can be controlled from the motifs entry point.



Classification based on reference elements

To assign cell types to individual cells in the TooManyPeaks tree, TooManyPeaks can use peaks from pseudo-bulk scATAC-seq or bulk ATAC-seq data from FACS-purified cells as reference cis-regulatory elements. Here, we annotated each murine bone marrow and spleen cells (Cusanovich et al., 2018) based on reference cis-regulatory elements. We generated reference cis-regulatory elements of 92 phenotypically defined FACS-sorted progenitor and differentiated hematopoietic cell types by analyzing their bulk ATACseg. To this end, fastq files for 186 samples were obtained from ImmGenn GSE100738 (Yoshida et al., 2019), and aligned to mm9 genome with BWA (version 0.7.13) (Li and Durbin, 2009) with parameters bwa aln -q 5 -l 32 -k 2 -t 6, after trimming with Trim Galore (version 0.4.1) with parameters -q 15-phred33-gzip-stringency 5 -e 0.1-length 20. Reads mapped to contigs, ENCODE blacklist, and marked as duplicates by Picard (version 2.1.0) were discarded and the remaining reads were used for peak calling and creating

Reproducible peaks in ATAC-seg replicates were identified following an implementation of ENCODE Irreproducible Discovery Rate (IDR) pipeline. Peaks in true replicates, pseudoreplicates, and pooled samples were identified using MACS (version 2.0.9) (Zhang et al., 2008) with parameters -p 1E-5 -g mm9-nomodel-format = BAM-bw = 300-keep-dup = 1. IDR cutoffs for true replicates, pseudoreplicates, and pooled samples were 0.05, 0.05 and 0.005 respectively. Replicates with Np/Nt 2 and N1/N2 2 were considered reproducible. The resulting ATAC-seq peaks were used as reference cis-regulatory elements of 92 phenotypically defined progenitor and differentiated hematopoietic cell types.

Given a set of observations (cells) $O = \{1...m\}$, a set of features (regions or genes) $F = \{1...n\}$, a set of reference regulatory elements (pseudo-bulk scATAC-seq or bulk ATAC-seq FACS-purified populations) R = 1...r, an $m \times n$ observation feature matrix **M** and a new $r \times n$ reference matrix **R**, TooManyPeaks first normalizes each row in **M** and **R** such that for some $m \times n$ matrix **X**,

$$p(\mathbf{X})(i,j) = \mathbf{e}_i^{-1} \mathbf{X}(i,j),$$
 (Equation 1)

where $e_i = \sqrt{\sum_{k=1}^{n} \mathbf{X}^2(i,k)}$ is the Euclidean norm of **X** row *i*. Then we can generate a new matrix **S** representing a bipartite graph of relationships between the observations in **M** and bulk populations in **R** with

$$\mathbf{S} = p(\mathbf{M})p(\mathbf{R})^T,$$
 (Equation 2)

where $-1 \le \mathbf{S}(i,j) \le 1$ is the score (cosine similarity) of relatedness between observation $1 \le i \le m$ to reference $1 \le j \le r$, with higher score indicating higher relatedness. Then the set of cell-type assignments A of length m by maximum score is defined by

$$\mathbf{A}_i = \max_{j \in R} \mathbf{S}(i, j).$$
 (Equation 3)

Clustering benchmarks

We adapted the clustering benchmark for scRNA-seq as previously described (Schwartz et al., 2020) to scATAC-seq. Briefly, using TooManyPeaks, APEC (Li et al., 2020), Cicero (Pliner et al., 2018), CisTopic (Bravo González-Blas et al., 2019), CisTopic with Louvain, Cusanovich2018 (Cusanovich et al., 2018), EpiScanpy (Danese et al., 2019), Signac (Stuart et al., 2020), and SnapATAC (Fang et al., 2021), we clustered separately two datasets of phenotypically defined cells within bone marrow and blood samples profiled using 10x Genomics (Satpathy et al., 2019) (starting from a cell-by-peak file generated by TooManyPeaks to keep cells consistent) or Fluidigm C1 (Buenrostro et al., 2018) (starting from peaks as given in the dataset) scATAC-seq platforms. To increase the robustness of our benchmark, we additionally clustered a simulated bone marrow dataset with a moderate noise level of 0.2 (Chen et al., 2019). As Cicero and CisTopic focused on generating features, we also ensured a version of CisTopic using Louvain clustering from Signac instead of densityClust. We based this clustering benchmark on the assumption that similar cell types should cluster together. As such, we used purity (Manning et al., 2008), entropy (Tan et al., 2019), mutual information (Kvålseth, 2017), adjusted rand index (ARI), homogeneity (Rosenberg and Hirschberg, 2007), and residual average Gini index (RAGI) (Chen et al., 2019) to compare clustering performances between algorithms. In summary, entropy and homogeneity assess the extent of cell-type label diversity within clusters. Purity evaluates the extent of the dominant cell-type labels within the clusters. RAGI evaluates cluster-specificity of enrichment for marker accessible elements. Finally, NMI measures dependency of information of the cell-type labels given the cluster labels. RAGI requires gene activities as well as a list of known marker genes and housekeeping genes, so we used Cicero to generate the gene activity matrix and the gene lists originally given with RAGI's introduction (Chen et al., 2019).

