Targeting Cbx3/HP1γ Induces LEF-1 and IL-21R to Promote Tumor-Infiltrating CD8 T-Cell Persistence

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Abstract
Checkpoint blockade can reverse CD8+ T-cell functional exhaustion, and TCF-1 is essential for this process. However, identifying mechanisms that can prevent functional senescence and potentiate CD8+ T-cell persistence in checkpoint blockade non-responsive tumors remains a challenge. We demonstrate that targeting Cbx3/HP1γ causes augmented transcription initiation, chromatin remodeling at Lef1 and Il21r leading to increased transcriptional activity at these loci. Mechanistic studies show LEF-1 and IL-21R are required for Cbx3/HP1γ-deficient CD8+ effector T cells to persist resulting in improved control of ovarian cancer, melanoma and neuroblastoma in preclinical models. Cbx3/HP1γ-deficient CD8+ T cells enhanced persistence in the TME facilitates remodeling of the chemokine/receptor landscape that ensures their optimal tumor invasion at the expense of CD4+ Tregs. Thus, CD8+ T cells heightened effector function consequent to Cbx3/HP1γ deficiency may be distinct from functional reactivation by checkpoint blockade, implicating Cbx3/HP1γ as a viable cancer T-cell-based therapy target for resistant, non-responsive solid tumors.

Keywords
Cbx3/HP1γ, LEF-1, IL-21R or IL-21 receptor, tumor-infiltrating CD8+ T-cell persistence, adoptive T-cell therapy (ACT), effector memory/progenitor memory CD8+ T cells, ovarian cancer, neuroblastoma, melanoma.

Introduction
Under persistent antigen exposure as in cancer or chronic infections, CD8+ effector T cells can enter an altered differentiation program known as T-cell exhaustion (T_ex)1. In the tumor microenvironment (TME), CD8+ progenitor T_ex cells express unique transcription factors as well as those shared with effector and memory cells including the high mobility group (HMG) transcription factor, T-cell factor 1 (TCF-1, encoded by Tcf7). They have reduced effector capacity but can be reinvigorated by check point blockade to proliferate and differentiate into terminally exhausted cells having reduced proliferative capacity yet can efficiently kill tumor cells through perforin 1 (PRF1), granzyme B (GrB) and interferon γ (INFγ)2, 3, 4. TCF-1 is implicated in the conversion of CD8+ progenitor T_ex to terminally differentiated T_ex5, 6, 7, 8, 9. However, given that only ~20% of all solid tumors and primarily those harboring high mutation loads respond to checkpoint blockade, in addition to the emergence of checkpoint blockade resistance, there is a
need to identify novel targets that can restore the effector and persistence capacity of exhausted CD8+ T cells in tumors harboring low mutation loads and do not respond to checkpoint blockade. In humans and mice, TCF-1 and the related lymphoid enhancer-binding factor 1 (LEF-1) are functionally linked during T-cell development in the thymus and the generation of precursor and functional memory CD8+ as well as CD4+ T cells\textsuperscript{10, 11, 12, 13, 14, 15, 16, 17, 18}. Nevertheless, unlike TCF-1, a role for LEF-1 in tumor response has not been documented.

IL-21R is widely expressed on various innate and adaptive immune cell-lineages including activated CD8+ T, CD4+ T\textsubscript{FH} and NK cells. During the course of a chronic viral infection or under IL-2-deprived conditions IL-21R signaling is critical for preventing CD8+ T-cell exhaustion\textsuperscript{19, 20}. In acute viral infections, IL-21R signaling is essential for the proliferation and survival of activated CD8+ T cells as well as the generation of long-lived memory cells\textsuperscript{21, 22, 23}. However, the function of IL-21R signaling in cancer is controversial and not completely understood\textsuperscript{24, 25, 26, 27, 28}.

Here, we demonstrate that targeting \textit{Cbx3/HP1\gamma} promotes increased/sustained expression of LEF-1 and IL-21R in CD8+ effector T cells. \textit{Cbx3/HP1\gamma} deficiency leads to augmented transcription initiation and chromatin remodeling at \textit{Lef1} and \textit{Il21r} resulting in increased transcriptional activity at these loci. Genetic ablation of \textit{Lef1} and \textit{Il21r} in \textit{Cbx3/HP1\gamma}-deficient mice causes a loss of CD8+ effector T cells accompanied by the reduction of \textit{Ifng, Gzmb} and \textit{Prf1} expression leading to uncontrolled growth of ovarian, melanoma and neuroblastoma tumors. By contrast, \textit{Tcf7} deletion does not affect the effector function of \textit{Cbx3/HP1\gamma}-deficient CD8+ T cells. Our data establish that LEF-1 and IL-21R are required for \textit{Cbx3/HP1\gamma}-deficient CD8+ T cells to expand, persist and maintain their effector capacity in tumors whereas TCF-1 is insufficient. Our findings together with those of others\textsuperscript{29, 30, 31} illustrate the complex mechanisms governing CD8+ T-cell effector differentiation and function within a given TME. They underscore the need for continuing exploration of novel targets that can reverse CD8+ T-cell dysfunction in tumors.

