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Antigen Dominance Hierarchies Shape TCF1+ Progenitor CD8 T Cell Phenotypes in Tumors

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INCLUSION AND DIVERSITY STATEMENT

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M.L.B, A.M.C., G.E.C., G.G., S.E.B., A.B., D.C., J.M.S., A.L., A.M.J., W.L.H., A.R., S.S., and T.J. designed the study; M.L.B., G.E.C., T.K., S.Z.T., A.L.B., A.G. and I.D.K. performed the mouse experiments in the lab of T.J.; M.L.B., G.E.C., M.H., W.L.H., and P.M.K.W. performed the mouse scRNA-seq in the labs of T.J. and A.R.; A.M.C., S.E.B. and A.B. performed the scRNA-seq computational analyses; G.G. and C.C.R. performed the t-CyCIF in the lab of S.S.; F.S.H, M.J.M. and M.H. provided melanoma samples; M.L.B. wrote the manuscript with input from other authors.

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We worked to ensure sex balance in the selection of non-human subjects. One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in science.

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SUMMARY

CD8 T cell responses against different tumor neoantigens occur simultaneously, yet little is known about the interplay between responses and its impact on T cell function and tumor control. In mouse lung adenocarcinoma, we found that immunodominance is established in tumors, wherein CD8 T cell expansion is predominantly driven by the antigen that most stably binds MHC. T cells responding to subdominant antigens were enriched for a TCF1+ progenitor phenotype correlated with response to immune checkpoint blockade (ICB) therapy. However, the subdominant T cell response did not preferentially benefit from ICB due to a dysfunctional subset of TCF1+ cells marked by CCR6 and Tc17 differentiation. Analysis of human samples and sequencing datasets revealed that CCR6+ TCF1+ cells exist across human cancers and are not correlated with ICB response. Vaccination eliminated CCR6+ TCF1+ cells and dramatically improved the subdominant response, highlighting a strategy to optimally engage concurrent neoantigen responses against tumors.

Graphical Abstract



IN BRIEF

Analysis of antigen dominance hierarchy and dynamics in a lung cancer model highlights the opportunity of optimizing T cell responses against subdominant neoantigens for better checkpoint blockade response through vaccination that adjusts T cell progenitor subpopulations

Keywords

TCF1; neoantigen; immunodominance; CCR6; Tc17; vaccine; lung cancer

INTRODUCTION

Accumulating evidence indicates that peptides derived from mutated proteins presented on major histocompatibility (MHC) molecules, termed neoantigens, drive productive T cell responses to tumors (Schumacher et al., 2019). Neoantigen-specific CD8 T cells expand in response to immune checkpoint blockade (ICB) therapies (e.g. anti-PD1/anti-PDL1) and patient responses correlate with high neoantigen and/or mutational burden (Keenan et al., 2019). Additionally, adoptive T cell therapies targeting neoantigens can promote meaningful tumor regression (Tran et al., 2017; Zacharakis et al., 2018). These observations provide strong rationale for development of immunotherapies directly targeting neoantigens and, in particular, have fueled recent large-scale efforts to develop neoantigen-targeted cancer vaccines (Hollingsworth and Jansen, 2019).

To identify therapeutically actionable neoantigens for immunotherapies, computational pipelines have been developed to predict immunogenic epitopes from exome sequencing of patient tumors (Peters et al., 2020; Wells et al., 2020). While these algorithms identify tens to hundreds of potential neoantigens, the vast majority of these epitopes do not elicit a detectable T cell response. Studies profiling T cell reactivity against predicted neoantigens have generally identified only a few, and sometimes zero, bona fide T cell-reactive epitopes across most cancer types (Linette et al., 2019; McGranahan et al., 2016; Scheper et al., 2019; Simoni et al., 2018). If this low frequency of T cell reactivity reflects the number of neoantigens capable of eliciting an anti-tumor response, this could pose a significant challenge in the selection of relevant neoantigens for immunotherapies.

Poor T cell reactivity against computationally predicted neoantigens may be in part attributable to shortcomings of prediction algorithms or detection limits of T cell assays. There are also plausible biological explanations, including selection against tumor cells expressing immunogenic neoantigens, resulting in loss of T cell-reactive antigens over time (i.e. immunoediting). Immunoediting has been observed in mouse models (DuPage et al., 2012; Matsushita et al., 2012; Milo et al., 2018) and in conjunction with clinical response to ICB and adoptive T cell therapies (Anagnostou et al., 2017; Verdegaal et al., 2016). While it is difficult to quantify the extent of immunoediting during tumor progression in humans (Rosenthal et al., 2019), evidence in mice suggests it may depend on the tumor type or tissue context (DuPage et al., 2011; DuPage et al., 2012; Matsushita et al., 2012). In addition, antigen immunodominance may be established during anti-tumor immune responses (Schreiber et al., 2002). As observed in acute viral infection, one or two epitopes may saturate the T cell response and suppress responses to additional antigens (Yewdell,

2006). Supporting this idea, healthy donor T cells can respond to tumor antigens that did not elicit a response from endogenous tumor infiltrating T cells, suggesting that responses against some antigens are depleted or suppressed during tumor progression (Stronen et al., 2016). Further, vaccines against melanoma neoantigens generate *de novo* responses, highlighting a degree of unrealized neoantigen immunogenicity in tumors (Carreno et al., 2015; Ott et al., 2017; Sahin et al., 2017). Thus, T cell responses against some neoantigens might outcompete others and narrow the T cell response to fewer epitopes.

To study the dynamics of anti-tumor T cell responses, our group previously engineered a Kras/p53-driven (KP) autochthonous mouse model of lung adenocarcinoma to express tumor-specific neoantigens (DuPage et al., 2011; DuPage et al., 2009). Tumors are initiated from single transformed cells *in situ* and develop over approximately five months, allowing ample time for tumor-immune crosstalk and longitudinal sampling of the T cell response at different tumor stages. Two model CD8 T cell neoantigens are expressed in tumor cells as a fusion to Luciferase: SIINFEKL (SIIN) from chicken ovalbumin and a synthetic peptide, SIYRYYGL (SIY) (termed LucOS). In this KP LucOS model, the CD8 T cell response initially slows tumor progression, but exhausts over time. Importantly, neoantigen and MHC expression is sustained in advanced tumors, indicating mechanisms other than immunoediting hinder the T cell response.

In the LucOS model, SIIN and SIY are concurrently expressed, providing an experimental model to study the interaction of distinct neoantigen-specific CD8 T cell compartments in the context of an anti-tumor response. Here, we find an antigen dominance hierarchy is established between SIIN and SIY in lung tumors that limits expansion of the subdominant CD8 T cell response and represses differentiation from a progenitor to an exhausted cell state. Progenitor CD8 T cells are correlated with response to ICB in mice and humans (Philip et al., 2021), yet their biology remains poorly understood. We uncover a previously undescribed subpopulation of progenitor cells that can arise from antigen subdominance and correlates with impaired functionality and poor response to ICB. Therapeutic vaccination eliminates this subpopulation and greatly expands the subdominant response, highlighting vaccination as a strategy to optimally engage simultaneous CD8 T cell responses against multiple neoantigens in tumors.

RESULTS

Longitudinal Analysis Uncovers Heterogeneity Between CD8 T Cell Responses to Different Neoantigens in Lung Adenocarcinoma

To investigate how CD8 T cell responses to tumor neoantigens may interact, we longitudinally characterized SIIN- versus SIY-specific CD8 T cells in tumors initiated by intratracheal instillation of lentivirus containing LucOS and *Cre* recombinase in *Kras*^{LSL-G12D/+;} $p53^{fl/fl}$ mice (Figure 1A; Dupage et al., 2011). At the peak of the response (5 weeks), we observed that SIIN drove substantially greater CD8 T cell expansion compared to SIY in the tumor-bearing lung (Figures 1B, 1C, S1A and S1B). The SIIN response contracted sharply thereafter, whereas the SIY response persisted better over time, resulting in similar numbers of SIIN- and SIY-specific cells by 12 weeks. Both populations increasingly displayed hallmarks of dysfunction over time, exhibiting progressive decline in

proliferation (Ki67) and upregulation of multiple inhibitory receptors (PD1, LAG3, TIM3; Figure 1D, 1E, S1C and S1D). Notably, inhibitory receptor expression was lower on SIYspecific cells at 5 and 8 weeks compared to SIIN-specific cells, suggesting SIY-specific cells exhibit slower kinetics of dysfunction. A greater proportion of SIY-specific cells also expressed a marker of long-lived T cells, IL-7R, which may contribute to their persistence (Figure 1F). 30% of SIIN-specific cells produced IFN γ compared to 15% of SIY-specific cells at 5 weeks, indicating that SIIN-specific cells have greater effector function early in the response (Figure 1G). However, few cells were IFN γ + TNF α +, or "polyfunctional", a phenotype associated with better tumor control (Spranger et al., 2014). Additionally, only a small proportion of cells expressed the cytotoxicity-associated molecule Granzyme B (GZMB; SIIN 7% versus SIY 4%; Figure 1H). These data are consistent with both SIINand SIY-specific CD8 T cells having poor anti-tumor activity overall; however, SIIN-specific cells have greater effector function compared to SIY-specific cells.

Single-Cell RNA-Sequencing Reveals Enrichment of a TCF1+ Progenitor Cell Phenotype Amongst SIY-Specific CD8 T Cells

To further investigate phenotypic differences between SIIN- and SIY-specific CD8 T cells, we performed single-cell RNA-sequencing (scRNA-seq) with paired T cell receptor (TCR) sequencing at 5 weeks post-tumor initiation. Comparison of transcriptional profiles by uniform manifold approximation and projection (UMAP) and unsupervised clustering revealed similar distribution of SIIN- and SIY-specific cells; however, some clusters were significantly enriched for cells directed against one antigen versus the other (Figures 2A–2C; Table S3). To explore this transcriptional heterogeneity, we classified single cells using "ProjecTILs" T cell atlases (Andreatta et al., 2021) derived from mouse lymphocytic choriomeningitis virus (LCMV) infection and B16 melanoma and MC38 colorectal carcinoma tumor infiltrating lymphocytes (TILS; Figures 2D and S2A; Table S1). Consistent with Figure 1, SIIN-specific cells were enriched for "effector" and "exhausted" T cell signatures, while SIY-specific cells were enriched for signatures of less differentiated "naïve" and "memory precursor" states. Further, cluster 7, which was almost exclusively composed of SIY-specific cells, displayed hallmarks of suppressed activation, including poor clonal expansion and high expression of AY036118.1 (ETS-related transcription factor 1, ERF1), a transcriptional repressor of *c-myc* and cell proliferation (Figures S2C, S2E and S4B; Verykokakis et al., 2007). Intriguingly, SIY-specific cells were also enriched for signatures of "progenitor" cells, a population marked by stem-like ability to replenish more cytotoxic, but terminally-exhausted cells (Figure 2D and Table S3; Utzschneider et al., 2016). Recent studies have shown ICB acts to promote differentiation of progenitor cells, rather than to reverse dysfunction of terminally-exhausted cells (Kurtulus et al., 2019; Miller et al., 2019; Sade-Feldman et al., 2019; Siddiqui et al., 2019); hence, progenitor cells appear to be the therapeutically relevant target of ICB therapy. Progenitor cells are commonly defined by expression of the transcription factor T cell factor 1 (TCF1) and the absence of terminal exhaustion markers, such as Havcr2 (TIM3; Miller et al., 2019). SIY-specific cells were preferentially assigned to clusters 4 and 8 (C4 and C8) that were enriched for progenitor gene signatures and expressed *Tcf7*(encoding TCF1) and other progenitor markers, including II7r (Figures 2C-2E and S2C; Tables S1 and S3). Conversely,

SIIN-specific cells were preferentially assigned to clusters C2 and C3, which expressed markers of cytotoxicity and exhaustion, including *Gzmb* and *Havcr2*.

The proximity of clusters C4 and C8 to C2 and C3 on the UMAP suggests transcriptional similarity between these populations, consistent with the previously proposed lineage relationship between progenitor and exhausted cells (Figure 2B; Siddiqui et al., 2019). We further examined this relationship by analyzing the distribution of TCR clonotypes (5 cells) with at least one cell assigned to progenitor clusters C4 or C8 (Figure 2F). Unsupervised hierarchical clustering largely segregated SIIN and SIY clonotypes, revealing antigen-specific distribution patterns across clusters of cell states (Figure 2F). SIIN clonotypes were distributed across most clusters, but were enriched for cells assigned to exhausted cluster C2 (Figures 2F and S4C). This supports a lineage relationship between progenitor and exhausted cell states and suggests SIIN clonotypes containing progenitor cells are well-progressed on the path to exhaustion. In contrast, both progenitor cellcontaining and total SIY clonotypes were biased in cell distribution to progenitor clusters C4 and C8 (Figures 2F, S4A and S4C), suggesting SIY-specific cells are skewed towards a less differentiated state. Further, while SIIN and SIY clonotype numbers were comparable (2) cells: SIIN 153, SIY 149), we observed that SIY clonotypes were smaller in size, indicating that clonal expansion may be repressed amongst the SIY response (Figures S4B).

Flow cytometric analysis confirmed that SIY-specific cells were enriched for a progenitor cell phenotype, referred to hereafter as "TCF1+ progenitor" and defined as CD8 α + CD44+ TCF1+ TIM3– cells (Figure 2G). Consistent with previous reports (Miller et al., 2019; Siddiqui et al., 2019), the majority of these cells expressed the inhibitory receptor PD1 (Figure S2F). Few expressed GZMB, indicating they are generally not cytotoxic, but they were more proliferative than TCF1– cells and produced IFN γ and TNF α (Figure 2G, 2H and S2G). SIY-specific cells were enriched for TCF1+ progenitor cells compared to SIIN-specific cells at 5 and 8 weeks; however, their proportion became similar by 12 weeks (Figure 2I). The percentage of TCF1+ cells increased over time, but the absolute number decreased, mirroring the overall contraction of the CD8 T cell response (Figure 2I). Altogether, these data indicate that SIY-specific cells are enriched for a less differentiated, TCF1+ progenitor state early in the response to KP LucOS lung tumors, while SIIN-specific cells differentiate more rapidly to an exhausted cell state.

