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*J Clin Invest.* 2022;132(17):e153437. [https://doi.org/10.1172/JCI153437](https://doi.org/10.1172/JCI153437).

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Tumor-intrinsic PRC2 inactivation drives a context-dependent immune-desert microenvironment and is sensitized by immunogenic viruses

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Our studies also point to intratumoral delivery of immunogenic viruses as an initial therapeutic strategy to modulate the immune checkpoint blockade (ICB) has demonstrated clinical success in “inflamed” tumors with substantial T cell infiltrates, but tumors with an immune–desert tumor microenvironment (TME) fail to benefit. The tumor cell–intrinsic molecular mechanisms of the immune–desert phenotype remain poorly understood. Here, we demonstrated that inactivation of the polycomb-repressive complex 2 (PRC2) core components embryonic ectoderm development (EED) or suppressor of zeste 12 homolog (SUZ12), a prevalent genetic event in malignant peripheral nerve sheath tumors (MPNSTs) and sporadically in other cancers, drove a context-dependent immune-desert TME. PRC2 inactivation reprogrammed the chromatin landscape that led to a cell-autonomous shift from primed baseline signaling-dependent cellular responses (e.g., IFN-γ signaling) to PRC2-regulated developmental and cellular differentiation transcriptional programs. Further, PRC2 inactivation led to diminished tumor immune infiltrates through reduced chemokine production and impaired antigen presentation and T cell priming, resulting in primary resistance to ICB. Intratumoral delivery of inactivated modified vaccinia virus Ankara (MVA) enhanced tumor immune infiltrates and sensitized PRC2-loss tumors to ICB. Our results identify molecular mechanisms of PRC2 inactivation-mediated, context-dependent epigenetic reprogramming that underlie the immune-desert phenotype in cancer. Our studies also point to intratumoral delivery of immunogenic viruses as an initial therapeutic strategy to modulate the immune-desert TME and capitalize on the clinical benefit of ICB.

Introduction

Polycomb-repressive complex 2 (PRC2), consisting of core components of enhancer of zeste homolog 1/-2 (EZH1/2), embryonic ectoderm development (EED), and suppressor of zeste 12 homolog (SUZ12), establishes and maintains H3K27me2/3 in the genome and regulates chromatin structure, transcription, cellular stemness, and differentiation (1). PRC2 is a context-dependent tumor suppressor, whose core components are frequently inactivated genetically or epigenetically in various cancer types, including malignant peripheral nerve sheath tumors (MPNSTs) (2, 3), melanoma (2), myeloid disorders (4, 5), T cell acute lymphocytic leukemia (ALL) (6), early T cell precursor ALL (7), pediatric gliomas (8–11), invasive breast cancer (12), and other cancers. Among all cancer types, high-grade MPNST, a group of aggressive soft tissue sarcomas with no effective therapies, has the highest prevalence of complete loss of PRC2 function through biallelic inactivation of the PRC2 core components EED or SUZ12 (2, 3, 13–15).

There is increasing evidence demonstrating the critical context-dependent role of PRC2 in regulating immune cell identity and function, including cytokotic CD8+ T cell repression (16–21), CD4+ T helper cell repression (22–24), and Treg activation (25–27) through regulation of cell lineage–specific gene expression. Therefore, selectively targeting PRC2 in immune cells may modulate the
tumor microenvironment (TME) and tumor responses to immunotherapy (28). Beyond immune cells, PRC2 in cancer cells has been shown to maintain bivalency at MHC class I (MHC-I) antigen-processing genes and silence MHC-I expression in selective MHC-I+ cancers (e.g., small cell lung cancer, neuroblastoma); targeting PRC2 can potentially enhance antitumor immunity by increasing MHC-I antigen presentation in this setting (29). However, it remains unclear how tumor-intrinsic PRC2 inactivation affects the tumor immune microenvironment and whether PRC2 has similar regulation of MHC-I in MHC-I+ cancers.