\textbf{Results}

\textit{Cbx3/HP1\gamma} deficiency modulates LEF-1 and IL-21R expression in CD8+ effector T cells. Previously we showed that germline deletion of the histone code reader \textit{Cbx3/HP1\gamma} impairs lymphoid-tissue germinal center (GC) reaction and high-affinity antibody response against a thymus (T)-dependent antigen (Ag) in a CD8+ T-cell-intrinsic manner\textsuperscript{32}. \textit{Cbx3/HP1\gamma} deficiency releases the effector capacity of CD8+ T cells to control tumor growth\textsuperscript{33}. However, it is not understood why \textit{Cbx3/HP1\gamma}-deficient CD8+ effector T cells are not subjected to functional
senescence and can persist in tumors. To ensure that Cbx3/HP1γ deletion was restricted to CD8⁺ T cells, all experiments were performed using the CD8α-Cre strain [34] crossed with our Cbx3/HP1γ-floxed mice (resulting animals were Cbx3/HP1γfl/fl and Cbx3/HP1γfl/+, collectively designated as Cbx3/HP1γ-deficient). Cre recombinase from this CD8α-Cre strain was active in CD8⁺CD4⁻ T cells not in CD4⁺CD8⁻ or CD11b⁺ cells (Extended Data Fig. 1a). Conditional ablation of Cbx3/HP1γ in CD8⁺ T cells resulted in a near or complete loss of protein expression and phosphorylation (Extended Data Fig. 1b). To determine mechanisms conferring persistence on Cbx3/HP1γ-deficient CD8⁺ effector T cells, ChIP-Seq data was analyzed (Extended Data Table 1) [33]. In wild-type (wt) differentiated CD8⁺ T cells, Cbx3/HP1γ was bound to the 5' untranslated region (UTR) surrounding transcriptional start sites (TSSs) of Lef1 (Fig. 1a,b) and Il21r (Fig. 1d,e). Western immunoblots showed an induction of LEF-1 expression in differentiated control CD8⁺ T cells; however, LEF-1 expression was further elevated in Cbx3/HP1γ-deficient CD8⁺ effector T cells (Fig. 1c). The level of LEF-1 detected in control and deficient thymocytes was similar (Extended Data Fig. 1c). In the absence of exogenous IL-21, differentiated Cbx3/HP1γ-deficient CD8⁺ T cells expressed more IL-21R protein and transcript compared to control cells (Fig. 1f-h). There was a steady expansion of Cbx3/HP1γ-deficient CD8⁺IL-21R⁺ T cells after differentiation while the growth of control cells remained low throughout the differentiation period (Fig. 1f,i). IL-21R was not detected on thymocytes from control or Cbx3/HP1γ-deficient mice (Extended Data Fig. 1d). In vitro-differentiated Cbx3/HP1γ-deficient CD8⁺ T cells expressed more Bcl6 and Tbx21 (T-bet) mRNAs compared to control cells (Extended Data Fig. 1e) [35]. By contrast Eomes transcript and protein levels were reduced (Extended Data Fig. 1e,f) [36]. The effector capacity of Cbx3/HP1γ-deficient CD8⁺ T cells was enhanced as evidenced by an upregulation of Prf1, Gzmb and Ifng mRNA expression (Extended Data Fig. 1g). As a result, more cleaved caspase 3 (CC3) was detected in B16 melanoma tumor cells co-cultured with Cbx3/HP1γ-deficient CD8⁺ effector T cells compared to control co-cultures (Extended Data Fig. 1h). Cbx3/HP1γ deficiency in CD4⁺ T cells did not enhance tumor killing. Our results establish that Cbx3/HP1γ deficiency induces the sustained increase of LEF-1 and IL-21R in CD8⁺ effector T cells, which are endowed with an enhanced effector capacity. IL-21R elevated expression likely provides signals mediating Cbx3/HP1γ-deficient CD8⁺ effector T-cell expansion in the absence of exogenous IL-21.
**Cbx3/HP1γ regulates Lef1 and Il21r transcription initiation and chromatin remodeling.** The rate of gene expression is governed in part by RNA Pol II initiation, elongation and/or chromatin remodeling. Pol II is phosphorylated at serine 5 (Pol II S5) during initiation of transcription while phosphorylation at serine 2 (Pol II S2) generally indicates chromatin remodeling concomitant with transcriptional elongation. To test whether alterations in transcriptional initiation, elongation and/or chromatin remodeling caused increased LEF-1 and IL-21R expression in Cbx3/HP1γ-deficient CD8+ effector T cells, ChIP-qPCR experiments were performed using ChIP-tested antibodies specific for Pol II S2 or S5, and H3K9me3. These results revealed augmented levels of Pol II S5 in or around transcription start sites (TSSs) of Lef1 and Il21r loci in Cbx3/HP1γ-deficient CD8+ effector T cells compared to control cells (Fig. 2a,b). Similar levels of Pol II S2 density were observed at both loci in control and deficient CD8+ T cells (Fig. 2c,d). The levels of H3K9me3 deposition at Lef1 locus was unaltered (Fig. 2e). However, there was a general loss of H3K9me3 around the TSS of Il21r locus in Cbx3/HP1γ-deficient CD8+ effector T cells (Fig. 2f). Thus, genetic deletion of Cbx3/HP1γ in CD8+ effector T cells results in enhanced transcription initiation at Lef1 and Il21r loci, with chromatin remodeling activity taking place in an extended region around the TSS of the Il21r locus. These changes likely underpinned the increased and sustained transcriptional activity of Lef1 and Il21r.

**Cbx3/HP1γ-deficient CD8+ effector T cells can persist and cause tumor rejection.** The increased/sustained expression of LEF-1 and IL-21R together with the enhanced effector capacity exhibited by Cbx3/HP1γ-deficient CD8+ effector T cells suggest they can persist to control tumor development. Thus, the ability of Cbx3/HP1γ-deficient CD8+ T cells to eradicate solid tumors was evaluated using the mouse ID8 or MOSE-L-TICv ovarian, B16 melanoma and NB-9464 neuroblastoma (NBL) tumor models. Ovarian and NBL tumors have low mutation rates, no clear defining tumor-associated antigens (TAAs) and are minimally responsive to checkpoint blockade. Accordingly, syngeneic ID8 or MOSE-L-TICv ovarian tumor cells were injected intraperitoneally (IP) into control and Cbx3/HP1γ-deficient mice. Mice were monitored until abdominal distension was visible, which indicated increased ascites and mirrored metastasis observed in humans. Tumor growth and production of ascites were inhibited in Cbx3/HP1γ-deficient mice compared to controls or mice ectopically expressing Cbx3/HP1γ driven by the human T-cell-restricted Cd2 promoter (Cbx3/HP1γTg) (Fig. 3a,b and Extended Data Fig. 2a,b). As
a result, Cbx3/HP1γ-deficient mice lived longer than controls (Fig. 3c). Cbx3/HP1γ-deficient mice were equally effective in reducing B16 melanoma (Fig. 3d) and NB-9464 NBL tumor burden (Fig. 3f), leading to their increased survival rate compared to controls (Fig. 3e,g). On day 120 after ID8 injection, there was an enrichment of CD8+NKG2D+ effector T cells and a decrease in CD4+CD25+FOXP3+ regulatory T cells (Tregs) in ascites from Cbx3/HP1γ-deficient mice compared to control or Cbx3/HP1γTg animals (Fig. 3h,i and Extended Data Fig. 2c,d). Similarly, enrichment of CD8+NKG2D+ effector T cells and decrease in CD4+CD25+FOXP3+ Tregs were observed in B16 melanoma as well as NBL tumors (Fig. 3j-m and Extended Data Fig. 2e-h). To show that Cbx3/HP1γ-deficient CD8+ effector T cells alone can persist and cause tumor rejection, CD8+ T cells (CD45.2+) were differentiated in vitro. On day 5 after differentiation, CD8+ T cells were collected for adoptive T-cell therapy in tumor bearing congenic B6.SJL (CD45.1+) recipients (Extended Data Fig. 3a). Treatment with Cbx3/HP1γ-deficient CD8+ effector T cells alone resulted in a statistically significant decrease in B16 melanoma and NBL tumor burden compared to treatment with control cells (Extended Data Fig. 3b,c). Within B16 melanoma tumors, there was an exclusive enrichment of transferred Cbx3/HP1γ-deficient CD8+NKG2D+ effector T cells, not endogenous nor control Cbx3/HP1γ-sufficient CD8+ effector T cells (Extended Data Fig. 3d). Contaminating donor CD4+NKG2D+ and NK1.1+NKG2D+ T cells were not detected in tumors or spleens indicating that they were not the source of tumor killing (Extended Data Fig. 3e-h). Inhibition of Cbx3/HP1γ-deficient CD8+ effector T cells engagement with tumor cells using a blocking/non-depleting anti-NKG2D antibody resulted in uncontrolled tumor growth and decreased survival of treated animals (Extended Data Fig. 4a-d), accompanied by the reduction of CD8+NKG2D+ effector T cells in tumors, to a level similar to that of untreated control mice (Extended Data Fig. 4e,f). NKG2D blockade did not affect CD4+NKG2D+ and NK1.1+NKG2D+ T-cell frequencies (Extended Data Fig. 4g,h). Comparable frequencies of NK1.1+NKG2D+ or CD4+NKG2D+ T cells was recovered from tumors of control, Cbx3/HP1γTg and Cbx3/HP1γ-deficient mice (Extended Data Fig. 5). Expression of Ifng, Gzmb and Prf1 was elevated in B16 melanoma and NBL tumors excised from Cbx3/HP1γ-deficient mice compared to controls or Cbx3/HP1γTg animals (Extended Data Fig. 6a-f). TUNEL staining revealed more apoptotic tumor cells (brown) in melanoma and NBL tumors from Cbx3/HP1γ-deficient mice compared to controls (Fig. 3n,o). Genetic ablation of Prf1 in Cbx3/HP1γ-deficient animals led to uncontrolled B16
melanoma growth and decreased tumor-cell apoptosis while Ifng deletion resulted in unchecked NBL tumor development in Cbx3/HP1γ-deficient mice (Extended Data Fig. 6g-i). Together, our results demonstrate that Cbx3/HP1γ-deficient CD8+ effector T cells expressing NKG2D can persist and cause tumor rejection, irrespective of tumor mutation status. The engagement of NKG2D with its ligands expressed on tumor cells likely facilitates the tumor killing activity of Cbx3/HP1γ-deficient CD8+ effector T cells through the enhanced production of PRF1, GrB and INF-γ.