Purity is based on the frequency of the most abundant class (e.g., cell type) in a cluster. Let $\Omega = \{\omega_1, \omega_2, ..., \omega_K\}$ be the set of cluster. ters and $\mathbb{C} = \{c_1, c_2, ..., c_J\}$ be the set of classes. Then purity is defined as

$$\operatorname{purity}(\Omega,\mathbb{C}) = \frac{1}{N} \sum_{k} \max_{j} |\omega_{k} \cap c_{j}|,$$

where N is the total number of cells, ω_k is the set of cells in cluster k, and c_i is the set of cells in class j (Manning et al., 2008). This measure ranges from 0, poor clustering, to 1, perfect clustering.

Entropy as a measure of cluster accuracy uses Shannon entropy (Shannon, 1948) to measure the expected amount of information from the clusters. The entropy of each cluster k is defined by

Cell Reports Report



$$H(\omega_k) = \sum_j \frac{|\omega_{kj}|}{|\omega_k|} \log \frac{|\omega_{kj}|}{|\omega_k|},$$

where ω_{kj} is the set of cells from $\omega_k \cap c_j$. Then the entropy for the entire clustering is (Tan et al., 2019)

$$\mathsf{entropy}(\varOmega,\mathbb{C}) = \sum_{k} \frac{|\omega_k|}{N} H(\omega_k).$$

Here, lower entropy of a clustering indicates higher accuracy.

Normalized mutual information (NMI) measures the normalized dependency of the class labels on the cluster labels, or the amount of information about the class labels gained when the cluster labels are given. Mutual information is defined by

$$I(Q; \mathbb{C}) = \sum_{k} \sum_{i} \frac{\left|\omega_{k} \cap c_{j}\right|}{N} \log \frac{N\left|\omega_{k} \cap c_{j}\right|}{\left|\omega_{k}\right|\left|c_{j}\right|}.$$

To compare mutual information across clusterings, $I(\Omega; \mathbb{C})$ is normalized to the interval [0, 1]. As $I(\Omega; \mathbb{C})$ is bounded by min $[H(\Omega), I]$ $H(\mathbb{C})$], where

$$H(\Omega) = -\sum_{k} \frac{|\omega_{k}|}{N} \log \frac{|\omega_{k}|}{N}$$

is the entropy of Ω along with the analogous $H(\mathbb{C})$, total normalization NMI can be defined by

$$\mathsf{NMI}(\varOmega,\mathbb{C}) = \frac{\mathit{I}(\varOmega;\mathbb{C})}{\mathsf{min}[\mathit{H}(\varOmega),\mathit{H}(\mathbb{C})]},$$

where higher values indicate more accurate clustering based on C (Kvålseth, 2017).

Homogeneity makes the assumption that clusterings assign all members within a single cluster a single label. Therefore, the label distribution within a single cluster should result in zero entropy. Thus, the perfect case of homogeneity would be the Shannon entropy of $H(\mathbb{C}|\Omega) = 0$. Then, instead of the raw entropy, homogeneity produces the normalized entropy by the maximum reduction in entropy from the clustering, namely $H(\mathbb{C})$. As 1 would be desirable as a maximum rather than 0, homogeneity is thus defined as (Rosenberg and Hirschberg, 2007)

$$h = \begin{cases} 1 & \text{if } H(\mathbb{C}|\varOmega) = 0 \\ 1 - \frac{H(\mathbb{C}|\varOmega)}{H(\mathbb{C})} & \text{otherwise} \end{cases}$$

where

$$H(\mathbb{C}|\Omega) = -\sum_{k} \sum_{j} \frac{|\omega_{k}j|}{N} \log \frac{|\omega_{k}j|}{\sum_{j} |\omega_{k}j|}.$$

Adjusted Rand Index (ARI) is calculated based on the number of pairings between two data clusterings, then adjusted for chance (Hubert and Arabie, 1985). Specifically, we first compute the Rand index

$$RI = \frac{TP + TN}{TP + FP + FN + TN},$$

where TP and FP is the number of true or false positives respectively, while TN and FN is the number of true or false negatives respectively, based on cells in the clustering pairs. Then, we can define the adjustment for chance as (Hubert and Arabie, 1985)

$$ARI = \frac{RI - \text{Expected } RI}{\text{max}RI - \text{Expected } RI}$$

For single-cell clustering accuracy, this measure requires a "ground truth" which was based on the given labels from each published dataset which defines coarse labels which some algorithms, such as TooManyPeaks, attempt to further delineate. As such, TooManyPeaks tends to perform poorly when using this type of measure with ambiguous "ground truth" clusterings.