Here, using both human MPNST tissues and engineered PRC2-loss murine models, we demonstrated that tumor-intrinsic PRC2 loss promoted immune evasion and an immune-desert TME through epigenetic reprogramming and consequent deficiency in antigen presentation, chemokine production, and IFN-γ signaling, as well as primary resistance to immune checkpoint blockade (ICB). Further, we demonstrated that intratumoral delivery of the inactivated immunogenic modified vaccinia virus Ankara (MVA) enhanced tumor immunity, altered the immune-desert TME, and sensitized the PRC2-loss tumors to ICB therapy. These studies indicate that genetic inactivation of PRC2 can be used as a biomarker for resistance to ICB therapy in selective cancers and that engineered immunogenic viruses can be used to enhance tumor immunity and serve as an initial immunomodulatory strategy to overcome the cold TME in PRC2-loss tumors.

Results

Tumor-intrinsic PRC2 loss is associated with an immune-desert TME in MPNST and other cancers. To characterize the role of PRC2 inactivation in cancer pathogenesis, we analyzed the transcriptomes of 41 histologically confirmed high-grade human MPNST tumor samples, consisting of both wild-type PRC2 (referred to hereafter as PRC2-wt) and PRC2-loss samples. PRC2 loss in MPNSTs was confirmed by the loss of H3K27me3 immunostaining and/or genetic inactivation of EED or SUZ12 by MSK-IMPACT (MSK - integrated mutation profiling of actionable cancer targets) (30) (Supplemental Figure 1A and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI153437DS1). Principal component analysis (PCA), which detects sources of variation, showed that the MPNST samples were readily separated by PRC2 status in the first principal component (PCI) (Supplemental Figure 1B). We generated a gene set composed of genes that were differentially expressed between PRC2-loss and PRC2-wt samples. Hierarchical clustering based on these genes robustly separated the PRC2-loss and PRC2-wt MPNSTs, with the majority of the most differentially expressed genes upregulated in PRC2-loss compared with PRC2-wt MPNSTs, consistent with the role of PRC2 in transcriptional repression (Supplemental Figure 1C). Consistently, gene set enrichment analysis (GSEA) showed that the most enriched pathways and gene sets in PRC2-loss MPNSTs included the PRC2 modules and H3K27me3 target genes, organ development and morphogenesis, neuron cell fate specification, and WNT signaling gene sets (3, 31–33) (Figure 1A, Supplemental Figure 1D, and Supplemental Table 2). A distinct smaller subset of genes was consistently downregulated in PRC2-loss compared with PRC2-wt MPNSTs. Remarkably, nearly all of these gene sets were associated with immune function, including both innate and adaptive immune response pathway genes, T and B cell receptor signaling pathway genes, and antigen-binding and presentation genes (Figure 1A, Supplemental Figure 1C, and Supplemental Table 2).

We next performed IHC immunostaining for several established markers of distinct immune subclasses in human MPNST tumor tissue, including CD45 (pan-leukocyte), CD3 (T cells, CD4+ and CD8+ subsets), CD68 (macrophages/monocytes), and CD20 (B cells). Quantification of each immune subset revealed that, compared with PRC2-wt, the PRC2-loss MPNSTs were associated with significant reductions in CD45+ leukocytes (Figure 1, B and C), CD3+, CD4+, and CD8+ T cells, and CD68+ macrophages/monocytes (Figure 1, D and E). Although not significantly different, CD20+ B cells were only marginally present in both PRC2-wt and PRC2-loss MPNSTs (Figure 1, D and E). These data corroborate the transcriptome results of diminished tumor immune infiltrates (Supplemental Figure 1E). To further corroborate that PRC2 inactivation in various cancer contexts was associated with a cold TME, we identified available archival tumor tissues with confirmed PRC2 inactivation through EED or SUZ12 loss-of-function mutations by MSK-IMPACT and loss of H3K27me3 immunostaining by IHC (Supplemental Table 3 and Supplemental Figure 1F). IHC of CD45+ leukocytes demonstrated that these PRC2-loss tumors had low levels of tumor immune infiltrates that were comparable to those of PRC2-loss MPNSTs (Supplemental Figure 1F and Figure 1C). These results suggest that tumor cell-intrinsic PRC2 inactivation may exclude immune infiltrates and drive an immune-desert TME in various cancer contexts.

