A unified atlas of CD8 T cell dysfunctional states in cancer and infection

Graphical abstract



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In brief

Pritykin et al. reveal a shared path of T cells toward dysfunction and the early bifurcation of functional and dysfunctional T cell states from a TCF1⁺ progenitor by implementing a unified analysis of transcriptional and epigenetic states in T cells in cancer and infection.

Highlights

- Meta-analysis of chromatin states in T cells responding to cancer and infection
- Chronically activated T cells follow a common path toward dysfunction
- Cell-state-specific transcription factors define T cell
 differentiation trajectories
- Functional and dysfunctional T cells emerge from similar TCF1⁺ progenitor states



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A unified atlas of CD8 T cell dysfunctional states in cancer and infection

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SUMMARY

CD8 T cells play an essential role in defense against viral and bacterial infections and in tumor immunity. Deciphering T cell loss of functionality is complicated by the conspicuous heterogeneity of CD8 T cell states described across experimental and clinical settings. By carrying out a unified analysis of over 300 assay for transposase-accessible chromatin sequencing (ATAC-seq) and RNA sequencing (RNA-seq) experiments from 12 studies of CD8 T cells in cancer and infection, we defined a shared differentiation trajectory toward dysfunction and its underlying transcriptional drivers and revealed a universal early bifurcation of functional and dysfunctional T cell states across models. Experimental dissection of acute and chronic viral infection using single-cell ATAC (scATAC)-seq and allele-specific single-cell RNA (scRNA)-seq identified state-specific drivers and captured the emergence of similar TCF1⁺ progenitor-like populations at an early branch point, at which functional and dysfunctional T cells diverge. Our atlas of CD8 T cell states will facilitate mechanistic studies of T cell immunity and translational efforts.

INTRODUCTION

Upon completing their differentiation in the thymus, mature naive T lymphocytes enter the periphery and recirculate through secondary lymphoid organs, where, upon an encounter with a cognate antigen in the presence of co-stimulatory molecules, they become activated, expand, and differentiate into effector or memory T cells. These cells then take up residence in lymphoid and non-lymphoid organs, where they exert their immune functions. In contrast, chronic or suboptimal antigenic stimulation (e.g., in the absence of co-stimulation) can result in a state of hypo-responsiveness or anergy (Murphy and Weaver, 2016). Over the last decade, this simple textbook view has evolved into a markedly more nuanced and complex picture of T cell differentiation, with a plethora of seemingly distinct states emerging from a large number of studies in mice and man (Fan and Rudensky, 2016; Jameson and Masopust, 2018; Kumar et al., 2018). CD8 T cells, whose function is essential for defense against viral and bacterial infections and for tumor immunity, serve as a case study in this regard. Phenotypic and functional analyses of CD8 T cells in acute and chronic viral infections,

cancer, transplantation, and "self" tolerance in both experimental animal models and in human patients have offered numerous descriptions of activated effector, long-lived central, and short-lived effector memory cells and their precursors, as well as an array of CD8 T cell states with perturbed functionalities dubbed anergic, exhausted, and reversibly or irreversibly dysfunctional. Recent characterization of a small subset of exhausted/dysfunctional cells, named "stem-cell-like" or progenitor cells and capable of self-renewal, adds further complexity to the topography of CD8 T cell activation and differentiation (McLane et al., 2019; Hashimoto et al., 2018; Blank et al., 2019).

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Studies of CD8 T cell dysfunctional states, besides being highly significant for understanding basic mechanisms of adaptive immunity, have attracted particular attention due to the realization that prevention or reversal of CD8 T cell dysfunction can serve as a potent strategy for the treatment of both solid organ and hematologic malignancies and chronic infections. Inefficient mobilization of endogenous CD8 T cell responses or a failure to engage them in cancer patients in response to checkpoint blockade inhibitors, as well as disease resistance or relapse following adoptive CD8 T cell therapies including CAR (chimeric



antigen receptor) T cells, has been attributed to a large degree to dysfunction or exhaustion of tumor- and virus-specific CD8 T cells (Ribas and Wolchok, 2018; Sharma et al., 2017; Kallies et al., 2020). Transcriptional and chromatin features associated with these states have been extensively explored through the analyses of epigenomes and transcriptomes of isolated subsets of functional and dysfunctional CD8 T cells using DNase I hypersensitive sites sequencing (DHS-seq), assay for transposaseaccessible chromatin using sequencing (ATAC-seq), and RNA sequencing (RNA-seq) and through single-cell transcriptomics and proteomics (Pauken et al., 2016; Sen et al., 2016; Scott-Browne et al., 2016; Philip et al., 2017; Mognol et al., 2017; Chen et al., 2019a, 2019c; Scharer et al., 2017; Utzschneider et al., 2016; Man et al., 2017; Miller et al., 2019; Brummelman et al., 2018; Bengsch et al., 2018; Yao et al., 2019; Beltra et al., 2020; Yu et al., 2017). These studies have significantly advanced the knowledge of CD8 T cell differentiation and highlighted pronounced changes in T cell chromatin states. However, the remarkable heterogeneity of CD8 T cell states revealed by these genome-wide analyses in diverse experimental and clinical settings poses a major problem of distinguishing between common and context-specific features of differentiation toward dysfunction and underlying regulatory mechanisms. For example, in chronic viral infection, T cells encounter antigen in an inflammatory and stimulatory context and have been described as progressing through an effector state prior to differentiating to a state often called exhaustion (Wherry and Kurachi, 2015). Meanwhile, a two-step process of differentiation from a reversible to an irreversible dysfunctional state was reported in studies, including our own, in the setting of early tumorigenesis, where naive tumor-specific T cells encounter antigen in a non-inflammatory setting that may result in inadequate priming or activation (Philip et al., 2017; Philip and Schietinger, 2019). Therefore, in addition to inconsistent terminology (i.e., exhaustion versus dysfunction), it is unclear how to reconcile different models of progression to dysfunction (the term we will use here) and how these differentiation programs give rise to two distinct states at late time points of chronic antigen exposure: self-renewing dysfunctional progenitors and terminally dysfunctional cells.

A vexing obstacle in addressing these issues has been the inability to directly compare genome-wide data from different studies due to technical sources of variation-including sample preparation, sequencing quality, batch effects, and cell numbers-making meaningful integration of massive amounts of data generated in mouse and human studies problematic. To address this challenge, we carried out a uniform reprocessing and a statistically principled batch effect correction approach to over 300 chromatin accessibility (ATAC-seq) and gene expression (RNA-seq) datasets generated in 12 independent studies of CD8 T cell states observed across experimental mouse models of acute and chronic infection and tumors. Our analysis revealed a universal signature of chromatin accessibility changes in the progression to terminal dysfunction in both tumors and chronic infection, implying early commitment to a dysfunctional fate in all settings of chronic antigen exposure. The chromatin state observed at early time points during the development of dysfunction was similar to that of dysfunctional progenitor cells found in late time points in infection and tumor

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models. Motif-based regression modeling of this unified chromatin accessibility compendium enabled inference of state-specific transcription factor (TF) activities and implicated new factors in the progression to terminal dysfunction. This bulk-level T cell state analysis suggested a universal early bifurcation of functional and dysfunctional T cell activation states across models of cancer and chronic infection.

We further characterized this early branch point by carrying out single-cell analysis of CD8 T cell populations in the context of acute and chronic lymphocytic choriomeningitis virus (LCMV) infection and observed a TCF1⁺ progenitor-like population resembling the memory precursor effector cell (MPEC) population in acute infection. Regression modeling of single-cell ATAC (scATAC)-seq clusters enabled refined association of T cell functional states with the activity of TFs, whose causal role was established through a comprehensive single-cell RNA (scRNA)-seq atlas in T cell populations from F1 hybrid mice combining evolutionary distant genomes of laboratory and wild-derived mouse strains.

Together, these results provide new insights into the early emergence of a progenitor-like population in response to chronic antigen stimulation that appears to precede the establishment of dysfunction-committed progenitor cells, elucidating recent reports (Utzschneider et al., 2020; Chen et al., 2019c) with rich single-cell characterizations. Our unified atlas of CD8 T cell chromatin and expression states across mouse models and single-cell analyses of the bifurcation between functional and dysfunctional T cell responses will provide a valuable resource to the community and facilitate further mechanistic and translational studies.

RESULTS

Dysfunctional T cells in tumors and chronic infection share a common epigenetic and transcriptional state space

We collected 166 chromatin accessibility (ATAC-seq) datasets (Figure 1A; Table S1) from 10 recent studies on CD8 T cell function in mouse models of infection and cancer. These encompassed T cells in settings of acute bacterial infection, acute and chronic viral infection, including memory precursor and tissue-resident memory cells, and tumor-infiltrating lymphocytes in hepatocarcinoma and melanoma models. The data included endogenous T cells and adoptively transferred CAR-T cells; T cells with and without treatment with anti-PD1 immunotherapy; and progenitor T cell populations isolated from various models of chronic antigen stimulation (Pauken et al., 2016; Sen et al., 2016; Scott-Browne et al., 2016; Scharer et al., 2017; Mognol et al., 2017; Philip et al., 2017; Chen et al., 2019a; Miller et al., 2019; Milner et al., 2017; Yu et al., 2017). We performed a uniform processing of these data to construct a high-resolution atlas of 129,799 reproducible chromatin accessibility peaks across CD8 T cell states; these peaks were further split into 221,054 subpeaks and associated to genes (Method details; Table S2).

Expectedly, principal-component analysis (PCA) of peak read counts clustered samples by data source (Figure S1A). We therefore applied a generalized linear model (GLM) accounting for data source and functional state (naive, functional,



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Figure 1. Dysfunctional CD8 T cells in tumors and chronic infection share a common chromatin state space

(A) Snapshot of the ATAC-seq compendium near the Pdcd1 locus (light gray bars, peaks; dark gray bars, peak summit regions).

(B) PCA of batch-effect-corrected ATAC-seq signal in peak summit regions (functional cell state, color; data source, shape).

(C) First principal component (PC1) of PCA for batch-effect-corrected ATAC-seq signal in dysfunctional T cells from different studies (shown separately below) for peaks differentially accessible between progenitor and terminally dysfunctional cells in chronic LCMV infection.

(D) Differential accessibility and differential expression between progenitor and terminally dysfunctional T cells in chronic LCMV infection and in melanoma, and between early and late states of dysfunction in hepatocarcinoma progression. Left: batch-effect-corrected ATAC-seq signal log2 fold change for peaks of significantly differentially accessible genes; right: log2 fold change of RNA-seq gene expression (color for significantly decreased/increased individual peaks or genes; FDR < 0.05).

See also Figures S1 and S2 and Tables S1, S2, and S3.

dysfunctional; see Method details). After this batch effect correction, distances between functionally related samples from different studies decreased (Figure S1B), and samples readily clustered into broad functional categories regardless of the data source (Figure 1B). A more conservative GLM correction that did not explicitly model differences between functional



and dysfunctional cells produced similar results (Figure S1C). Furthermore, naive cells across studies clustered together in the PCA plot, while effector and memory cells (as well as preactivated cells injected in melanoma-bearing mice, but ignorant of the tumor antigen) formed a cluster of functional cells. Interestingly, cells profiled at day 4 in acute infection (Scharer et al., 2017) were positioned between clusters of naive and functional cells, suggesting their intermediate state. Strikingly, the chromatin states of dysfunctional tumor-infiltrating T cells from different tumor models and T cells in chronic viral infection formed a distinct cluster, suggesting a universal program of T cell dysfunction across models and immune challenges. A similar analysis of 136 gene expression (RNA-seq) datasets from eight studies (Pauken et al., 2016; Scott-Browne et al., 2016; Man et al., 2017; Utzschneider et al., 2016; Miller et al., 2019; Mognol et al., 2017; Philip et al., 2017; Chen et al., 2019a) showed consistent results (Figure S1D; Tables S1 and S2). Differential expression between functional and dysfunctional cells was significantly correlated with differential accessibility (Figure S1E). This confirmed that dysfunctional T cells are epigenetically and transcriptionally similar in the settings of chronic infection and across tumor models.

Genes with the strongest differential accessibility between functional and dysfunctional cells at their promoter, intronic, and nearby intergenic peaks (Method details) included wellknown markers of T cell activation, cytotoxicity, adhesion, and apoptosis, as well as those encoding key TFs, cytokines and cytokine receptors, and other cell-surface and intracellular proteins (Figure S1F; Table S2). Consistent with global differential accessibility and expression correlation (Figure S1E), differential accessibility of individual genes was often associated with significant differential expression (Figure S1F). Differential accessibility between naive or memory and dysfunctional CD8 T cells in mouse models was significantly associated with orthologous changes in human donors and cancer patients, suggesting generalizability to the human context (Figure S1G; Table S3; Method details).

Thus, our integrative accessibility and expression analyses across our compendium (Table S2) identified a high-confidence universal epigenetic and transcriptional gene signature of T cell dysfunction.

T cell temporal progression in tumors and chronic infection mirrors the state change from progenitor to terminal dysfunction

Dysfunctional T cells profiled as early as days 5–8 (d5–8) after antigen encounter in tumor or chronic viral infection models clustered closely with terminally dysfunctional cells profiled at d22–35 rather than with effector or memory cells (Figures 1B and S1D). Cells characterized as progenitor dysfunctional cells in different studies also clustered with terminally dysfunctional cells. This suggested that T cells adopt a dysfunctional rather than an effector chromatin state as early as d5 after antigen encounter.

We next compared chromatin states of different dysfunctional T cell subsets across models and immune challenges. Differentially accessible peaks between progenitor and terminally dysfunctional cells, including endogenous or transferred CAR-T cells in melanoma or dysfunctional cells in chronic viral infec-

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tion, were consistent between models and displayed concordant changes between early (d5-8) and late (d22-35) states of dysfunction in chronic viral infection and hepatocarcinoma tumorigenesis (Figures 1C and S2A-S2C; Table S2). Changes in accessibility between early and late dysfunctional states, and between progenitor and terminally dysfunctional states, correlated with changes in gene expression (Figure S2D). Importantly, T cells at early time points in all mouse models were similar in bulk chromatin state to progenitor dysfunctional cells, a subpopulation sorted from late time points in chronic viral infection or tumors. Differential accessibility between progenitor and terminally dysfunctional CD8 T cells in the melanoma mouse model was significantly associated with that in tumor infiltrating lymphocytes from melanoma patients (Sade-Feldman et al., 2018), again supporting generalizability to human data (Figure S2E; Table S3; Method details). These analyses suggested a common axis of differentiation of T cell dysfunction across models and immune challenges.