Enrichment of TCF1+ Progenitor Cells in the SIY Response is Driven by an Antigen Dominance Hierarchy

Having observed that SIIN drove a larger CD8 T cell response than SIY, we hypothesized that competition between the T cell responses might underlie enrichment of TCF1+ progenitor cells amongst the SIY response. To test this, we engineered lentiviruses to express SIIN-only or SIY-only as fusions to luciferase (LucSIIN and LucSIY; Figure 3A). Strikingly, in LucSIY mice, SIY-specific cells expanded similarly to SIIN-specific cells in LucOS and LucSIIN mice (Figures 3A, 3B and S3A). Furthermore, enrichment for TCF1+ progenitor cells in the SIY response was no longer observed (Figures 3C and S3B). A higher proportion of SIY-specific cells in LucSIY mice expressed markers of effector memory and exhausted cells (CX3CR1, GZMB, PD1, LAG3, TIM3; Figures S3C–S3F).

Thus, the SIY response in LucSIY mice more closely resembled the response against SIIN in LucOS and LucSIIN mice. These data demonstrate that an antigen dominance hierarchy can form in tumors and this significantly influences the phenotype of cells responding to the subdominant antigen. Competition with SIIN promotes a TCF1+ progenitor phenotype amongst SIY-specific cells and hinders differentiation to effector and exhausted cell states.

In acute viral infection, several factors underlie antigen immunodominance, including: relative stability of peptide-MHC (pMHC) complexes, number of pMHC complexes on the surface of antigen-presenting cells, and functional avidity of the T cell repertoire for each antigen (Yewdell, 2006). SIINFEKL is known to bind H-2K^b MHC molecules with greater affinity and stability than SIYRYYGL (Eisen et al., 2012); thus, we hypothesized that differential MHC binding might contribute to the observed dominance hierarchy. To test this, we created lentiviruses expressing SIINYEKL (Y5), a point mutant of SIIN that binds H-2K^b molecules with reduced stability (K_{off}) and affinity (K_d) (Figure 3D–3F and S3G; Howarth et al., 2004). The Y5 mutation conserves key TCR contact residues and is largely cross reactive with the SIIN T cell precursor pool (Figure 3G and S3H; Bentzen et al., 2018). By comparing SIIN and Y5, we can decouple the contribution of pMHC binding and TCR repertoire/avidity to antigen dominance. Remarkably, when Y5 and SIY were expressed together (LucY5S), we found the antigen dominance hierarchy was reversed compared to LucOS, favoring SIY as the immunodominant antigen (Figures 3D and 3H). SIY-specific T cells expanded substantially more than Y5-specific cells and the Y5 response was enriched for TCF1+ progenitor cells (Figures 3H and 3I). When Y5 was expressed alone (LucY5), expansion of Y5-specific cells rebounded and enrichment for TCF1+ progenitor cells was no longer observed. These results demonstrate that differential pMHC binding can establish an antigen dominance hierarchy in cancer, and subdominant responses, irrespective of the antigen, are enriched for a TCF1+ progenitor cell phenotype. Y5 bound H-2K^b with higher affinity, but lower stability than SIY (Figure 3E and 3F), suggesting pMHC stability plays a larger role than affinity in establishing Y5 subdominance.

To evaluate the dynamics between antigens with similar MHC binding properties, we expressed two neoantigens derived from point mutations in murine methylcholanthrene (MCA)-induced sarcoma, mALG8 and mLAMA4 (Figure S3K; Gubin et al., 2014). Compared to mLAMA4, mALG8 binds H-2K^b with similar, but slightly lower affinity and stability (Figures S3I and S3J). Expression of mALG8 or mLAMA4 alone (LucA and LucL, respectively) in tumors resulted in responses of equivalent magnitude (Figure S3L). However, when co-expressed (LucAL), neither response became dominant in terms of T cell expansion, but both decreased in magnitude compared to the single-antigen setting (Figure S3L). The proportion of TCF1+ progenitor cells was also increased for both responses (Figure S3M), trending towards greater enrichment for the mALG8 response, the slightly weaker MHC binder. Thus, antigens with similar MHC binding properties are largely co-dominant, but their competition can drive enrichment for a TCF1+ progenitor phenotype, similar to subdominant antigens.

Subdominant Antigen-Specific T Cells Do Not Preferentially Benefit from anti-PD1/CTLA4 Therapy

Given TCF1+ progenitor cells play a key role in CD8 T cell responses to ICB therapy, we investigated whether SIY-specific cells in LucOS mice might preferentially benefit from ICB. In response to one week of anti-PD-1/CTLA-4 treatment, SIIN- and SIY-specific T cell expansion was greatest at 5 weeks, diminished at 8 weeks and absent by 12 weeks, mirroring the kinetics of T cell dysfunction and contraction of TCF1+ progenitor cells described above (Figures 4A and 4B). ICB treatment shifted cells away from a TCF1+ TIM3– phenotype and towards a TCF1– TIM3+ phenotype, consistent with increased differentiation to an exhausted state (Figure 4C). The absolute number of TCF1+ TIM3– cells increased, despite their decreased proportion, indicating that TCF1+ progenitor cells also expand with ICB (Figure 4D). ICB treatment also increased cell proliferation (Ki67) and cytotoxicity (GZMB; Figures 4E and 4F). Surprisingly, despite being enriched for TCF1+ progenitor cells, SIY-specific cells did not respond better to ICB compared to SIIN-specific cells. A somewhat greater shift in TCF1+ TIM3– to TCF– TIM3+ cells was observed, but did not translate into greater T cell expansion or increased effector functions (Figures 4B–4F).

Recent studies in chronic LCMV infection and cancer have identified the chemokine receptor CX3CR1 as a marker of cells recently differentiated from TCF1+ progenitor cells (Hudson et al., 2019; Zander et al., 2019). CX3CR1 expression correlates with cytotoxic function and depletion of CX3CR1+ cells results in a loss of disease control. CX3CR1 was expressed on a subset of SIIN- and SIY-specific TCF1+ cells and likely marks cells that are actively differentiating (Figure 4G). CX3CR1 expression was lower on SIY-specific TCF1+ cells and SIY-specific cells overall, suggesting SIY cells are repressed in their differentiation from a progenitor to an exhausted cell state (Figure 4G). ICB treatment was able to rescue much of this deficit, consistent with ICB promoting differentiation of progenitor cells (Figure 4G). Similarly, expression of SIY alone in LucSIY mice rescued CX3CR1+ cells to SIIN levels, which suggests that competition with the SIIN response represses SIY cell differentiation (Figure S3D). Notably, however, ICB was insufficient to fully rescue differentiation of SIY-specific cells, as the SIY response matched, but did not exceed the SIIN response, disproportionate with the enrichment of TCF1+ cells amongst the SIY response.

The Subdominant CD8 T Cell Response is Enriched for a CCR6+ TCF1+ Progenitor Cell Subset with a Tc17 Differentiation Trajectory

One possible explanation for a worse than expected SIY response to ICB is that SIY-specific TCF1+ progenitor cells are intrinsically less functional than their SIIN counterparts. To explore potential heterogeneity in progenitor cell phenotype or differentiation, we performed a separate analysis of progenitor clusters C4 and C8 and exhausted clusters C2 and C3 from the scRNA-seq data (Figure 5A). Both C4 and C8 were enriched for a number of genes previously associated with TCF1+ progenitor cells (e.g. *Tcf7, Xcl1, Slamf6, Ccr7)*, but were localized distally on the UMAP and were characterized by unique gene signatures (Figure 5B; Table S3). C8 was marked by genes associated with memory T cells, including *Klf2, S1pr1* and *ll7r*, while C4 expressed markers of T cell dysfunction, tolerance and anergy, and

most notably, showed strong enrichment for a gene signature of Tc17 cells (Figure 5B, 5C, S5A and S5B; Tables S2 and S3).

Tc17 cells, marked by expression of the chemokine receptor CCR6, are an IL17A-producing CD8 T cell subset commonly associated with autoimmune inflammation (Srenathan et al., 2016). *Ccr6* was highly expressed in progenitor cluster C4 and overlapped with *Tcf7* expression; however, cells exhibiting other hallmarks of Tc17 cells, including *Rorc* (RORγT) and *II17a* expression, were predominantly found adjacent to *Tcf7*-expressing cells within C4 (Figure 5D). Monocle3 analysis of putative lineage trajectories (Cao et al., 2019; Trapnell et al., 2014) predicted a trajectory connecting *Tcf7/Ccr6*-expressing cells with the Tc17 population (Figure 5D). Further, unsupervised clustering of TCR clonotypes revealed a group of five clonotypes, clonotype cluster 7, that predominantly contained cells spanning both the *Tcf7/Ccr6* and *Rorc/II17a* regions of C4 (Figure 5E, S4A and S4D). Only a small number of TCR clonotypes expressed *II17a* and might suggest a subset of TCRs promote Tc17 differentiation (Figure S4A). However, the majority of these clonotypes expressed *II17a* in a small proportion of cells and were distributed across multiple clusters, indicating Tc17-associated clonotypes also give rise to other cell states.

Flow cytometry analyses confirmed the presence of CCR6+ SIIN and SIY-specific cells and revealed enrichment amongst the SIY response (Figure 5F). The majority of CCR6+ cells expressed TCF1 as well as SLAMF6, another marker used to define TCF1+ progenitor cells (Figure 5G and S5D; Miller et al., 2019). Most CCR6+ cells also expressed RORyT and about half of these expressed TCF1 (Figure 5G). SIY-specific cells produced more IL17A compared to SIIN-specific cells, especially amongst the RORyT+ and CCR6+ populations, indicating increased Tc17 differentiation of SIY-specific cells (Figure 5H). Comparing functionality of CCR6+ TCF1+ versus CCR6- TCF1+ cells, we found both populations proliferated similarly (Ki67), but GZMB expression was largely restricted to a small proportion of CCR6- TCF1+ cells (Figures S5E and S5F). CCR6+ TCF1+ cells had higher expression of tolerance/anergy markers (CD200, EGR2, CD83), inhibitory receptors (PD1, LAG3, TIGIT) and costimulatory receptors often upregulated on exhausted cells (ICOS, OX40) compared to CCR6- TCF1+ cells (Figure S5C). CCR6+ TCF1+ cells also expressed higher levels of TOX, a transcription factor closely tied with T cell dysfunction (Scott et al., 2019). Altogether, these results uncover previously undescribed heterogeneity amongst TCF1+ progenitor cells. High expression of markers of dysfunction/tolerance, low GZMB expression and Tc17 differentiation suggest CCR6+ TCF1+ cells represent an unconventional TCF1+ population with reduced functionality.

Remarkably, the CCR6+ subset made up nearly 40% of SIY-specific TCF1+ progenitor cells, accounting for much of the enrichment of TCF1+ progenitor cells amongst the SIY response at 5 and 8 weeks (Figures 5I and S5G). The proportion of CCR6+ TCF1+ progenitor cells increased over time for the SIIN response, correlating with the kinetics of dysfunction and loss of ICB response (Figure 5I and S5G). Thus, a higher proportion of CCR6+ TCF1+ versus CCR6- TCF1+ progenitor cells correlates with worse ICB response and suggests the CCR6+ subset has reduced capacity to respond to ICB. This may explain why SIY-specific cells do not respond better than SIIN-specific cells to ICB despite enrichment for TCF1+ progenitor cells. Consistent with this, the proportion of

CCR6+ TCF1+ cells was reduced after one week of anti-PD1/CTLA4 ICB treatment (Figure 5J). Since total TCF1+ cell numbers increase with ICB response (Figure 4D), this indicates that ICB preferentially expands or recruits CCR6– TCF1+ cells. Additionally, little to no expression of CX3CR1 was observed on CCR6+ TCF1+ cells, while up to 20% of CCR6– TCF1+ cells expressed CX3CR1 (Figure 5K and S5H). This suggests CCR6+ TCF1+ cells are impaired in conventional differentiation to a cytotoxic state.

To assess the functionality of CCR6+ TCF1+ derived Tc17 cells, we utilized an IL17A reporter allele to permanently mark cells that have expressed IL17A with Tomato fluorescence (Figure 5L; Hirota et al., 2011). Consistent with the flow cytometry data, the majority of SIY-specific CCR6+ and RORyT+ cells expressed Tomato (Figure S5I). Around 30% of TCF1+ cells also expressed Tomato and harbored most of the CCR6+ population (Figure S5I). Strikingly, Tomato+ cells expressed no GZMB, even following one week of anti-PD-1/CTLA4 ICB therapy (Figure 5L). This indicates cells that have expressed IL17A do not differentiate into cytotoxic cells. Further, the proportion of Tomato+ cells was reduced following ICB treatment and suggests ICB reduces differentiation down the Tc17 pathway (Figure S5J). Altogether, these data suggest that CCR6+ TCF1+ and Tc17 cells constitute functionally inferior cell states that contribute poorly to therapeutic response to ICB.

Therapeutic Vaccination Breaks Antigen Dominance and Eliminates the CCR6+ Subset of TCF1+ Progenitor Cells

CCR6 upregulation and Tc17 differentiation in autoimmune diseases is driven by reactivity to self-antigens (Srenathan et al., 2016). Hence, we hypothesized that suboptimal T cell priming conditions might induce this phenotype in tumors and sought to improve T cell priming conditions by therapeutic vaccination of LucOS mice with SIIN and SIY long peptides (Figure 6A). Vaccination vastly improved the SIIN- and SIY-specific CD8 T cell responses in the tumor-bearing lung, increasing T cell expansion, proliferation (Ki67) and cytotoxicity (GZMB) (Figure 6B-6D). Greater fold expansion was observed for the SIY versus SIIN response, suggesting SIY-specific cells preferentially benefit from vaccination. Spatial profiling by tissue-based cyclic immunofluorescence (t-CyCIF) showed increased infiltration of CD8 T cells into tumors post-vaccination, with greater proportions expressing Ki67 and GZMB (Figures 6E-6G, S6B and S6C). This correlated with decreased tumor size, particularly when immune cell infiltrate was excluded (Figure 6F and S6A). The effects of vaccination were antigen-specific, with no contribution of adjuvant alone (Figure S6D). Interestingly, little response was observed in the lung 7 days after the initial vaccine dose (Figures 6H and S6E). However, antigen-specific T cells were expanded in the blood, inguinal lymph nodes and spleen, especially for the SIY response (Figure 6I and S6F). These data indicate that vaccination acts predominantly by increasing priming and/or expanding activated cells in the periphery rather than boosting the existing response in the lung.