Antigen presentation and IFN-γ signaling are suppressed in PRC2-loss MPNSTs. To understand how tumor-intrinsic PRC2 loss leads to an immune-desert TME, we further characterized the transcriptomes of PRC2-loss versus PRC2-wt MPNSTs and focused on the initiating steps of the antitumor immune response (34). Among the genes downregulated by PRC2-loss compared with PRC2-wt tumors, antigen processing and presentation was one of the most negatively enriched gene sets by GSEA (Figure 2A and Supplemental Table 2). IFN-γ is an established key regulator of antigen presentation and chemokine for immune cell recruitment (35, 36). Consistently, IFN-γ signaling was impaired in PRC2-loss tumors, including cellular responses to IFN-γ, IFN-responsive genes, and regulation of IFN-γ production (Figure 2A). Expression levels of the tumor cell-autonomous gene IFNγR1 and downstream signaling axis and effector genes, including JAK1, JAK2, and IRF1, were all significantly lower in PRC2-loss tumors than in PRC2-wt tumors (P < 0.05; Figure 2B). In addition, the expression of IFNG encoding IFN-γ, which is usually produced by immune cells, was markedly low in most PRC2-loss tumors (Figure 2B), consistent with the PRC2-loss-associated immune-desert phenotype. Overall, expression of the IFNG/IFNGR1/JAK signaling axis genes was markedly reduced in PRC2-loss compared with PRC2-wt tumors. Consequently, accompanying the PRC2 loss, the expression of genes relevant to antigen processing and presentation, including MHC-I (e.g., HLA-A and B2M), MHC-II (e.g., CD74 and HLA-DMα), and antigen processing (e.g., TAP1) (Figure 2B), was also decreased, suggesting decreased tumor immunogenicity in PRC2-loss compared with PRC2-wt tumors. We further validated the decreased protein expression of MHC-I, B2M, and MHC-II by IHC in PRC2-loss tumors using an MPNST tissue microarray (TMA) (Figure 2, C
NSG mice, gave rise to high-grade MPNSTs with histological (e.g., monotonous spindle cell morphology, herringbone pattern, and fascicular growth) and immunostaining (e.g., H3K27me3 and Ki67 staining) features resembling those of high-grade human MPNST (13) (Supplemental Figure 2A). Nucleoplasmic and chromatin fractionation demonstrated that the global decrease in H3K27me3 and the increase in H3K27ac occurred on chromatin in SUZ12-loss tumor cells (Supplemental Figure 2B). These characterizations combined with the loss of H3K27me3 immunostaining in PRC2-loss M3 cell line–derived MPNST tumors validated the model system for mechanistic studies.

We speculated that PRC2 loss may alter the chromatin context and directly affect the transcriptional regulation of genes related to immune signaling and responses. We first examined the impact of PRC2 loss on genome-wide distribution of chromatin accessibility by the assay for transposase-accessible chromatin using sequencing (ATAC-Seq) and PRC2-relevant chromatin marks by ChIP-Seq and CUT&RUN in PRC2-isogenic M3 cells. Globally, compared with PRC2-wt, loss of PRC2 not only led to a marked increase and decrease in H3K27me3, H3K27me2, and H3K27me1 marks and a reciprocal global increase in the H3K27ac mark in PRC2-isogenic human MPNST cells (sgCon vs. sgSUZ12) (Figure 3A). The PRC2-isogenic M3 cells, when orthotopically transplanted into the sciatic nerve pockets of immunodeficient and D). Furthermore, the key chemokines responsible for immune cell recruitment, including CXCL9/10 (P < 0.01) and CCL2/3/4/5 (P < 0.05), were significantly lower in PRC2-loss tumors than in PRC2-wt tumors (Figure 2B). These data further posit that the tumor cell–intrinsic PRC2-loss–associated immune-desert phenotype is driven by impaired antigen presentation and diminished IFN-γ signaling in tumors.