Residual Average Gini Index (RAGI) is a recently proposed measure to define accuracy based on accessibility between known housekeeping genes and marker genes (Chen et al., 2019). This method first requires a gene activity matrix generated from the accessibility data, which we created using Cicero. Next, we calculate the mean accessibility values for all cells in each cluster. We use the Gini index on this vector of values based on either housekeeping genes or marker genes, both of which we used as previously reported (Chen et al., 2019). The Gini index (Gini, 1997) measures dispersion based on inequality among values in a distribution. Briefly, if x_i is the mean accessibility of i of all cells in a cluster, then the Gini index of n accessibility sites would be





$$G = \frac{\sum_{i=1}^{n} \sum_{j=1}^{n} |x_i - x_j|}{2\sum_{i=1}^{n} \sum_{j=1}^{n} x_j}.$$

Then, we define the RAGI value as the difference in Gini index means between the housekeeping and marker genes for a clustering (Chen et al., 2019).

We used all algorithms with either default parameters as outlined in their function definitions or associated vignettes throughout the entire study which includes preprocessing with Seurat (Butler et al., 2018), with the exception of: knn = False in epi.pp.neighbors followed by episcanpy.tl.diffmap and another neighbor identification round for EpiScanpy and PAGA for visualizations to avoid low nearest neighbor errors as suggested by the Scanpy vignettes, and RunSVD with 3 dimensions to have fewer dimensions than topics in CisTopic with Louvain in the rare population benchmarks to also avoid an error. Furthermore, while the recommended latent semantic indexing (LSI, analogous to LSA) transformation through Signac was done to maintain a standard between Cicero and CisTopic with Louvain clusterings, topics were directly inputted into Signac UMAP for visualization of CisTopic with Louvain to avoid UMAP artifacts. All UMAP projections of the same data from different tools used the same seed.

Rare population benchmarks

We adapted the rare population benchmark for scRNA-seq as previously described to scATAC-seq (Schwartz et al., 2020). To this end, we generated ten random datasets each from two immune cell datasets using subsampling. The first set of ten samples included 1000 cells each with one common B cell population (ranging from 900 to 990 cells), one rare CD8+T population (5 to 50 cells), and one rare T regulatory cell (Treg) population (5 to 50 cells) (starting from sequencing fragments or peaks, depending on which algorithm accepts which format as all cells were included) (Satpathy et al., 2019). The second set of ten samples included 500 cells each with one frequent common myeloid progenitor population (400 to 450 cells), one rare monocyte population (5 to 25 cells), and one rare plasmacytoid dendritic cell population (5 to 25 cells), with fewer cells due to a smaller dataset (starting from peaks as given in the dataset) (Buenrostro et al., 2018). Additionally, we benchmarked on synthetic data generated using simATAC (Navidi et al., 2021), where each of the common and two rare populations were generated from different seeds.

To quantify these benchmarks, we calculated a contingency table of the fraction of pairwise labels. For all rare cell pairs, we called a true pair if the two cells were of the same cell type (e.g., a Treg with another Treg or a CD8+ T cell with another CD8+ T), while we assigned a false pair if the two cells were of different cell types (e.g., a Treg with a CD8+ T cell). Then, the measure for accuracy in this benchmark was the fraction of true pairs in all pairs.

Timing benchmark

We ran each algorithm three times on a dataset of 2,954 cells (Buenrostro et al., 2018) using a machine with Ubuntu 20.04, 512GiB Memory, Intel® Xeon® CPU E5-2670 v3 @ 2.30GHz, 2 physical processors 24 cores, and 48 threads.

T-ALL scATAC-seq statistical analyses

We used the Kruskal-Wallis test with the Benjamini-Hochberg method for multiple-hypothesis correction (Benjamini and Hochberg, 1995) for differential accessibility between populations normalized by total sequence fragment. For the differential expression analysis, we used edgeR (Robinson et al., 2010) for normalization and the Benjamini-Hochberg multiple-hypothesis correction with quasi-likelihood (QL) F-test p value.

T-ALL scATAC-seq motif analyses

We used HOMER findMotifsGenome.pl (Heinz et al., 2010) on the differential accessibility list of resistant-like / other parental cells, keeping peaks that were considered significant at q < 0.05. This process generated a list HOMER identified as known motifs differential between these subpopulations. To understand the ontology of these motifs, we then performed a Metascape (Zhou et al., 2019) analysis on the motifs significant at q < 0.05. To identify putative regulatory elements that correlated with differential expression, we intersected these found elements with the differential gene expression between resistant-like and non-resistant-like parental cells of this system (Schwartz et al., 2020).

In order to identify motifs for putative regulatory elements at the LINC00977 locus, we used FIMO (Bailey et al., 2009) on the LINC00977 peak using the JASPAR reference database (Sandelin et al., 2004). To identify regulatory elements correlating with differential gene expression as with the global analysis above, we intersected this candidate list with the differential gene expression between resistant-like and non-resistant-like parental cells (Schwartz et al., 2020).

Statistical parameter definitions

Definitions of statistical parameters such as *n* and box-plot notations are defined in their respective figure legends.