Many individual genes displayed similar expression and chromatin accessibility changes across models along this common differentiation axis (Figure 1D; Table S2). For example, genes encoding markers of terminal dysfunction Entpd1 (Cd39), 2B4 (Cd244), and Cd38 were significantly more accessible in terminal dysfunction and late tumor-specific dysfunction, while genes encoding progenitor cell markers Cxcr5, Slamf6, and IL7Ra chain were significantly more accessible in progenitor and early dysfunctional cells. Additional genes encoding cell-surface proteins, such as CD9, CD200, TNFRSF25 (DR3), CD83, and CD69, were significantly more accessible and expressed in progenitor cells and could serve as candidate progenitor cell markers. The locus of the TF Id2 was more accessible in terminal/late dysfunction, while Id3, Tcf7, Lef1, Nfkb1, Pou2f2, and Pou6f1 were more accessible in progenitor or early dysfunctional cells, implicating multiple TFs in the establishment and maintenance of these states. In most cases, differential accessibility at individual genes in early/progenitor versus late/dvsfunctional states was associated with differential expression, consistent with overall correlation of differential expression and accessibility (Figure S2D). However, our joint analysis did recover contextspecific features, including accessibility and expression patterns associated with T cell activation in chronic viral infection compared to tumor contexts (Figure S2D).

Tox and *lkzf2*, whose heightened expression was previously associated with terminal dysfunction, were indeed significantly more accessible and highly expressed in terminal than in early T cell dysfunction during hepatocarcinoma tumorigenesis; however, both loci had many peaks significantly more accessible in progenitor cells, and lkzf2 was significantly overexpressed in progenitor cells both in chronic LCMV infection and melanoma, suggesting the activity of these TFs in progenitor cells (Figure 1D).

Thus, our analysis identified universal markers and TFs associated with T cell differentiation toward dysfunction across models and organisms.

Coordinated activity of many TFs characterizes functional and dysfunctional T cell states

We next sought to identify TFs associated with different T cell activation and differentiation states, particularly the progenitor

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Figure 2. Predictive modeling identifies TFs associated with progenitor dysfunctional CD8 T cells

(A) Schematic of the negative binomial GLM to infer TF associations with chromatin accessibility.

(B) Inferred TF motif coefficients (with the highest variance across conditions, Z score row normalized) from each sample, consolidated between replicates across studies (median over replicates; error bars, 95% confidence intervals around median).

(C) Inferred TF motif coefficients with the highest variance across dysfunctional states (Z score row normalized).

(D) Network of predicted regulatory interactions between TFs in progenitor dysfunctional T cells (binding motifs not available for TOX and IKZF2, yellow). See also Figure S3.

cells. We analyzed TF targets, as predicted by TF binding motifs, and used negative binomial regression with ridge regularization to predict absolute levels of chromatin accessibility (ATAC-seq read coverage) in peaks from TF motif scores for each sample (Figures 2A, S3A, and S3B; Method details). For this analysis, we focused on 105 motifs for TFs expressed in CD8 T cells after grouping TFs with indistinguishable motifs. Using coefficients of the regression models, we estimated the effect on accessibility of each TF in each sample. This allowed us to map chromatin accessibility profiles into a lower-dimensional inferred TF activity space, largely preserving the relationships between samples (Figure S3C).

In order to identify TFs most strongly associated with T cell functional states, we consolidated the inferred TF activities across replicates and studies (Figure 2B). For example, as expected, the Eomes/Tbx21 motif was associated with low



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accessibility in naive cells and high accessibility in all antigenexperienced cells, and the Lef1/Tcf7 motif was associated with high accessibility in naive cells and low accessibility in terminally dysfunctional cells (Figure 2B). Nr4a, Nfkb, Nfat, Pou, and AP1 family motifs were strongly associated with high accessibility in early dysfunctional and progenitor cells, as opposed to terminally dysfunctional, naïve, or functional cells, while Zfp143 and Zfp523 motifs were associated with high accessibility in terminal dysfunction (Figures 2B and 2C); this included both previously (Chen et al., 2019a; Philip et al., 2017; Doering et al., 2012; Im et al., 2016; Beltra et al., 2020) and newly identified factors. Interestingly, Ctcf motifs were associated with differential accessibility between conditions (Figures 2B and 2C), suggesting changes in chromatin looping between T cell functional states (Johanson et al., 2019; Splinter et al., 2006); consistently, we observed a significant relationship between differential accessibility and differential expression when associating ATAC-seq peaks to genes using chromatin loops defined using published Hi-C data in naive CD8 T cells (He et al., 2016) instead of our default approach (Figure S3D; Method details). Overall, motifbased regression modeling of ATAC-seq data was a powerful approach for associating candidate TFs with T cell functional states.

Inferred TF activities across all antigen-experienced cells were highly correlated, potentially suggesting coordinated regulatory programs in different functional states; importantly, correlated TFs generally had different motifs and were bound at different sites (Figure S3E). Many TFs expressed in progenitor dysfunctional cells had multiple predicted binding sites in loci encoding other TFs, including Tox and Ikzf2, for which we did not have a motif, and thus could potentially regulate their expression (Figure 2D). This suggested that the coordinated and hierarchically organized activity of a broad range of TFs may be required for establishing and maintaining T cell functional states, as opposed to a single "master regulator."

Single-cell chromatin accessibility analysis reveals the early emergence of a progenitor-like T cell population

Our integrative analysis of bulk ATAC-seq data found an early (d5–8) divergence of CD8 T cells between responses to acute and chronic immune challenges (Figure 1B) and showed that



plastic and reprogrammable cells early (d7–8) in the development of T cell dysfunction were more similar to immunotherapy-responsive progenitor dysfunctional cells identified at later (d20–35) time points than to terminally dysfunctional cells (Figure 1C). To better characterize chromatin states of the early divergence between functional and dysfunctional fates, we performed scATAC-seq analysis of the total splenic CD8 T cell compartment in mice at d7 upon infection with LCMV Armstrong (Arm), resulting in acute infection, and LCMV clone 13 (Cl13), resulting in chronic infection (Figure 3A).

We first constructed a combined atlas of 189,281 chromatin accessibility peaks including both the bulk ATAC-seq data peak atlas and 59,482 newly identified scATAC-seq peaks (Table S4). Quantification and filtering yielded 4,767 and 5,865 cells, with 12,598 and 13,195 median reads per cell, initially isolated from Arm- and Cl13-infected mice, respectively. Normalization using term frequency-inverse document frequency (TF-IDF), dimensionality reduction using PCA followed by uniform manifold approximation and projection (UMAP), and Louvain clustering (Method details) suggested that d7 responses to infection with the two LCMV clones were heterogeneous and overlapping (Figures 3B, 3C, S4A, and S4B).

To characterize functional features of d7 cell populations, we scored scATAC-seq data with epigenetic signatures derived from bulk ATAC-seq data (Figures 3D, S4C, and S4D; Method details) and used single-sample gene set enrichment analysis (ssGSEA) to associate scATAC-seq cell clusters with the bulk ATAC-seq data compendium (Figure 3E; Method details). We found that cells in clusters 0, 7, 9, 11, 12, and 14 were likely a mixture of naive and background-activated memory cells not specific to the LCMV antigen, while clusters 1–6, 8, 10, and 13 likely consisted of LCMV responding cells. Some clusters showed no strong bias between LCMV clones, but we also found differences (e.g., in clusters 1 and 6) (Figures 3E, 3F, and S4E), suggesting a clear divergence between responses to Arm and Cl13 as early as d7 post-infection.

To validate and refine our bulk TF analyses (Figures 2 and 1D), we applied the same motif-based predictive modeling to the pseudo-bulk signal aggregated from each scATAC-seq cluster, separately for acute and chronic infection. This identified TFs

Figure 3. scATAC-seq analysis reveals heterogeneity and overlapping but divergent CD8 T cell responses to acute and chronic immune challenges

(A) Experimental setup.

(B) UMAP of TF-IDF-transformed scATAC-seq data.

(C) Louvain clustering of scATAC-seq data.

(D) Single-cell heatmap showing the naive cell signature from bulk ATAC-seq data.

(E) Association by ssGSEA of batch-effect-corrected bulk ATAC-seq data with normalized cluster-aggregated scATAC-seq signal in each of the two samples (Z score row normalized).

(F) Fraction of cells in each sample that belong to each cluster (restricted to clusters 1–8, 10, and 13).

(G) Inferred TF motif coefficients (with the highest variance, Z score row normalized) for cluster-aggregated scATAC-seq signal (for clusters 1–8, 10, and 13) in each of the two samples.

(H) Progenitor dysfunctional signature derived from bulk ATAC-seq data scored in scATAC-seq data for cells in clusters 1–8, 10, and 13 (single-cell heatmap, top; violin plot for cluster-aggregated signal, bottom).

(I) Genome browser tracks for selected peaks. Bulk ATAC-seq for progenitor and terminally dysfunctional cells and for terminal MPECs. Cluster-aggregated scATAC-seq for clusters 1–8 and 10.

(J) Progenitor dysfunctional and MPEC signatures derived from bulk ATAC-seq data scored in cells from cluster 10 in acute and chronic infection (Mann-Whitney U test).

See also Figures S4 and S5 and Table S4.



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most strongly associated with functional states in early response to acute or chronic infection (Figures 3G and S4F).

Cells in cluster 10, when compared with bulk ATAC-seq data, were most similar to progenitor dysfunctional cells profiled at late chronic infection (Figures 3E and 3H; Method details), suggesting the presence of a progenitor-like state early in infection by Cl13 (Utzschneider et al., 2020). Surprisingly, cluster 10 also contained a small number of cells from acute infection that were similar to memory precursor cells (Joshi et al., 2007; Vodnala et al., 2019) based on comparison with bulk ATAC-seq data (Figures 3F, 3H, S4E, S5A, and S5B). Cluster 10 cells from acute and chronic infection displayed similar genomewide chromatin accessibility profiles (and thus clustered together); TF activities via enrichment of Lef1/Tcf7, Nr4a, Nfkb, Nfat, Pou, and AP1 family motifs (Figure 3G); and accessibility at genes important for T cell activation and function, including progenitor marker Cxcr5 and dysfunction marker Tox (Figures 3I and S5A-S5C), consistent with bulk ATAC-seq analysis (Figures 1D, 2B, and 2C). This shared cluster suggested that progenitor-like cells, by analogy to memory precursor cells, are established as early as at d7, as recently reported (Miller et al., 2019; Beltra et al., 2020; Utzschneider et al., 2020). Furthermore, employing epigenetic signatures from mice enabled a characterization of progenitor dysfunctional T cells in human tumors using scATAC-seg data (Satpathy et al., 2019) (Figures S5D-S5G; Method details). However, we also observed differential accessibility between the progenitor-like cells in chronic and acute infection (Figures 3J and S5C) and potentially different TF activity (Figure 3G), suggesting an even earlier divergence between functional and dysfunctional fates.

By complementing our bulk ATAC-seq analysis across models, scATAC-seq analysis characterized the early divergence between CD8 T cell chromatin states in response to acute and chronic immune challenges, identified progenitor-like subpopulations in acute and chronic infection, and implicated cellstate-specific TFs.

Single-cell transcriptional analysis confirms a progenitor/precursor T cell population in early response to both acute and chronic infection

To further explore the heterogeneity and temporal progression of CD8 T cell responses at single-cell resolution and to validate our state-specific TF predictions through allele-specific analyses, we performed scRNA-seq analysis of CD8 T cells from hybrid mice generated upon breeding C57BL/6J and SPRET/EiJ mice (F1[B6xSpret]) at three time points during acute and chronic



LCMV infection (Figures 4A and S6). This approach takes advantage of widespread natural genetic variation (~40 million SNPs) between the evolutionary distant parental genomes and assessing its effect on gene expression and chromatin accessibility. Virus-specific T cells were profiled before infection (d0) and upon activation at d7 and d40 during the acute LCMV infection, and activated CD62L-negative cells were profiled at d7 and d35 during the chronic LCMV infection. Cells isolated from chronically infected B6 mice at d35 were used as a control. Bulk ATAC-seq profiling of CD8 T cells at d0, d7, and d60 upon acute LCMV infection in F1(B6xSpret) mice (van der Veeken et al., 2019) confirmed their consistency with the counterparts from B6 mice (Figure S7A).

After scRNA-seq filtering steps (Method details), our dataset consisted of 9,822 genes profiled in 24,400 cells, ranging from 3,419 to 4,824 cells per sample, with median 1,147 expressed genes per cell. UMAP embedding and Louvain clustering revealed a well-separated naive state and diverging phenotypic arms in acute and chronic responses with heterogeneity within each sample and gradients of expression of well-known T cell function markers potentially reflecting differentiation trajectories (Figures 4B–4D, S7B, and S7C). Cells at d35 in chronic infection from B6 and F1(B6xSpret) mice were highly similar (Figures 4B-4D, S7D, and S7E). Dimension reduction using other well-known methods produced similar results (Figures S7F-S7H).

We used a combination of complementary approaches to functionalize the scRNA-seq data (Figures 4E, 4F, S8, and S9A; Table S5; Method details). Our analysis identified cell clusters encompassing archetypal CD8 T cell subsets including naive T cells, memory precursor cells, short-lived effector cells, central memory cells, effector memory cells, terminally dysfunctional cells, and their precursors. This cellular spectrum is similar to, but more detailed than, previously published scRNA-seq datasets of CD8 T cell responses in acute or chronic infection (Chen et al., 2019c; Miller et al., 2019; Yao et al., 2019; Kurd et al., 2020; Milner et al., 2020), showing that these cell states are consistently generated across independent studies and in mice of distinct genetic backgrounds.