**PRC2 loss reprograms the chromatin landscape and suppresses a subset of IFN-responsive genes.** To examine the role of PRC2 loss in MPNST while minimizing cell line–specific confounding factors, we generated and validated PRC2-isogenic human MPNST cells using CRISPR/Cas9-mediated knockout of the PRC2 core component SUZ12 in a PRC2-wt, NF1−/−CDKN2A−/− M3 cell line derived from a human NF1–associated MPNST (13). SUZ12 loss led to a global reduction of the H3K27me3 and H3K27me2, and H3K27me1 marks and a reciprocal global increase in the H3K27ac mark in PRC2-isogenic human MPNST cells (sgCon vs. sgSUZ12) (Figure 3A). The PRC2-isogenic M3 cells, when orthotopically transplanted into the sciatic nerve pockets of immunodeficient NSG mice, gave rise to high-grade MPNSTs with histological (e.g., monotonous spindle cell morphology, herringbone pattern, and fascicular growth) and immunostaining (e.g., H3K27me3 and Ki67 staining) features resembling those of high-grade human MPNST (13) (Supplemental Figure 2A). Nucleoplasmic and chromatin fractionation demonstrated that the global decrease in H3K27me3 and the increase in H3K27ac occurred on chromatin in SUZ12-loss tumor cells (Supplemental Figure 2B). These characterizations combined with the loss of H3K27me3 immunostaining in PRC2-loss M3 cell line–derived MPNST tumors validated the model system for mechanistic studies.

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also a marked decrease in chromatin accessibility at 15,346 (16% of all ATAC peaks) and 20,099 (21% of all ATAC peaks) genomic loci, respectively (Figure 3B). The substantially changed chromatin accessibility regions (increased or decreased) were relatively similarly distributed at promoter (7.7%, 8.3%) and nonpromoter regions, including distal regulatory (59.9%, 68.8%) and intergenic regions (32.4%, 22.9%) (Supplemental Figure 2C). The markedly increased ATAC peaks by PRC2 loss overlapped more with H3K27me3-enriched loci (10.5%) compared to the decreased ATAC peaks (2.4%) (Supplemental Figure 2D), consistent with the PRC2 function in chromatin compaction and transcription repression. Importantly, the integration of differential ATAC peaks with multiple histone modifications showed that the increased/decreased H3K27ac peaks were well correlated with open/closed chromatin accessibility changes at both promoter and nonpromoter regions as a result of PRC2 loss (Figure 3C and Supplemental Figure 2, E and F). The global distribution of H3K36me2 and H3K36me3 chromatin marks that had been previously described to interact with PRC2 and H3K27me3 (37–42) were not obviously affected, especially surrounding the H3K27me3-enriched regions in the PRC2-wt context (Supplemental Figure 2, F–H).