Cell Reports, Volume 36

Supplemental information

TooManyPeaks identifies
drug-resistant-specific regulatory elements
from single-cell leukemic epigenomes

Gregory W. Schwartz, Yeqiao Zhou, Jelena Petrovic, Warren S. Pear, and Robert B. Faryabi

Figure S1 Α 0.8-0.8 0.8 Entropy 0.0 Homogeneity 0.6 0.6 0.6 0.6 0.6 0.0 0.4 Purity 0.4 ₩ 0.4 0.2 0.2 0.2 0.2 0.2 0.0 0.0 0.0 *\$07075505500550* Bins Bins Bins В 0.8 0.8 0.8 0.8 0.8 Entropy 0 0.0 Homogeneity 0.6 - 0.2 - 0.2 - 0.2 Anuty 0.6 0.6 0.6 0.6 0.4 0.4 ₩ 0.4 0.4 0.2 0.2 0.2 0.2 0.2 0.0 0.0 0.0 70 80 LSA 70 80 LSA 70 80 LSA 70 80 LSA 90 100 90 100 50 90 100 70 80 LSA 90 100 60 90 100 60 50 60 60 50 60 50 60 70 8 LSA 80 90 100 С D E TooManyPeaks with Cicero Gene Activity TooManyPeaks with Cistopic Topic CLP TooManyPeaks default CMP GMP GMP1-Low 2 HSC LMPP MCP MEP Gini Difference 7k) M PUTIS MPP Other Mono pDC Buenrostro et al. 2018 (C1 Fluidigm) TooManyPeaks with CisTopic Topic TooManyPeaks with Cicero Gene Activity G Н CLP TooManyPeaks with Cicero Gene Activity TooManyPeaks with Cistopic Topic HSC/MPP LMPP TooManyPeaks default Memory B Naive 0.75 ONaive CD4 T1 Plasma cell 0.50 OPre-B 0.25 0.00 Gini Difference M Puris Entropy 7p, Satpathy et al. 2019 TooManyPeaks with CisTopic Topic TooManyPeaks with Cicero Gene Activity (10X Genomics) 25,000 20,000 Time (s) 15,000

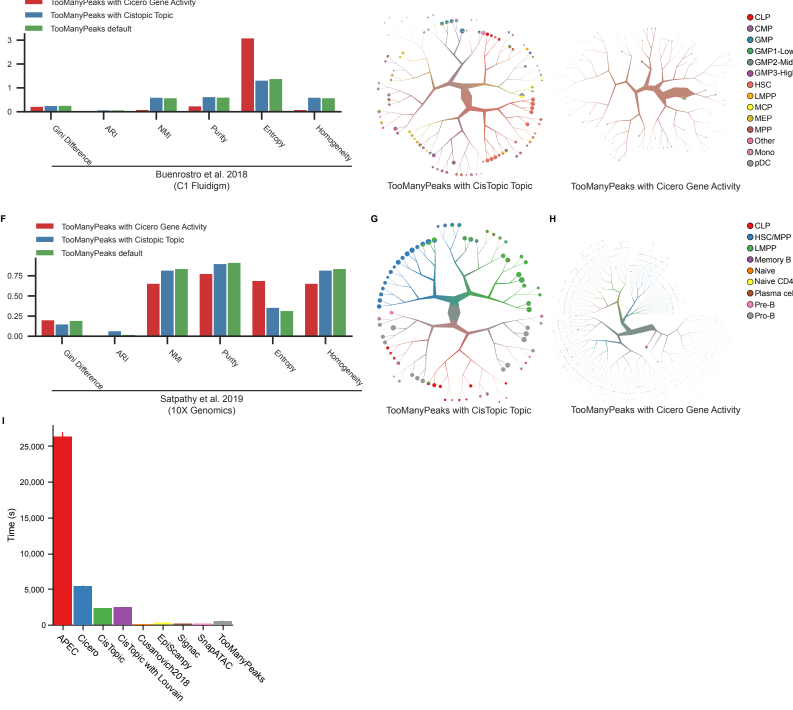


Figure S1: Comparison of TooManyPeaks for varying parameters, feature definition algorithms, and timing. (**A-B**) Comparison of TooManyPeaks across varying parameters on a human peripheral blood cell population (n = 44,814 cells) (Satpathy et al., 2019). Performance was measured using a clustering benchmark with, from left to right, lower entropy, higher purity, higher normalized mutual information (NMI), higher adjusted Rand index (ARI), and higher homogeneity showing more accurate clustering by varying either the number of bins (A) or the number of LSA dimensions (B). (**C-H**) Running TooManyPeaks using features defined from other scATAC-seq algorithms. (C, F) Clustering benchmark quantification from either the tree generated using CisTopic (D, G) or Cicero (E, H) on CD34⁺ hematopoietic progenitor cells profiled using 10x Genomics (n = 7,771 cells) (Satpathy et al., 2019) (C-E), or Fluidigm C1 (n = 2,954 cells) (Buenrostro et al., 2018) (F-H). (I) Timing benchmark of scATAC-seq algorithms. Running time of algorithms were compared using a human bone marrow profiled with Fluidigm C1 (n = 2,954 cells) (Buenrostro et al., 2018). Bar plots (mean plus / minus standard error) quantifies timing for 3 independent runs using a machine with Ubuntu 20.04, 512GiB Memory, Intel[®] Xeon[®] CPU E5-2670 v3 @ 2.30GHz, 2 physical processors 24 cores, and 48 threads. For an unbiased comparison, default or suggested filterings and parameters were used for all algorithms unless otherwise noted (see STAR Methods). Related to Figure 1.