Consistent with our scATAC-seq analysis (Figure 3), our scRNA-seq analysis identified a subpopulation of cells (cluster 9) (Figure 4G) resembling the previously characterized progenitor dysfunctional cells based on the enrichment of signature genes such as Id3, Tcf7, Slamf6, and Cxcr5 (Figures 4E-4H and S9B) and comparison with bulk RNA-seq from sorted subpopulations (Figure S9A). Many genes significantly more accessible and expressed in progenitor dysfunctional cells in bulk

Figure 4. scRNA-seq analysis uncovers phenotypic heterogeneity of CD8 T cell response and cell-state-specific TF expression (A) Experimental setup.

(C) Louvain clustering of scRNA-seq data.

- (G) MAGIC-imputed gene expression in individual samples.
- (H) Log2 fold change of expression in individual samples of selected genes in cluster 9 cells versus all other non-naive cells.
- (I) Cells at d7 in acute and chronic infection separately and their cluster composition, within overall UMAP.
- (J) MAGIC-imputed expression of selected genes encoding TFs.

See also Figures S6-S11 and Table S5.

⁽B) UMAP of normalized scRNA-seq data.

⁽D) Barplot of the number of cells in each cluster from each sample.

⁽E) Cluster-aggregated scRNA-seq gene expression in each cluster for differentially expressed genes between clusters (Z score row normalized).

⁽F) MAGIC-imputed gene expression for selected genes (MAGIC, Markov Affinity-based Graph Imputation of Cells [van Dijk et al., 2018]).



RNA-seq analysis and bulk and scATAC-seq analysis (Figures 1D, 3I, and S5) were enriched in scRNA-seq cluster 9. Thus, cluster 9 consisted of progenitor and progenitor-like cells with a mixed gene expression profile resembling naive, recently activated, dysfunctional, and memory cells.

Finally, we wanted to validate the similarity of progenitor-like cells in early chronic infection and memory precursor cells, their counterpart cells in acute infection, as observed in scATAC-seq analysis (Figures 3E–3I, S4E, S5A, and S5B). Indeed, scRNA-seq cluster 9 contained cells from all non-naive samples (Figures 4D and S9C). Despite the limited resolution of differential expression analysis for small subsets of cells, many critical marker genes were consistently overexpressed in this cluster when considered independently in d7 acute, d7 chronic, or d35 chronic infection, including TFs and putative drivers of chromatin accessibility changes such as Tox, Pou2f2, and Ikzf2; the most overexpressed gene in this cluster, Xcl1; previously described progenitor markers Cxcr5 and Slamf6; and memory precursor marker II7r (Figures 4G, 4H, and S9D). This again confirmed the emergence of progenitor-like cells early in chronic infection, together with a small number of similar cells in acute infection that clustered with progenitor dysfunctional cells, consistent with recent reports (Chen et al., 2019b; Utzschneider et al., 2020). We independently confirmed the presence of this CD8 T cell subpopulation at d7 in acute infection by reanalyzing three recently published scRNA-seq datasets (Yao et al., 2019; Kurd et al., 2020; Chen et al., 2019c) and verified enrichment of genes Tox, Ikzf2, and Cxcr5 in this subpopulation (Figure S10). Trajectory analysis using RNA velocity, although noisy, suggested that progenitor cells arise from less differentiated cells that are only sparsely captured in our dataset (Figure S11).

Thus, our scRNA-seq analysis provided a comprehensive atlas of CD8 T cell functional and dysfunctional states and confirmed the existence of progenitor/progenitor-like cell populations with similar transcriptional profiles in acute and chronic infection.

scRNA-seq characterizes overlapping but divergent CD8 T cell responses to acute and chronic viral infection and cell-state-specific TF expression

We next sought to characterize the divergence of TF drivers of acute and chronic immune responses using our scRNA-seq dataset. Consistent with our scATAC-seq analysis (Figure 3), cells were already highly heterogeneous and markedly different transcriptionally at d7 in acute and chronic infection (Figure 4I; Table S5). Thus, although acute infection predominantly drives CD8 T cell response along the effector cell trajectory, a small fraction of responding cells differentiate toward memory or even potentially along the dysfunctional trajectory. Conversely, chronic infection infrequently drives cells into an effector cell trajectory, consistent with previous reports (Chen et al., 2019c). Together, our observations suggest that both chronic and acute infection give rise to cells at similar functional, activation, and differentiation states, but these cells accumulate at different proportions depending on the challenge.

Cell-state-specific enrichment of TF expression (Figures 4J and S7C) was suggestive of cell-state-specific function, consistent with observations from bulk and scATAC-seq data (Figures

1D, 2B, 2C, and 3G). For example, we identified Gabpa as effector specific; Rora as memory specific; factors Tcf1, Nr4a1, Nfat5, Id3, Pou2f2, Pou6f1, Nfkb1, and Batf as specific for progenitor dysfunctional state; and Zfp143 and Zfp523 as specific for terminal dysfunction. Thus, our scRNA-seq data support the cell-state-specific expression of TF drivers described in our bulk and scATAC analyses.

Allele-specific analysis validates cell-state-specific TF activity

To complement our observations of cell-state-specific TF activities from bulk and single-cell ATAC-seq and RNA-seq data analyses, we exploited our scRNA-seq profiling in F1 hybrid mice via allele-specific analysis of gene expression and TF binding, following and extending our previous studies (van der Veeken et al., 2019, 2020). Indeed, we identified thousands of genes with significant allele-specific expression in many scRNA-seq clusters (Figure S12A), including many genes involved in T cell activation and function (Figure 5A). Most genes were consistently imbalanced in their expression toward B6 or Spret allele across clusters, while a few genes, including Gzmk or Fas, were significantly imbalanced in a cluster-dependent manner. Allelic imbalance of gene expression between B6 and Spret was correlated with differential expression between B6 and F1(B6xSpret) mice, supporting the accuracy of our allelic imbalance estimates (Figure S12B).

To gain evidence for the causal role of TFs in regulating gene expression, we looked for an association between allele-specific TF binding, as predicted by motif analysis, and allele-specific expression (Figures 5B and 5C). For each TF motif match in the B6 or Spret sequence around an ATAC-seq peak summit, we calculated the TF motif binding imbalance as the difference in motif log-odds scores between B6 and Spret, Z score-normalized. Many genes (e.g., Gzmk) (Figure 5C) had many TFs with predicted binding imbalance at their promoter and enhancer ATAC-seq peaks. We expected that genes with only a few potential TF regulators would yield stronger association between allele-specific TF binding and allele-specific expression. Accordingly, when we restricted to genes with a single TF with predicted allele-specific binding, this allele-specific binding was concordant with allele-specific expression (Figure 5D). Considering genes with at most 20 ATAC-seq peaks, we found that imbalanced binding of certain TF motifs was significantly associated with gene expression imbalance. For example, genes with stronger TCF1/LEF1 promoter binding to the Spret allele were significantly more imbalanced in their expression toward Spret than were genes with stronger binding of the same motif in B6 (Figure 5E). This suggests that TCF1/LEF1 binding at promoters is associated with activation of expression. Overall, this approach associated TF binding with activation or repression of gene expression for multiple factors (Figures 5F and S12C-S12E).

TCF1 was reported as a critical marker of progenitor dysfunctional cells and is required for their generation during chronic infection (Im et al., 2016; Utzschneider et al., 2016). To experimentally validate that TCF1 binding is associated with activation of gene expression, we mapped 3,325 TCF1 bound sites by CU-T&RUN analysis in progenitor dysfunctional cells from

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Figure 5. Allele-specific scRNA-seq analysis reveals cis-regulation of gene expression by TFs

(A) Allelic imbalance of gene expression in scRNA-seq data clusters (color, p < 0.05, Mann-Whitney U test).

(B) Schematic for analysis of association between allelic imbalance of motif-based TF binding and gene expression.

(C) Left: ATAC-seq from B6/Spret F1 mice at Gzmk locus and examples of peaks with B6-specific (green), Spret-specific (brown) and ambiguous (gray) ATAC-seq signal and allele-specific predicted TF binding. Right: allelic specificity of Gzmk expression with significant B6-specific (green) and Spret-specific (brown) expression highlighted.

(D) Predicted allelic imbalance of TF binding (difference between TF motif log odds scores, *Z* score normalized) for genes with significant allele-specific expression in any of the scRNA-seq clusters (for genes with a single imbalanced motif); p-value, Mann-Whitney U test.

(E and F) Allele-specific expression analysis of scRNA-seq data. CDF plots: allelic imbalance between B6 and Spret of normalized cluster-aggregated scRNA-seq signal, for genes predicted to be bound more strongly in B6 or Spret using sequence motif analysis in promoter peak summit regions. Barplots: summary of the above analysis for each TF motif over clusters (black bars, Kolmogorov-Smirnov, p < 0.05; CDF, cumulative distribution function).

(G) Examples of allele-specific TCF1 CUT&RUN binding sites in progenitor dysfunctional cells in the established chronic infection.

(H) Allelic imbalance between B6 and Spret of normalized scRNA-seq signal in cluster 9 at d35 in chronic infection, for genes bound by TCF1 more strongly in B6 or Spret as measured by CUT&RUN in progenitor dysfunctional cells (p-value, Kolmogorov-Smirnov).

See also Figures S12 and S13 and Table S6.



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Figure 6. Progenitor CD8 T cells from chronic infection transferred to acute infection are committed to dysfunction (A) Setup for the adoptive cell transfer experiment.

(B and C) Flow cytometry of selected genes for progenitor cells before transfer (B) and after transfer and expansion (C) in acute infection. (D) Quantification of flow cytometry results for donor cells after transfer and expansion.

(E) Left: UMAP of all scRNA-seq data including the expanded donor cells; right: UMAP showing individual samples within the overall map.

(F) New Louvain clustering of all scRNA-seq data, with the barplot showing the number of cells in each cluster from each sample.

(G) Comparison of new clustering with the previous one (Figure 4C). Shown is a fraction of cells in clusters t0-t23 that belong to previously obtained clusters 0-21. (H) Fraction of cells in each sample that belong to the progenitor cluster t4.

(I) Cluster composition of samples. Heatmap shows for each cluster what fraction of cells (excluding naive) that cluster occupies in each sample. See also Table S5.

F1(B6xSpret) hybrid mice at d35 upon Cl13 infection (Figure S13A; Table S6). Direct TCF1 binding sites were strongly enriched for the TCF1 motif (Figure S13B) and included many T cell

activation genes and progenitor markers (Figures S13C and S13D), including some targets with allele-specific TCF1 binding such as Tox, Tox2, Ifng, and Nfkb1 (Figures 5G and S13E).

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Consistent with previous observations and our predictions (Figure 2), accessibility levels at nearly all TCF1 targets were much higher in progenitor cells than in terminally dysfunctional cells (Figure S13F). Surprisingly, these accessibility changes were not associated with changes in expression of nearby genes (Figure S13F). However, allele-specific TCF1 binding was significantly associated with allele-specific expression of nearby genes in progenitor dysfunctional cells in established chronic infection (Figure 5H), consistent with a causal role for TCF1 in activation of gene expression. Furthermore, allele-specific motif enrichment analysis in CUT&RUN-defined TCF1 binding sites revealed TCF1 binding co-factors, such as Runx and Nfkb family factors (Figure S13G), consistent with allele-agnostic co-factor analysis (Figure S13B).

Thus, our allele-specific analysis in the hybrid genome revealed *cis*-regulatory effects of TF binding on gene expression in CD8 T cells that are missed with standard approaches.

Progenitor cells differentiate to terminal dysfunction under acute viral challenge

Adoptive transfer experiments using sorted progenitor dysfunctional cells have demonstrated their potential to self-renew and give rise to terminally dysfunctional cells in models of melanoma and chronic infection (Miller et al., 2019; Im et al., 2016; Beltra et al., 2020). Since single-cell chromatin and expression analyses showed that dysfunctional progenitors resemble progenitor-like cells at the plastic stage of T cell dysfunction development as well as the MPEC population in acute infection, we asked whether progenitor cells retain the potential to give rise to effector cells. We therefore transferred fluorescence-activated cell sorting (FACS) identified progenitor dysfunctional cells isolated from chronically infected mice on d35 post-infection into congenically marked recipients infected with LCMV Arm (Figures 6A and 6B; Method details). The transferred cells were profiled at d7 post-infection using flow cytometry and scRNA-seq. Flow cytometry revealed that the transferred CD8 T cells proliferated and persisted (Figures 6C and 6D). Furthermore, scRNA-seq analysis showed that these cells exhibited heterogeneous gene expression states (Figures 6E-6I). Despite the acute infection setting, the phenotypic space defined by transferred cells after expansion most strongly overlapped with that of cells in chronic infection settings. Clustering analysis of the combined scRNA-seq data (Figures 6F and 6G) confirmed that the expanded transferred cells were most similar in cluster composition and cluster-wise differential expression to cells at d7 in chronic infection (Figure 6I; Table S5). The expanded population lacked effector cells (clusters t3, t10, and t11; Figures 6F and 6I) suggesting the transferred population did not contain naive or memory cells. Importantly, progenitor cells persisted upon transfer, confirming their self-renewal capacity (cluster t4; Figure 6H). Consistent with a recent report (Utzschneider et al., 2020), our singlecell analyses suggest that the progenitor cells observed in chronic infection (d35) were already committed to a dysfunctional fate.

DISCUSSION

Our unified analysis of bulk ATAC-seq and RNA-seq data across diverse studies of T cell response to chronic antigen exposure defined a universal program of progression to terminal dysfunc-



tion. By joint analysis of data from numerous resources, we gained statistical power to robustly identify individual chromatin-accessible sites, patterns of accessibility and expression changes common across experimental models, and TFs whose binding motifs explain global accessibility patterns in distinct chromatin states in T cell differentiation.

Given the universality of T cell dysfunction program across mouse models, we then focused on the LCMV infection as a model to elucidate the early divergence in chromatin and transcriptional states between dysfunctional and functional T cell responses at single-cell resolution using scATAC-seq and scRNA-seq. We found that a subset of antigen-specific T cells in chronic LCMV infection are appropriately activated and become effectors at d7, while most progress along a dysfunctional trajectory. Conversely, a small number of cells in acute LCMV infection had progressed to a differentiated effector memory state and potentially also to a dysfunctional state by d7. These findings, consistent with previous smaller-scale scRNAseq studies of functional or dysfunctional T cells, enable a more nuanced comparison of acute versus chronic immune responses, where T cells can commit to multiple fates at differing frequencies.