H3K27ac is an established chromatin mark preferentially enriched at active promoters and distal regulatory enhancers (13, 43, 44). Despite the increase in the total amount of chromatin-bound H3K27ac with PRC2 loss, we found relatively balanced gains and losses in the genome when regions enriched with H3K27ac were compared (Figure 3D). While the markedly increased H3K27ac peaks mainly localized to distal regulatory and intergenic regions and overlapped with H3K27me3 peaks, the markedly decreased H3K27ac peaks mainly localized to promoters and existing superenhancers (SEs) in PRC2-wt control tumor cells, with minimal overlap with H3K27me3 (Figure 3D and Supplemental Figure 2I). Consistently, the increased H3K27ac peaks in the PRC2/H3K27me3 loss context showed a trend toward closer distances to the nearest H3K27me3 peaks compared with the decreased H3K27ac peaks, while their distances to H3K36me3 peaks were reversed (Supplemental Figure 2J). Interestingly, we observed that with PRC2/H3K27me3 loss, a diffuse H3K27ac signal spread into the H3K27me3-enriched regions in the PRC2-wt context, without enrichment of specific genomic loci, leading to a mild overall increase in the baseline H3K27ac signal (e.g., loci 1–4) (Supplemental Figure 2, G and H). Notably, the decreased H3K27ac peaks localized to genes with higher baseline expression levels compared with increased H3K27ac peaks (Figure 3F). Moreover, H3K27ac changes were well correlated with transcriptome changes at both promoter and nonpromoter regions (Figure 3F).
Figure 3. PRC2 loss reprograms the chromatin landscape and suppresses a subset of IFN-γ-responsive genes. (A) Immunoblots of the indicated proteins in PRC2-isogenic human MPNST cells. (B) Volcano plot of chromatin accessibility changes by ATAC-Seq. Red dots represent markedly changed ATAC peaks (FDR q < 0.1, fold change ≥1.5). (C) Density plot of histone modifications by ChIP-Seq, centered on markedly increased (left) and decreased (right) ATAC peaks by SUZ12 knockout in M3 cells. Promoter: transcriptional start site (TSS) ± 2 kb; nonpromoter: rest of the genomic regions other than promoters, including distal regulatory enhancers and intergenic regions. (D) Distribution of differential H3K27ac peaks across different genomic regions in PRC2-isogenic M3 cells (FDR q < 0.05, fold change ≥2), including promoter (TSS ± 2 kb), distal regulatory (–50 kb from the TSS to the transcriptional end site [TES] + 5 kb), and intergenic (nonpromoter, nondistal regulatory) regions and SEs. (E) Violin plots of mRNA baseline expression of genes associated with PRC2 loss induced significantly increased (SigUP) and decreased (SigDN) H3K27ac at their respective loci in M3 sgCon cells. (F) Correlation of transcriptome and H3K27ac enrichment changes at promoter (TSS ± 2 kb) and nonpromoter (FDR q < 0.05, fold change ≥2) regions. Blue and gray dots represent peaks mapped to genes with (DiffExpGene) and without (nonDiffExpGene) significant transcriptome changes, respectively. (G) HOMER motif analysis of PRC2-loss-associated significantly decreased and increased H3K27ac peaks in PRC2-isogenic M3 cells. (H) ChIP-Seq and ATAC-Seq profiles at the loci of representative IFN-γ-responsive genes, e.g., CCL2, in PRC2-isogenic M3 cells. Pink indicates an H3K27ac change; yellow indicates an H3K27me3 change.
Motif analysis revealed significant enrichment of transcription factor binding motifs associated with immune signaling pathways and responses (e.g., the IFN signaling-associated IRF family and ISRE motifs) only in the decreased, but not the increased, H3K27ac peaks associated with PRC2 loss (Figure 3G and Supplemental Table 4). In addition, we observed that a subset of IFN-γ-responsive gene loci were directly affected by PRC2 loss. For example, H3K27ac enrichment at the SE locus of the monocyte chemotactic protein CCL2 was significantly (P > 0.05, fold change > 2) diminished by PRC2 loss (sgSUZ12) compared with controls (Figure 3H), whereas H3K27ac enrichment at other IFN-γ target gene loci (e.g., IRF1, CD274) was unchanged (Supplemental Figure 2K). We observed that many decreased and increased SE regions by PRC2 loss were preferentially flanked by broad enrichment of H3K27me3 in the genome of PRC2-wt tumor cells (Figure 3H and Supplemental Figure 2L). Further, inhibition of the main histone acetyl transferase (CBP/P300) for H3K27ac by A-485, or of the binding of bromo-ure 2L). Further, inhibition of the main histone acetyl transferase (CBP/P300) for H3K27ac by A-485, or of the binding of bromodomain (BRD) proteins to H3K27ac by JQ1, suppressed IFN-γ-responsive CCL2 gene expression in PRC2-wt M3 cells (Supplemental Figure 2M). These data indicated that CCL2 expression in the PRC2-wt context required H3K27ac modification at the CCL2 locus and that the diminished expression of CCL2 in the PRC2-loss context was a direct consequence of the decreased H3K27ac enrichment at the CCL2 locus. These data suggest that PRC2 loss affects the global distribution of H3K27ac and reprograms the genome-wide chromatin context of both the promoter and enhancer landscapes, which in turn alters signaling-dependent transcriptional responses (Supplemental Figure 2N).