Comparison of the SIIN/SIY pooled vaccine to SIIN-only and SIY-only vaccines demonstrated that the benefits of vaccination were restricted to the targeted antigen (Figure 6J and S6G). However, the SIY-only vaccine was able to increase the magnitude and

functionality of the SIY response to a similar extent as the SIIN/SIY vaccine (Figure 6J and S6G). This indicates that targeting a subdominant antigen alone with vaccination can be sufficient to rescue the subdominant response to the level of the dominant response. The SIY response remained enriched for TCF1+ progenitor cells compared to the SIIN response after vaccination against SIIN/SIY or SIY-only (Figure 6K and data not shown); however, CCR6+ TCF1+ progenitor cells were all but eliminated (Figure 6L and S6G). Amongst the remaining CCR6+ cells, fewer expressed ROR_YT, suggesting reduced propensity for Tc17 differentiation (Figure 6M). Altogether, these results provide strong evidence that CCR6 expression amongst TCF1+ cells and Tc17 differentiation is driven by suboptimal T cell priming. Moreover, they demonstrate that vaccination can "break" antigen dominance, improving the overall contribution and quality of the subdominant T cell response.

We further assessed whether vaccination could break the antigen dominance hierarchy between SIY and Y5 in LucY5S mice. Surprisingly, vaccination expanded only the dominant SIY response while subdominant Y5-specific cells remained low in number (Figure S6H). Still, Y5-specific cells benefitted functionally, with similar increases in GZMB and Ki67 expression and depletion of CCR6+ TCF1+ cells compared to SIY-specific cells (Figure S6H). Because Y5 is much less stable on MHC compared to SIY and SIIN (see Figure 3F), this suggests a minimum threshold of pMHC stability may be required to overcome antigen dominance in response to vaccination. Still, the functional enhancement of Y5-specific cells indicates that inclusion of unstable MHC binders in vaccines may still be beneficial overall.

CCR6+ TCF1+ Progenitor and Tc17 Cells are Found Across Human Cancers

To extend our findings to human cancer, we scored clusters from the mouse scRNA-seq dataset for enrichment of gene signatures (Table S4) derived from three published human CD8 T cell scRNA-seq datasets spanning lung, melanoma, colon, endometrial and renal cancers (Guo et al., 2018; Sade-Feldman et al., 2019; Wu et al., 2020). The mouse TCF1+ progenitor cluster, C8, was strongly enriched for gene signatures from clusters associated with progenitor and memory cell phenotypes, most notably two clusters described as containing TCF1+ progenitor cells in melanoma (Sade-Feldman CD8 4- and CD8 6-Memory/Effector) but also Guo CD8 C2-CD28 and C3-CX3CR1 and Wu 8.6.KLRB1 (Figures 7A and S7A). In contrast, mouse cluster C4, containing CCR6+ TCF1+ cells, showed weak to no enrichment for these signatures and was instead enriched for signatures of T cell exhaustion, including Sade-Feldman CD8 1-Exhaustion/CellCycle and Guo CD8_C6_LAYN (Figures 7A and S7A; Table S4). In a reciprocal analysis, we scored the tumor CD8 T cell subset of the Wu et al. pan-cancer dataset with gene signatures of mouse clusters C8 and C4 and found strong enrichment of C8 in the "progenitor region" of the UMAP, marked by TCF7 expression and cells assigned to the published progenitorlike cluster, 8.6.KLRB1 (Figures 7B and S7B). The C4 signature was weakly, but more broadly enriched across the UMAP, including in regions with HAVCR2 expression and cells assigned to published clusters with exhausted phenotypes, 8.3.Trm and 8.5.Mitosis (Figures 7B and S7B). These results suggest that both C8 and C4 cell populations resemble populations in human cancer; however, the C8 population is better aligned with progenitor cell states described in humans.

In the reanalysis of the Wu et al. dataset, we further identified cells co-expressing TCF7 and CCR6, some of which also expressed RORC, indicating that CCR6+ TCF1+ cells are indeed found in human cancer (Figure 7C; Table S6). IL17A-expressing cells, some co-expressing TCF7, CCR6 and/or RORC, were also found (Figure 7C; Table S6). While CCR6+ TCF7+ cells were predominantly localized to the progenitor region of the UMAP, IL17A-expressing cells were distally localized in the region expressing HAVCR2 and exhibiting a more differentiated phenotype. We identified similar cell populations with analogous localization patterns in four additional scRNA-seq datasets from individual cancer types, including lung cancer, melanoma and basal cell carcinoma (Figure S7C; Guo et al., 2018, Sade-Feldman et al., 2019, Tirosh et al., 2016, Yost et al., 2019). We further validated the presence of CCR6+ TCF1+ cells in human tumor tissue by t-CyCIF imaging and found these cells made up 2% of CD8 T cells on average across early-stage lung adenocarcinoma and metastatic melanoma samples (Figures 7D and S7D; Tables S5 and S7). Our observations in mouse lung adenocarcinoma indicate that while only a small percentage of TCF1+ cells are antigen-experienced, the majority of CCR6+ cells are contained within the activated, tumor antigen-specific response (Figure S7G). Hence, despite comprising a small proportion of total CD8 T cells, CCR6+ TCF1+ cells likely constitute a larger fraction of tumor-reactive CD8 T cells compared to CCR6- TCF1+ cells in humans.

Finally, we directly investigated a correlation between CCR6+ TCF1+ cells and response to ICB by reanalyzing the CD8 T cell subset of the Sade-Feldman et al. melanoma dataset from patients treated with ICB (Sade-Feldman et al., 2019). Consistent with published observations, cells derived from responder versus non-responder patients segregated spatially on the UMAP, with responder cells colocalizing with *TCF7* expression and progenitor-associated clusters CD8_4 thru CD8_6 (Figure 7G and S7C). Scoring the dataset with signatures derived from *CCR6+ TCF7*+ versus *CCR6- TCF7*+ cells from the mouse scRNA-seq data revealed significant enrichment of the *CCR6- TCF7*+ signature in clusters CD8_4 and CD8_6 and cells from responder patients (Figures 7H–7J). In contrast, the *CCR6+ TCF7*+ signature was enriched in exhaustion-associated cluster CD8_1, and no significant enrichment was observed in responder cells. These data indicate that CCR6- TCF1+, and not CCR6+ TCF1+ cells, are positively correlated with patient response to ICB. This is consistent with our findings in mouse lung adenocarcinoma and suggests CCR6+ TCF1+ cells contribute poorly to ICB response in human patients.

DISCUSSION

Neoantigens can drive potent anti-tumor T cell responses, yet T cell reactivity against predicted neoantigens is starkly limited across most cancers (Schumacher et al., 2019). Our study demonstrates that antigen dominance hierarchies in tumors may contribute to this narrowly focused CD8 T cell response. In KP mouse lung adenocarcinomas expressing pairs of immunogenic neoantigens, we found that CD8 T cell responses against different neoantigens compete and can result in the establishment of an immunodominant antigen. This profoundly impacted the response against the subdominant antigen, suppressing T cell expansion, differentiation and effector function. These findings demonstrate that CD8 T cell responses against concurrently expressed tumor neoantigens are interdependent, and their competition can limit the overall diversity and effectiveness of the anti-tumor response.

Multiple factors may contribute to the establishment of antigen dominance hierarchies in tumors (Schreiber et al., 2002; Yewdell, 2006). Here, using the Y5 mutant of SIIN that poorly binds MHC, we demonstrated that pMHC binding plays a central role. It has previously been shown that only tumor peptides with high affinity for MHC are efficiently cross-presented (Engels et al., 2013); hence, weak pMHC binding may limit opportunities for good interactions with antigen presenting cells (APCs). The subdominant SIY and Y5 responses in KP LucOS and LucY5S mice, respectively, were rescued when these antigens were expressed alone, despite pMHC binding being unchanged. Thus, it may be that competition between CD8 T cell responses for interactions with APCs amplifies a deficit in MHC binding and results in formation of antigen dominance hierarchies.

Subdominant T cell responses in our model were associated with enrichment of TCF1+ progenitor cells that drive CD8 T cell responses to ICB therapy in mice and humans (Philip et al., 2021). However, we found that the subdominant SIY response in KP LucOS mice did not preferentially benefit from ICB therapy, possibly due to predominance of a dysfunctional subset of TCF1+ cells marked by CCR6 expression. Another recent study from our lab corroborates there being heterogeneity amongst TCF1+ cells and identifies a SLAMF6+ subset, that contains CCR6+ TCF1+ cells, as a population recently arrived from the tumor draining lymph node (Schenkel et al., in press Immunity). Analysis of human scRNA-seq datasets spanning multiple cancer types and imaging of human lung adenocarcinoma and melanoma samples indicates this heterogeneity is reflected across human cancers (this study and Schenkel et al., in press Immunity). Furthermore, while CCR6– TCF1+ cells were correlated with melanoma patient response to ICB therapy, no correlation was observed for CCR6+ TCF1+ cells. Therefore, CCR6+ TCF1+ cells appear to contribute poorly to ICB response in both mice and humans.

In KP LucOS mice, we identified a lineage trajectory from CCR6+ TCF1+ cells to a Tc17 population. Tc17 cells have previously been reported in cancer, but their role remains unclear. While Tc17 cells have been reported to mediate tumor control in some transplant tumor settings, their production of IL17A may also contribute to tumor-promoting inflammation (Chang et al., 2014; Srenathan et al., 2016; Zhao et al., 2020). Here, we found that CD8 T cells marked by an IL17A reporter did not express GZMB, even in response to ICB therapy, suggesting Tc17 cells do not contribute to cytotoxic responses against tumors. Furthermore, our lab previously showed that treatment of KP lung adenocarcinoma with a neutralizing antibody to IL17A significantly reduces tumor burden by inhibiting recruitment of tumor-promoting neutrophils (Jin et al., 2019). Similarly, in humans, tumor infiltration by myeloid derived suppressor cells correlates with accumulation of IL17A-producing cells (Wu et al., 2014; Zhuang et al., 2012). Thus, in addition to contributing poorly to ICB response, CCR6+ TCF1+ cell differentiation to a Tc17 phenotype may also promote tumorigenesis through IL17A production.

Tc17 differentiation is predominantly observed in autoimmune inflammation and is driven by autoreactive TCR signals (Srenathan et al., 2016). Hence, we hypothesized that CCR6+ TCF1+ cells might be derived from suboptimal TCR stimulation during priming and differentiation. Consistent with this, vaccination of KP LucOS mice initially expanded cells in the blood and peripheral lymphoid organs, especially for the subdominant SIY response.

TCF1+ cells were also significantly expanded in the lung following the vaccine boost; however, the CCR6+ subset was lost. Increased antigen presentation with vaccination likely alleviates the constraints of low affinity/stability pMHC binding, improving TCR stimulation and reducing differentiation of CCR6+ TCF1+ cells. An increasingly immunosuppressive tumor microenvironment may further contribute to poor TCR stimulation as tumors progress and could explain why SIIN-specific cells develop a CCR6+ TCF1+ phenotype over time.

Vaccination of KP LucOS mice dramatically improved the magnitude and functionality of tumor-reactive CD8 T cells, and the response to vaccination was correlated with a reduction in tumor burden. Likewise, early clinical studies have demonstrated the therapeutic potential of neoantigen vaccines in melanoma, where they improve T cell priming and tumor control (Hollingsworth and Jansen, 2019). Vaccination has shown a remarkable ability to elicit *de novo* T cell responses (Ott et al., 2017; Sahin et al., 2017) and, in combination with anti-PD1 therapy, also promotes epitope spreading to neoantigens not included in the vaccine (Ott et al., 2020). Therefore, vaccination against a range of predicted immunogenic neoantigens may be effective at priming new T cell responses even in cancers with few pre-treatment T cell-reactive epitopes. Our data suggest the presence of CCR6+ TCF1+ cells is indicative of subdominant and/or poorly primed T cell responses in tumors. Consequently, CCR6+ TCF1+ cells might mark tumors that would preferentially benefit from vaccination, and should be further explored as a biomarker of patient response to vaccine therapies.

pMHC affinity is used for identification of neoepitopes for cancer vaccines. High affinity neoepitopes are typically chosen (Ott et al., 2017) and may selectively target immunodominant T cell responses. Our results indicate subdominant T cell responses against some lower pMHC affinity neoantigens may experience greater benefit from vaccination and remain enriched for TCF1+ progenitor cells. Since TCF1+ cells repopulate the cytotoxic T cell pool and are ICB-responsive, engaging subdominant T cell responses through vaccination might result in more durable tumor control and better response to ICB. Overall, these findings provide rationale for evaluating the relative response to high versus low pMHC affinity/stability antigens in clinical trials of pooled neoantigen cancer vaccines, where subdominant antigens may contribute more to tumor control than previously realized.

Limitations of the Study

In this study, neoantigen pairs were expressed clonally in tumors and the expression level of each antigen was held constant. Human tumors are generally thought to be more heterogenous, containing variable numbers of neoantigens expressed at different levels and with varying clonality. These factors may influence hierarchical ordering of antigens and could be explored in future studies by modifying antigen number, expression level and clonal fraction.

In the mouse scRNA-seq analysis, we mapped a differentiation trajectory of CCR6+ TCF1+ cells to a Tc17 population that lacks cytotoxic capacity, even after ICB. However, as lineage tracing by adoptive transfer is technically intractable in the KP lung cancer model, we were unable to directly delineate differentiation of CCR6+ TCF1+ cells. Use of novel genetic alleles, such as a fate-mapping CCR6 reporter, or investigation in other models may shed

further light on the relative functionality and plasticity of CCR6+ TCF1+ cells in response to ICB.

STAR METHODS

RESOURCE AVAILABILITY

Lead Contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Tyler Jacks (tjacks@mit.edu).

Materials Availability—Plasmids generated in this study have been deposited to Addgene: Lenti-LucOS (Cat. No. 22777), Lenti-LucSIIN (Cat. No. 174043), Lenti-LucSIY (Cat. No. 174044), Lenti-LucY5 (Cat. No. 174046), Lenti-LucY5S (Cat. No. 174045), Lenti-LucAL (Cat. No. 174049), Lenti-LucA (Cat. No. 174047), Lenti-LucL (Cat. No. 174048).

Data and Code Availability

- The mouse scRNA-seq data has been deposited to GEO and is publicly available as of the date of publication. The accession number is listed in the key resources table. This paper additionally included analyses of publicly available human scRNA-seq datasets. The accession numbers for these datasets are listed in the key resources table.
- All original code for the mouse and human scRNA-seq analyses and the t-CYCIF imaging analyses has been deposited on GitHub and is publicly available as of the date of publication. The DOIs are listed in the key resources table.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice—*Kras*^{LSL-G12D/+};*p53*^{fl/fl} (KP) and IL17a^{Cre};Rosa26^{LSL-tdTomato} mice have been previously described (DuPage et al., 2011; Hirota et al., 2011; Madisen et al., 2010) and were on a C57BL/6 genetic background. Male and female KP mice were used in all experiments and experimental arms were gender and age-matched within 3 weeks. 8 week old, female IL17a^{Cre};Rosa26^{LSL-tdTomato} were used for the collection of bone marrow to reconstitute lethally irradiated KP mice. All studies were performed under an animal protocol approved by the Massachusetts Institute of Technology (MIT) Committee on Animal Care. Mice were assessed for morbidity according to guidelines set by the MIT Division of Comparative Medicine and were humanely sacrificed prior to natural expiration.