**Cbx3/HP1γ-deficient CD8+ T cells remodel the tumor chemokine/receptor landscape.** CD8+ effector T cells homing to tumors is mediated primarily by the interaction between CCL2/CCR2 and CXCL9/CXCL10/CXCR3 chemokine/receptor pairs while CD4+ Tregs trafficking to tumors is induced mostly by CCL28/CCR10 and CXCL12/CXCR440, 41, 42, 43, 44. Chemokines are generally produced by tumor-associated myeloid cells including dendritic cells (DCs) and macrophages (MØs) whereas their cognate receptors are expressed on T cells as well as other immune cells. It is proposed that changes in the tumor chemokine/receptor landscape influence anti-tumor responses; however, precise mechanisms through which this occurs has not been clearly established. RT-qPCR assays demonstrated that Ccl2/Ccr2 and Cxcl9/Cxcl10/Cxcr3 levels were higher in B16 melanoma tumors from Cbx3/HP1γ-deficient mice than those from controls (Fig. 4a,b). In NBL tumors from Cbx3/HP1γ-deficient mice, Ccl2/Ccr2 and Cxcl10 remained similar to those of controls whereas Cxcl9/Cxcr3 levels were elevated (Fig. 4c,d). Compared to tumors from controls, Ccr10 and Cxcl12/Cxcr4 levels were decreased in B16 melanoma tumors from Cbx3/HP1γ-deficient mice (Fig. 4e,f). In NBL tumors from Cbx3/HP1γ-deficient mice, the level of Ccl28/Ccr10 and Cxcl12 expression was reduced (Fig. 4g,h). These observations indicate that homing of Cbx3/HP1γ-deficient CD8+ effector T cells into B16 melanoma tumors requires CCL2 and CXCL9/CXCL10 present in the TME and their receptors CCR2 and CXCR3 expressed on CD8+ T cells. However, Cbx3/HP1γ-deficient CD8+ effector T cells likely depend on CXCL9/CXCR3 interaction to invade NBL tumors. Changes in the levels of CD4+ Tregs chemokines and receptors suggest that CCR10 and CXCL12/CXCR4 regulate homing of CD4+ Tregs into B16 melanoma tumors while invasion of NBL tumors by CD4+ Tregs is governed by CCL28/CCR10 and CXCL12. In short, the presence of Cbx3/HP1γ-deficient CD8+ effector T cells in the TME mediates remodeling of the chemokine/receptor landscape that favors their optimal trafficking into tumors at the expense of CD4+ Tregs.
**LEF-1 and IL-21R are indispensable for halting tumor growth.** To establish that LEF-1 and IL-21R contribute to the control of tumor growth by Chx3/HP1γ-deficient CD8+ effector T cells, compound mutant mice were created (Extended Data Fig. 7a). In the Chx3-Lef1-deficient mouse, Lef1 and Chx3/HP1γ deletion was restricted to CD8+ T cells; in the Chx3-Il21r-deficient mouse, Chx3/HP1γ ablation was restricted to CD8+ T cells while Il21r was deleted in all tissues. LEF-1 and IL-21R expression was abolished in deficient CD8+ T cells compared to control cells (Fig. 5a,b). Loss of one Lef1 or Il21r allele showed a decrease in protein levels that were almost completely abrogated upon deletion of both alleles. Progenitor T-cell development proceeded normally in the thymus of compound mutant animals (Extended Data Fig. 7b,c). Mesenteric lymph nodes (mLNs) of compound mutant mice displayed normal frequencies of mature CD8+ and CD4+ T cells (Extended Data Fig. 7d,e). Normal ratios of naive (CD44+CD62L+), effector (CD44+CD62L−) and memory (CD44+CD62L+) T cells were observed in the mLNs of compound mutant animals (Extended Data Fig. 7f,g). In the mLNs, IL-21R expression was restricted to the CD8+ effector (CD44+CD62L−) T-cell population and was also dependent on gene dosage (Extended Data Fig. 7h). IL-21R was not detected on any CD4+ T-cell populations (Extended Data Fig. 7i). To determine LEF-1 contribution to tumor control, ID8 (ovarian), B16 (melanoma) or NB-9464 (NBL) tumor cells were injected into Chx3-Lef1-deficient and control mice. Lef1 genetic ablation resulted in unrestrained ovarian, melanoma and NBL tumor growth (Fig. 5c,d,f). Tumor burden was higher than that observed for Chx3/HP1γ-deficient mice, but similar to controls. Because Il21r was deleted in all tissues, adoptive transfer of differentiated CD8+ T cells from Chx3-Il21r-deficient and control mice was performed. Melanoma and NBL growth proceeded unchecked in tumor-bearing B6.SJL (CD45.1+) congenic mice treated with control or Chx3-Il21r-deficient CD8+ effector T cells (CD45.2+) compared to animals receiving Chx3/HP1γ-deficient CD8+ effector T cells (CD45.2+) (Fig. 5e,g). Our results showing that Chx3-Lef1-deficient mice cannot control tumor growth indicates that TCF-1, which is intact in our models, may not be sufficient in restraining tumor development. To assess TCF-1 role in tumor control in our model, we created Chx3-Tcf7-deficient mice in which both genes were conditionally abrogated in CD8+ T cells (Extended Data Fig. 8a). Unlike Lef1 and Il21r, deletion of both Tcf7 alleles was necessary to achieve near complete abrogation of TCF-1 expression (Fig. 5h). Growth of all three tumors was halted in Chx3-Tcf7- and Chx3/HP1γ-deficient mice compared to controls (Fig. 5i-k), Chx3-Lef1-deficient mice or those receiving Chx3-Il21r-deficient CD8+ effector T cells (Fig. 5c-g). Our
data establish that LEF-1 and IL-21R are required for the optimal control of ovarian, melanoma and NBL tumor growth consequent to Cbx3/HP1γ deficiency while TCF-1 is insufficient. Moreover, LEF-1 expression in control CD8+ effector T cells is induced (Fig. 1c), yet they are incapable of persisting in tumors, suggesting there may be a threshold of expression level required for LEF-1 to function effectively. IL-21R expression is confined to the CD8+ effector population and its loss following deletion suggest that IL-21R function is essential to this population, which is critical for halting tumor development.