Figure S2

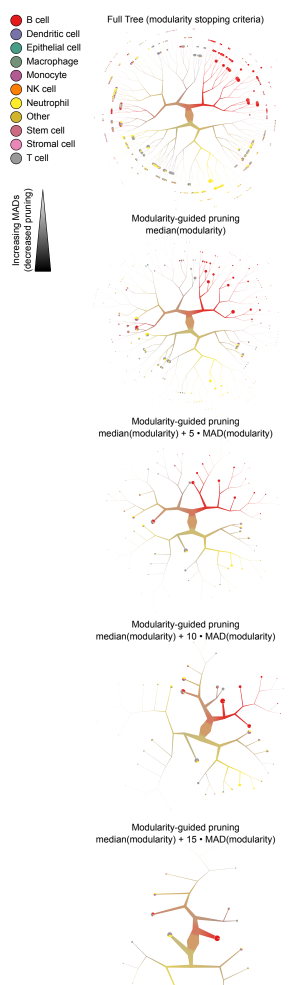


Figure S2: TooManyPeaks provides a number of pruning options, including a modularity-guided pruning, to control the tree depth (Schwartz et al., 2020). Example of controlling the depth of the TooManyPeaks tree with modularity stopping criteria of mouse spleen and bone marrow cells with modularity-guided pruning when the threshold is set to no pruning or median(modularity) plus 0, 5, 10, and 15 \times MAD(modularity) (from top to bottom in that order). MAD: median absolute deviations. Related to Figure 2.