Lentiviral Tumor Induction— 2.5×10^4 PFU of lentivirus containing *Cre* recombinase and model neoantigens were injected intratracheally into KP mice as previously described (Dupage et al., 2011, Dupage et al., 2009). Mice were at least 8 weeks old at the time of injection. Details of the lentiviral constructs and lentivirus production are included below. Mice were randomized post-infection for therapy trials. **Bone Marrow Chimeras**—Bone marrow was harvested from the femur and tibia of 8 week-old female IL17a^{Cre};Rosa26^{LSL-tdTomato} mice into sterile RPMI 1640 media. Total bone marrow was pelleted by centrifugation at 1200 rpm for 5 minutes, resuspended in PBS and filtered through a 70 μ m cell strainer. 1×10⁷ cells in 200 ul PBS were injected retroorbitally into lethally irradiated KP mice (two doses of 550 rad, three hours apart). 6 weeks after bone marrow transfer, mice were injected intratracheally with Lenti-LucOS lentivirus as described above.

METHOD DETAILS

Lentiviral Constructs and Lentivirus Production—The LucOS lentiviral construct was previously described (Dupage et al., 2011). The LucSIIN and LucSIY vectors were generated by deleting the 24 nucleotides of SIINFEKL or SIYRYYGL coding sequence, respectively, from the LucOS vector. The LucY5S vector was generated by creating a point mutation at amino acid position 5 in SIINFEKL in the LucOS construct. To make the LucY5 vector, the 24 nucleotides of SIYRYYGL coding sequence were deleted from LucY5S. The LucA and LucL vectors were generated by fusing the mALG8 or mLAMA4 neoepitope sequence plus approximately 6 flanking amino acids to luciferase. These sequences were cloned in tandom to create LucAL, with care to avoid creation of additional potentially immunogenic epitopes.

Lentiviruses were produced in 293FS* cells by transfection with the lentiviral constructs above, psPAX2 and VSV-G packaging plasmids at a 4:3:1 ratio using Mirus TransIT LT1 (MirusBio). At 48- and 72-hours post-transfection, viral supernatant was collected, passed through 0.45 μ m filters and concentrated by ultracentrifugation (25,000 rpm for 2 hours at 4°C). Concentrated lentivirus was resuspended in a 1:1 solution of Opti-MEM and HBSS and stored at -80°C. Viral titers were determined using the GreenGo 3TZ cell line as previously described (Sanchez-Rivera et al., 2014).

Anti-PD1/anti-CTLA4 Therapy—KP LucOS mice were treated for one week with *Invivo*MAb anti-PD1 (29F.1A12; BioXCell) and *Invivo*MAb anti-CTLA4 (9H10; BioXCell) monoclonal antibodies or *Invivo* isotype controls (Rag IgG2a, 2A3; Syrian Hamster, polyclonal; BioXCell). An initial dose of 200 µg each of anti-PD1 and anti-CTLA4 (or isotype control) was injected i.p. at day 0, followed by 200 µg anti-PD1 and 100 µg anti-CTLA (or isotype control) at day 3 and 6. All doses were delivered i.p. in 200 µl of PBS. Mice were analyzed one day following the last dose.

Therapeutic Vaccination—KP LucOS mice were vaccinated via subcutaneous tailbased injection with 30 amino acid peptides containing SIINFEKL, SIYRYYGL and/or SIINYEKL (10 nmol; New England Peptide) and cyclic di-GMP adjuvant (0.25 mg/ml; Invitrogen) 6 weeks post-tumor initiation. A booster injection was given 14 days later and mice were euthanized 9 weeks post-tumor initiation. All doses were delivered in two 50 µl boluses and control mice were injected with PBS. The long peptide sequences used were: SMLVLLPDEVSGLEQLE<u>SIINFEKL</u>TEWTS, GRCVGSEQLE<u>SIYRYYGL</u>LLKERSEQKLIS and SMLVLLPDEVSGLEQLESIINYEKLTEWTS (New England Peptide).

For assessment of SIIN-specific TCR cross-reactivity with Y5, wild-type mice were vaccinated against SIINFEKL as described above.

Tissue Collection and Flow Cytometry—To distinguish lung tissue-resident versus circulating immune cells, mice were injected retroorbitally with a fluorescently-conjugated anti-CD45 antibody (PE-CF594 or AlexaFluor780; 30-F11; BD Bioscience) 2-3 minutes prior to euthanasia (Anderson et al., 2014). Circulating CD8 T cells staining positive for the CD45 antibody were excluded from our analyses. Lung tissue was collected and minced by hand with spring scissors and incubated in 125 U/mL collagenase IV (Worthington Biochemical) and 40 U/mL DNase I (Sigma-Aldrich) for 30 minutes at 37°C. Following incubation, the tissue was dissociated using the m lung 2.0.1 protocol on a gentleMACS Dissociator using gentleMACS C tubes (Miltenyi Biotec) and passed through a 70 µm cell strainer. Spleen and lymph nodes were dissociated through a 70 µm cell strainer into RPMI 1640 media containing 1% heat-inactivated fetal bovine serum. Cell suspensions were pelleted by centrifugation (1200 rpm for 5 minutes), resuspended in 1X RBC Lysis Buffer (eBioscience) and incubated on ice for 10 minutes to lyse red blood cells. Pellets were resuspended in PBS and single cell suspensions were transferred to a 96-well U-bottom plate. Cells were then stained with a fixable viability dye to exclude dead cells (20 minutes on ice; Zombie Dye; Invitrogen; Tonbo Ghost Dye; Tonbo Biosciences) and resuspended in FACS buffer (1% heat-inactivated FBS in PBS) and stained with the following surface antibodies for 15-30 minutes on ice: CD8a (53-6.7), CCR6 (29-2L17), CX3CR1 (SA011F11), CXCR3 (CXCR3-173), CD44 (IM7), IL7R (A7R34), LAG3 (C9B7W), PD1 (RMP1-30), SLAMF6 (13G3), TIGIT (1G9), TIM3 (RMT3-23), OX40 (OX-86), ICOS (7E.17G9), CD200 (OX-90), CD83 (Michel-19) purchased from ThermoFisher Scientific, BD Biosciences or Biolegend (see Key Resources Table). Cells were simultaneously stained with H-2K^b peptide-MHC tetramers specific to SIINFEKL, SIYRYYGL, SIINYEKL, mALG8 or mLAMA4 (monomer, NIH Tetramer Core Facility; PE and APC streptavidin, Invitrogen). For intracellular staining, cells were fixed for 1 hour at room temperature using the eBioscience Fixation/Permeabilization Kit (ThermoFisher Scientific). Cells were then stained overnight at at 4°C with the following antibodies: TCF1/ TCF7 (C63D9), RORyT (B2D), TBET (eBio4B10), Granzyme B (GB11), Ki67 (B56), TOX (REA473), EGR2 (erongr2) purchased from Cell Signaling Technology, ThermoFisher Scientific, BD Biosciences, Biolegend or Miltenyi Biotec (see Key Resources Table). Samples were analyzed on a BD Biosciences LSR Fortessa or LSR II Flow Cytometry Analyzer.

For assaying cytokine production, T cells were enriched in lung samples by depletion of tumor cells and myeloid cells. Cells were incubated with 2 µg each of purified Ly-6G, EpCAM and F4/80 antibodies (Biolegend; 4°C for 20 minutes), then 125 µl of sheep anti-rat Dynabeads (Invitrogen; 4°C for 30 minutes with rotation). Note, to avoid depleting intravascularly labeled CD45+ cells, a mouse anti-mouse CD45.2 antibody (Brilliant Violet 510, Biolegend) was used for the intravascular stain. A Dynabeads magnet (Invitrogen) was applied to remove the beads and the T cell-containing supernatant was transferred to a clean tube, washed with 10 mL FACS buffer and pelleted by centrifugation (1200 rpm for 5 minutes). Pellets were resuspended in T-cell media (RPMI 1640 containing 10% FBS,

10 mM HEPES, 1 mM sodium pyruvate, 1X MEM Non-essential amino acids, 2 mM L-glutamine, 0.275 mM beta-mercaptoethanol, and 50 U/mL penicillin-streptomycin) and transferred to a 96-well U-bottom plate. To assay IFNY and TNF α production, samples were pelleted by centrifugation (1200 rpm for 5 minutes) and were resuspended in T cell media plus 1X Monensin (Biolegend), 1X Golgi Plug (BD Bioscience) and SIINFEKL or SIYRYYGL peptide (167 nM; New England Peptide). For IL17A staining, cells were resuspended in T cell media containing PMA (2.5 ng/ml; EMD Millipore) and Ionomycin (1 μ M; Sigma-Aldrich), 1X Monensin and 1X Golgi Plug. Samples were then incubated at 37°C for 4–5 hours. Ten percent of each sample was aliquoted and left unstimulated as a control. Cells were subjected to viability staining, surface staining and fixation as described above and stained with the following antibodies overnight at 4°C: IFN- γ (XMG1.2), TNF α (MP6-XT22) and IL17A (17B7) from ThermoFisher Scientific, BD Biosciences or Biolegend (see Key Resources Table).

scRNA-seq on Mouse CD8 T Cells—Lung single cell suspensions were prepared from 10 mice as described above. Dead cells were removed with an EasySep Dead Cell Removal Kit (Annexin V; StemCell). Cells were then stained with 1 µg TotalSeq-C hashtag antibodies 1-10 (M1/42, 30-F11; Biolegend) and surface antibodies against CD44 (IM7), CD8a (53-6.7) and H-2K^b-SIINFEKL and H-2K^b-SIYRYYGL tetramers for 30 minutes on ice. FACS sorting was performed on a FACS Aria III (BD Biosciences) and CD44+ CD8a+ SIINFEKL+ and CD44+ CD8a+ SIYRYYGL+ T cells were sorted into separate tubes, counted and resuspended in PBS (no calcium or magnesium) with 0.04% BSA. Approximately 65,000 SIINFEKL+ cells and 14,000 SIYRYYGL+ cells were loaded across two channels of a Chromium single-cell 5' chip (10X Genomics) according to manufacturer's instructions. Single cells were partitioned into droplets with gel beads to form emulsions, after which cellular lysis and barcoded reverse transcription of mRNA was performed. Paired 5' gene expression, hashtag barcode and V(D)J libraries were prepared using the following kits from 10X Genomics and protocols provided by the manufacturer: Chromium Next GEM Single Cell 5' Library and Gel Bead Kit v1.1; Chromium Single Cell V(D)J Enrichment Kit, Mouse T Cell. RNA expression libraries were sequenced individually using HiSeq X (Illumina). VDJ and hashtag barcode libraries were sequenced with HiSeq 2500 (Illumina).

Peptide Stabilization of H-2K^b on RMA-S Cells—TAP-deficient RMA-S cells were cultured in RPMI 1640 media (10% fetal bovine serum, 2 mM L-glutamine, 10 mM Hepes and 50 U/µl Penicillin-Streptomycin) at 26°C overnight (16 hours) to increase expression of empty MHC molecules on the cell surface. Cells were then plated at 100,000 cells per well in a 96-well U-bottom plate (150 µl per well). 10-fold serial dilutions of peptides (10 mM in DMSO stock) were generated in RPMI media at 4X final concentration in a 96-well U-bottom plate. 50 µl of peptide was added to the RMA-S cells, bringing the final volume to 200 µl. Cells were incubated with peptide for 2 hours at 26°C to allow for peptide binding followed by 1 hour at 37°C to degrade empty MHC molecules. Each condition was plated in triplicate. Cells were washed in FACS buffer (1% heat-inactivated FBS in PBS) and stained with an APC-conjugated H-2K^b (AF6-88.5.5.3; ThermoFisher Scientific) antibody for 15 minutes on ice. Plates were assayed on a BD Biosciences LSR II flow cytometry analyzer.

Differential Scanning Fluorimetry Assay for pMHC Thermal Melting—Empty peptide-receptive H-2K^b molecules (Saini et al., 2019) were diluted to 200 µg/ml in 50 mM HEPES pH 7.5 containing 1X SYPRO Orange dye. 18 µl of this solution was combined with 2 µl of 100 µM peptide (diluted in PBS) in triplicate in a 384 well quantitative PCR plate. The plate was incubated on ice for 10 minutes to allow for peptide-MHC interaction. The plate was then assayed on a Roche LightCycler 480 instrument. Fluorescence was acquired with excitation at 465 nM and emission at 580 nM (SYPRO Orange filter), Samples were equilibrated at 25 °C for 1 min and then heated with a gradient of 0.1 °C/s from 25 °C to 95 °C with 6 fluorescence readings for every 1 °C. Melting curves were normalized to the minimum and maximum fluorescence value and the inverse of the first derivative (–dF/dT) was then calculated. The minimum value of –dF/dT was taken at the T_m.

Histology and H&E Staining—Tumor-bearing lung lobes and a portion of the spleen were fixed in 4% paraformaldehyde overnight and embedded in paraffin. Hematoxylin and eosin stain (H&E stain) was performed with a standard method by the Hope Babette Tang Histology Facility at Koch Institute. Digitally scanned images of H&E slides were obtained with an Aperio ScanScope AT2 at 20X magnification and tumor area was annotated using Aperio ImageScope software.

t-CyCIF Staining and Imaging—The t-CyCIF experimental protocol was conducted as previously described (Du et al., 2019; Lin et al., 2018). In brief, the mouse and human lung adenocarcinoma and human melanoma FFPE slides were baked at 60°C for 30 min, dewaxed using Bond Dewax Solution (Leica Biosystems) at 72°C, and antigen retrieval was performed with Epitope Retrieval 1 Solution (Leica Biosystems) at 100°C for 20 minutes using the BOND RX Automated IHC/ISH Stainer (Leica Biosystems). All antibodies were diluted in Odyssey Intercept Buffer (plus Hoechst 33342 0.25 µg/mL; LI-COR Biosciences) and incubated overnight at 4°C in the dark. See the Key Resources Table for the complete list of antibodies. Slides were coverslipped using 20–50% glycerol solution (Sigma-Aldrich) in PBS. Images were taken using DAPI, FITC, Cy3, and Cy5 channels on the RareCyte CyteFinder Instrument (20x/0.75NA objective lens). After imaging, the fluorophores were inactivated with photobleaching solution (4.5% H2O2 and 20 mM NaOH in PBS) for 45 minutes under LED ligfFigurehts.

t-CyCIF Image Processing—The image processing of tissue cyclic immunofluorescence is organized in the following steps, each of which is described in detail below:

- i. the software ASHLAR is used to stitch, register, and correct for image acquisition artifacts (using the BaSiC algorithm). The output of ASHLAR is a single pyramid ome.tiff file for each region imaged;
- the ome.tiff file is re-cut into tiles (typically 5000×5000 pixels) containing only the highest resolution image for all channels. One random cropped image (250×250 pixels) per tile is outputted for segmentation training (using ImageJ/ Fiji);

- **iii.** using the ilastik software the labelling of nuclear, cytoplasmic and background areas are trained on the cropped images. Based on the user training the Ilastik software outputs a 3-color RGB image with label probabilities;
- the RBG probability images are thresholded and watershed in MATLAB to segment the nuclear area. The cytoplasmic measurements are derived by dilating the nuclear mask;
- v. single-cell measurements are extracted for each channel (cell pixel median and mean for both nuclear and cytoplasmic area) as well as morphological measurements of area, solidity, and cell coordinates location.