In our single-cell analyses, we found a TCF1⁺ progenitor-like cell population emerging as early as d7 post-infection in both chronic and acute infection. This population formed the shared cluster across the two infection settings in both scATAC-seq and scRNA-seq data and therefore displayed the strongest transcriptional and epigenomic similarity among all the cells profiled, in contrast with recent analysis of sorted populations (Utzschneider et al., 2020). Accordingly, in both settings, we found the same genes enriched in expression and chromatin accessibility at multiple promoter and enhancer peaks in this population, including Tcf7, Xcl1, Cxcl10, Ccr7, Slamf6, Id3, and Cxcr5, but also Tox and Ikzf2, which had previously been associated with terminally dysfunctional T cells. Notably, multiple lines of evidence implicated several TFs as strongly associated with the progenitor and progenitor-like population (Figures 1D, 2B, 2C, 3G, 4J, 5E, 5F, S5B, and S9B), including the previously described TCF1 and the newly identified OCT2 (encoded by Pou2f2). However, differential accessibility analysis and motif regression modeling, despite being supported by analysis of relatively small cell numbers, identified subtle differences between cells from chronic and acute samples in this progenitorlike subpopulation, with cells from acute infection potentially representing an IL7R^{hi} MPEC subset and those from chronic infection forming a nascent progenitor reservoir (Figures 3G, 3J, and S5C). These findings may suggest an earlier divergence between progenitor and MPEC populations at the level of chromatin state and gene expression.

The presence of cells from both chronic and acute viral infection in the scRNA-seq cluster containing TCF1⁺ PD1⁺ progenitor cells and in the scATAC-seq progenitor-like cluster led us to test the plasticity of the progenitor cell population. Using an adoptive cell transfer experiment together with scRNA-seq profiling and mapping these data to the comprehensive scRNA-seq atlas of CD8 T cell responses that we constructed, we confirmed that progenitor cells isolated from an established chronic viral infection were committed to a dysfunctional fate. While transferred



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progenitor cells could proliferate in a new host under acute immune challenge, they displayed the single-cell phenotypic profile of a chronic immune response. These results are consistent with earlier adoptive T cell transfer studies showing that by d15 postantigen encounter, T cells that had responded to chronic LCMV infection lost their ability to form functional memory under acute LCMV challenge (Angelosanto et al., 2012) and with similar experiments documenting loss of plasticity for tumor-specific T cells in an autochthonous liver cancer model (Schietinger et al., 2016). However, these earlier studies also confirmed that at early time points in settings of chronic antigen stimulation, antigen-specific T cells-or at least a subpopulation of these cells-retained the potential to mount a functional response and form memory cells. Our single-cell analyses of T cells under chronic immune challenge suggest this transient plasticity resides within the early progenitor-like population, consistent with recent phenotypic studies (Utzschneider et al., 2020).

Trajectory analysis using RNA velocity remains noisy. Further studies of the origins and regulatory mechanisms of maintenance and differentiation potential of the progenitor and progenitor-like cell populations are warranted, (e.g., using heritable bar codes for tracking cell fates within a TCR clonal population over time [Wagner and Klein, 2020] and at time points earlier than d7 upon T cell activation). Performing these studies in F1 hybrid mice can help further interrogate regulatory mechanisms of gene expression by linking allele-specific analysis of TF motif occurrences (or allele-specific TF occupancy) to allelic imbalance of target gene expression.

In conclusion, our datasets and analyses lay the groundwork for resolving the fundamental questions of T cell differentiation programs. We expect that our unified T cell atlas will provide a valuable resource both for basic T cell biologists and for cancer immunologists seeking to therapeutically target the transcriptional programs underlying progression to fixed dysfunction.

Limitations of the study

Sorted populations subjected to bulk profiling are still a mixture of the purer subpopulations we characterized with single-cell analysis, complicating interpretation of the bulk analysis. While our batch-effect correction succeeded in finding strong similarities and some differences between models, generating all data in a controlled manner within the same lab remains the gold standard for differential analyses. We limited our single-cell analyses to the LCVM model to enable comparison of acute and chronic responses; however, similar single-cell analyses in a tumor model would directly address whether the underlying distribution over T cell states differs in cancer versus chronic infection. T cell differentiation is a particularly difficult setting for computational trajectory analysis: subpopulations rapidly expand or collapse, and overall differences between functional states can be subtle. While we provide results of RNA velocity analysis, we believe that direct experimental determination with cell tagging approaches will be needed to resolve differentiation trajectories.

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
- Mice
- METHOD DETAILS
 - Construction of ATAC-seq peak atlas
 - Batch effect correction for ATAC-seq
 - Differential accessibility analysis
 - Transcription factor motif analysis
 - O Predictive modeling of transcription factor activity
 - O RNA-seq data analysis
 - Comparison of ATAC-seq and RNA-seq using Hi-C contacts
 - Single-cell ATAC-seq experiments
 - Single-cell ATAC-seq data preprocessing, dimensionality reduction, clustering
 - Comparison of scATAC-seq with bulk ATAC-seq data compendium
 - Isolation of cells for scRNA-seq
 - Adoptive transfer of progenitor dysfunctional cells
 - Single-cell barcoding, library preparation and sequencing
 - Single-cell RNA-seq data preprocessing, dimensionality reduction, clustering
 - Overview of functional annotation of scRNA-seq data
 - Differential expression analysis of scRNA-seq data
 - Comparison of scRNA-seq with bulk RNA-seq data compendium
 - Analysis of scRNA-seq data from adoptive cell transfer experiment
 - Allele-specific scRNA-seq expression
 - Allele-specific predicted TF binding
 - CUT&RUN experiments
 - Identification of transcription factor binding sites in CUT&RUN data
 - Allele-specific analysis of CUT&RUN data
 - Construction of human ATAC-seq peak atlas and scA-TAC-seq count matrix
 - Analysis of human scATAC-seq data
 - Comparison of differential accessibility between mouse and human
- RNA velocity analysis
- QUANTIFICATION AND STATISTICAL ANALYSIS

SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

Conceptualization, Y.P., J.v.d.V., A.Y.R., and C.S.L.; methodology, Y.P., J.v.d.V., A.R.P., Y.Z., M.S., L.M., D.P., A.Y.R., and C.S.L.; software, Y.P., A.R.P., Y.Z., M.S., and C.S.L.; formal analysis, Y.P., A.R.P., and C.S.L.; investigation, Y.P., J.v.d.V., A.R.P., Y.Z., M.S., L.M., D.P., A.Y.R., and C.S.L.; writing – original draft, Y.P. and C.S.L.; writing – review & editing, Y.P., J.v.d.V., A.Y.R., and C.S.L.; supervision, Y.P., J.v.d.V., A.Y.R., and C.S.L.; funding acquisition, Y.P., D.P., A.Y.R., and C.S.L.

DECLARATION OF INTERESTS

A.Y.R. is an SAB member; has equity in Sonoma Biotherapeutics, RAPT Therapeutics, Surface Oncology, and Vedanta Biosciences; and is a co-inventor or has IP licensed to Takeda that is unrelated to the content of this study.

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STAR*METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
CD8α - Brilliant Violet 605	BioLegend	Cat#100744; RRID: AB_2562609
TCRβ - PE-eFluor 610	ThermoFisher	Cat#61-5961-82; RRID: 2574644
CD44 - APC	Tonbo Bioscience	Cat#20-0441-U100; RRID: AB_2621572
CD62L - APC-eFluor780	ThermoFisher	Cat#47-0621-82; RRID: AB_1603256
NK1.1 - eFluor 450	ThermoFisher	Cat#48-5941-82; RRID: AB_2043877
CD4 - PerCP-Cyanine5.5	Tonbo Bioscience	Cat#65-0042-U100; RRID: AB_2621876
PD-1 - FITC	ThermoFisher	Cat#11-9985-82; RRID: AB_465472
CD39 - PerCP-eFluor 710	ThermoFisher	Cat#46-0391-80; RRID: AB_10717513
CD73 - eFluor 450	ThermoFisher	Cat#48-0731-82; RRID: AB_10853356
CD9 - PE	BioLegend	Cat#124805; RRID: AB_1279327
CD38 - APC	ThermoFisher	Cat#17-0381-82; RRID: AB_469382
Rabbit anti-TCF1 monoclonal	Cell Signaling Technology	Cat#2203S; RRID: AB_2199302
anti-mouse CD4	BIOXCELL	Cat#BE0003-1; RRID: AB_1107636
Bacterial and virus strains		
LCMV Armstrong and Clone 13	Produced in-house	N/A
Chemicals, peptides, and recombinant proteins		
D(b) NP396 LCMV Tetramers	NIH Tetramer Core Facility	N/A
Agencourt AMPure XP - PCR Purification	Beckman Coulter	Cat#A63881
Ghost Dye Live/Dead stain	Tonbo Biosciences	Cat#13-0870-T100
Protein A/G-MNase	Produced in-house	N/A
Complete mini EDTA-free Protease Inhibitor	Sigma-Aldrich	Cat#11836153001
Critical commercial assays		
eBioscience Foxp3/Transcription Factor Staining Buffer Set	Thermo Fisher	Cat#00-5523-00
MinElute Reaction Cleanup Kit	QIAGEN	Cat#28206
KAPA Hyper Prep Kit	Kapa Biosystems	Cat#KK8504
KAPA UDI adaptor kit	Kapa Biosystems	Cat#KK8727
Deposited data		
scATAC-seq, scRNA-seq, CUT&RUN	This paper	NCBI GEO: GSE164978
ATAC-seq, RNA-seq	NCBI GEO	Accession numbers listed in Table S1
Experimental models: Organisms/strains		
Spret/EiJ	JAX	001146
CD45.1	JAX	002014
C57BL/6J	JAX	000664
Software and algorithms		
Flowjo	Tree Star	https://www.flowjo.com
Prism	GraphPad	https://www.graphpad.com/scientific- software/prism/
Illustrator	Adobe	N/A
fastq-dump	NCBI SRA Toolkit	https://trace.ncbi.nlm.nih.gov/Traces/sra/ sra.cgi?view=toolkit_doc
Bowtie2	Langmead and Salzberg, 2012	http://bowtie-bio.sourceforge.net/bowtie2
SAMtools	Li et al., 2009	http://www.htslib.org/

Resource



Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
MACS2	Zhang et al., 2008	https://github.com/macs3-project/MACS
IDR	Li et al., 2011	https://github.com/nboley/idr
Rsubread	Liao et al., 2019	https://bioconductor.org/packages/ release/bioc/html/Rsubread.html
deepTools	Ramírez et al., 2016	https://github.com/deeptools/deepTools
DESeq2	Love et al., 2014	https://bioconductor.org/packages/ release/bioc/html/DESeq2.html
gprofiler2	Raudvere et al., 2019	https://cran.r-project.org/web/packages/ gprofiler2/index.html
HOMER	Heinz et al., 2010	http://homer.ucsd.edu/homer/
FIMO	Grant et al., 2011	https://meme-suite.org/meme/doc/ fimo.html
mpath	Wang et al., 2014	https://cran.r-project.org/web/packages/ mpath/index.html
HISAT2	Kim et al., 2019	http://daehwankimlab.github.io/hisat2/
HiC-Pro	Servant et al., 2015	https://github.com/nservant/HiC-Pro
HiC-DC	Carty et al., 2017	https://bitbucket.org/leslielab/hic.dc
cellranger-atac	10X Genomics	https://support.10xgenomics.com/single- cell-atac/software/downloads/latest
scanpy	Wolf et al., 2018	https://github.com/theislab/scanpy
GSVA	Hänzelmann et al., 2013	https://www.bioconductor.org/packages/ release/bioc/html/GSVA.html
MAGIC	Van Dijk et al., 2018	https://github.com/ KrishnaswamyLab/MAGIC
STAR	Dobin et al., 2013	https://github.com/alexdobin/STAR
cellranger	10X Genomics	https://support.10xgenomics.com/single- cell-gene-expression/software/downloads/ latest
velocyto	La Manno et al., 2018	http://velocyto.org/
scvelo	Bergen et al., 2020	https://scvelo.readthedocs.io/

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Christina Leslie (cleslie@cbio.mskcc.org).

Materials availability

This study did not generate new unique reagents.

Data and code availability

The code for computational analysis and additional data visualizations are available at https://bitbucket.org/leslielab/cd8t_atlas. The accession number for the new scATAC-seq, scRNA-seq and CUT&RUN data reported in this paper is NCBI GEO: GSE164978. Previously published datasets reanalyzed in this study are listed in Table S1.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

Animals were housed at the Memorial Sloan Kettering Cancer Center (MSKCC) animal facility under specific pathogen free (SPF) conditions on a 12-hour light/dark cycle under ambient conditions with free access to food and water. All studies were performed under protocol 08-10-023 and approved by the MSKCC Institutional Animal Care and Use Committee. Mice used in this study had no previous history of experimentation or exposure to drugs. Male Spret/EiJ mice were purchased from Jackson laboratory and bred to female CD45.1 mice on a C57BL/6 genetic background. Adult (at least 8 weeks old) male and female F1 offspring were used for experiments as indicated.