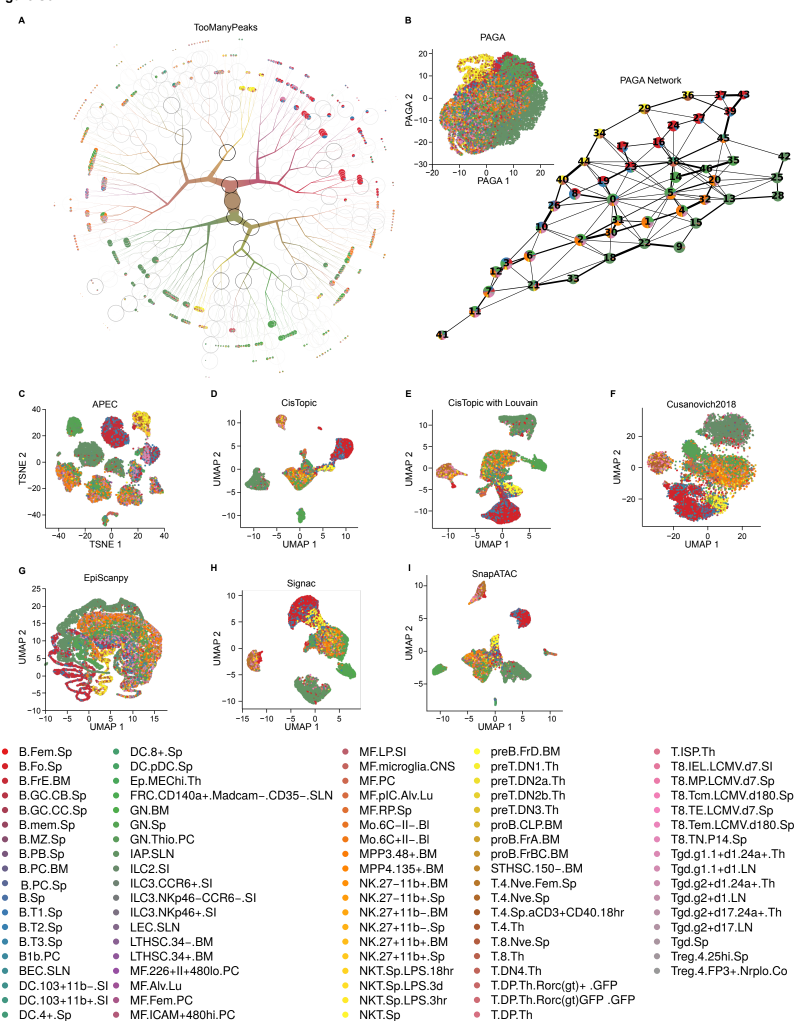


Figure S3: Stratification and annotation of murine marrow and spleen cells. The TooManyPeaks algorithm for cell-type annotation based on reference cis-regulatory elements was used to predict cell lineages in mouse bone marrow and spleen ($n=16,749\,\text{cells}$) (Cusanovich et al., 2018). Reference cis-regulatory elements of 92 phenotypically defined progenitor and differentiated hematopoietic cell types are generated from the analyses of bulk ATAC-seq in FACS-sorted cells (Yoshida et al., 2019). (A) TooManyPeaks with modularity stopping criteria and no pruning shows all 92 progenitor and differentiated hematopoietic cell types. At each bipartitioining, darker circle circumference represents higher modularity. (B-I) PAGA-initiated UMAP (B, left panel) or PAGA network (B, right panel), t-SNE output of APEC (C), as well as UMAP outputs of CisTopic (D), CisTopic with Louvain (E), Cusanovich2018 (F), EpiScanpy (G), Signac (H), and SnapATAC (I) are colored by assigned cell-type labels. For an unbiased comparison, each projection used the corresponding package's UMAP or t-SNE implementation. Related to Figure 2.

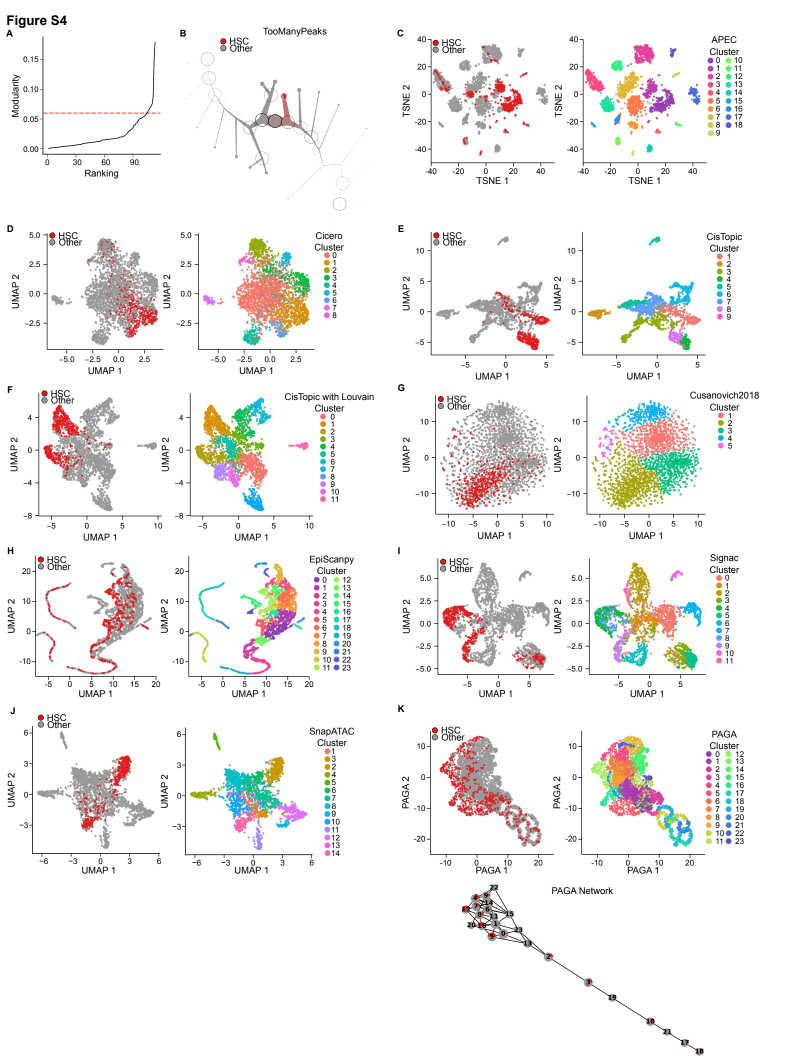


Figure S4: Comparing localization of hematopoietic stem cells (HSC) in the TooManyPeaks tree and UMAP or t-SNE outputs of human bone marrow profiled with Fluidigm C1 (Buenrostro et al., 2018). **(A)** The modularity-guided pruning threshold is set based on the beginning of the left plateau (marked by dashed red line) in the ranked modularity curve of the human bone marrow TooManyPeaks tree nodes (n=2,954 cells). **(B)** HSC cells are colored on the pruned TooManyPeaks tree per (A). At each bipartitioining, darker circle circumference represents higher modularity. **(C-K)** HSC cells are colored on UMAP or t-SNE outputs (red dots, left panel) generated by APEC (C), Cicero (D), CisTopic (E), CisTopic with Louvain (F), Cusanovich2018 (G), EpiScanpy (H), Signac (I), SnapATAC (J), and PAGA (K) initiated UMAP (top two panels) or PAGA network (bottom panels). Coordinates from the UMAP or t-SNE outputs (C-K, left panels) colored by each algorithm cluster label (C-K, right panels) fails to clearly localize HSC cells. For an unbiased comparison, each projection used the corresponding package's UMAP or t-SNE implementation. Moreover, default or suggested filters and parameters were used for all algorithms unless otherwise noted (see STAR Methods). Related to Figure 2.