BaSiC: The BaSiC ImageJ plugin tool was used to perform background and shading correction of the original images (Peng et al., 2017). The BaSiC algorithm calculates the flatfield, the change in effective illumination across an image, and the darkfield, which captures the camera offset and thermal noise. The dark field correction image is subtracted from the original image, and the result is divided by the flatfield image correction to obtain the final image.

ASHLAR: Alignment by Simultaneous Harmonization of Layer/Adjacency Registration (ASHLAR) is used to stitch together image tiles and register image tiles in subsequent layers to those in the first layer (Lin et al., 2018; Rashid et al., 2019). For the first image layer, neighboring image tiles are aligned to one another via a phase correlation algorithm that corrects for local state positioning error. A similar method is applied for subsequent layers to align tiles to their corresponding tile in the first layer. ASHLAR outputs an OME-TIFF file containing a multi-channel mosaic of the full image across all imaging cycles. Full codes available at: https://github.com/labsyspharm/ashlar.

<u>Ilastik:</u> Ilastik is a machine learning based bioimage analysis tool that is used to obtain nuclear and cytoplasmic segmentation masks from OME-TIFF files (Berg et al., 2019). For increased processing speed, randomly selected 250×250 pixel regions from the original OME-TIFF are used as training data. ilastik's interactive user interface allows the user to provide training annotations on the cropped regions. Users are presented with a subset of the channels stacked images and label pixels as either nuclear area, cytoplasmic area, or background area. The annotations are used to train non-linear classifiers that are applied to the entire image to obtain probability masks describing the probabilities of each pixel belonging to the nuclear, cytoplasmic, or background area. A MATLAB (version 2018a) script uses these masks to construct binary masks for nuclear and cytoplasmic area.

QUANTIFICATION AND STATISTICAL ANALYSIS

Single-cell RNA Sequencing Analysis

Mouse CD8 scRNA-seq Data Processing, Cell Clustering, and Differential Expression Analysis: Cell Ranger, version 3.1.0 (Zheng et al., 2017), was used to process raw sequencing data for RNA expression, VDJ, and Cell Hashing libraries from mouse CD8 scRNA sequencing experiments. RNA expression data was aligned to the GRCm38/mm10 reference mouse transcriptome (version 3.0.0) and VDJ sequencing data was aligned to the

prebuilt mouse (GRCm38/mm10) VDJ reference supplied by 10X Genomics (version 3.1.0; Zheng et al., 2017). Cell Ranger estimated 6,472 and 3,646 cells with 15,939 and 14,834 genes detected for RNA expression libraries of SIIN- and SIY- specific CD8 T-cells at a sequencing depth of 80.4% and 76.9%, respectively. VDJ libraries captured an estimated 4,713 and 2,705 cells with 3,857 and 2,196 of those cells containing productive V-J Spanning Pair for SIIN- and SIY- specific CD8 T-cell libraries, respectively. Cell Hashing libraries were estimated to capture 6,073 and 3,319 cells at a sequencing saturation of 10.3% and 13.2% for SIIN- and SIY- specific CD8 T Cell libraries, respectively. Cell Hashing data were demultiplexed with Seurat, version 4.0.0 (Butler et al., 2018; Stoeckius et al., 2018), using counts normalized by centered-log ratio (CLR) transformation at a positive quantile threshold of 0.98 to infer which mouse each cell was harvested from.

Seurat (version 4.0.0) was used to normalize, scale, select variable features, and perform differential gene expression analysis. To filter out dying cells with poor data quality and remove probable doublets, individual cells with <100 or >4000 detected genes, with >20,000 reads, or with >5% of reads aligned to mitochondrial genome in RNA expression libraries were removed from downstream analyses. Thereafter, any remaining cells, called as doublets, were also removed from downstream analyses. RNA expression data were normalized by total expression per cell and natural log transformed with a scale factor of 10,000. Expression for all genes in SIIN- and SIY- libraries were first centered by subtracting average expression of each gene and subsequently scaled by dividing gene expression levels by their standard deviations. SIIN- and SIY- libraries were then merged in Seurat ("merge.data = TRUE"). For each cell passing quality control thresholds, metadata assignments for V(D)J clonotypes and cell hashing (mouse of origin) were made using *Python* with *Pandas* and *Numpy* (Harris et al., 2020; McKinney, 2010).

The top 2,000 variable genes used for downstream dimensionality reductions were selected in Seurat using the *vst* method (Butler et al., 2018). Principal component analysis (PCA) was performed for the first 50 principal components (PCs). An estimation of PCs to include for downstream analyses was performed with the elbow method and JackStraw analysis. Based on this estimate, and in manual consideration of gene features of PCs, the first 30 PCs were selected for downstream analyses. A shared nearest neighbor graph (SNN, k = 20) was constructed with 30 PCs and used to perform Louvain clustering (default parameters, resolution)(Meo et al., 2011). The data was then visually projected in 2-dimensional space by using the Uniform Manifold Approximation and Projection (UMAP) algorithm (Butler et al., 2018) McInnes et al., 2018) in Seurat (default parameters). Differential gene expression was performed between all clusters using the default Wilcoxon Ranked Sum test (FindAllMarkers, min.pct = 0.25) (Table S3). The top 10 differentially expressed genes for each cluster are represented in a heatmap produced with ComplexHeatmap (cluster_columns = FALSE, cluster_rows = FALSE)(Figure S2D). Enrichment for SIIN or SIY was calculated by hypergeometric test (phyper, Stats R package; alpha = 0.05).

Gene expression scores were projected on UMAPs with Seurat using the FeaturePlot function (order = TRUE). UMAP embeddings that also depict gene expression scores were generated using the plot_cells function from Monocle3.

Subsetted C2, C3, C4, C8 Analysis and Trajectory Inference: A separate analysis of cells in the mouse CD8 scRNA-seq dataset assigned to C2, C3, C4 and C8 was performed with Monocle3, version 0.2.3.0 (Cao et al., 2019; Qiu et al., 2017; Trapnell et al., 2014). The raw RNA expression data for these cells were subsetted in Seurat and exported to Monocle3 for dimensionality reduction, clustering, and trajectory analyses. Preprocessing of raw expression data (normalization and PCA) was performed for 20 PCs, using all other default parameters. Further dimensionality reduction was performed with the UMAP algorithm (default parameters, except: umap.n_neighbors = 30L, umap.fast_sgd = FALSE, preprocess_method = "PCA"; McInnes et al., 2018). Leiden clustering (Tragg et al., 2018) was performed to group cells into clusters for trajectory analyses (resolution = 0.001) with otherwise default parameters. For trajectory analyses, a principal graph was learned and plotted in UMAP space (learn_graph, use_partition = FALSE, rann.k = 20). Monocle3 UMAP coordinates were exported to Seurat for gene expression and signature analysis. Differentially expressed genes were calculated in Seurat, as described above.

Gene lists reflecting CD8 T cell states and functionality were curated from a combination of the differentially expressed genes between Seurat clusters and literature review. Clusters C2, C3, C4 and C8 were scored for expression of these genes by the same methodology of Monocle3's plot_genes_by_group function. Gene expression scores were then scaled for each cluster and visualized in heatmaps produced with ComplexHeatmap in R (cluster_columns = FALSE, cluster_rows = FALSE) (Figure 5B).

ProjecTILs T Cell Subtype Classification and Annotation: The mouse CD8 scRNA-seq data was aligned to the reference tumour-infiltrating lymphocytes (TIL) atlas and reference lymphocytic choriomeningitis virus (LCMV)-specific CD8 T cell atlas using the ProjecTILs R package (version 0.5.1; Andreatta et al., 2021). A normalized expression matrix (Seurat logNorm data) was provided as input and non-T cells were filtered using TILPRED-1.0 before alignment and batch-effect correction. Additional parameters for "make.projection" were kept as the default. Cell states were predicted for each cell using a nearest-neighbor algorithm (ProjecTILs "cellstate.predict"; Table S1) and overlaid onto the original mouse UMAP with TILPRED filtered cells set as "NA". Cell states of interest were highlighted on UMAP plots using Seurat (Butler et al., 2018), with cells of interest plotted last. Cell state annotations were used "as-is" (without a confidence threshold). We further validated these assignments by applying a confidence score threshold of 0.5 for calling a cell state and confirmed that cell state clusters remain the same.

Enrichment of SIY and SIIN cells labeled as TIL (or LCMV) progenitor or exhausted states was evaluated using a hypergeometric test (phyper, Stats R package; alpha = 0.05). Cells annotated as progenitor or exhausted were tested similarly for enrichment in the mouse clusters.

<u>Gene Signature Scoring and UMAP Projection:</u> Scoring of published mouse gene signatures was performed with Seurat (version 4.0.0; Butler et al., 2018) using the AddModuleScore function, which calculates mean expression level for each signature subtracted by aggregated expression of control signatures for each cell. Signatures were filtered to exclude genes not detected in the query dataset. For projection of the TIL/LCMV

progenitor exhausted and terminally exhausted signatures (Miller et al, 2019), deletional tolerance signature (Parish et al., 2009) and anergy signature (Safford et al., 2005), published gene signatures were filtered for genes captured in the mouse scRNA seq dataset (Table S2). Cells were then scored for each resultant signature using the AddModuleScore function in Seurat and highlighted on the Monocle3-based mouse UMAP of the mouse cluster C2, C3, C4 and C8 analysis.

For projection of the Tc17 versus Tc1 gene signatures from Linehan et al. (Linehan et al., 2018), raw RNA-seq sequence FASTQ files for Tc17 and Tc1 cells from the skin of Imiquimod-treated or *S. epidermidis*-infected mice were obtained from the Sequence Read Archive (SRA accession PRJNA419368; Imiquimod Tc17 runs: SRX3425166, SRX3425165, SRX3425145; Imiquimod Tc1 runs: SRX3425153, SRX3425146; *S. epidermidis* Tc17 runs: SRX3425152, SRX3425151, SRX3425149; *S. epidermidis* Tc1 runs: SRX3425150, SRX3425148, SRX3425149; *S. epidermidis* Tc1 runs: SRX3425150, SRX3425148, SRX3425147). Sequences were aligned to the mouse genome (NCBI37/mm9 assembly) using Bowtie, v. 1.2.3 (Langmead et al., 2009), and feature counts were quantified using rsem, v. 1.3.1 (Li and Dewey, 2011), and UCSC mm9 genome annotation (genome.ucsc.edu). Pairwise gene expression differential analyses were conducted with the DESeq2 package, v. 1.16.1 (Love et al., 2014), in R (v. 3.6.0; r-project.org). The Tc17 vs Tc1 (*S. epidermidis* or Imiquimod; Table S2) signatures were scored using Seurat (AddModuleScore with default parameters) and highlighted on the Monocle3-based mouse UMAP of the mouse cluster C2, C3, C4 and C8 analysis.

CCR6+ TCF7+ and *CCR6– TCF7*+ signatures were derived from the mouse scRNA-seq data by identifying differentially expressed genes using Seurat (FindMarkers with default parameters) with a fold change > 2 and adjusted p-value < 0.05 (Table S3). *Ccr6+ Tcf7*+ marker genes were defined by comparing cells from cluster C4 expressing both *Ccr6* and *Tcf7* (log-normalized expression count > 0) to that of all other cells in the dataset except those in cluster C4. To define *Ccr6– Tcf7*+ marker genes, cells from cluster C8 expressing *Tcf7* but not *Ccr6* were compared to the expression of all other cells in the dataset except those in cluster C8. Mouse symbols were translated to human symbols using the Broad GSEA chip file "Mouse Gene Symbol Remapping to Human Orthologs MSigDB.v7.2.chip" (Liberzon et al., 2011; Subramanian et al., 2005), subsequently scored in the Sade-Feldman dataset and visualized with UMAPs using Seurat.

TCR Clonotype Analysis: Cells with productive TCR sequences in the mouse CD8 scRNA-seq dataset were assigned to clonotypes by Cell Ranger. While the vast majority (581 of 652) of clonotypes appear to originate exclusively from a single mouse, a minority (22 of 652) of clonotypes appear to have one or more cells that originate from multiple mice. These clonotypes were identified in Python with *Pandas* and excluded from downstream analyses. Most of the clonotypes detected by Cell Ranger contain less than 5 cells (548 of 652), which may reflect their underlying biological abundance (i.e. degree of clonal expansion). However, it is impossible to decouple this underlying biology from artifacts of technical dropout in these lowly-abundant clonotypes. As such, clonotypes containing less than 5 cells were removed in most analyses, resulting in a total of 103 clonotypes.

In order to understand how clonotypes were distributed across the Seurat clusters (C0– C10), a matrix was generated that quantifies the number of cells in each clonotype assigned to each Seurat cluster. To permit comparison of clonotypes of different sizes, this matrix was normalized to reflect the fraction of cells in each clonotype. To cluster clonotypes with similar distributions across Seurat clusters, Euclidean distance was first calculated between clonotypes in the normalized clonotype-cluster matrix. Visualizations of the clonotype-cluster matrix were made with ComplexHeatmap. Rows (clonotypes) in this heatmap were ordered by hierarchical clustering (method = "ward.D") of default pairwise euclidean distances. Columns (clusters) were ordered by hierarchical clustering (method = "ward.D") of pairwise Jaccard distance. Subsequently, complete linkage hierarchical clustering (hclust, R, method = "ward.D") was performed on these distance calculations. A total of 12 clonotype clusters were generated by cutting the resulting dendrogram (h = 0.6, cutree, R).