**LEF-1 and IL-21R are required for Cbx3/HP1γ-deficient CD8+ effector T cells to persist in tumors.** The inability of Cbx3-Left1- and Cbx3-II21r-deficient mice to halt tumor development suggests that LEF-1 and IL-21R are required for the persistence of CD8+ effector T cells in tumors. Analyses of tumors from Cbx3-Left1-deficient mice and those receiving Cbx3-II21r-deficient CD8+ T cells (hereafter simplified as Cbx3-II21r-deficient) revealed that the frequency of CD8+ effector T cells was reduced to control levels in all tumors (Fig. 6a-e). By contrast, tumors from Cbx3-Tcf7- and Cbx3/HP1γ-deficient animals exhibited similar CD8+ effector T-cell frequencies that were elevated over control mice (Fig. 6f-h). The proportion of CD8+ T cells exhibiting progenitor (CD8+TIM3+CXCR5+) or terminal exhausted (CD8+TIM3+CXCR5+) phenotype was not perturbed (Extended Data Fig. 8b). The frequency of NK1.1+NKG2D+ and CD4+NKG2D+ T cells in all tumor types was low and similar to that of controls (Extended Data Fig. 8c). Myeloid and B-cell populations were not altered. Our data underscore the fact that LEF-1 and IL-21R are required for the persistence of Cbx3/HP1γ-deficient CD8+ effector T cells in tumors with varied mutation loads and are checkpoint blockade responsive (B16 melanoma) or non-responsive (ovarian and NBL). In the absence of checkpoint blockade, TCF-1 is not sufficient in conferring persistence on CD8+ effector T cells.

**LEF-1 and IL-21R are required to maintain effector activity in tumors.** To confirm that the loss of CD8+ effector T cells will lead to diminished effector activity in tumors from Cbx3-Left1- and Cbx3-II21r-deficient mice, RT-qPCR assays were done using tumor RNA samples from mutant and control mice. In melanoma tumors from Cbx3-Left1- and Cbx3-II21r-deficient mice, Prf1 and Gzmβ transcript levels were reduced to controls, which were lower than those in tumors from Cbx3/HP1γ-deficient mice (Fig. 7a-d). Prf1, Gzmβ and Ifng expression in NBL tumors from Cbx3-Left1- and Cbx3-II21r-deficient mice was diminished compared to Cbx3/HP1γ-deficient mice but similar to that of controls (Fig. 7e-j). By contrast, Prf1, Gzmβ or Ifng transcripts were readily
detected in B16 and NBL tumors from Cbx3/HP1γ- and Cbx3-Tcf7-deficient mice (Fig. 7a,b and e-g). Thus, our findings show that LEF-1 and IL-21R, not TCF-1, are indispensable for maintaining Prf1, Gzmb and Ifng presence in tumors. The loss of CD8+ effector T cells likely contributes to the observed blunted effector activity observed in tumors from Cbx3-Lef1- and Cbx3-II21r-deficient mice that in turn results in unchecked tumor growth.

Discussion
We establish that depletion of Cbx3/HP1γ in CD8+ T cells simultaneously induces the elevated, sustained expression of factors conferring both persistence and heightened effector capacity on Cbx3/HP1γ-deficient CD8+ T cells that in turn enables them to control the growth of diverse tumor types irrespective of mutation loads. Once in the TME, Cbx3/HP1γ-deficient CD8+ T cells can remodel the chemokine/receptor landscape that favors their optimal trafficking into tumors at the expense of CD4+ Tregs thus eliminating immune suppression. Cbx3/HP1γ deficiency allows for a higher rate of transcription initiation and chromatin remodeling at Lef1 and Il21r loci. Consequently, Cbx3/HP1γ-deficient CD8+ T cells express elevated levels of LEF-1 and IL-21R that halt functional senescence and enable their persistence in tumors without the assistance of checkpoint blockade. By contrast, Tcf7 deletion does not affect the effector function of Cbx3/HP1γ-deficient CD8+ T cells. Thus, CD8+ T cells heightened effector function consequent to Cbx3/HP1γ deficiency may be distinct to functional reactivation by checkpoint blockade, implicating Cbx3/HP1γ as a viable cancer T-cell-based therapy target for resistant, non-responsive solid tumors.

To date, no essential functions for LEF-1 have been identified in CD8+ effector T cells despite Lef1 expression being detected in human and mouse stem-like tumor infiltrating lymphocyte (TIL) subsets that also express Tcf7. Here, we present compelling evidence that LEF-1 has a critical and non-redundant function in tumor rejection. The exclusive increase of Cbx3/HP1γ-deficient CD8+ effector T cells in tumors and their subsequent disappearance when Lef1 is ablated suggest that LEF-1 is required for the persistence of Cbx3/HP1γ-deficient CD8+ effector T cells in varied tumor types. Our results showing tumor growth is successfully inhibited in Cbx3-Tcf7-deficient mice indicates that TCF-1 function may be insufficient. Our data are distinct to those found with checkpoint blockade where TCF-1 activity is implicated in the rejuvenation of tumor CD8+ TEx cells. Instead, our findings are consistent with those reported by Carr et. al. wherein
they show LEF-1 playing primary roles in the post selection and effector fate differentiation of iNKT2 cells in the thymus. Similarly, Reya and colleagues have shown that LEF-1 is required for progenitor B-cell proliferation and survival through Wnt signaling. In both models, LEF-1 exhibits a more dominant role whereas TCF-1 is not sufficient for the differentiation and expansion of the immune-cell populations examined. Thus, there may be specific conditions wherein LEF-1 function is dominant and sufficient for the development and differentiation of various immune cells while that of TCF-1 is not required.