Figure S5

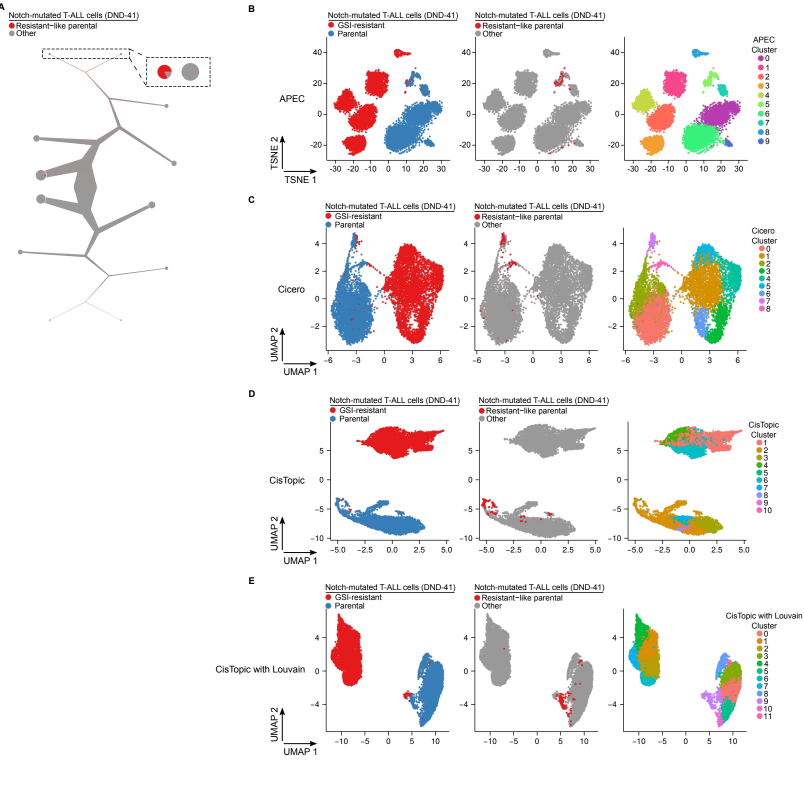


Figure S5: Comparisons of DND-41 T-ALL scATAC-seq data visualization. **(A)** The TooManyPeaks tree colored by resistant-like parental cells. **(B-E)** projection outputs of APEC (B), Cicero (C), CisTopic (D), and CisTopic with Louvain (E) colored by resistance status (left), resistant-like parental cells as defined by TooManyPeaks (middle), or algorithm cluster assignment (right) (n = 7,989 cells). For an unbiased comparison, each projection used the corresponding package's UMAP or t-SNE implementation. Moreover, default or suggested filters and parameters were used for all algorithms unless otherwise noted (see STAR Methods). Related to Figure 3.

Figure S6

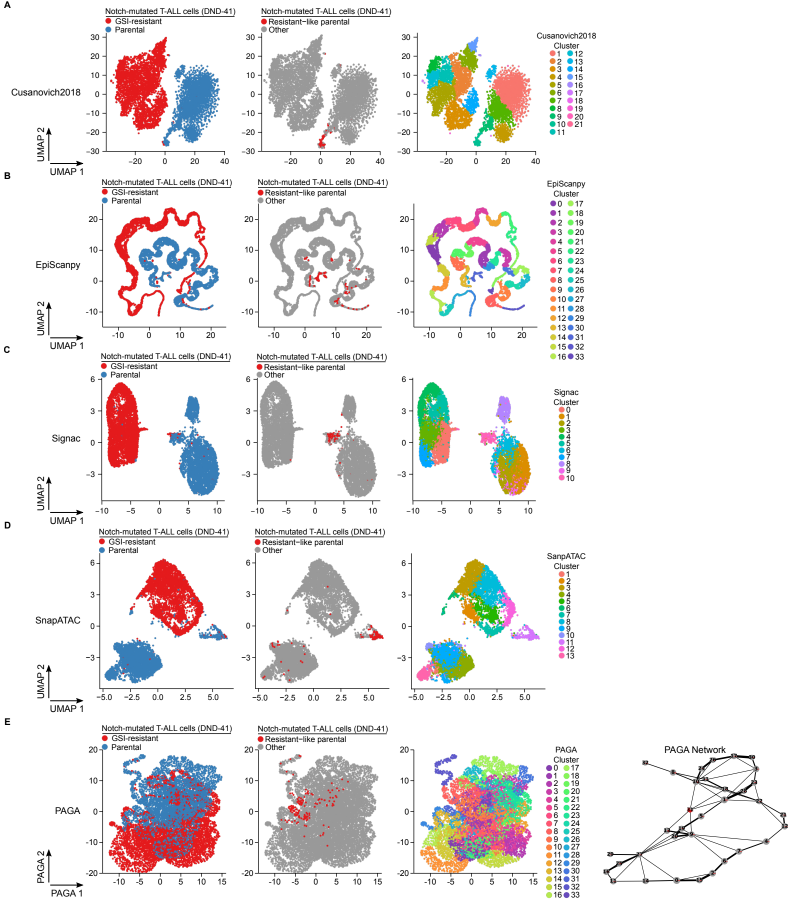
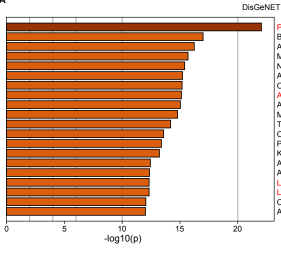