Molecular Cell Resource

METHOD DETAILS

Construction of ATAC-seq peak atlas

ATAC-seq data from multiple previous studies were downloaded from GEO using fastq-dump from NCBI SRA Toolkit (https://trace. ncbi.nlm.nih.gov/Traces/sra/sra.cgi?view=toolkit_doc). Functional annotation of samples (naive, effector, memory, dysfunctional, etc.) was obtained from each study (Table S1). Reads were aligned to the mouse genome mm10.GRCm38 using bowtie2 v2.3.4.3 (Langmead and Salzberg, 2012). Uniquely aligned reads were extracted using SAMtools v1.9 (Li et al., 2009). Peaks were called using MACS2 v2.1.1.20160309 (Zhang et al., 2008). For each data source, peaks were called using all samples from all replicates combined. Then IDR (Li et al., 2011) was used to identify reproducible peaks between at least one pair of replicates of the same condition. The union of all such peaks formed the peak library for each study. Then peaks identified in each study were all combined into a common list, by taking the union of all genomic positions covered by peaks from different studies using function reduce(c()) in R (R v3.4.0 was used in all our analysis except when otherwise noted). Peaks were further split into subpeaks around summits of signal, using a custom script based on the signal processing package https://bitbucket.org/leslielab/biosignals. Peak summit regions were then filtered based on ATAC-seq read coverage of 150bp regions centered at the summits, by including only the peak summit regions with at least 100 reads in at least one sample and at least 5 reads on average across all samples within any of the three categories (naive, functional, and dysfunctional). Peaks that included at least one of such summit region were included in the final atlas. This resulted in an atlas of 129799 chromatin accessibility peaks for CD8 T cells. Peaks were annotated using GENCODE vM14 (Frankish et al., 2019). Each peak was associated with the closest gene, if this gene was within 50Kb in genomic coordinates. Peaks were annotated by applying the following sequence of rules: peaks were classified as promoter peaks if within 2Kb from a transcription start site of any annotated transcript; otherwise as exonic if overlapping with any exon of any annotated transcript; otherwise as intronic if within a gene body of any annotated gene; or finally as intergenic if within 50Kb of a gene. All the remaining peaks were left unclassified. Reads from each sample were counted in peaks and 150bp genomic regions centered around peak summits using Rsubread v1.32.4 (Liao et al., 2019). For visualization, bigWig files and count matrices in bins around peak summits were produced using deep-Tools (Ramírez et al., 2016). Snapshots of ATAC-seq signal at selected genomic regions were obtained from UCSC Genome Browser (Kent et al., 2002).

Batch effect correction for ATAC-seq

DESeq2 v1.22.2 (Love et al., 2014) was used to fit multi-factorial models to ATAC-seq read counts in peaks or peak summits. The main model included two factors where one factor represented the source of the data, and the other factor represented the functional annotation of the sample, broadly defined as naive, functional, and dysfunctional. For PCA analysis and visualization, batch effect-corrected values were used. To do the batch effect correction, log2FC values associated with the data source factor were extracted from the model for all peaks and used to correct the original counts, by dividing them by the exponentiated log2FC values.

The alternative variant of the model included a two-valued functional annotation factor, where one value was for naive samples, and the other for all antigen-experienced cells, including all functional and dysfunctional cells. This alternative variant of the model produced results similar to the main model (Figure S1C).

In order to determine whether memory precursor cell samples and tissue-resident memory cell samples were similar to functional or dysfunctional cells, a variant of this alternative model with a two-valued functional annotation factor was used. For this, differentially accessible peaks were detected between functional and dysfunctional cells but excluding memory precursor and tissue-resident memory cell samples. Then PCA analysis for all samples was run on ATAC-seq read counts restricted to these differentially accessible peaks, revealing that memory precursor and tissue-resident memory cells are more similar to other functional effector and memory cells.

Differential accessibility analysis

DESeq2 v1.22.2 was used to perform differential accessibility analysis between selected pairs of conditions. For this, original counts were used, and the model included factors for functional annotation, cell state, data source, and/or type of immune challenge, where appropriate. For each gene, overall differential accessibility of its peaks between a certain pair of cell states was assessed as follows. A Mann-Whitney U test was used to compare the vector of all peak accessibility log2 fold changes between these two cell states with the vector of log2 fold changes only for the peaks associated with the gene. Then the resulting p values (one for each gene) were adjusted for multiple hypothesis testing (multiple tested genes) using q-value. This analysis was visualized with a scatterplot where the x axis shows the log2 fold change of gene expression, the y axis shows mean accessibility log2 fold change of the peaks associated with a gene, black indicates significant differential expression (FDR < 0.05), and red and blue indicate significant overall differential accessibility of peaks associated with a gene (q < 0.01), only for significantly differentially expressed genes. For functional analysis, the most significantly differentially accessible genes (q < 0.01) were selected, split into those with positive and negative average differential accessibility across the gene's peaks, and functional enrichment assessed for these genes using gprofiler2 with the background of all expressed genes (Raudvere et al., 2019). Genes significantly more accessible in terminally dysfunctional cells in thronic infection as compared with terminally dysfunctional cells in the tumors (Figure S2D) were most significantly enriched for GO terms such as "immune system process" (rank 1, p < 3e-19) and "T cell activation" (rank 4, p < 6e-14). Genes significantly

Resource



more accessible in endogenous than in CAR-T dysfunctional cells (Figure S2D) were often associated with terminal dysfunction as evidenced by higher accessibility at loci of important dysfunction marker genes (Tox, Havcr2, Entpd1, Tigit).

Transcription factor motif analysis

Transcription factor (TF) binding motifs for *Mus musculus* were downloaded from CIS-BP version 1.02 (Weirauch et al., 2014) via the web interface (compressed archive Mus_musculus_2016_06_01_2-46_pm.zip). For the DNA sequences in 150bp-wide regions around peak summits, script findMotifsGenome.pl from HOMER suite (Heinz et al., 2010) was run with parameters "mm10 -len 8,10,12 -size given -S 100 -N 1000000 -bits -p 10 -cache 1000" in order to identify the significance of presence of each motif in the sequences of the peaks as compared with the background sequences. We limited the analysis to motifs corresponding to expressed TFs, defined as those with at least 200 library-size normalized RNA-seq reads in at least one condition in at least one study. We focused only on the motifs present in at most 50% of the peak summit region sequences. Furthermore, we detected *de novo* motifs using HOMER in the same sequences and associated them with TFs by similarity with the CIS-BP motifs using script compareMotifs.pl with parameters "-reduceThresh 0.7 -matchThresh 0.9." The most significant motif per TF, either from the database or identified *de novo*, was selected for further analysis (with potentially multiple TFs associated with the same motif), if it had HOMER p < 0.001. This resulted in 113 motifs. Furthermore, we merged motifs that had correlation of motif occurrences in the peak summits above 0.75. This resulted in the list of 105 motifs corresponding to 204 TFs for further analysis. FIMO version 4.11.2 (Grant et al., 2011) was used to search for motif matches in 150bp regions around the peak summits. Matches with p < 1e–3 were chosen as significant.

Predictive modeling of transcription factor activity

To infer cell-state-specific TF activities, we performed a supervised modeling of chromatin accessibility data based on TF motif occurrences. We formed a feature matrix that consisted of TF motif match predictions in regions around peak summits. Each value in the matrix was a sum of FIMO log-odds scores of a TF motif occurrence in a peak summit region. For each ATAC-seq sample, we performed negative binomial generalized linear regression modeling of the batch-effect corrected chromatin accessibility values y in peak summit regions using this feature matrix, with ridge regularization, using function cv.glmreg() from R package mpath (Wang et al., 2014) with parameters family = "negbin," alpha = 0, theta = 1, nfolds = 5, maxit = 20000, thresh = 1e-5. We limited the regression analysis to peak summit regions with at least 10 batch-effect corrected reads on average across all samples. We identified a hyperparameter multiplier a of the ridge regularization penalty term using 5-fold cross-validation; for this we ran the regression with 30 values of a formed by multiplying the mean of y by a vector of values from 10^{-8} to 10^3 (with equidistant log₁₀ values) and choosing the value of a that maximizes the Spearman correlation between the observed and predicted values of y (Figure S3B). Then we used the coefficients of this regression as a proxy for TF activity scores and used them for downstream analysis; for this, we limited our downstream analysis to results of regression only for those ATAC-seq samples where the fit converged for at least 4 values of the hyperparameter a.

RNA-seq data analysis

RNA-seq data from multiple previous studies were downloaded from GEO using fastq-dump. Functional annotation of samples (naive, effector, memory, dysfunctional, etc.) was obtained from each study (Table S1). Reads were aligned to the mouse genome mm10.GRCm38 using hisat2 v2.1.0 (Kim et al., 2019). Reads from each sample were counted in genes annotated by GENCODE vM14 using Rsubread v1.32.4. Batch effect correction and differential expression analysis were performed the same as for ATAC-seq data.

Comparison of ATAC-seq and RNA-seq using Hi-C contacts

Published Hi-C data from naive CD8 T cells (He et al., 2016) was used to define associations of ATAC-seq peaks to genes and compare it with our default peak-to-gene association (see Construction of ATAC-seq peak atlas), using a computational experiment comparing ATAC-seq and RNA-seq data. Hi-C read pairs were processed using HiC-Pro pipeline (version 2.11.1) (Servant et al., 2015). Reads were trimmed to 30nt and mapped to the mm10 reference genome. We used valid pairs output in cis from HiC-Pro that were mapped uniquely to the genome. Valid pairs from replicates were merged and binned into 10kb bins. HiC-DC (Carty et al., 2017) was used to assign statistical significance to each interaction bin. Significant chromatin loops between 10Kb bins were defined at q < 0.05 and with support by more than 20 Hi-C valid pairs, resulting in 22,337 significant loops. Median distance between loop ends was 180Kb, and only 5% of loops had distance between ends shorter than 50Kb. For control, non-significant loops were defined as those with q-value between 0.1 and 0.2 and supported by not more than 10 Hi-C valid pairs, resulting in 118,497 loops. Peaks were associated to genes using these loops as follows. For a loop between 10Kb-long genomic regions A and B, a peak was associated to a gene if the peak was within 5Kb window from A and another peak at a promoter of the gene was within 5Kb from B. Then differential gene expression (RNA-seq log2FC) between naive and effector cells was compared between genes associated with at least one significantly more accessible peak in naive than in effector cells and in effector than in naive cells (FDR < 0.05), for different methods of association of peaks to genes: default associations, associations based on significant loops, and associations based on non-significant loops (Figure S3D). The same analysis was performed for comparison between naive and dysfunctional cells (Figure S3D).



Single-cell ATAC-seq experiments

Four male C57BL/6J mice per group were infected with either LCMV Armstrong (2x10⁵ p.f.u. via intraperitoneal injection) or LCMV Clone 13 (2x10⁶ p.f.u. via retroorbital injection). On day 7 post-infection, total CD8 T cells from pooled spleens were double sorted by flow cytometry and prepared for single cell ATAC-seq analysis.

scATAC-seq was performed using Chromium instrument (10x Genomics) and Single Cell ATAC Reagent Kits (Chemistry v1). The suspension of cells was processed following the User Guide (CG000168 Rev A). Briefly, cells were lysed in bulk, washed and resulting nuclei suspension treated with ATAC reagents provided in the kit. Approximately, 8000 nuclei per sample were loaded onto micro-fluidics Chip E and encapsulated with DNA barcodes and reaction mix. Following emulsion-PCR the resulting material was purified and subjected to 12-cycles of indexing PCR. The indexed scATAC-Seq libraries were double-size selected using SPRI beads and sequenced on an Illumina NovaSeq 6000 instrument (Read 1 – 50 cycles, i7 index – 8 cycles, i5 index – 16 cycles, and Read 2 – 50 cycles) at 180M reads per sample.

Single-cell ATAC-seq data preprocessing, dimensionality reduction, clustering

The data was preprocessed with cellranger-atac from 10X. Then using the BAM files with read alignments for the two samples produced by cellranger-atac, peaks were detected using MACS2. An extended list of peaks was formed by combining the previously constructed bulk ATAC-seq peaks with the non-overlapping newly identified peaks. The cellranger-atac tool was then rerun with this extended peak list as input to obtain a count matrix for each of the two samples. Cells with a log library size less than 3.5 and peaks that were accessible in less than 4 cells were excluded from the analysis. The count matrix was then binarized and transformed using term frequency-inverse document frequency (TF-IDF, scikit-learn package v0.20.3, https://scikit-learn.org/) for normalization, followed by PCA and UMAP for dimensionality reduction and visualization. Cells were clustered by the Louvain clustering method. Analysis and visualizations were performed using python package scanpy v1.4.4 (Wolf et al., 2018).

Comparison of scATAC-seq with bulk ATAC-seq data compendium

Signature peak sets for progenitor, naive, effector, MPEC, and terminal dysfunction were derived from bulk ATAC-seq analysis (Table S2) and then used to score cells by taking the average normalized counts of the peak set and subtracting the average normalized counts of a reference peak set. The naive cell signature was defined as peaks significantly more accessible in naive cells in naive versus functional and naive versus dysfunctional cell comparisons (using thresholds baseMean > 50, FDR < 0.001, log2 fold change < -2 for both comparisons). The effector cell signature was defined as peaks significantly more accessible in effector cells in naive versus effector (baseMean > 50, FDR < 0.01, log2 fold change > 1), effector versus memory (baseMean > 10, FDR < 0.05, log2 fold change < 0), and effector versus early dysfunctional (baseMean > 10, FDR < 0.05, log2 fold change < 0) cell comparisons. The memory cell signature was defined as peaks significantly more accessible in memory cells in effector versus memory (baseMean > 50, FDR < 0.001, log2 fold change > 0.5) and memory versus terminally dysfunctional (baseMean > 50, FDR < 0.001, log2 fold change < -2) cell comparisons. The MPEC signature was defined as peaks significantly more accessible in MPEC versus effector cells (baseMean > 50, FDR < 0.001, log2 fold change > 1). The progenitor dysfunctional cell signature was defined as peaks significantly more accessible in progenitor versus terminally dysfunctional cells (baseMean > 100, FDR < 0.001, log2 fold change < -2). The terminally dysfunctional cell signature was defined as peaks significantly more accessible in dysfunctional cells in naive versus dysfunctional (baseMean > 10, FDR < 0.05, log2 fold change > 0), memory versus terminally dysfunctional (baseMean > 10, FDR < 0.05, log2 fold change > 0), progenitor versus terminally dysfunctional (baseMean > 50, FDR < 0.01, log2 fold change > 1), effector versus early dysfunctional (log2 fold change > 0) cell comparisons.

BAM files were generated with aligned scATAC-seq reads corresponding to cells from each cluster, further split between the two samples. Pseudobulk counts were obtained from these BAM files for the extended peak list using Rsubread v1.32.4. Limited only to clusters 1-6, 8, 10, 13 that contained activated virus-specific cells, differential accessibility analysis was then run using these pseudobulk counts for each of the clusters, within each of the two samples, versus all the remaining clusters split between the two samples. The most significantly differentially open peaks formed a signature set of peaks for each cluster split between the two samples. Then ssGSEA (Barbie et al., 2009) was run for these signatures against batch-effect corrected library-size normalized bulk ATAC-seq counts using the package GSVA v1.30.0 (Hänzelmann et al., 2013).