Our observations that IL-21R+CD8+ effector T cells increase in number after differentiation and are lost upon Il21r deletion indicate that IL-21R provides survival as well as expansion signals to these cells. This is in keeping with work showing that, in humans and mice, IL-21R has an essential role in regulating viral infections and tumor immunity. During the course of a chronic viral infection or under IL-2-deprived conditions, IL-21R signaling is critical for preventing CD8+ T-cell exhaustion. In acute viral infections, IL-21R signaling is essential for the proliferation and survival of activated CD8+ T cells as well as the generation of long-lived memory cells. In these models, IL-21R can activate the STAT1/STAT3 signaling pathways, which subsequently upregulate pro-survival factors BCL-2 and BCL-XL and downregulate TRAIL. Moreover, several lines of evidence indicate that IL-21R is involved in tumor immunity. First, high expression of IL-21R on CD8+ T cells in tumors correlates positively with overall survival and lack of tumor recurrence in hepatocellular carcinoma (HCC) patients. Second, in mice, IL-21R signaling reduces accumulation of myeloid derived suppressor cells (MDSCs) in the TME to control rapid HCC growth and maintains an immunological memory response to tumor re-challenge. Third, newly diagnosed HER2+ breast cancer patients with higher Il21r expression may have a reduced risk of distant relapse when treated with trastuzumab (anti-HER2/ErB2 mAb) in combination with chemotherapy; IL-21R expression on CD8+ effector T cells, not NK cells, is required for optimal anti-ErB2 mAb efficacy. Our results underscore IL-21R critical function in providing signals that induce the expansion of CD8+ effector T cells to promote their anti-tumor capacity.

Our results suggest a model whereby Cbx3/HP1γ normally restrains the effector and persistence potential of CD8+ T cells, which will eventually succumb to functional inactivation and apoptosis. Removal of the Cbx3/HP1γ restraint allows CD8+ T cells to differentiate into long-lived effector cells armed with an intrinsic, heightened killing and persistence capacity to control tumor growth; LEF-1 and IL-21R are required while TCF-1 is insufficient. Our model does not preclude the
possibility that LEF-1 and IL-21R may also be implicated in CD8+ T-cell responses downstream of checkpoint blockade. Since mouse and human Chx3/HP1γ peptides are 100% identical, we suspect that human Chx3/HP1γ will behave identically to the mouse ortholog: Chx3/HP1γ deficiency would have the same effects on human CD8+ T cells. Furthermore, ovarian and neuroblastoma are tumors with low mutation rates that do not respond to checkpoint blockade, yet their growth is effectively controlled by Chx3/HP1γ-deficient CD8+ effector T cells. Thus, our findings provide a rationale for targeting Chx3/HP1γ in human T cells to treat tumors harboring low mutation loads and do not respond to checkpoint inhibitors.

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Author Contributions
T.H.T. N.H. and P.T.L. conceived the project and wrote the original draft with input and guidance from K.M.E., J.L.D., E.M.S., A.B., H.H.X., P.B.S., T.H.T., P.T.L., N.H. and N.K.T. designed and performed experiments. P.T.L., N.H., A.B., H.H.X., P.B.S. and T.H.T. analyzed data and wrote the final manuscript.

Declaration of Interests
The authors declare no competing interests.

Figure Legends
Figure 1. Chx3/HP1γ deficiency modulates LEF-1 and IL-21R expression in CD8+ effector T cells. a, Schematic of mouse Lef1 locus. -2 and 12 indicated positions of first and last primer pairs, respectively. b, Read-density tracks of Chx3/HP1γ peaks across Lef1 identified by ChIP-Seq using chromatin from wt day 5 differentiated CD8+ T cells (as in c, pooled from 3 mice); y axis: number of reads per million mapped per 25-bp window. c, LEF-1 expression was assessed by Western
immunoblot using total lysates from differentiated control and Cbx3/HP1γ-deficient CD8+ T cells; 3/28+IL2: CD8+ T cells differentiated for 5 days with plate-bound anti-CD3 and anti-CD28 plus IL-2; C: control (CD8α-Cre or wt); fl/+: Cbx3fl/fl; fl/fl: Cbx3fl/fl; CD8+ T-cell-restricted deletion of Cbx3/HP1γ using the CD8α-Cre mouse. d, Schematic of mouse Il21r locus. -4 and 4: positions of first and last pair of primers, respectively. e, Read-density tracks of Cbx3/HP1γ peaks across Il21r identified by ChIP-Seq as in (b). f, IL-21R expression on differentiated CD8+ T cells was determined by flow cytometry; left panels depicted percent CD8+IL-21R+ cells from each culture; right panels indicated IL-21R levels on the same cells; numbers: percent cells; representative of 3 experiments; n = 3 for each genotype. g, Graphs represented IL-21R mean fluorescence intensity (MFI) from (f). h, Relative expression of Il21r, normalized to Gapdh, in day 5 differentiated CD8+ T cells was evaluated by RT-qPCR using RNA from cells obtained in (f); Graphpad unpaired student t-test: **p≤0.01; representative of 3 experiments. i, Number of CD8+IL-21R+ T cells in cultures were calculated from (f); Graphpad unpaired student t-test: *p≤0.05, **p≤0.01.

**Figure 2.** Cbx3/HP1γ regulates Lef1 and Il21r transcription initiation and chromatin remodeling. a,b, Levels of Pol II S5 bound to Lef1 (a) and Il21r (b) were quantified by ChIP-qPCR using chromatin from day 5 differentiated CD8+ T cells. TSS: transcription start site, numbers on X axis: positions of primers along the two loci; 150 bp products amplified by Lef1 or Il21r primers. c,d, Levels of Pol II S2 bound to Lef1 (c) and Il21r (d) were quantified as in (a,b). e,f, H3K9me3 deposition on Lef1 (e) and Il21r (f) was quantified as in (a,b). Statistics: Graphpad unpaired student t-test: *p≤0.05, **p≤0.01, ***p≤0.001, ****p≤0.0001; representative of 4 independent ChIPs using pooled chromatin from 3 mice.