Decreased chromatin accessibility for IFN-γ-responsive gene loci in PRC2-loss MPNST cells. One of the fundamental functions of the chromatin context is to prime the cells for the signaling-dependent transcriptional response. Reasoning that an altered local chromatin context mediated by PRC2 loss may change the transcriptional output, we next examined the transcriptome changes of IFN-γ-responsive genes in response to IFN-γ stimulation in PRC2-isogenic MPNST cells. Stimulation with 10 ng/mL exogenous IFN-γ for 24 hours had no significant impact on the levels of H3K27me3 modification or SUZ12/EED mRNA expression in PRC2-isogenic M3 cells (Supplemental Figure 3, A and B). PCA of ATAC-Seq replicates under various conditions demonstrated robust clustering of replicates and separation of samples based on PRC2 status (PC1) and IFN-γ stimulation (PC2) (Figure 4A). K-means clusters 2, 6, and 7 were most representative of increased chromatin accessibility in response to IFN-γ stimulation in the PRC2-loss (sgSUZ12) and PRC2-wt (sgCon) contexts, respectively (Figure 4B). Consistently, de novo motif analysis of the K-means clusters showed that only clusters 2, 6, and 7 identified IFN-γ-stimulation-related motifs (e.g., IRF8 and PU.1:IRF8) but with differential significance. The IFN-γ-related motif IRF8 was the top-most enriched motif of cluster 7, with a P value of 1 x 10-12,99, consistent with a primed chromatin context in response to IFN-γ stimulation through IRF activation in the PRC2-wt context. In contrast, the top-most enriched motif for cluster 2 was the GCN4/AP1 motif (P = 1 x 10-1227), whereas the classic IFN-γ-signaling-relevant motif IRF8/PU.1 was less significantly enriched (P = 1 x 10-25) (Figure 4B and Supplemental Table 5), suggesting a shift from IRF signaling to GCN4/AP1 signaling in a PRC2-loss-primed chromatin context. Moreover, gene ontology (GO) analysis also revealed substantial differences between the PRC2-loss representative cluster 2 and the PRC2-wt representative cluster 7 of chromatin accessibility changes induced by IFN-γ. GO analysis revealed that cluster 7 was most enriched for immune response-related signaling pathways, including pathways for antigen presentation and IFN signaling (Figure 4C), whereas cluster 2 was most enriched for development-related pathways, e.g., cellular differentiation, organ development, and PRC2 and H3K27me3 target pathways (Figure 4D).

Nevertheless, the transcriptome change tracked changes in the local chromatin context. In response to IFN-γ stimulation, the transcription of IFN-γ response genes (e.g., IRF1 and CD274) was unchanged when the local chromatin context was unperturbed (Figure 4E and Supplemental Figure 2K), and the transcriptional activation of IFN-γ response genes (e.g., CCL2, CD74, CIITA, and HLA-DRA) was significantly blunted (Figure 4F and Supplemental Figure 3C) when the gene loci were associated with a decrease in chromatin accessibility and H3K27ac signals as a result of PRC2 loss (Figure 3H and Supplemental Figure 2E). Further, the blunted transcriptional activation of these genes was most pronounced when the IFN-γ source was low (Figure 4F and Supplemental Figure 3C). We also restored EED in an EED-mutant human MPNST cell line sNF96.2 and validated the restoration of PRC2 function by H3K27me3 using a doxycycline-inducible EED system (Supplemental Figure 3, D and E). Reexpression of wild-type EED (PRC2 restoration) in sNF96.2 cells enhanced the expression of IFN-γ-responsive genes such as CCL2 and CD74 that were otherwise downregulated as a result of PRC2 loss (Supplemental Figure 3F). These observations suggest that PRC2 loss reprograms the steady-state primed chromatin context and leads to a blunted IFN-γ response in tumor cells.