Cells assigned to each clonotype were scored for expression of *Gzmb, Havcr2, Cx3cr1, Tcf7, Ccr6,* and *II17a* by counting the number of cells with gene expression values > 0.5. To test enrichment of these genes, we performed a hypergeometric test (phyper, Stats R package; alpha = 0.05). To test enrichment of clonotypes specific for SIIN and SIY in C4+C8 and C2, we performed a 2-dimensional 2-sample KS test and visualized differences in proportion of cells with an empirical cumulative distribution plot. Additionally, to provide a metric for how equally distributed each clonotype is across clusters, we calculated the Gini index (Hurley, 2009) for each clonotype.

To visualize the location of cells from large and small clonotypes in UMAP space, we assigned each cell a value equivalent to the size of the clonotype assigned to each cell and plotted this feature in Seurat.

Enrichment of Human Signatures in Mouse Clusters: Marker genes for published CD8 T cell cluster signatures from Wu et al., Supplemental Table 4 (Wu et al., 2020), Guo et al., Supplemental Table 3 (Guo et al., 2018), and et al., Supplemental Table 4 (Sade-Feldman et al., 2019) were translated from human symbols to mouse symbols using the Mouse Genome Informatics (MGI) mouse/human orthology with phenotype annotations obtained on November 16, 2020 (Baldarelli et al., 2020) [http://www.informatics.jax.org/downloads/ reports/HMD HumanPhenotype.rpt]. For ambiguous mappings (i.e. human genes mapping to more than one mouse gene), we first prioritized the lowercase equivalent of the human gene, if available. For cases where lowercase equivalents were not an option, we checked to see if one of the possible translations was in the mouse dataset and if so, then we arbitrarily selected the first translation (but avoided symbols with the Gm* prefix if another option was available). For cases where neither the lowercase equivalent nor a match in the mouse dataset was found, the first translation was kept as a placeholder. In addition, genes without a mapping were dropped and not included in the final lists. Signature gene lists used for downstream analysis are provided in Table S4. Human signatures were scored in the mouse dataset using Seurat (AddModuleScore with default parameters) and visualized on the mouse Seurat-based UMAP (Figure S7A).

Heatmaps illustrating enrichment of human signatures in mouse clusters (Figure 7A) were generated in R using heatmap.2 and data was preprocessed using a custom script. For each human signature and mouse cluster, a Wilcoxon rank-sum test was performed to test for differential enrichment of a signature across clusters (i.e. a set of signature scores for a given cluster compared to signature scores across all other clusters) similar to DeTomaso et al. (DeTomaso et al., 2019). To reduce overall noise and highlight enrichment for clusters that included "sufficient" high-scoring cells, significance testing was conditioned according to distributional (i.e. quantile) cutoffs. Differential enrichment between sets of signature scores was only tested if the following conditions were met: the upper 2nd percentile of scores of cells outside the cluster, (ii) the upper 2nd percentile of scores of cells across the signature, and (iii) the upper 40th percentile of scores all signatures (i.e. a full matrix of signature scores per cell per cluster).

Analysis of External Human scRNA-seq Datasets

GSE139555 Wu et al.: Original cluster assignments for cells in the Wu et al. dataset (Wu et al., 2020) were retrieved from GEO (accession GSE139555; GSE139555_tcell_metadata.txt). Additionally, utility functions within the software library released in the GEO record (GSE139555 software.tar.gz) were used for extracting raw count tables with code updates to subset the dataset to contain cells assigned to CD8 T cell clusters in the original study (8.1.Teff, 8.2.Tem, 8.3a.Trm, 8.3b.Trm, 8.3c.Trm, 8.4.Chrom, 8.5.Mitosis, 8.6.KLRB1). Raw count tables for CD8 T cells per tumor sample, ignoring cells from normal adjacent tissue and peripheral blood samples across all patients, were then used in subsequent analyses. Counts per tumor sample were normalized using Seurat v3.2 SCTransform (Butler et al., 2018; Hafemeister and Satija, 2019), while regressing out percentage mitochondrial content per cell. Samples were integrated in a hierarchical fashion (per patient, then across patients) using the Seurat reference-based workflow (Stuart et al., 2019). Choice of reference sample did not affect result interpretation. PCA (Principal Component Analysis with top 30 components used for downstream analyses) dimensionality reduction and UMAP (Uniform Manifold Approximation and Projection) embeddings were generated using the integrated dataset.

For projection of mouse scRNA-seq cluster gene signatures onto the Wu analysis, the top 20 positive marker genes for each mouse cluster (C4 and C8, clustering resolution 0.7) were identified using Seurat (FindMarkers) with a fold change cutoff of 1.2 and adjusted p-value less than 0.05. Mouse symbols were translated to human symbols using the Broad GSEA chip file (MSigDB v7.2; https://data.broadinstitute.org/gsea-msigdb/msigdb/annotations_versioned/ Mouse_Gene_Symbol_Remapping_Human_Orthologs_MSigDB.v7.2.chip)(Liberzon et al., 2011; Subramanian et al., 2005), subsequently scored in the Wu dataset and visualized on the Wu UMAP using Seurat. Signature gene lists used for downstream analysis (Table S3) reflect the collection of unique genes for which an ortholog has been mapped in human.

<u>GSE120575 Sade-Feldman et al.</u>: Log2-TPM counts were retrieved from GEO along with patient characteristics, including response to immune checkpoint blockade, and processed

with Seurat v3.0.0 (Hafemeister and Satija, 2019). A Seurat object was created such that the counts slot and data slot were populated with TPM counts and log-TPM counts, respectively. Only cells assigned to published CD8 T cell clusters (Table S2; Sade-Feldman et al., 2019) were selected and used for downstream analyses. Highly variable genes were identified and the data was scaled (while regressing out percentage mitochondrial content per cell and the number of detected reads per cell) followed by dimensionality reduction with PCA and UMAP (using 24 principal components) using Seurat's standard preprocessing procedure.

GSE99254 Guo et al.: Raw counts were retrieved from GEO (accession GSE99254). Cells from seven CD8 T cell clusters in the published analysis were retained for downstream analyses. This subset was processed using the SCT normalization flow of Seurat v3.0.0 (Hafemeister and Satija, 2019). Following dimensionality reduction using PCA, 12 principal components were used for downstream analyses including UMAP embeddings.

GSE72056 Tirosh et al.: Log2(TPM/10 + 1) counts were retrieved from GEO along with patient metadata and processed with Seurat v3.2 (Hafemeister and Satija, 2019). A Seurat object was created such that the counts slot and data slot were populated with TPM counts and log-TPM counts, respectively. The dataset was filtered to retain T cells based on metadata annotation and further filtered to retain only CD8 T cells by filtering on expression of CD8A (> 0) and CD4 (= 0). Only six patient samples with 50 or more CD8 T cells were selected and used for downstream analyses and integrated using Seurat's reciprocal PCA (RPCA) integration flow. 30 principal components were used for downstream UMAP dimensionality reduction and data embedding.

GSE123813 Yost et al.: Raw counts were retrieved from GEO along with patient information and processed with Seurat v3.2 (Hafemeister and Satija, 2019). The dataset was filtered to retain basal cell carcinoma (BCC) T cells based on metadata annotation and further filtered to retain only CD8 T cells by filtering on expression of CD8A (> 0) and CD4 (= 0). A hierarchical integration flow based on SCT normalization was used, similar to the one described for Wu et al. above (GSE139555). Pre- and post-treatment samples per patient were integrated in the first round and cross-patient integration was performed using Seurat's reference-based integration workflow. The code was largely based on that used for the GSE139555 analysis (described above). 30 principal components were used for downstream analyses.

For the analyses of GSE139555, GSE120575, GSE99254, GSE72056, GSE123813 above, the Seurat default "RNA" assay with log-normalized counts was used for feature plots, marker gene identification, and all expression-based analyses. For single gene and co-expression analyses, log-normalized expression count thresholds of 0.0 (raw count datasets GSE139555, GSE99254, GSE123813) and 0.5 (TPM datasets GSE72056, GSE120575) were used for considering a gene as expressed in a given cell.

Identification of mucosal associated invariant T (MAIT) cells: In the human scRNA-seq datasets we identified a population of mucosal associated invariant T (MAIT) cells that phenotypically resemble Tc17 cells (Toubal et al., 2019). MAIT cells express invariant TCRs that recognize antigen presented on the MHC-related molecule MR1. Cells with

these TCRs were assigned to background (i.e. not highlighted) in the *TCF7/CCR6/RORC/ IL17A* expression plots and excluded from statistical analyses of CCR6+/– TCF7+ signature enrichment in Figures 7C, 7I, 7J and S7C. Importantly, MAIT cells are also found in mice, but do not express CD8 and are not found amongst SIIN- or SIY-specific T cells that are MHC I K^b-restricted in KP LucOS lung tumors.

For Wu et al., clonotype assignments were retrieved from GEO record GSE139555. MAIT cells were identified based on their invariant TCR using the following logic: (alpha_v = "TRAV1-2") AND one of (\$alpha_j = "TRAJ33") OR (\$alpha_j = "TRAJ12") OR (\$alpha_j = "TRAJ20") and assigned to the background set (not highlighted) in co-expression analysis (Toubal et al., 2019). For Sade-Feldman et al., clonotype assignments were requested and obtained from the authors (private communication). MAIT cells were identified using the logic described above. For Guo et al., cells identified as belonging to a MAIT cluster (CD8-C7-SLC4A10) in the published analysis (Guo et al., 2018) were tagged as MAIT cells. For Tirosh et al., we obtained raw scRNAseq FASTQ files through the Broad Institute Data Used Oversight System (duos.broadinstitute.org). This data was processed using MiXCR's v3.0.13 "analyze shotgun" workflow (Bolotin et al., 2017) to catalog the T cell receptor repertoire. No cells satisfying the MAIT clonotype assignments described above were detected in this set. For Yost et al., clonotype assignments were requested and obtained from the authors (private communication). MAIT cells were identified using the logic described above were detected in this set. For Yost et al., clonotype assignments were requested and obtained from the authors (private communication). MAIT cells were identified using the logic described above.

Lists of MAIT cells per dataset are included in Table S6. Using a relaxed approach, cells without clonotype information were assumed to be non-MAIT. Overall, MAIT cells were not highlighted in co-expression plots and were dropped from CCR6+/– TCF7+ signature enrichment analyses.

Enrichment of CCR6+/– TCF7+ signatures in GSE120575 (Sade-Feldman et al.): A hypergeometric test (phyper, Stats R package; alpha = 0.05) was used to test for enrichment of cells from published clusters in the set of cells with high *CCR6+ TCF7+* or *CCR6– TCF7+* signature scores (standardized signature score > 2). Similarly, cells annotated as responders/non-responders were tested for enrichment in the set of cells with high *CCR6+/ TCF7+* signature scores (standardized signature score > 2). Response categories were determined by RESIST criteria: responders, complete or partial response; non-responders, stable or progressive disease (Sade-Feldman et al., 2019). MAIT cells were removed from the dataset prior to calculating proportions.

t-CyCIF Data Analysis Workflow—The data analysis is divided in a set of preprocessing steps in which data from different tissues is i) log2-transformed and aggregated together, ii) filtered for image analysis errors, and iii) normalized on a channel-by-channel basis across the entire data from a single experiment. All the steps are performed in MATLAB.

Data Aggregation: The image processing workflow outputs one ome.tiff image and one data file (.mat) for each tissue area imaged. The data matrices from each .mat file are concatenated into a single matrix for each metric measured (median/mean, nuclear/

cytoplasmic) into a single structure ("AggrResults"). The morphological data (i.e., area, solidity, and centroid coordinates) is concatenated into a single structure ("MorpResults"), which also contains the indexing vector to keep track of the tissue of origin within the dataset.

Data Filtering: Single cells are filtered to identify and potentially exclude from subsequent analysis errors in segmentation and cells lost through the rounds of imaging. Two types of criteria are used to filter cells: morphological criteria based on cell object segmented area, which are applied to all the rounds for the cell object, and DAPI-based criteria which are applied to the DAPI measurement for each imaging round. The latter corrects for cell loss during cycling and computational misalignment, which are both round specific.

Morphological filtering criteria are: 1) nuclear area within a user-input range; 2) cytoplasmic area within a user-input range; 3) nuclear object solidity above a user-input threshold. DAPI-based criteria are: 1) nuclear DAPI measurement above a user-input threshold; 2) ratio between nuclear and cytoplasmic DAPI measurement above a user-input threshold. The filter information for the criteria is allocated to a logical (0–1) structure 'Filter', which is used to select the cells to analyze in the further analysis by indexing. The threshold selection is dataset dependent and is performed by data inspection. The values used in each dataset are available with the codes used for data analysis in the github repository https://github.com/santagatalab/2021_Burger_et_al_CyCIF_codes.

Data Normalization: Each channel distribution is normalized by probability density function (pdf) centering and rescaling. The aim is to center the distribution of the log2 fluorescent signal at 0 and rescale the width of the distribution to be able to compare across channels. The data is first log-transformed (base 2). The standard normalization is performed using a 2-component Gaussian mixture model, each model capturing the negative and the positive cell population. If the 2-component model fails to approximate the channel distribution, two other strategies are attempted: i) a 3-component model is used assuming the components with the two highest means are the negative and positive distribution (i.e., discarding the lowest component) or ii) the user selects a percentage 'x' of assumed positive cells and a single Gaussian distribution fit is performed on the remainder of the data to capture the negative distribution. The single Gaussian fit is then used as the lower component in a 2-component model to estimate the distribution of the positive population. The strategy chosen for each channel in each dataset is available in the github repository https://github.com/santagatalab/2021_Burger_et_al_CyCIF_codes. The "add_coeff" is defined as the intersection of the negative and positive distributions. The "mult_coeff" is defined as the difference between the mean of the negative and positive distributions. The full distribution is normalized by subtracting the add coeff and dividing by the mult_coeff. The normalization is performed on the nuclear and cytoplasmic single-cell, single-channel distributions individually. The data preprocessing workflow is performed on all datasets. The individual analyses used in the paper are performed only in select datasets as follows.

<u>Cell Type Calling:</u> Cells from tissue-based experiments are classified into lineage compartments by cell type markers, by gating on the sign of the normalized values of

cell type markers. Cells are scored by increasingly deeper lineage layers, starting with separating them into epithelial, immune and stromal compartments, and subdividing the immune population into lymphoid lineages. The cell types identified and the markers used to do so are included in Table S7. If a cell was called as more than one single cell-type, this is defined as a conflict. The conflicts are resolved by comparing the markers that triggered each of the cell type calls and assigning the cell type with the highest marker level. If the markers are within 10% of each other, the cell is deemed "not classifiable" for the specific layer.