**Figure 3.** Cbx3/HP1γ-deficient CD8+ effector T cells can persist and cause tumor rejection. a, Ascites from control (CD8α-Cre or wt), Cbx3Tg, Cbx3fl/+ and Cbx3fl/fl mice (X axis) were visualized on day 120 after intraperitoneal injection of mouse ID8 ovarian tumor cells; Cbx3Tg: Cbx3/HP1γ T-cell-restricted expression driven by the human Cd2 promoter; Cbx3fl/+ and Cbx3fl/fl: CD8+ T-cell-restricted deletion of Cbx3/HP1γ using the CD8α-Cre mouse. b, Ovarian ascites (vol: volume) of mice in (a) were measured and graphed; bars: group median; Graphpad unpaired student t-test: **p≤0.01, ***p≤0.001; each symbol = one mouse; n = 5-8. c, Survival curves of ovarian tumor-bearing mice were determined; Graphpad log-rank (Mantel-Cox) test; n = 5-8. d, B16 tumor cells were injected subcutaneously (sc) and growth was assessed starting on day 14
after tumor-cell injection then every 2 days through day 20; Graphpad two-way ANOVA: **p ≤ 0.01, ****p ≤ 0.0001; n = 5. e, Survival curves of B16 melanoma tumor-bearing mice.; Graphpad log-rank (Mantel-Cox) test; n = 5. f, NB-9464 tumor cells were injected sc, NBL tumor growth was determined every 2 days starting on day 22 through day 28; Graphpad two-way ANOVA: ****p ≤ 0.0001; n = 5. g, Survival curves of NBL tumor-bearing mice; Graphpad log-rank (Mantel-Cox) test; n = 5. h, Frequencies of CD8+ NKG2D+ T cells in ovarian ascites, B16 and NBL tumors from control (CD8α-Cre or wt), Cbx3Tg, Cbx3fl/+ and Cbx3fl/fl mice; data were extracted from flow analysis (Extended Data Fig. 2c,e,g); bars: group median; Graphpad unpaired student t-test: **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001; each symbol = one mouse; n = 5-8. i, j, Frequencies of CD4+ FOXP3+ Tregs in ovarian ascites, B16 and NBL tumors (Extended Data Fig. 2d,f,h); bars: group median; Graphpad unpaired student t-test: *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, ****p ≤ 0.0001; each symbol = one mouse; n = 5-8 for each genotype. n, o, Apoptotic cells (brown) in B16 melanoma and NBL tumor sections were identified by TUNEL staining; representative of 4 tumors from each mouse strain.

**Figure 4.** Cbx3/HP1γ-deficient CD8+ T cells remodel the tumor chemokine/receptor landscape. a,b, Relative expression of chemokines and receptors mediating CD8+ T cells trafficking into B16 tumors were measured by RT-qPCR using RNA of day 5 differentiated CD8+ T cells from control and Cbx3/HP1γ-deficient CD8+ T cells as in figure 1c, results were normalized to Gapdh. c,d, Relative expression of chemokines and receptors mediating CD8+ T cells trafficking into NBL tumors. e,f, Relative expression of chemokines and receptors inducing CD4+ Tregs homing into B16 tumors normalized to Gapdh. g,h, Relative expression of chemokines and receptors inducing CD4+ Tregs homing into NBL tumors. X axis: mouse strains; Graphpad unpaired student t-test: *p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001; ns: not significant; n = 4 tumors from each mouse strain.

**Figure 5.** LEF-1 and IL-21R are indispensable for halting tumor growth. a, Western blot of LEF-1 expression was done using total lysates of day 5 differentiated CD8+ T cells; C: Cbx3/HP1γ-deficient; Lef1fl/fl and Lef1fl/fl: Cbx3-Lef1-deficient. b, IL-21R expression was evaluated by flow; Cbx3fl/fl: Cbx3/HP1γ-deficient, Il21r+/- and Il21r-/-: Cbx3-Il21r-deficient. Ovarian ascites were measured on day 125 after tumor injection; bars: group median; Graphpad unpaired student t-test: **p ≤ 0.01, ****p ≤ 0.0001; each symbol = one mouse; n = 3-5. Cbx3fl/flLef1fl/fl: Cbx3-Lef1-.
Ifng deficient tumors tumor RNA from indicated mice. Figure 7. 5 student t-test; *p≤0.0001; each symbol = mouse; for each genotype n = 4-7. e, On day 10 after injection of tumor cells, tumor bearing B6SJ/L mice were treated with in vitro-differentiated CD8+ T cells from control, Cbx3fl/fl or Cbx3fl/fl,Il21rΔ/Δ (Cbx3-Il21r-deficient); ***p≤0.001, n = 3-4 recipients. f, NBL tumor burden was determined starting on day 25 after injection of tumor cells then every 2 days until day 33; Graphpad two-way ANOVA: ***p≤0.001; n = 2-5. g, On day 14 after NBL injection, tumor bearing B6SJ/L mice were treated with in vitro-differentiated CD8+ T cells from control, Cbx3fl/fl or Cbx3fl/fl,Il21rΔ/Δ (Cbx3-Il21r-deficient); ****p≤0.0001, n = 3-4 recipients. h, Western blot of TCF-1 expression was done using total lysates of day 5 differentiated CD8+ T cells; C: Cbx3/HP1γ-deficient; Tcf7fl/+ and Tcf7fl/fl; Cbx3-Tcf7-deficient. i-k, Ovarian ascites (i), B16 tumor size (j) and NBL tumor volume (k) were measured in control, Cbx3/HP1γ-deficient and Cbx3-Tcf7-deficient mice; Graphpad unpaired student t-test: *p≤0.05, **p≤0.01, ***p≤0.001; n = 3-4 per genotype.

Figure 6. LEF-1 and IL-21R are required for Cbx3/HP1γ-deficient CD8+ effector T cells to persist in tumors. a,b,c, Frequencies of CD8+NKG2D+ T cells in ovarian ascites (a), B16 (b) and NBL (e) tumors were determined using data extracted from flow analysis. d,e, Frequencies of CD8+NKG2D+ T cells (gated on CD45.2+ populations) in B16 melanoma (d) and NBL (e) tumors from B6.SJL mice treated with in vitro-differentiated control, Cbx3/HP1γ- or Cbx3-Lef1-deficient CD8+ effector T cells on day 10 and day 14 after injection of B16 and NBL tumor cells, respectively. f-h, ovarian ascites (f), B16 tumor size (g) and NBL tumor volume (h) of control, Cbx3/HP1γ- or Cbx3-Tcf7-deficient mice. Statistics: bars: group median; Graphpad unpaired student t-test: *p≤0.05, **p≤0.01, ***p≤0.001; each symbol = one mouse; n = 3-5.

Figure 7. LEF-1 and IL-21R are required to maintain effector activity in tumors. a,b, Prf1 and Gzmb relative expression in day 18 B16 melanoma tumors was quantified by RT-qPCR using tumor RNA from indicated mice. c,d, Prf1 and Gzmb relative expression in day 18 B16 melanoma tumors from B6.SJL mice treated with in vitro-differentiated control, Cbx3/HP1γ- or Cbx3-Lef1-deficient CD8+ effector T cells on day 10 after injection of B16 tumor cells. e-g, Prf1, Gzmb and Ifng relative expression in day 33 NBL tumors. h-j, Prf1, Gzmb and Ifng relative expression in day 33 NBL tumors from B6.SJL mice treated with in vitro-differentiated control, Cbx3/HP1γ- or
Cbx3-Lef1-deficient CD8+ effector T cells on day 14 after injection of NBL tumor cells. Statistics: Graphpad unpaired student t-test: *p≤0.05, **p≤0.01, ***p≤0.001, ****p≤0.0001. n = 3 tumors from each mouse strain; representative of 3 experiments.