Figure S6: Comparisons of DND-41 T-ALL scATAC-seq data visualization, continued from Figure S5. **(A-D)** projection outputs of Cusanovich2018 (A), EpiScanpy (B), Signac (C), and SnapATAC (D) colored by resistance status (left), resistant-like parental cells as defined by TooManyPeaks (middle), or algorithm cluster assignment (right). **(E)** Projection output of PAGA-initiated UMAP colored by resistant status (left), resistant-like parental (second from left), cluster assignment (second from right). Right panel shows PAGA network colored by resistant-like parental cells (n = 7,989 cells). For an unbiased comparison, each projection used the corresponding package's UMAP or t-SNE implementation. Moreover, default or suggested filters and parameters were used for all algorithms unless otherwise noted (see STAR Methods). Related to Figure 3.



Precursor T-Cell Lymphoblastic Leukemia-Lymphoma Burkitt Lymphoma

Anaplasia

Malignant neoplasm of soft tissue

Nephroblastoma Adult Burkitt Lymphoma Childhood Burkitt Lymphoma

Adult T-Cell Lymphoma/Leukemia

Adult Classical Hodgkin Lymphoma

Myeloid Leukemia

Tumor Initiation

Classical Hodgkin's Lymphoma

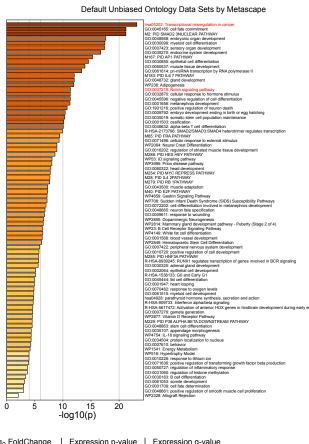
Progression of prostate cancer

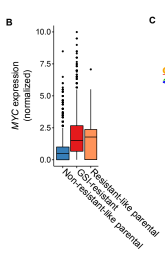
Ki-1+ Anaplastic Large Cell Lymphoma Acute Promyelocytic Leukemia

Aortic Aneurysm, Abdominal

Leukemia, T-Cell Lymphoid leukemia

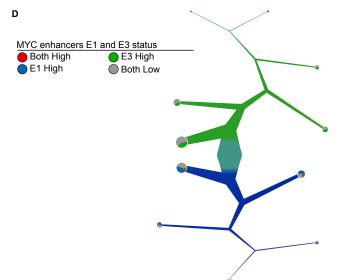
Congenital chromosomal disease Acute Erythroblastic Leukemia





EASCACGTGS	
STCCCAGGGG	ı

Motif	Motif p-value	Motif q-value	log ₂ FoldChange	Expression p-value	Expression q-value	
с-Мус	1e-6	0.00	1.08	5.67e-6	1.94e-3	
EBF1(EBF)	1e-24	0.00	0.856	0.0228	0.397	



Ε

ASA TOM C
C_AUT_G_T_

Motif	Motif p-value	Motif q-value	log ₂ FoldChange	Expression p-value	Expression q-value	
TCF7	1.33e-5	0.0306	-2.63	2.45e-7	9.30e-5	
GMEB1	-0.987	0.0283	-0.987	0.0283	0.350	

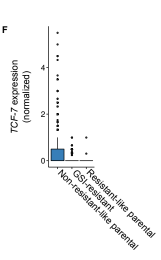


Figure S7: MYC and TCF-1 potentially bind differentially accessible elements of resistant-like parental cells. (**A**) Metascape analysis with DisGeNET database (left) and Ontology gene sets (right) showing that transcription factors with enriched binding motifs at the differential accessible elements of resistant-like parental cells are associated with pathways involved in T cell development and malignancies (highlighted in red, see STAR Methods). (**B**) *MYC* expression levels in GSI-resistant, resistant-like parental, and non-resistant-like parental cells identified in the TooManyCells tree (n = 7,371 cells). (**C**) Transcription factors from HOMER with both significant upregulation and more accessible binding motif recognition sites in resistant-like parental compared to non-resistant-like parental cells. (**D**) TooManyPeaks tree of parental and GSI-resistant DND-41 cells as in Figure 3A showing the accessibility of the Notch-dependent and Notch-independent *MYC* enhancers (n = 7,989 cells). (**E**) Transcription factors from JASPAR (Sandelin et al., 2004) with both significantly lower expression and accessible binding motif recognition sites at the *LINC00977*-proximal enhancer E2 of the resistant-like compared to non-resistant-like parental cells. (**f**) *TCF-7* expression levels in GSI-resistant, resistant-like parental, and non-resistant-like parental cells identified in the TooManyCells tree (Schwartz et al., 2020) (n = 7,371 cells). Related to Figure 3.