Identification of CCR6+ TCF1+ CD8+ T Cells: The quantification of positive cells was performed on a single channel basis for CD8, CCR6 and TCF1. For CD8 and TCF1 signal the cells were called positive based on the normalization described above. However, in the lung cancer tissues imaged, CD20 positive B cells expressed high levels of CCR6, consistent with the current literature (Kadomoto et al., 2020). The variability of B cells between individual tissues and the spatial proximity between T cells and B cells can potentially lead to overestimation of CCR6 positivity. In order to avoid potential biases we modified the protocol in three ways: 1) the CCR6 protein was normalized at a single tissue basis to match the negative signal distribution, 2) when calculating the CCR6+ CD8+ cells (independent of TCF1 status) we discarded all cells closer than 32.5 μ m (100 pixels), and 3) we imaged the CCR6 protein twice in the first and last round of the experiment.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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DECLARATION OF INTERESTS

T.J. is on the Board of Directors of Amgen and ThermoFisher Scientific, a co-founder of Dragonfly Therapeutics and T2 Biosystems, an SAB member of Dragonfly Therapeutics, SQZ Biotech, and Skyhawk Therapeutics and President of Break Through Cancer. T.J. lab receives funding from the Johnson & Johnson Lung Cancer Initiative and The Lustgarten Foundation for Pancreatic Cancer Research, but this funding did not support this research. A.R. is a founder and equity holder of Celsius Therapeutics, an equity holder in Immunitas Therapeutics and until Aug. 31, 2020 was an SAB member of Syros Pharmaceuticals, Neogene Therapeutics, Asimov and ThermoFisher Scientific. From Aug. 1, 2020, A.R. is an employee of Genentech (Roche Group). S.S. is a consultant for RareCyte, Inc. A.M.C. interned at Skyhawk Therapeutics. None of these affiliations represent a conflict of interest with respect to the design or execution of this study or interpretation of data presented in this manuscript.

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HIGHLIGHTS

- Peptide-MHC binding property underlies antigen dominance hierarchy in a lung cancer model
- T cells responding to subdominant antigens are enriched in TCF1+ progenitors
- CCR6+ TCF1+ progenitor cells contribute poorly to checkpoint blockade response
- Vaccination can eliminate CCR6+ TCF1+ cells and improve the subdominant response





(A) KP LucOS genetically engineered mouse model of lung adenocarcinoma expressing tumor-specific neoantigens SIINFEKL (SIIN) and SIYRYYGL (SIY).

(B-C) Percentage (B) and absolute number (C) of SIIN- and SIY-specific CD8+ CD44+ T cells in tumor-bearing lung tissue by H-2K^b peptide-MHC tetramer staining and flow cytometry.

(D-H) Percentage of SIIN- and SIY-specific CD8 T cells expressing Ki67 (D), PD1, LAG3 and TIM3 (E), IL7R (F), IFNy and TNFa (G) and GZMB (H) by flow cytometry. (B-H) are representative of 3 independent experiments per time point, n 5 mice per group. Results here and in the following figures are expressed as mean + SD. Statistics were calculated by two-tailed Student's t test: ns = not significant, * p < 0.05, ** p < 0.01, *** p < 0.001.

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Figure 2. Single-Cell RNA-Sequencing Reveals Enrichment of a TCF1+ Progenitor Cell Phenotype Amongst SIY-Specific CD8 T Cells

(A) UMAP embedding of scRNA-seq 5' gene expression data comparing SIIN- and SIY-specific CD8 T cells at 5 weeks post-tumor initiation. n = 10 mice, 4,023 SIIN and 1,861 SIY cells.

(B) Unsupervised clustering of the scRNA-seq data in (A).

(C) Assignment of SIIN- versus SIY-specific cells to the clusters in (B). Brackets indicate significant enrichment (p < 0.05) for SIIN- or SIY-specific cells.

(D) Classification of individual cells from (A) using the ProjecTILs pipeline (Andreatta et al., 2021). SIY-specific cells are enriched for a progenitor phenotype (TIL p = 0.002; LCMV p = 2.51E-15); SIIN-specific cells are enriched for an exhausted phenotype (TIL p = 4.01E-17; LCMV p = 1.52E-10).

(E) Log-normalized expression of genes associated with progenitor (*Tcf7 and II7r*) or exhausted *Gzmb and Havcr2*) cell states.

(F) Heatmap depicting proportions of TCR clonotypes (rows) assigned to each cluster (columns), for clonotypes 5 cells with 1 cell assigned to TCF1+ progenitor clusters C4 or C8.

(G-H) Percentage of SIIN- versus SIY-specific cells expressing TCF1, TIM3 and GZMB (G) and the proportion of SIY-specific TCF1+ TIM3– and TCF1– cells expressing IFN γ and TNFa (H) at 5 weeks by flow cytometry.

(I) Percentage and absolute number of SIIN- and SIY-specific TCF1+ TIM3- cells over time by flow cytometry.

(G-I) are representative of 3 independent experiments, n 5 mice per group.



Figure 3. Enrichment of TCF1+ Progenitor Cells in the SIY Response is Driven by an Antigen Dominance Hierarchy

(A) Lentiviral constructs expressing SIIN and SIY (LucOS), SIIN-only (LucSIIN) and SIY-only (LucSIY) and representative flow cytometry plots of antigen-specific cells in the lung 5–6 weeks post-tumor initiation.

(B and C) Percentage of SIIN- and SIY-specific cells of CD8 T cells (B) and the proportion that are TCF1+ TIM3– (C) in LucOS, LucSIIN and LucSIY mice at 5–6 weeks by flow cytometry.

(D) Lentiviral constructs expressing SIINYEKL (Y5) in combination with SIY (LucY5S) or alone (LucY5) and representative flow cytometry plots of antigen-specific cells in the lung 5–6 weeks post-tumor initiation.

(E) pMHC affinity for each antigen assessed by flow cytometric quantification of $H-2K^b$ stabilization on TAP-deficient RMA-S cells. Dotted lines = 50% binding concentration, which approximates the affinity (K_d) of each peptide for $H-2K^b$.

(F) pMHC complex stability (K_{off}) for each antigen measured by Differential Scanning Fluorimetry of pMHC thermal melting. Dotted lines = melting temperature (Tm) of each pMHC complex. Displayed Tm values are averaged from 3 experiments.

(G) TCR cross-reactivity measured by SIIN and Y5 tetramer co-staining of CD8 T cells from spleens of SIIN-vaccinated mice. Percentage of tetramer+ cells stained by both tetramers is graphed.

(H-I) Percentage of Y5-, SIIN- and SIY-specific cells of CD8 T cells (H) and the proportion that are TCF1+ TIM3– (I) at 5–6 weeks in LucOS, LucY5S, LucY5 and LucSIIN mice. (A-F) and (H-I) are representative of 3 independent experiments, n 5 mice per group. (E and G) are representative of 2 independent experiments.

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Figure 4. Subdominant Antigen-Specific T Cells Do Not Preferentially Benefit from anti-PD1/CTLA4 Therapy

(A) Time course of anti-PD1/CTLA4 therapy initiated at 5, 8 or 12 weeks post-tumor initiation, comprised of 3 doses (arrows) of isotype control (Iso) or anti-PD1/CTLA4 (PC) antibodies in LucOS mice.

(B-G) Absolute number of SIIN- versus SIY-specific CD8 T cells (B), the proportion expressing TCF1+ TIM3- versus TCF1- TIM3+ (C), the absolute number of TCF1+ TIM3- cells (D), and the proportion expressing Ki67 (E), GZMB (F) and CX3CR1 (TCF1+ and total; G) assessed by flow cytometry in LucOS mice following treatment with Iso or PC starting at 8 weeks unless otherwise indicated. Representative of 3 independent experiments, n 5 mice per group.



Figure 5. The Subdominant CD8 T Cell Response is Enriched for a CCR6+ TCF1+ Progenitor Cell Subset with a Tc17 Differentiation Trajectory

(A) UMAP embedding of clusters C2, C3, C4 and C8 from the scRNA-seq data in Figure 2B and log-normalized expression of *Tcf*7 and *Havcr2*.

(B) Expression (mean log(expression +1)) of genes associated with the indicated

classifications across clusters C2, C3, C4 and C8.

(C) Scoring of cells from (A) for enrichment of a gene signature differentially upregulated in Tc17 versus Tc1 CD8 T cells in the skin of mice infected with *S. epidermidis* (Linehan et al., 2018; Table S2).

(D) Monocle3 lineage trajectories overlaid with UMAP visualizations of *Ccr6, Rorc* and *II17a* expression.

(E) TCR clonotype cluster 7 (see Figures S4A and S4D) highlighted on the UMAP plots from (A).

(F-G) Percentage of SIIN- and SIY-specific cells expressing CCR6 (F) and the proportion of these cells expressing TCF1, SLAMF6 and ROR γ T (G) at 5 weeks by flow cytometry. CCR6+ ROR γ T+ cells are subsetted by TCF1 expression.

(H) Percentage of SIIN- and SIY-specific cells expressing IL17A at 5 weeks and flow cytometry plots showing co-expression of IL17A with CCR6, RORyT and TCF1 on SIY-specific cells.

(I-J) Percentage of SIIN- and SIY-specific TCF1+ TIM3– cells expressing CCR6 over time and following one week of treatment with isotype (Iso) or anti-PD1/CTLA4 (PC) antibodies starting at 5 weeks.

(K) Percentage of SIIN- and SIY-specific TCF1+ cells that express CX3CR1 subsetted by CCR6 expression following Iso or PC treatment at 5 weeks.

(L) Irradiated KP mice were reconstituted with IL17a^{Cre} × Rosa26^{LSL-tdTomato} bone marrow and LucOS tumors were initiated. Flow cytometric analysis of the percentage of SIY-specific cells expressing Tomato (marking current/prior IL17A expression) following one week of Iso or PC treatment starting at 5 weeks is shown. Data are compiled from 2 independent experiments. Data in (F-K) are representative of 3 independent experiments, n 5 mice per group.



Figure 6. Therapeutic Vaccination Breaks Antigen Dominance and Eliminates the CCR6+ Subset of TCF1+ Progenitor Cells

(A) SIIN and SIY long-peptide and cyclic di-GMP vaccine. Mice were vaccinated at 6 weeks (s.c. tail base), boosted at 8 weeks and analyzed 9 weeks post-tumor initiation.(B-D) Absolute number of SIIN- and SIY-specific cells (B) and the percentage expressing Ki67 (C) and GZMB (D) in PBS control (Ctrl) versus vaccine (Vax) treated mice by flow cytometry.

(E-G) Tissue-based cyclic immunofluorescence (t-CyCIF) imaging of lung lobes of Ctrl versus Vax-treated mice (E) and quantification of CD8 T cell infiltration and tumor size (F) and the percentage of Ki67 and GZMB positive CD8 T cells in tumors (G). Results are averaged from all tumors from two lung lobes per mouse.

(H-I) Absolute number of SIIN- and SIY-specific cells in the lung (H) and the percentage in the blood (I) of Ctrl versus Vax treated mice 7 days after the initial vaccine dose.

(J) Absolute number of SIIN- and SIY-specific cells in mice treated with Ctrl, Vax, SIINonly vaccine or SIY-only vaccine as in (A).

(K-L) Percentage of SIIN- and SIY-specific cells that are TCF1+TIM3-(K) and the proportion of these cells that express CCR6 (L) in Ctrl versus Vax treated mice by flow cytometry.

(M) Proportion of CCR6+ SIIN- and SIY-specific cells expressing RORyT in Ctrl versus Vax treated mice by flow cytometry.

(B-D) and (J-M) are representative of 3 independent experiments, n 5 mice per group. (H-I) are compiled from 2 independent experiments.



Figure 7. CCR6+ TCF1+ and Tc17 Cells are Found Across Human Cancers

(A) Enrichment of CD8 gene signatures from three human cancer scRNA-seq datasets (Table S4) in the mouse scRNA-seq clusters from Figure 2B. Human clusters, right; mouse clusters, top. Colored boxes indicate significant enrichment, p = 0.05.

(B-C) UMAP of tumor CD8 T cells from a human pan-cancer scRNA-seq dataset (Wu et al., 2020) showing enrichment of gene signatures derived from mouse progenitor clusters C4 and C8 (B) and expression of *TCF7*, *HAVCR2*, co-expression of *TCF7*, *CCR6 and RORC* and co-expression of *IL17A* with *TCF7*, *CCR6* and/or *RORC* (C).

(D-F) t-CYCIF images of human lung adenocarcinoma and metastatic melanoma samples (D-E) and quantification of CD8 T cells expressing TCF1 and CCR6 (F).

(G) UMAP embedding of CD8 T cells from an anti-PD1 and/or anti-CTLA4 melanoma scRNA-seq dataset (Sade-Feldman et al., 2018) showing cell assignment to published clusters CD8_1 thru CD8_6 and responder versus non-responder patients.

(H) Scoring of cells in (G) for enrichment of gene signatures (Table S3) derived from mouse Ccr6-Tcf7+ and Ccr6+Tcf7+ cells.

(I-J) Enrichment of the gene signatures from (H) in clusters CD8_1 thru CD8_6 (I) and responders versus non-responders (J).

In (C) and (I-J), mucosal invariant T (MAIT) cells that phenotypically resemble Tc17 cells were excluded from the analysis based on TCR sequence (Table S6; STAR methods).