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Methods

Mice

Cbx3/HP1γ floxed mice were generated and provided by Dr. Prim B. Singh. These mice were backcrossed to C57BL/6 for twelve generations; they bred and developed normally before and after Cre-mediated deletion. The Cbx3/HP1γ transgenic line was generated by inserting a 1 kb fragment of mouse Cbx3/HP1γ cDNA, derived from activated CD8+ T cells, into EcoR1/Sma1 cloning sites of the VA-hCD2 vector provided by Dr. Dimitris Kioussis. The purified Cbx3/HP1γ-VA-hCD2 DNA construct was injected into C57BL/6 pronuclei by Dr. Lina Du at the fee-based Dana-Farber/Harvard Cancer Center (DF/HCC) Transgenic Mouse Core. Two founder transgenic lines were produced. The Lef1 floxed mice were generated as described and provided by Dr. Hai-Hui Xue. The following mouse lines were purchased from The Jackson Laboratory: Tcf7 floxed (B6(Cg)-Tcf7tm1Hlx/J)4, Il21r knock out (B6.129-Il21r<sup>tm1Kopf/J</sup>)5, CD8α-Cre transgenic (C57BL/6-Tg(Cd8a-cre)1Itan/J)6, ROSA26 (B6;129-Gt(Rosa)26Sor<sup>tm2Shao/J</sup>)7, Prf1 knock out (C57BL/6-Prf1<sup>tm1Sdz/J</sup>)8, and Ifng knock out (B6.129S7-Ifng<sup>tm1Ts/J</sup>)9. The B6.SJL line (B6.SJL-Ptprc<sup>a</sup>/BoyAiTac) was purchased from Taconic. All mice were maintained in specific pathogen-free (SPF) conditions. All mouse protocols were approved by the BIDMC Institutional Animal Care and Use Committee. All experiments were performed in accordance with relevant guidelines and regulations.

Tumor cell lines.

The mouse syngeneic ID8 ovarian tumor line was obtained from Dr. Katherine F. Roby under an MTA executed by the University of Kansas Medical Center. Dr. Eva M. Schmelz provided the mouse syngeneic MOSE-L<sub>TIC</sub> ovarian tumor line<sup>11,12</sup>. The NB9464 neuroblastoma cell line was a kind gift from Dr. Crystal L. MacKall (Stanford University). The B16-F10 melanoma cell line was purchased from ATCC and tested negative for mycoplasma.

Antibodies

All flow cytometry fluorochrome-conjugated antibodies were purchased from BioLegend, eBiosciences or BD Biosciences. The following Western blot and ChIP-tested antibodies were purchased from Cell Signaling: tri-methyl-histone H3 (Lys9) (D4W1U) rabbit mAb #13969, phospho-Rpb1 CTD (Ser2) (E1Z3G) rabbit mAb #13499, phospho-Rpb1 CTD (Ser5) (D9N51) rabbit mAb #13523, LEF-1 (C12A5) rabbit mAb #2230, cleaved caspase-3 (Asp175) antibody #9661 and phospho-HP1γ (Ser83) Ab #2600. The Western blot and ChIP-tested mouse anti-HP1γ
mAb (clone 42s2) was purchased from Millipore Sigma. The rabbit polyclonal to TBR2/Eomes (ab23345) was purchased from Abcam. The rat mAb to mouse granzyme B (16G6) was from Invitrogen/ThermoFisher Scientific.

**Tumor induction**

**Ovarian cancer**

Mice were injected intraperitoneally (IP) with ID8 (5 x 10^6/mouse in 200 µl PBS) or MOSE-L_LTCV (1 x 10^4/mouse in 200 µl PBS). On day 120 (ID8) or 36 (MOSE-L_LTCV), when abdominal distension was visible, mice were euthanized, ascites were collected, and volume measured using syringes fitted with 18-gauge needles.

**Melanoma**

Mice were implanted subcutaneously with B16 tumor cells (1 x 10^5/mouse in 100µL PBS). Using a digital caliber, B16 tumor size was measured and calculated on day 14 and at 2-day or 3-day intervals until day 20 depending on the size of the tumor, at which time mice were euthanized for analysis.

**Neuroblastoma**

Mice were implanted subcutaneously with NB-9464 tumor cells (1 x 10^6/mouse in 100µL PBS). Using a digital caliber, NB-9464 tumor volume was measured and calculated (W x L x 0.4) starting on day 22 and at 2-day intervals until day 30 depending on the size of the tumor, at which time mice were euthanized for analysis.

**Fluorescence-activated cell sorting or flow cytometry (FACS)**

FACS was performed on the BD 5-laser LSR II or the Beckman Coulter CytoFLEX LX. Analysis was performed with FlowJo software (Tree Star, Inc.). Tumors and ascites were harvested, minced (NBL and melanoma), and cell-suspensions were filtered through 70µm cell-strainers (Fisherbrand). Cells were stained for appropriate surface markers as indicated in figures.

**In vitro differentiation of CD8^+^ T cells**

CD8^+^CD44^- T cells were purified from spleen and peripheral lymph nodes using the MojoSort™ Mouse CD8 Naïve T-Cell Isolation Kit (BioLegend, #480044) followed by CD25-depletion using the mouse CD25 MicroBead Kit (Miltenyl Biotec, #130-091-072), all according manufacturers’ protocols. Naïve CD8^+^ T cells (1 x 10^6/mL) were activated with plate-bound anti-CD3 (clone 145-2C11, 0.25 µg/ml, BioLegend) and anti-CD28 (clone 37.51, 0.5 µg/ml,
BioLegend) in T-cell medium (high glucose DMEM, 10% FBS, penicillin/streptomycin, non-essential amino acids, HEPES, L-glutamate and sodium pyruvate) at 37°C in 10% CO₂ for 2 days. Cells were then removed from CD3/CD28 activation and re-cultured (5 × 10⁵/ml) in differentiation medium (T-cell medium supplemented with 10 IU/ml rhIL-2 from NCI) at 37°C in 10% CO₂. Cells were then sub-cultured every day in differentiation medium at 5 × 10⁵/ml each sub-culture. On day 5, cells were harvested and used for Western blots, ChIP-qPCR, qPCR or adoptive T-cell therapy.

**Adoptive T-cell transfer**

*In vitro*-generated donor CD8⁺ effector T cells (CD45.2⁺) were prepared as above. On day 0, B6.SJL (CD45.1⁺) mice were implanted subcutaneously with NB-9464 (1 x 10⁶/mouse in 100µL PBS) or B16 (1 x 10⁵/mouse in 100µL PBS) tumor cells. On day 10 (B16) or 14 (NB-9464) after tumor induction, mice were treated with donor CD8⁺ effector T cells (3 x 10⁶/mouse in 100µL PBS) via tail vein intravenous injection. B16 tumor size and NB-9464 tumor volume were measured and calculated on indicated dates.