Engineered PRC2 loss recapitulates the diminished IFN-γ signaling and the cold TME in both MPNSTs and breast cancer murine models. To evaluate the impact of tumor cell–intrinsic PRC2 inactivation on the TME in vivo, we generated a histologically confirmed Nf1–/– Cdkn2a/b–/– murine MPNST tumor–derived cell line (SKP605) from skin-derived precursors (SKPs) of C57BL/6J mice (45, 46) (Supplemental Figure 4, A and B). Using CRISPR/Cas9-mediated knock-out of the PRC2 core component Eed, we generated and validated PRC2-isogenic murine MPNST cells (SKP605, sgCon vs. sgEed) amenable for orthotopic and syngeneic transplantation into immunocompetent C57BL/6J mice (Figure 5, A and B, and Supplemental Figure 4C). The orthotopically transplanted PRC2-loss (sgEed) tumors exhibited accelerated growth in the sciatic nerve pockets of C57BL/6J mice compared with PRC2-wt (sgCon) tumors (Figure 5, C and D, and Supplemental Figure 4D). The expression levels of immune cell recruitment chemokines, e.g., Ccl2 and Cxcl10, and the lymphocyte activation cytokine Il2 were significantly decreased in PRC2-loss compared with PRC2-wt tumors (Supplemental Figure 4E), indicating decreased immune cell infiltration. Profiling of tumor immune infiltrates by FACS demonstrated a significant reduction in CD45+ leukocytes in PRC2-loss tumors compared with PRC2-wt tumors (Figure 5E and Supplemental Figure 4F). Further, we observed a reduction of tumor immune infiltrates in PRC2-loss tumors across all major subclasses of immune cells, including MHCII+CD11c+ DCs, TCRβ+ T cells, B220+ B cells, and, to a lesser extent, F4/80+CD11b+ macrophages (Figure 5F, Supplemental Figure 4G, and Supplemental Figure 5), phenocopying the cold TME of human PRC2-loss MPNSTs. Importantly, the IFN-γ- and TNF-α–mediated separation of samples based on PRC2 status (PC1) and IFN-γ stimulation or MPNST cells. Stimulation with 10 ng/mL exogenous IFN-γ stimulated the global distribution of H3K27ac and reprograms the genome-wide chromatin context of both the promoter and enhancer landscapes, which in turn alters signaling-dependent transcriptional responses (Supplemental Figure 2N).
CD4+ T cells were both significantly reduced in PRC2-loss tumors (Figure 5G and Supplemental Figure 4H). The immune infiltration differences between PRC2-isogenic SKP605 tumors were further confirmed by IHC of CD45+ leukocytes and various subclasses of immune cells (Supplemental Figure 4I). These data suggest that PRC2 inactivation in MPNSTs led to diminished recruitment of tumor immune infiltrates and a reduction of functional T cells, which together contributed to a cold TME.

To evaluate whether PRC2 loss has a similar effect on the TME in other cancer types, we used CRISPR/Cas9-mediated knock-out of the PRC2 core components Eed or Suz12 and generated a PRC2-isogenic murine mammary tumor model (AT3, sgCon vs. sgEed or sgSuz12) amenable for syngeneic transplantation into C57BL/6J mice (Supplemental Figure 4I). These data suggest that PRC2 inactivation in MPNSTs led to diminished recruitment of tumor immune infiltrates and a reduction of functional T cells, which together contributed to a cold TME.

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antigen presentation–related genes (e.g., Tap1, Tap2), and MHC-I (e.g., H2-k1, H2-q4, H2-23) and MHC-II (e.g., H2-t10 and H2-aa) genes in PRC2-loss versus PRC2-wt tumors (Figure 6D), indicating impaired tumor immunogenicity by tumor cell–intrinsic PRC2 inactivation. The immune recruitment chemokines were also markedly impaired tumor immunogenicity by tumor cell–intrinsic PRC2 inactivation. The immune recruitment chemokines were also markedly decreased in PRC2-loss compared with PRC2-wt AT3 tumors (e.g., Cxcl9, Cxcl10, Ccl5) (Figure 6D). Therefore, engineered PRC2-loss murine mammary tumors phenocopied the transcriptome changes and recapitulated the impaired immunogenicity of PRC2-loss human MPNSTs, irrespective of tumor lineage.