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
In VivomAb PD1	BioXCell	Clone 29F.1A12; Cat# BE0273; RRID: AB_2687796
In VivomAb CTLA4	BioXCell	Clone 9H10; Cat# BE0131; RRID: AB_10950184
<i>In Vivo</i> mAb rat IgG2a	BioXCell	Clone 2A3; Cat# BE0089; RRID: AB_1107769
In VivomAb polyclonal Syrian Hamster IgG	BioXCell	Cat# BE0087; RRID: AB_1107782
FC: CD8a	BD Biosciences	Clone 53-6.7; Cat# 563786, 612759; RRID: AB_2732919, AB_2870090
FC: CCR6 (CD196)	BioLegend	Clone 29-2L17; Cat# 129822; RRID: AB_2687019
FC: CX3CR1	BioLegend	Clone SA011F11; Cat# 149031; RRID: AB_2565939
FC: CXCR3 (CD183)	BD Biosciences	Clone CXCR3-173; Cat# 562937, 741032; RRID: AB_2687551, AB_2740650
FC: CD44	BD Biosciences; BioLegend	Clone IM7; Cat# 563736, 612799, 103057, 103059; RRID: AB_2738395, AB_2870126, AB_2564214, AB_2571953
FC: CD45	ThermoFisher Scientific; BD Biosciences	Clone 30-F11; Cat# 47-0451-80, 562420; RRID: AB_1548790, AB_11154401
FC: CD83	BioLegend	Clone Michel-19; Cat# 121506; RRID: AB_572009
FC: CD200	BD Biosciences	Clone OX-90; Cat# 565546; RRID: AB_2739288
FC: EGR2	ThermoFisher Scientific	Clone erongr2; Cat# 17-6691-80; RRID: AB_11150966
FC: Granzyme B	BioLegend, BD Biosciences	Clone GB11; Cat# 515408, 560213; RRID: AB_2562196, AB_1645453
FC: H-2K ^b	ThermoFisher Scientific	Clone AF6-88.5.5.3; Cat# 17-5958-82; RRID: AB_1311280
FC: ICOS (CD278)	ThermoFisher Scientific	Clone 7E.17G9; Cat# 46-9942-80; RRID: AB_2744727
FC: IFN-γ	BioLegend	Clone XMG1.2; Cat# 505822; RRID: AB_961359
FC: IL17A	ThermoFisher Scientific	Clone 17B7; Cat# 11-7177-81; RRID: AB_763581
FC: IL7R (CD127)	ThermoFisher Scientific	Clone A7R34; Cat# 25-1271-82; RRID: AB_469649
FC: Ki67	BD Biosciences	Clone B56; Cat# 561277, 563756; RRID: AB_10611571, AB_2732007
FC: LAG3 (CD233)	BD Biosciences, ThermoFisher Scientific; BioLegend	Clone C9B7W; Cat# 741820, 46-2231-82, 11-2231-80, 125219; RRID: AB_11151334, AB_2572483, AB_2566571
FC: OX40 (CD134)	BioLegend	Clone OX-86; Cat# 119419; RRID: AB_2687175
FC: PD1 (CD279)	BioLegend	Clone RMP1-30; Cat# 109110, 109121; RRID: AB_572017, AB_2687080

REAGENT or RESOURCE	SOURCE	IDENTIFIER
FC: RORŸT	ThermoFisher Scientific	Clone B2D; Cat# 25-6981-82, 17-6981-82; RRID: AB_2784671, AB_2573254
FC: SLAMF6 (LY-108)	BD Biosciences	Clone 13G3; Cat# 742272; RRID: AB_2871448
FC: TCF1/TCF7	Cell Signaling Technology	Clone C63D9; Cat# 6444, 6709, 9066; RRID: AB_2797627, AB_2797631, AB_2797696
FC: TIGIT	BD Biosciences	Clone 1G9; Cat# 565474; RRID: AB_2739254
FC: TIM3	BioLegend	Clone RMT3-23; Cat# 119721, 119725; RRID: AB_2616907, AB_2716066
FC: TNFa	BioLegend	Clone MP6-XT22; Cat# 506306; RRID: AB_315427
FC: TOX	Miltenyi Biotec	Clone REA473; Cat# 130-118-335; RRID: AB_2751485
FC: H-2Kb SIINFEKL monomer	NIH Tetramer Core Facility	Custom
FC: H-2Kb SIYRYYGL monomer	NIH Tetramer Core Facility	Custom
FC: H-2Kb SIINYEKL monomer	NIH Tetramer Core Facility	Custom
FC: H-2Kb mALG8 monomer	NIH Tetramer Core Facility	Custom
FC: H-2Kb mLAMA4 monomer	NIH Tetramer Core Facility	Custom
Hashtag: TotalSeq-C anti-mouse	BioLegend	Clones M1/42, 30-F11; Cat# 155861, 155863, 155865, 155867, 155869, 155871, 155873, 155875, 155877, 155879; RRID: AB_2800693, AB_2800694, AB_2800695, AB_2800696, AB_2800697, AB_2819910, AB_2819911, AB_2819912, AB_2819913, AB_2819914
tCycif: anti-mouse B220 (CD45R)	ThermoFisher Scientific	Clone RA3-6B2; Cat# 41-0452-80; RRID: AB_2573598
tCycif: anti-mouse CD4	ThermoFisher Scientific	Clone 4SM95; Cat# 41-9766-82; RRID: AB_2573637
tCycif: anti-mouse CD8a	Cell Signaling Technology	Clone D4W2Z; Cat# 98941S; RRID: AB_2756376
tCycif: anti-mouse FOXP3	ThermoFisher Scientific	Clone FJK-16 s; Cat# 11-5773-82; RRID: AB_465243
tCycif: anti-human a.SMA	Abcam	Clone EPR5368; Cat# ab202509; RRID: AB_2868435
tCycif: anti-human CD4	R&D Systems	Cat# FAB8165G; RRID: AB_2728839
tCycif: anti-human CD8a	ThermoFisher Scientific	Clone AMC908; Cat# 50-0008-80; RRID: AB_2574148
tCycif: anti-human CD20	ThermoFisher Scientific	Clone L26; Cat# 50-0202-80; RRID: AB_11151691
tCycif: anti-human CD45	BioLegend	Clone HI30; Cat# 304008; RRID: AB_314396
tCycif: anti-human CCR6	Abcam	Clone EPR22259; Cat# ab243852; RRID: AB_2860033
tCycif: anti-human FOXP3	eBioscience	Clone 236A/E7; Cat# 41-4777-80; RRID: AB_2573608
tCycif: anti-human TCF1	Cell Signaling Technology	Clone C63D9; Cat# 6444S; RRID: AB_2797627

REAGENT or RESOURCE	SOURCE	IDENTIFIER
tCycif: anti-human TTF1 (also reacts with mouse)	Abcam	Clone EPR595(2); Cat# ab206726; RRID: AB_2857980
Purification: Ly6G Rat Monoclonal	BioLegend	Clone 1A8; Cat# 127602; RRID: AB_1089180
Purification: EpCam (CD326) Rat Monoclonal	BioLegend	Clone G8.8; Cat# 118202; RRID: AB_1089027
Purification: F4/80 Rat Monoclonal	BioLegend	Clone BM8; Cat# 123102; RRID: AB_893506
Biological samples		
Human formalin fixed paraffin embedded tissue samples from lung adenocarcinoma cases	Partners Healthcare Institutional Review Board at Brigham Health, Boston, MA, USA	Excess tissue, discarded tissue protocol number 2018P001627
Human formalin fixed paraffin embedded tissue samples from metastatic melanoma cases	Specialized Histopathology Services - Longwood Core Brigham and Womens Hospital, Boston, MA, USA	Project: 05-042 - T Jacks Lab slides
Chemicals, peptides, and recombinant proteins		
Collagenase IV	Worthington Biochemical	Cat# LS004189
DNase I	Sigma-Aldrich	Cat# 10104159001
Monensin	BioLegend	Cat# 420701
Golgi Plug	BD Biosciences	Cat# 555028
Phorbol-12-myristate-13-acetate (PMA)	EMD Millipore	Cat# 524400
Ionomycin Calcium Salt	Sigma	Cat# 13909
Streptavidin, allophycocyanin conjugate	Invitrogen	Cat# \$32362
Streptavidin, R-phycoerythrin conjugate	Invitrogen	Cat# S21388
Mirus TransIT LT1	Mirus Bio	Cat# MIR 2300
SYPRO Orange Protein Gel Stain	Thermo Fisher Scientific	Cat# \$6651
Cyclic-di-GMP	Invitrogen	Cat# tlrl-nacdg
SIINFEKL peptide	New England Peptide	Custom
SIYRYYGL peptide	New England Peptide	Custom
SIINYEKL peptide	New England Peptide	Custom
ITYTWTRL peptide (mALG8)	Genscript	Custom
VGFNFRTL peptide (mLAMA4)	Genscript	Custom
SMLVLLPDEVSGLEQLESIINYEKLTEWTS	New England Peptide	Custom
SMLVLLPDEVSGLEQLESIINFEKLTEWTS peptide	New England Peptide	Custom
GRCVGSEQLESIYRYYGLLLKERSEQKLIS peptide	New England Peptide	Custom
Empty peptide-receptive H-2K ^b molecules	Saini et al., 2019	Generated in-house
Zombie Fixable Viability Kit	BioLegend	Cat# 423102
Ghost Dye	Tonbo Biosciences	Cat# 13-0865-T500
Critical commercial assays		
Intracellular Fixation & Permeabilization Buffer Set Kit	ThermoFisher Scientific	Cat# 88-8824-00
Dynabead Sheep Anti-Rat IgG	ThermoFisher Scientific	Cat#11035
Chromium Next GEM Single Cell 5' Library and Gel Bead Kit	10X Genomics	Cat# 000167
Chromium Single Cell V(D) I Enrichment Kit Mouse T Cell	10X Genomics	Cat#1000071

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Easysep Dead Cell Removal (Annexin V) Kit	StemCell	Cat#17899
Deposited data		
Mouse KP LucOS SIIN versus SIY scRNA-seq	GEO	GEO: GSE164177
Sade-Feldman et al. scRNA-seq dataset	Sade-Feldman et al., 2019	GEO: GSE120575
Wu et al. scRNA-seq dataset	(Wu et al., 2020)	GEO: GSE139555
Guo et al. scRNA-seq dataset	Guo et al., 2018	GEO: GSE99254
Yost et al. scRNA-seq dataset	(Yost et al., 2019)	GEO: GSE123813
Tirosh et al. scRNA-seq dataset	Tirosh et al., 2016	GEO: GSE72056; Broad Data Use Oversight System: DUOS-000002
Mouse and human single cell RNA-sequencing analysis code	This Paper	GitHub: https://doi.org/10.5281/ zenodo.5110770
t-CYCIF analysis code	This Paper	GitHub: https://doi.org/10.5281/ zenodo.5104869
Experimental models: Cell lines		
GreenGo 3TZ for lentiviral titering	Sánchez-Rivera et al., 2014	Available upon request
293FS* viral packaging cell line	This paper	Available upon request
RMA-S TAP-deficient cell line	Schumacher et al., 1990	Available upon request
Experimental models: Organisms/strains		
Mouse: B6.129S4-Krastm4Tyj/J	Jackson Laboratories	Stock No: 008179
Mouse: B6.129P2- <i>Trp53</i> tm1Btn/J	Jackson Laboratories	Stock No: 008462
Mouse: II17atm1.1(icre)Stck/J	Jackson Laboratories	Stock No: 016879
Mouse: B6.Cg-Gt(ROSA)26Sortm14(CAG-tdTomato)Hze/J	Jackson Laboratories	Stock No: 007914
Recombinant DNA		
Plasmid: Lenti-LucOS	DuPage et al., 2011	Addgene Cat# 22777
Plasmid: Lenti-LucSIY	This Paper	Addgene Cat# 174044
Plasmid: Lenti-LucSIIN	This Paper	Addgene Cat# 174043
Plasmid: Lenti-LucY5S	This Paper	Addgene Cat# 174045
Plasmid: Lenti-LucY5	This Paper	Addgene Cat# 174046
Plasmid: Lenti-LucA	This paper	Addgene Cat# 174047
Plasmid: Lenti-LucL	This paper	Addgene Cat# 174048
Plasmid: Lenti-LucAL	This paper	Addgene Cat# 174049
Software and algorithms		
FlowJo	FlowJo	Version 10.7.1 https://www.flowjo.com/ solutions/flowjo/downloads
Aperio ImageScope	Leica Biosystems	Version 12 https:// www.leicabiosystems.com
ImageJ	NIH	https://imagej.nih.gov/ij/
ImageJ BaSiC Plugin	Peng et al., 2017	https://www.helmholtz- muenchen.de/icb/research/groups/marr- lab/software/basic/index.html
ASHLAR	The Python Package Index	https://pypi.org/project/ashlar/
ilastik	Berg et al., 2019	https://www.ilastik.org/download.html
MATLAB	MathWorks	version 2018a https://www.mathworks.com/ products/matlab.html

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REAGENT or RESOURCE	SOURCE	IDENTIFIER
R/Rstudio	Rstudio	version 1.2.5042 https:// www.rstudio.com/
Seurat	Butler et al., 2018	version 4.0.0 https://cran.r-project.org/web/ packages/Seurat/index.html
CellRanger	10X Genomics	version 3.1.0 https://support.10xgenomics.com/single- cell-gene-expression/software/overview/ welcome
ComplexHeatmap	CRAN Gu et al., 2016	version 2.4.3 https:// bioconductor.org/packages/release/bioc/ html/ComplexHeatmap.html
Tidyr	CRAN	version 1.1.2 https://tidyr.tidyverse.org, https://github.com/tidyverse/tidyr
Cowplot	CRAN	version 1.1.0 https://wilkelab.org/ cowplot/
GGplot2	CRAN	version 3.2.2 https:// ggplot2.tidyverse.org, https:// github.com/tidyverse/ggplot2
GridExtra	CRAN	version 2.3 https://cran.r-project.org/web/ packages/gridExtra/index.html
Plyr	CRAN	version 1.8.6 https://cran.r-project.org/web/ packages/plyr/index.html
Dplyr	CRAN	version 1.0.2 https://cran.r-project.org/web/ packages/dplyr/index.html
Monocle3	Trapnell et al., 2014	version 0.2.3.0 https://cole-trapnell- lab.github.io/monocle3/
Patchwork	CRAN	version 1.1.1 https://github.com/ thomasp85/patchwork
Viridis	CRAN	version 0.5.1 https://github.com/ sjmgarnier/viridis
Circlize	CRAN	Version: 0.4.11 https://github.com/ jokergoo/circlize
Stringr	CRAN	Version: 1.4.0 https:// stringr.tidyverse.org, https://github.com/ tidyverse/stringr
Philentropy	CRAN	Version: 0.4.0 https://github.com/ drostlab/philentropy
Dendextend	CRAN	Version: 1.14.0 https://cran.r-project.org/web/ packages/dendextend/index.html
FactoEXTRA	CRAN	Version: 1.0.7 http://www.sthda.com/ english/rpkgs/factoextra
ProjecTILs	Andreatta et al., 2021	v.0.5.1 https://github.com/carmonalab/ ProjecTILs
gplots	Warnes, 2015	v.3.1.0 https://cran.r-project.org/web/ packages/gplots/index.html
RColorBrewer	Neuwirth, 2014	v.1.1-2 https://cran.r-project.org/web/ packages/RColorBrewer/index.html