Isolation of cells for scRNA-seq

For isolation of CD8 T cells responding to acute infection, B6/Spret F1 mice, where B6 stands for C57BL/6J and Spret stands for SPRET/EiJ, were infected with 2x10⁵ p.f.u. of LCMV Armstrong via intraperitoneal injection or left uninfected. Splenocytes were stained with a cocktail of fluorescent antibodies, NP396 tetramer, and viability dye to mark dead cells, and sorted using flow cytometry. Naive CD8 T cells were isolated from pooled spleens of 2 uninfected mice as CD44-CD62L+ CD8 T cells. Activated and memory effector cells were isolated as NP396+ CD8 T cells from pooled splenocytes of 2 or 3 mice on day 7 or day 40 post-infection, respectively and used as input for scRNA-seq analysis.

For isolation of cells responding to chronic infection, B6/Spret F1 or B6 mice were infected with LCMV clone 13 by retroorbital injection of 4x10⁶ p.f.u. of virus. Mice were depleted of CD4 T cells by injection of 200ug aCD4 antibody (BioXCell cat: BE0003-1) one day prior to and one day post-infection. Cells were isolated as live CD62L- CD8 T cells from spleens of 3 pooled mice per group and used as input for scRNA-seq analysis.



Resource

Adoptive transfer of progenitor dysfunctional cells

Four C57BL/6J mice were infected with LCMV Clone 13 (2x10⁶ p.f.u. via retroorbital injection). On day 35 post-infection, splenocytes were pooled and stained using a cocktail of fluorescently labeled antibodies and Ghostdye Violet 510 (Tonbo cat: 13-0870-T100) to distinguish dead cells. Progenitor dysfunctional cells were isolated by flow cytometry-based sorting as PD1+, CD73+, CD39- CD8 T cells. This population was also CD9+ and CD38 low. 50,000 progenitor dysfunctional cells/recipient were transferred into 4 CD45.1/ CD45.2 mice via retroorbital injection. Recipients were infected with LCMV Armstrong (2x10⁵ p.f.u. via intraperitoneal injection). On day 7 post-infection donor cells were re-isolated based on expression of congenic markers and prepared for scRNA-seq.

Single-cell barcoding, library preparation and sequencing

Single-cell suspensions were loaded on 10x Genomics Chromium instrument and encapsulated in microfluidic droplets with barcoded DNA hydrogel beads and RT reagents from Chromium Single Cell 3' Reagent kit (v3). The cDNA synthesis/barcoding was performed following manufacturer's instructions: 53°C for 45 min followed by heat inactivation at 85°C for 5 min. The barcoded-cDNA was purified and PCR-amplified and prepared for sequencing according to the Single Cell 3'Reagent kit 3 User Guide (CG000183; Rev B). The DNA sequencing was performed on the Illumina NovaSeq 6000 instrument (R1 read - 26 cycles, R2 read - 70 cycles or higher, and index read – 8 cycles), aiming for \sim 200 million reads per \sim 5,000 single cells.

Single-cell RNA-seq data preprocessing, dimensionality reduction, clustering

We analyzed scRNA-seq data from the six samples labeled "acute_d00," "acute_d07," "acute_d40," "chronic_d07," "chronic_d35," "chronic_d35_b6." Reads were aligned to the combined B6 and Spret genome using hisat2 v2.1.0. For this, files Mus_musculus.GRCm38.dna.toplevel.fa.gz and Mus spretus spreteij.SPRET EiJ v1.dna.toplevel.fa.gz with genomic sequences of B6 and Spret mice, respectively, were obtained from NCBI FTP server, and hisat2 index for the chromosomes from both files was constructed. Using the BAM files of scRNA-seq read alignment, UMI counts for each gene in B6 and Spret genomes were obtained using a custom script, by overlapping read alignments with exonic annotations of genes in either B6 or Spret and counting UMI corrected using the method from seqc (Azizi et al., 2018). For this, files Mus_musculus.GRCm38.91.gtf.gz and Mus_spretus_spreteij.SPRE-T EiJ v1.86.qtf.gz with gene annotations for B6 and Spret genomes, respectively, were obtained from NCBI FTP server. The resulting read count tables were used for downstream analysis in scanpy v1.3.7. Cells with less than 100 genes with positive counts or cells with more than 40000 total positive UMI counts were filtered out, and genes with less than 500 positive counts across cells from all samples were filtered out. Cells with high expression of B cell genes (likely contamination) were filtered out (15 cells). Only proteincoding genes were included in the analysis, and furthermore ribosomal genes were excluded from the analysis. Based on inspection of the distribution of the number of genes detected per cell in each sample, cells from sample "acute_d00" with log₁₀ gene count less than 2.85, from sample "acute_d07" with log₁₀ gene count less than 2.9, from sample "acute_d40" with log₁₀ gene count less than 2.9, from sample "chronic_d07" with log₁₀ gene count less than 2.9, from sample "chronic_d35" with log₁₀ gene count less than 2.8, from sample "chronic_d35_b6" with log₁₀ gene count less than 2.8 were excluded from the analysis. Then the count matrix was filtered again to include only cells with at least 500 genes and genes with at least 100 counts. This resulted in a read count table of 9822 genes in 24400 cells across the six samples.

Counts for 9124 genes present in both B6 and Spret annotations were used for normalization, dimensionality reduction, and clustering. For normalization, the total count in each cell was calculated, excluding the top 50 genes with the highest total count across all cells, and then each count divided by the total count per cell and multiplied by the median of total counts per cell. For dimensionality reduction and visualization, principal component analysis (PCA) was run for the normalized counts, the first 50 principal components (PCs) were selected, and the kNN (nearest neighbor) graph was built for k = 50 nearest neighbors per cell using Euclidian metric. The kNN graph was then used to construct three-dimensional uniform manifold approximation and projection (UMAP) with default parameters. The data was also visualized with t-distributed stochastic neighbor embedding (tSNE) using function "TSNE" from package sklearn.manifold with perplexity = 150 and otherwise default parameters applied to the first 50 PCs of normalized count matrix; with diffusion maps and force-directed atlas applied to the kNN graph. Louvain clustering was then applied to the kNN graph with resolution = 1.9 and otherwise default parameters. For visualization of gene expression, imputation algorithm MAGIC (Van Dijk et al., 2018) was applied to the normalized count matrix using package magic with parameters a = 15, k = 30, knn_dist = 'euclidean', $n_pca = 50$, random_state = 0, t = 3.

To assess the similarity of samples "chronic_d35" and "chronic_d35_b6," kNN graph analysis was performed. For each cell c and a sample S, the distance between c and S was calculated as the average distance in the kNN graph between c and cells from S, measured as the number of edges in the shortest path. This value was calculated for each cell c from samples "chronic_d35" and "chronic_d35_b6" against each of the samples, and shown as a boxplot (Figure S7E).

Overview of functional annotation of scRNA-seq data

We used a combination of complementary approaches in order to functionalize the scRNA-seq data. First, unbiased differential expression analysis identified genes enriched in each cluster (Figures 4E, S8A, and S8B; Method details). Second, visualizing the expression of the T cell signature genes (Figures 4F, S7C, and S8C) further helped to interpret the clusters and characterize subsets of cells. Clusters 0, 7, 19, dominated by cells profiled before infection (Figure S8D), were enriched for naive T cell marker genes Ccr7, Ccr9, Lef1, Tcf7. Gzmb was highly expressed in clusters 4, 6, 14, all dominated by cells at d7 upon acute infection, consistent with



Molecular Cell Resource

effector function. Cell proliferation and survival related genes Birc5, Stmn1, Cdca3 were enriched in clusters 4, 6, 8, 14, 17, 20, which were dominated by cells from early response (d7) to acute or chronic infection. Clusters 2, 10, 13 displayed high II7r expression and were dominated by cells from acute infection at d40, which were mostly memory cells. Clusters 1, 3, 11, 12, 13, 21 were dominated by cells at d35 in chronic infection and enriched for markers of dysfunction Pdcd1, Lag3, Entpd1, and Cd38. Clusters 5, 15 were dominated by cells from chronic infection at d7 and enriched for Lgals1, Lgals3, Mt1, Mt2. Cluster 18 consisted of dysfunctional cells enriched for Pdcd1, Lag3, and Entpd1 that also overexpressed Ifng, Ccl3, Ccl4, and Nr4a2 and thus were also highly activated. Clusters 9 and 16 overexpressed markers of progenitor cells Cxcr5, Slamf6, Tcf7, and Id3. Finally, association of our bulk RNA-seq compendium (Figure S1D) with scRNA-seq cluster-specific expression profiles using ssGSEA largely confirmed our cluster characterizations (Figure S9A).

Differential expression analysis of scRNA-seq data

Differential expression analysis between cells from groups A and B was performed as follows. Let x_G and y_G be vectors of normalized scRNA-seq expression values of the gene G in individual cells from groups A and B, respectively. Then log2 fold change (log2FC) of expression for G is estimated as $log_2((Y_G + c) / (X_G + c))$ where $X_G = mean(x_G)$, $Y_G = mean(y_G)$ are means of normalized expression of G in cells from groups A and B, respectively, and c is a corrective value defined as median over all values M_g where M_g is the mean expression of a gene g across all cells from all samples. Only genes with the absolute log2FC above a certain threshold were then tested for significance. For each such a gene G, the significance of the log2FC for G was estimated with the Mann-Whitney U test applied to normalized counts x_G and y_G for G in individual cells from groups A and B; the Bonferroni correction for multiple hypothesis testing was then applied to these p values.

Using this approach, differential expression analysis was performed for each cluster versus cells from all other clusters, testing for significance of all absolute log2FC values above 0.5. The same analysis was also repeated when excluding all naive cells defined as cells from sample "acute_d00" or clusters 0, 7, 19. Differential expression analysis was also performed for each pairwise comparison between clusters. Furthermore, for each sample, differential expression analysis was performed for each cluster versus cells from all other clusters when restricting the analysis only to cells from that sample.

Comparison of scRNA-seq with bulk RNA-seq data compendium

The most significantly differentially expressed genes in each scRNA-seq cluster (adjusted p < 0.001, log2FC > 0.8), defined as the union of at most 100 genes from the comparison of the cluster with all other clusters and at most 100 genes from the same comparison restricted to non-naive cells, formed a signature set of genes for this cluster. ssGSEA was then run for these signatures against batch-effect corrected library-size normalized bulk RNA-seq read counts using package GSVA v1.30.0.

Analysis of scRNA-seq data from adoptive cell transfer experiment

The six main scRNA-seq samples were combined with the sample "transfer_acute_d07" from the progenitor dysfunctional cells extracted from the established chronic infection and adoptively transferred and expanded under acute infection. The extended dataset was preprocessed in the same manner as the main dataset, resulting in a count matrix for 9274 genes in 28447 cells, including 4047 cells from the transfer sample. The analysis of this extended count matrix was performed in the same manner as with the main six samples, resulting in clusters t0–t23. This cell clustering, restricted to the cells from the main six samples, was compared with the previously obtained clusters, suggesting cluster t4 consisted of the progenitor and progenitor-like cells. The cluster composition of each sample, calculated as fraction of cells in each sample that belongs to each cluster, was also compared between samples, suggesting the sample "transfer_acute_d07" was most similar to the sample "chronic_d07."

Allele-specific scRNA-seq expression

For allele-specific analysis, scRNA-seq sequencing reads were re-aligned in allele-specific manner as previously described (van der Veeken et al., 2019; Crowley et al., 2015). Briefly, the genetic variants of Spret mice were obtained from the mouse genome project (Keane et al., 2011). A pseudo-Spret genome was built by modifying the reference genome with SNPs, insertions and deletions found in the wild-derived inbred strain. The RNA-seq reads were aligned to the reference and pseudo Spret genomes in parallel using STAR (Dobin et al., 2013) with the following parameter settings: "STAR-runMode alignReads-readFilesCommand zcat-outSAMtype BAM-outBAMcompression 6–outFilterMultimapNmax 1–outFilterMatchNmin 30–alignIntronMin 20–alignIntronMax 20000–alignEndsType Local." Next, the genomic coordinates of pseudo-genome aligned reads were converted back to the corresponding B6 coordinates. To determine the allelic origins of the reads, the mapping scores of the two alignments of each read were compared. We retained the alignment with the highest score and generated the final BAM files. In cases where the diploid genome alignment produced identical scores for both genomes, one of the alignments was selected randomly.

Spret- and B6-specific and ambiguous UMI counts of gene expression for each cell were obtained with custom scripts, in the same manner as in the main analysis (see Single-cell RNA-seq data preprocessing, dimensionality reduction, clustering). Further analysis was restricted only to cells and genes selected for the main scRNA-seq analysis. Library-size normalization for Spret- and B6-specific and ambiguous read counts was obtained applying the same scaling for each cell as in the main analysis. The allelic imbalance of expression of each gene in each scRNA-seq cluster was defined as $log2((w_1 + 0.5 * w_2) / (w_0 + 0.5 * w_2))$, where w_0 and w_1 are B6- and Spret-specific total library-size normalized counts for this gene in this cluster, respectively, and w_2 is the ambiguous count for this

Resource



gene in this cluster. Significance of the allelic imbalance was assessed by a Mann-Whitney U test applied to all B6-specific versus all Spret-specific library-size normalized gene expression estimates for this gene over cells in this cluster.

Allele-specific predicted TF binding

Spret sequences of bulk ATAC-seq peak summit regions, as defined above (see Construction of ATAC-seq peak atlas), were obtained by introducing genetic variants between B6 and Spret into B6 sequences of these regions. To estimate allelic specificity of predicted TF binding, FIMO was run for Spret sequences and rerun for B6 sequences of the peak summit regions with the same motif collection as described previously (see Transcription factor motif analysis) with a relaxed p value threshold of 5e-3. All matches at p < 1e-4 in either B6 or Spret sequence were selected, and allelic imbalance of predicted TF binding for each motif in each peak summit region was estimated as the difference between the total FIMO log odds score for all matches of this motif in Spret and in B6. Then the distribution of these values for this motif across all peaks was z-score-normalized.

Analysis of the association of allelic imbalance of predicted TF binding and of scRNA-seq gene expression was performed for each TF motif and each scRNA-seq cluster as follows. All predicted TF motif binding sites with substantially stronger predicted match at Spret or B6 sequence were selected as those with z-score above 0.2 or below -0.2, respectively. Then the scRNA-seq allelic imbalance in this cluster was compared between genes nearby B6-specific and Spret-specific predicted binding sites, with significance assessed by Kolmogorov-Smirnov test.