**Co-culturing effector cells with B16 tumor cells**

Target B16 tumor cells were plated in 12-well plates at a density of 5 x 10⁵ cells per well in 750µl of medium (high glucose DMEM, 10% FBS, penicillin/streptomycin, non-essential amino acids, HEPES, L-glutamate and sodium pyruvate). CD8⁺ or CD4⁺ effector T cells were added. For ratio of 1:1 (effector:target), 5 x 10⁵ CD8⁺ or CD4⁺ T cells: 5 x 10⁵ B16 cells were co-cultured in the same well; ratio 5:1, 2.5 x 10⁶ CD8⁺ or CD4⁺ T cells: 5 x 10⁵ B16 cells. Plates were incubated at 37°C in 5% CO₂ for 24 hours. Wells were washed to remove non-adherent T cells. Adherent B16 cells were collected and washed with 1 ml cold PBS. Pellets were resuspended in Radio-Immunoprecipitation Assay (RIPA) lysis buffer containing a protease inhibitor cocktail (Roche) and used immediately or stored at −80°C for further analysis.

**Western Blots**

Cells (1 x 10⁶) were lysed with RIPA buffer (Boston BioProducts) containing a protease inhibitor cocktail on ice for 30 minutes. Lysates were centrifuged at 13,000 rpm for 10 minutes at 4°C. Protein extracts were denatured at 95°C for 10 minutes, separated by SDS-PAGE, and transferred to PVDF membranes (EMD Millipore). Membranes were probes with primary antibodies. Proteins of interest were detected with HRP-conjugated secondary antibodies and the Pierce™ ECL
Western Blotting Substrate (ThermoFisher Scientific). Membranes were exposed with HyBlot CL Autoradiography films (Denville Scientific, Inc.), and developed with the Kodak X-OMAT 2000 Processor.

**Chromatin immunoprecipitation followed by qPCR (ChIP-qPCR) and ChIP followed by deep sequencing (ChIP-Seq)**

*ChIP-qPCR*

ChIP was performed using the SimpleChIP Kit #9003 according to manufacturer’s protocol (Cell Signaling). Briefly, 2 x 10^7 CD8⁺ T cells were used for each ChIP. Cells were fixed in 1% formaldehyde to cross-link proteins to DNA, then lysed with 500µL of lysis buffer containing 1X ChIP buffer and protease inhibitor cocktail. Chromatin was sheared using a Sonic Dismembrator (Fisher Scientific Model 120) at 120W-20kHz power, at 15 seconds per cycle, 45 seconds break in between, for 3 cycles total. Chromatin was subjected to immunoprecipitation using specific antibodies at 4°C overnight with rotation. Following incubation, ChIP grade protein G magnetic beads were added to each ChIP and incubated for 2 hours at 4°C with rotation. Samples were placed in a magnetic separation rack for 2 minutes each time and washed 3X with high and low salt buffers. Chromatin was eluted from antibody/protein G magnetic beads using 1X ChIP elution buffer at 65°C for 30 minutes. Eluted chromatin was collected and subjected to cross-link reversal using 5M NaCl and 20 mg/ml Proteinase K, incubated for 2 hours at 65°C. After reversal of protein-DNA cross-link, the DNA was purified using DNA purification spin columns and eluted with DNA elution buffer provided in the kit. Purified ChIP DNA was immediately used for qPCR using primers shown in Table S3. BioRad hard-shell PCR Plates (BioRad) were used. In each well 2µL of purified ChIP DNA was added in triplicates along with 18µl primer mixture, which consisted of 1µL forward primer (5µM), 1µL reverse primer (5µM), 6µL nuclease-free water, and 10µL of 2X QuantiNova SYBR Green PCR Master Mix (Qiagen) or SimpleChIP Universal qPCR Master Mix (Cell Signaling). The plate was centrifuged at 300 RCF for 1 minute and read in the BioRad CFX384 Real-Time System (BioRad). The following qPCR settings were used: initial denaturation 95°C for 3:00 minutes, denatured at 95°C for 0:15 minutes, annealing and extension at 60°C for 1:00 minute, GOTO step denature and extension for a total of 40 cycles. Quantitative PCR result was analyzed, and the IP efficiency was calculated per formula provided in Simple ChIP Kit #9003 (Cell Signaling). Primers are listed in Extended Data Table 3.
**ChIP-Seq**

Detailed experiments and analyses have been described previously\(^\text{14}\).

**Reverse transcription-quantitative polymerase chain reaction (RT-qPCR).**

The BioRad hard-shell PCR Plates (BioRad) were used. In each well 2.5\(\mu\)L of sample was added in triplicates along with 7.5\(\mu\)L of the primer mixture (Table S2), which consisted of 0.5\(\mu\)L forward primer (10\(\mu\)M), 0.5\(\mu\)L reverse primer (10\(\mu\)M), 1.7\(\mu\)L PCR grade water, and 5\(\mu\)L of Roche's LightCycler 480 SYBR Green I Master (Roche). The plate was centrifuged at 300 RCF for 1 minute and read in the BioRad CFX384 Real-Time System (BioRad). The qPCR settings: 95°C for 10:00 minutes, 95°C for 0:10 minutes, 62°C for 0:20 minutes, 72°C for 0:20 minutes (Read Plate), GOTO step 2 for 39 times, 92°C for 0:05 minutes, melt curve 65°C to 97°C, at increment of 0.5°C for 1:00 minute (Read Plate), 40°C for 0:10 minutes. Primers are listed in Extended Data Table 2.

**Immunohistochemistry (IHC)**

Tumor section processing and TUNEL staining were performed by HistoWiz (Brooklyn, NY). Briefly, all IHC staining is automated using BOND Rx. Slides were treated with Dewax, a solvent-based solution (Leica Biosystems). Tissue sections were fixed by adding 10% neutral buffered formalin for 15 minutes. Slides were treated with Proteinase K at 1:500 dilution. Tissue sections were re-fixed using 10% neutral buffered formalin. Tissue sections were then treated with the equilibration buffer and incubated for 12 minutes at room temperature (RT). Subsequently the TdT reaction mix was added and incubated for 60 minutes at 37°C. The TUNEL reaction was stopped by adding 2X SSC. Tissue slides were treated with Peroxide Block (3-4% Hydrogen Peroxide) followed by wash buffer. Streptavidin HRP was added and incubated for 30 minutes at RT. DAB was added onto the slides and incubated for 10 minutes. Counter-staining was done by adding hematoxylin. Slides were covered using the Sakura Tissue Tek strainer and cover slips then scanned at 40x using the Leica AT2 scanner.

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