CUT&RUN experiments

Terminal and progenitor dysfunctional cells were isolated from the spleens of 4 male B6/Spret F1 mice or B6 mice on day 35 postinfection, based on expression of cell surface markers (see Adoptive transfer of progenitor dysfunctional cells). Cells from 2 mice were pooled to generate biological duplicates with approximately 70,000 cells per replicate. CUT&RUN libraries were prepared as described (Skene and Henikoff, 2017) with the modifications described below. Because Concanavalin-A (ConA) is a well-known T cell mitogen, we avoided the use of ConA-coated beads for cell isolation and handling. 70k cells per replicate were collected in a V-bottom 96 well plate by centrifugation and washed in antibody buffer (buffer 1 (1x permeabilization buffer from eBioscience Foxp3/Transcription Factor Staining Buffer Set diluted in nuclease free water, 1X EDTA-free protease inhibitors, 0.5mM spermidine) containing 2mM EDTA). Cells were incubated with TCF1 antibody for 1h on ice. After 2 washes in buffer 1, cells were incubated with pA/G-MNase at 1:4000 dilution in buffer 1 for 1h at 4 degrees. Cells were washed twice in buffer 2 (0.05% (w/v) saponin, 1X EDTA-free protease inhibitors, 0.5mM spermidine in PBS) and resuspended in calcium buffer (buffer 2 containing 2mM CaCl2) to activate MNase. Following a 30 minute incubation on ice, 2x stop solution (20mM EDTA, 4mM EGTA in buffer 2) was added and cells were incubated for 10 minutes in a 37 degree incubator to release cleaved chromatin fragments. Supernatants were collected by centrifugation and DNA was extracted using a QIAGEN MinElute kit.

CUT&RUN libraries were prepared using the Kapa Hyper Prep Kit (Kapa Biosystems KK8504) and Kapa UDI Adaptor Kit (Kapa Biosystems KK8727) according to manufacturers protocol with the modifications described below. A-tailing temperature was reduced to 50 degrees to avoid melting of short DNA fragments and reaction time was increased to 1h to compensate for reduced enzyme activity as described by Liu et al. 2018. Following the adaptor ligation step, 3 consecutive rounds of Ampure purification were performed using a 1.4x bead to sample ratio to remove excess unligated adapters while retaining short adaptor-ligated fragments. Libraries were amplified for an average of 15 cycles using a 10 s 60°C annealing/extension step to enrich for shorter library fragments. Following amplification, libraries were purified using 3 consecutive rounds of Ampure purification with a 1.2x bead to sample ratio to remove amplified primer dimers while retaining short library fragments. A 0.5x Ampure purification step was included to remove large fragments prior to sequencing.

Identification of transcription factor binding sites in CUT&RUN data

CUT&RUN reads from B6 mouse samples were aligned to the genome using bowtie2 v2.3.4.3. Similar to RNA-seq, paired-end CU-T&RUN reads from F1 B6/Spret mouse samples were mapped to the diploid genome using STAR with the splicing alignment feature switched off. The command line was as follow: "STAR-runMode alignReads-genomeLoad NoSharedMemory-readFilesCommand zcat-outSAMtype BAM SortedByCoordinate-outFilterMultimapNmax 1-outFilterMatchNmin 40-outBAMcompression 6-outFilter-MatchNminOverLread 0.4-seedSearchStartLmax 20-alignIntronMax 1-alignEndsType Local." We only used the read pairs if their fragment length was between 50 to 500 bp.

For analysis, we used two control IgG samples and biological replicates for TCF1 CUT&RUN, including two samples from B6 mice and two samples from F1 B6/Spret mice. CUT&RUN and control reads were counted in 150bp windows around ATAC-seq peak summits. To estimate library sizes, control regions were obtained by shifting ATAC-seq peak summit regions by 2Kb in either direction, extending the shifted segments by 500bp preserving their center, and excluding those that overlapped with ATAC-seq peak summit regions, and then calculating control and CUT&RUN read counts in these control regions and using DESeq2 v1.22.2, as described previously (Konopacki et al., 2019). Differential CUT&RUN count analysis was then run using DESeq2, identifying 3325 TCF1 binding sites in the progenitor dysfunctional cells defined as ATAC-seq peak summit regions with significantly higher TCF1 CUT&RUN read counts than IgG control (adjusted p value < 0.1). To produce heatmaps of TCF1 CUT&RUN signal around binding sites, bigWig files and count matrices in bins around binding sites were produced using deepTools (Ramírez et al., 2016).



Motif enrichment in TCF1 binding sites was estimated as log2FC of frequency of TCF1 binding sites with significant TCF1 motif match (FIMO p < 1e-4) over such frequency for all ATAC-seq peak summit regions, with significance of enrichment estimated using hypergeometic test followed by Bonferroni correction for multiple hypothesis testing.

Molecular Cell

Resource

Allele-specific analysis of CUT&RUN data

For the TCF1 binding sites, allelic imbalance of TCF1 binding was defined as $log2((w_1 + 0.5 * w_2) / (w_0 + 0.5 * w_2))$, where w_0 and w_1 are B6- and Spret-specific CUT&RUN counts, respectively, and w_2 is the count of ambiguously aligned reads. For the CDF plots of allelic imbalance of TCF1 binding (Figure S13E), TCF1 binding sites with predicted stronger TCF1 motif batch in Spret or B6 were selected as those with TCF1 motif match imbalance z-score above 0.1 or below -0.1, respectively. For the allele-specific TCF1 co-factor analysis, Spret- or B6-specific TCF1 binding sites were detected as those with log2FC of CUT&RUN read count in Spret over B6 above 0.5 or below -0.5, respectively. Then for these sets of allele-specific TCF1 binding sites, for each motif, co-factor motif enrichment was calculated by comparing motif match imbalance z-scores against those in all ATAC-seq peak summit regions, with significance estimated using hypergeometric test. Motif match imbalance z-scores were then plotted as a boxplot (Figure S13G) for all the motifs that were detected as significant both for Spret- and B6-specific TCF1 CUT&RUN binding sites.

Construction of human ATAC-seq peak atlas and scATAC-seq count matrix

Bulk and single-cell ATAC-seq data for CD8 T cells in human donors and patients was obtained from three recent publications (Sade-Feldman et al., 2018; Philip et al., 2017; Satpathy et al., 2019). Bulk ATAC-seq data (Sade-Feldman et al., 2018; Philip et al., 2017) was processed in the same way as described above for mouse data. For scATAC-seq data (Satpathy et al., 2019), samples SU001_Tcell_Post2, SU001_Tcell_Post, SU006_Tcell_Pre, SU008_Tcell_Post, SU008_Tcell_Pre, SU009_Tcell_Post, SU009_Tcell_Pre were identified as those with large number of T cells profiled, according to metadata available at https://github.com/GreenleafLab/10x-scATAC-2019. Fragment data for these samples was obtained from GEO and mapped from hg19 to hg38 coordinates using liftOver tool from UCSC Genome Browser (Kent et al., 2002). Fragments associated with cells from T cell clusters 12-19 (Satpathy et al., 2019) were likely corresponding to CD8 T cells and therefore were selected for further analysis. Pseudo-reads of length 30nt were created for each start and end of the selected fragments, and MACS2 was run on this collection of pseudo-reads to identify accessibility peaks. These peaks were merged with those from bulk ATAC-seq fragment ends (corresponding to Tn5 cut sites) in these peaks were counted using Rsubread and findOverlaps(), respectively. Then peaks with at least total count of 50 in any of the three datasets (two bulk ATAC-seq datasets and one scATAC-seq dataset) were selected, forming an atlas of 161,140 peaks. Of them, 40,449 or 36,241 peaks overlapped at least one mouse atlas peak or peak summit, respectively, when mapped to human genome using liftOver().

Analysis of human scATAC-seq data

Count matrices for each of the seven human scATAC-seq samples obtained as described above were merged together, binarized, and transformed using the TF-IDF transform, as done with the mouse scATAC-seq data. The nearest neighbor graph was computed using the top 50 PCs and batch corrected by patient (SU001, SU006, SU008, SU009) using the batch balanced k-nearest neighbor procedure (Polański et al., 2020). Subsequent UMAP dimensionality reduction and Leiden clustering with a resolution of 0.8 were performed on the corrected kNN graph.

Comparison of differential accessibility between mouse and human

Differential accessibility in human bulk ATAC-seq data between naive and dysfunctional cells, central memory and dysfunctional cells, and progenitor and terminally dysfunctional cells was performed in the same manner as for mouse data. For comparison, differentially accessible peak summit regions in mice between naive and dysfunctional cells and between central memory and dysfunctional cells were selected as those with log2FoldChange > 1 and adjusted p < 0.001, between progenitor and terminally dysfunctional cells in melanoma as those with log2FoldChange > 0.5 and adjusted p < 0.001. The selected differentially accessible peaks were then mapped to human genome using liftOver() and overlapped with human peaks. For human scATAC-seq analysis, peak signatures obtained from mouse bulk ATAC-seq data were mapped to human genome using liftOver() and then scored in human scATAC-seq clusters, as described above for mouse scATAC-seq data analysis. For differential accessibility analysis between Leiden clusters 0 and 8, scATAC-seq counts were aggregated over all cells in these clusters in each sample separately, and then these pseudobulk counts compared between the two clusters treating each sample as a biological replicate (7 samples in each category) using DESeq2.

RNA velocity analysis

Spliced and unspliced read counts for each sample were obtained by running cellranger followed by velocyto (La Manno et al., 2018), using default mm10 genome. Then RNA velocity analysis for each sample was run using scvelo (Bergen et al., 2020). Only the cells that were used in the main scRNA-seq analysis after filtering were used for the velocity analysis. Naive cells (from clusters 0, 7, 19) were excluded from analysis of each sample. Genes for the analysis of each sample were selected using scvelo.pp.filter_genes() with min_shared_counts = 30. Furthermore, for each sample except "chronic_d35_b6," only those genes were used in subsequent analysis that had Spearman correlation at least 0.8 between spliced counts and previously obtained F1 genome counts used in

Resource



the main scRNA-seq analysis. RNA velocity analysis was run using function scvelo.tl.velocity() with mode = 'deterministic', implementing the approach from the original single-cell RNA velocity publication (La Manno et al., 2018). Genes labeled as velocity genes by scvelo.tl.velocity() were further filtered to include only those with R^2 of the fit (velocity_r2) above 0.3. This resulted in 36 genes for sample "acute_d07," 27 genes for "acute_d40," 72 genes for "chronic_d07," 22 genes for "chronic_d35" and 141 genes for "chronic_d35_b6" that were used in constructing velocity vectors for velocity graph construction and projection to 2-dimensional UMAP. Vector map visualizations were obtained using scvelo.pl.velocity_embedding_grid() with parameters arrow_size = 1, arrow_length = 3, density = 1, smooth = 0.7, min_mass = 3.

We suspect that the RNA velocity analysis in our data has a number of caveats. Insufficient sequencing coverage resulting in low read counts, particularly for unspliced reads, may have caused only a small number of genes to be selected for the velocity analysis. Furthermore, transcriptional differences across CD8 T cell functional and differentiation states captured in our scRNA-seq data may be smaller than those between different cell types previously analyzed with RNA velocity analysis (La Manno et al., 2018; Bergen et al., 2020). Finally, interpretation of the velocity vectors projected to 2-dimensional UMAP may be complicated, because of the oversimplification from reducing the data to only two dimensions and thus overlaying potential differentiation trajectories. Altogether, this may have contributed to noisier results of the analysis that are harder to interpret than those in previous publications (La Manno et al., 2018; Bergen et al., 2018; Bergen et al., 2018; Bergen et al., 2020). Nevertheless, we observe consistently across samples that velocity is directed toward cells on the right side of the UMAP, e.g., in cluster 1, which suggests these cells are more differentiated, consistent with their characterization as terminally dysfunctional.

QUANTIFICATION AND STATISTICAL ANALYSIS

All the details of quantifications and statistical analyses are fully described in the main text, figure legends, and Method details section.

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Supplemental information

A unified atlas of CD8 T cell dysfunctional

states in cancer and infection

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Figure S1. Universal program of CD8 T cell dysfunction (related to Figure 1). A. Principal component analysis (PCA) of library-size normalized ATAC-seq read counts in 150bp windows around peak summits (functional cell state shown by color, data source by symbol) without batch-effect correction. B. Distributions of distances between ATAC-seq vectors of read counts in 150bp windows around peak summits. Distances were calculated in the two-dimensional PCA space built using 20000 most variable peak summit read counts, for pairs of samples in the same functional category (naïve, functional, dysfunctional), separately for pairs of samples from the same study and from different studies, before and after batch effect correction. C. PCA of library-size normalized batch-effect corrected ATAC-seq read counts in 150bp windows around peak summits (labels as in A) using 10000 most variable peak summit read counts. Here GLM-based batch effect correction used a factor encoding the functional cell state using two values, for naïve cells and all other cells. D. PCA of librarysize normalized batch-effect corrected RNA-seg read counts for 1000 genes with most variable counts (functional cell state shown by color, data source by symbol). E. Scatter plot of differential expression (RNA-seq log2FC, x-axis) and differential accessibility (mean ATAC-seq log2FC over all peaks associated with a gene, y-axis) between functional and dysfunctional cells (computed as in D). Significantly differentially expressed genes shown as black dots. Significantly differentially accessible genes highlighted with red or blue color. F. "Diamond" plots of differential accessibility and differential expression between functional and dysfunctional cells for transcription factors and for genes associated with T cell activation, cytotoxicity, adhesion, and apoptosis. In each panel, left: library-size normalized batch-effect corrected ATAC-seq read count log2FC for peak summits (diamond shown in color for significantly decreased/increased, FDR < 0.05) of significantly differentially accessible genes; right: log2 fold change of RNA-seq gene expression for the same genes. G. Cumulative distribution function of differential accessibility (ATAC-seq log2FC) between naive and dysfunctional (left) or central memory and dysfunctional (right) CD8 T cells in human donors or cancer patients. Shown are peaks evolutionarily conserved when compared with peak summit regions in mice that were significantly more accessible in naïve or memory (blue) or dysfunctional (red) cells; background distribution is for all peaks identified in human cells that were conserved with mouse peak summit regions (black); count and p-value from Kolmogorov-Smirnov test against the background distribution.





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gene RNA-seg

5















Figure S2. Progression from early or progenitor to terminal dysfunctional state (related to Figure 1). A. Comparisons of differential accessibility results. Each plot is a scatter plot of log2FC of library-size normalized batch-effect corrected ATAC-seq read counts in peak summit regions for two differential accessibility analyses. Color (blue, red, dark gray) is used to highlight quadrants with peak summit regions significantly differentially accessible in both pairwise comparisons. Counts in corners indicate the number of such peak summits in each quadrant. Spearman's correlation is calculated over all such peak summits. B. Cumulative distribution function (CDF) of log-transformed quantilenormalized library-size normalized batch-effect corrected ATAC-seq read counts from various studies in 5940 peak summits significantly more accessible in progenitor than terminally dysfunctional T cells and in 2768 peak summits significantly more accessible in terminally than progenitor dysfunctional T cells in chronic LCMV infection (Kolmogorov-Smirnov p < 2e-16 for all comparisons). C. First principal component (PC1) of PCA for library-size normalized batch-effect corrected ATAC-seg read counts in dysfunctional T cells from different studies (see Figure 1C). PCA was calculated based on significantly differentially accessible peaks in different pairwise comparisons. For clarity, vertical random jiggle is added. D. Scatter plots of differential expression (RNA-seq log2FC, x-axis) and differential accessibility (mean ATAC-seq log2FC over all peaks associated with a gene, y-axis) between cell states. Significantly differentially expressed genes shown as black dots. Significantly differentially accessible genes highlighted with red or blue color. E. Cumulative distribution function of differential accessibility (ATAC-seq log2FC) between progenitor (TIM3- CD39-) and terminally (TIM3+ CD39+) dysfunctional TILs in human tumors. Shown are peaks evolutionarily conserved when compared with peak summit regions in mouse melanoma model that were significantly more accessible in progenitor (blue) or terminally dysfunctional (red) cells; background distribution is for all peaks identified in human cells that were conserved with mouse peak summit regions (black); count and p-value from Kolmogorov-Smirnov test against the background distribution.



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Figure S3. Predictive models of transcription factor association with chromatin accessibility (related to Figure 2). A. Schematic of the negative binomial generalized linear regression analysis to infer transcription factor (TF) associations with chromatin accessibility. B. Spearman correlations between library-size normalized batch-effect corrected ATAC-seq read counts and model predictions for each sample and each value of the regularization parameter. C. Inferred TF motif coefficients for regressing library-size normalized batch-effect corrected ATAC-seq read counts in each sample, for motifs corresponding to coefficients with the highest variance across samples, z-score normalized within each row. D. Top: Boxplots of log2FC of RNA-seq gene expression between naïve and effector cells for genes associated with at least one peak significantly more accessible in naïve than effector cells and associated with at least one peak significantly more accessible in effector than in naïve cells. Left: default peak-gene associations (peak associated to the closest gene in genomic coordinates, if this gene is within 50Kbp). Center: peak-gene associations defined using significant Hi-C contacts in naïve cells. Right: peak-gene associations defined using non-significant Hi-C contacts in naïve cells. Bottom: same analysis for naïve vs. dysfunctional cells. E. Spearman correlation of inferred TF motif coefficients across all antigen-experienced samples (left) and of TF motif match scores across the atlas of peak summit regions (right).



Figure S4. scATAC-seq analysis of CD8 T cells in acute and chronic infection (related to Figure 3). **A.** Barplot showing the number of cells in each cluster from each sample. **B.** Components 2 and 3 of the UMAP representation of TF-IDF-transformed scATAC-seq data with Louvain clusters. **C.** Heatmap for single cells in scATAC-seq data, separately for acute and chronic infection, showing the naïve cell signature derived from bulk ATAC-seq data. **D.** Violin plots for scores in scATAC-seq data clusters, separately for acute and chronic infection, of peak signatures (for naïve cells, memory cells) derived from bulk ATAC-seq data. **E.** Violin plots for scores in scATAC-seq data clusters 1-8, 10, 13, separately for acute and chronic infection, of peak signatures (for effector cells, memory precursor cells, terminally dysfunctional cells) derived from bulk ATAC-seq data. **F.** Inferred TF motif coefficients for scATAC-seq counts averaged over cells in each cluster in each of the two samples. Inferred coefficients with the highest variance are shown (z-score row normalized).



Figure S5. scATAC-seq analysis reveals a similar population of precursor/progenitor cells in acute and chronic infection (related to Figure 3). A. Genome browser tracks of ATAC-seq data for selected peaks. Bulk ATAC-seq for progenitor and terminally dysfunctional cells, and for terminal effector and memory precursor cells. Normalized aggregated single-cell ATAC-seq for cells in each cluster (for clusters 1-8, 10) in each of the two samples. For genes significantly more accessible in scATAC-seq cluster 10 as compared with other clusters. B. Differential accessibility of genes in cluster 10. Each panel shows normalized scATAC-seq pseudo-count (averaged over cells in each cluster in each sample) log2 fold change for peak summits of significantly differentially accessible genes; mean log2FC value is highlighted (with a shade of red) for genes that were overall differentially accessible. Shown are comparisons between cells in cluster 10 and cells in clusters 1-8, separately for acute and chronic infection, for genes highlighted in Figure 1E. C. Same as B, but for the most significantly differentially accessible genes between cells from acute and chronic infection within cluster 10. D. UMAP for scATAC-seq of CD8 T cells from human cancer patients (Satpathy et al. 2019). Top and bottom left: patient label and treatment status of each sample. Top right: clusters identified in the original publication. Bottom right: newly identified clusters (Methods). E. Comparison of new clusters with those from the original publication. Shown is a fraction of cells in each of the published clusters that belong to each of the new clusters 0-8. F. Violin plots for scores in new human scATAC-seq clusters 0-8 of peak signatures derived from bulk ATAC-seq data in mice. Shown are comparisons of all clusters (left) and of the cluster with the highest median score against the union of all other clusters (right). The results suggest a classification of cluster 8 as progenitor dysfunctional cells, and cluster 0 as terminally dysfunctional cells. G. Differential accessibility of genes between new human scATAC-seq clusters 0 and 8. Shown are normalized scATAC-seq pseudocount log2 fold changes for peaks of selected genes with at least one significantly differentially accessible peak (color, FDR < 0.1). Most genes show the trend similar to that in differential bulk and single-cell ATAC-seg signal between progenitor and terminally dysfunctional cells in mice, with only a few notable exceptions (e.g. KLRG1, SLAMF6, LEF1).

Figure S6 A Isola



Isolation of CD8 T cell subsets from LCMV Armstrong infected mice

В

PD-1

Isolation of CD8 T cell subsets from LCMV CI13 infected mice



CD127

CXCR3

Figure S6. Sorting for CD8 T cell subpopulations for scRNA-seq profiling (related to Figure 4). A-

B. Sorting strategy for isolating CD8 T cells responding to acute and chronic LCMV infection. **C.** Protein expression in cells isolated from chronic infection.



Figure S7. Dimensionality reduction and visualization of scRNA-seq data (related to Figure 4). A. Principal component analysis (PCA) of library-size normalized batch-effect corrected ATAC-seq read counts in 150bp windows around peak summits (functional cell state shown by color, data source by symbol), including samples from naïve, effector, and memory cells from hybrid F1(B6xSpret) mice. This analysis suggests that CD8 T cell functional states are similar between hybrid and B6 mice. **B.** Projections of three-dimensional UMAP representation of library-size normalized scRNA-seq data. **C.** Log-transformed gene expression (library-size normalized scRNA-seq UMI counts) on UMAP. **D.** Cluster composition of samples. Heatmap showing for each cluster what fraction of cells that cluster occupies in each sample. **E.** Boxplot of the kNN graph distances from cells in samples "chronic_d35" (left) and "chronic_d35_b6" (right) to cells in other samples. For each cell C, average distance in the kNN graph from C to cells in each sample was calculated, and distribution of these values across cells C was plotted. **F-H.** Results of dimensionality reduction methods applied to normalized scRNA-seq data.



Figure S8. Differential expression analysis of scRNA-seq data (related to Figure 4). A. Average scRNA-seq gene expression (library-size normalized UMI counts, z-score row normalized) across clusters for 261 differentially expressed genes obtained as the union of three overlapping gene sets: genes significantly overexpressed in a cluster as compared with a union of all other clusters (top 20 genes per cluster, log2FC > 0.7, adjusted p-value < 0.001, total 224 genes), genes significantly overexpressed in a non-naïve-cell cluster as compared with a union of all other non-naïve-cell clusters (top 20 genes per cluster, log2FC > 1.0, adjusted p-value < 0.001, total 188 genes), genes significantly differentially expressed between a pair of non-naïve-cell clusters (top 5 genes, log2FC > 1.5, adjusted p-value < 0.001, total 98 genes). **B.** Barplots of scRNA-seq gene expression log2 fold change values for top 30 genes significantly differentially over-expressed and under-expressed in each cluster. **C.** MAGIC-imputed gene expression for selected genes. **D.** Fraction of cells classified as naïve (clusters 0, 7, 19) in individual samples.



Figure S9. scRNA-seq cluster 9 consists of precursor/progenitor cells in acute and chronic infection (related to Figure 4). A. ssGSEA association of library size-normalized batch effect-corrected bulk RNA-seq data with library size-normalized scRNA-seq data averaged over cells in each cluster (z-score column normalized). Clusters 0, 7, 19 were most strongly associated with naïve cells, clusters 2, 10, 13 with memory cells, clusters 1, 3, 11, 12, 18, 21 with late and terminally dysfunctional cells, clusters 4, 5, 6, 8, 14, 15, 17, 20 with effector and early and progenitor dysfunctional cells, clusters 9 and 16 with progenitor cells and with early dysfunctional cells. **B.** Average scRNA-seq gene expression (library-size normalized UMI counts) in each cluster for genes significantly overexpressed in cluster 9 as compared with all other cells and as compared with all other cells except naïve cells (clusters 0, 7, 19). **C.** Fraction of cells that belong to cluster 9 in individual samples. **D.** MAGIC-imputed gene expression in individual samples.





D21Spleen 2







Figure S10. Analysis of published scRNA-seq data confirms presence of progenitor-like subset at d7 in acute LCVM infection (related to Figure 4). Enrichment of progenitor markers in a cluster of cells profiled at d7 in acute LCMV infection in three studies (Yao *et al.* 2019, Kurd *et al.* 2020, Chen *et al.* 2020) (A-C). Shown are significant (q < 0.01) log2 fold change values of expression in a selected cluster as compared with cells from all other clusters, as identified by diffxpy (barplot, rightmost column). Clusters occupying at least 1% of the total data set are labeled on UMAP (third column). UMAP and clustering of the whole dataset are shown (leftmost two columns).







Figure S11. RNA velocity analysis (related to Figure 4). A-E. Vector maps of RNA velocities (left) and cell cycle gene signature scores (right) are shown for scRNA-seq samples "acute_d07" (**A**), "acute_d40" (**B**), "chronic_d07" (**C**), "chronic_d35" (**D**), and "chronic_d35_b6" (**E**). **F-G.** Phase maps of unspliced and spliced read counts (left), RNA velocities (middle), and gene expression values (averaged over nearest neighbors in the kNN graph, right) for selected genes in cells from samples "acute_d07" (**F**) and "chronic_d35" (**G**).





cluster 3

154 134

0.0 0.5 1.0

scRNA-seq allelic imbalance log2(Spret / B6)

genes with stronger binding in Spret

genes with stronger binding in B6

p < 0.0

0.8

0.0

-1.0

E(x) 0.4





allelic imbalance difference

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Mef2a/d





0.3 0.4 0.5

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0.0 0.1 0.2 12

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-2 -1 ò 2 18

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Figure S12. Allele-specific analysis of TF binding and scRNA-seq data (related to Figure 5). A. Allelic imbalance of gene expression in clusters of scRNA-seq data (color, p < 0.05, Mann-Whitney U). **B.** Boxplots of differential expression (log2 fold change) between cells within the same scRNA-seq cluster profiled at d35 in chronic infection in B6 and in B6/Spret F1 mice, for genes with significant allelic imbalance towards B6 or Spret in B6/Spret F1 mice. Cluster number indicated on top of each boxplot. Significant difference between log2FC distributions (p < 2e-4 for clusters 2, 5, 10, 18, p < 2e-16 for all other clusters, Mann-Whitney U test) in all cases. **C-E.** Allelic specificity expression analysis of scRNA-seq data. CDF plots: allelic imbalance between B6 and Spret of library-size normalized scRNA-seq counts aggregated over cells in a selected cluster, for genes predicted to be bound more strongly in B6 or Spret using sequence motif analysis for each TF motif over all clusters; black bar indicates significant association (Kolmogorov-Smirnov p < 0.05).



Figure S13. CUT&RUN in progenitor dysfunctional CD8 T cells from B6/Spret F1 hybrid mice maps allele-specific TCF1 binding (related to Figure 5). A. Normalized TCF1 CUT&RUN signal in TCF1-bound ATAC-seq peak summits (library-size normalized read counts in 50bp bins in 3000bp window around peak summit, ordered by total signal). B. Log2 fold change enrichment of motif frequency in TCF1 CUT&RUN peaks relative to all ATAC-seq peak summits. C. Examples of TCF1 binding sites. D. Genome browser tracks of bulk ATAC-seq for progenitor and terminally dysfunctional cells and TCF1 CUT&RUN signal between B6 and Spret for all TCF1 targets (black) and for those predicted to be bound more strongly in B6 (purple) or Spret (orange) using sequence motif analysis. F. Top: CDF plots of library size-normalized batch effect-corrected ATAC-seq signal for TCF1-bound sites vs. all ATAC-seq peak summits. Bottom: CDF plots of library-size normalized batch-effect corrected RNA-seq gene expression for genes with TCF1 binding vs. all expressed genes. Shown is the comparison between progenitor and terminally dysfunctional cells in chronic LCMV infection. G. Imbalance of predicted TF binding using sequence motif analysis in peaks with stronger TCF1 binding in B6 or in Spret allele as identified by CUT&RUN.