We next analyzed the population change of infiltrating immune cells in explanted PRC2-isogenic syngeneically transplanted AT3 tumors (Supplemental Figure 5). We observed a significant reduction of CD45+ leukocytes in PRC2-loss compared with PRC2-wt tumors, which was confirmed by IHC (Figure 6E and Supplemental Figure 6, F and G). Similarly, the reduction of tumor immune infiltrates with PRC2 loss was seen across all major subclasses of immune cells, including MHC-II CD11c+ DCs, TCRβ+ T cells (both CD4+ and CD8+), F4/80+CD11b+ macrophages, and to a lesser extent B220+ B cells (Figure 6F and Supplemental Figure 6H). Importantly, we detected a significant reduction in functional IFN-γ+ T cells in both in CD4+ and CD8+ T cell subclasses (Figure 6G and Supplemental Figure 6J), as well as a reduction of TNFα+ and a trend toward a reduction of the granzyme B+ (GzmB+) CD8+ T cells in PRC2-loss tumors (Supplemental Figure 6, J and K). These observations indicate that PRC2 loss in tumors not only led to diminished numbers of T cells, but also significant functional impairment of T cells. Since functional T cells are the main source of IFN-γ, these data indicate that IFN-γ is probably diminished in PRC2-loss tumors, which can further amplify the immune evasion phenotype.

To further dissect how PRC2 loss drives an immune evasion phenotype, we specifically evaluated T cell priming in tumor draining lymph nodes (TDLNs), a critical initial step in the development of antitumor immunity (34, 47). We induced exogenous expression of the model antigen OVA and generated PRC2-isogenic AT3 OVA+ cells (sgCon vs. sgEed), orthotopically transplanted these cells into the mammary fat pads of C57BL/6J mice, and analyzed the OVA-specific CD8+ T cells in TdLNs (Supplemental Figure 6, L and M). The MHC-I OVA tetramer+ CD8+ T cells were significantly diminished in the TdLNs from mice bearing PRC2-loss OVA+ compared with PRC2-wt OVA+ AT3 tumors (Figure 6H and Supplemental Figure 6N). These data indicate that PRC2 loss in tumors suppressed the initial antigen cross-presentation by DCs and impaired tumor-specific CD8+ T cell priming. These events, combined with the decreased tumor-infiltrating DCs and macrophages as well as diminished expression of immune recruitment chemokines, collectively contributed to the immune-desert TME. The observations from the AT3 murine mammary tumor model also indicated that the PRC2-loss–mediated immune evasion was not restricted to the MPNST context but could be generalized to other cancer types as well.

Engineered PRC2-loss tumors confer primary resistance to ICB.

Since tumor cell–intrinsic PRC2 loss drives an immune-desert TME, we speculated that PRC2 loss might confer primary resistance to FDA-approved ICB immunotherapies, including anti–programmed cell death 1 (anti–PD-1) and anti–cytotoxic T
recruitment of CD45+ immune cells, including both CD4+ and CD8+ T cells, was significantly blunted in the PRC2-loss tumors; this was accompanied by a reduction of functional IFNγ+CD4+ and IFNγ+CD8+ T cells (Figure 7, E–G). These results demonstrated that PRC2-loss tumors were resistant to ICB therapy.

To evaluate the cellular and molecular components that mediate the ICB treatment responses in PRC2-wt AT3 (sgCon) tumors, we used depletion antibodies to selectively deplete CD4+ T cells, CD8+ T cells, or NK cells and used an IFNγ-blocking antibody to inhibit IFNγ signaling. We then examined the ICB treatment responses under these perturbations (Supplemental Figure 7D).

lymphocyte–associated protein 4 (anti-CTLA4) antibodies (48). We evaluated the therapeutic efficacy of combining anti-PD-1 and anti-CTLA4 antibodies in transplanted PRC2-isogenic AT3 tumor models (Figure 7A). Combined ICB was effective and significantly inhibited the growth of PRC2-wt AT3 tumors; in contrast, it failed to retard the growth of PRC2-loss AT3 tumors (Figure 7B). Moreover, the combined ICB treatment increased CD45+ immune infiltrates (Figure 7, C and D, and Supplemental Figure 7A), particularly TCRβ+CD4+ and TCRβ+CD8+ T cell infiltrates in the PRC2-wt tumors (Figure 7, E and F, and Supplemental Figure 7, B and C). However, in response to combined ICB treatment, the recruitment of CD45+ immune cells, including both CD4+ and CD8+ T cells, was significantly blunted in the PRC2-loss tumors; this was accompanied by a reduction of functional IFNγ+CD4+ and IFNγ+CD8+ T cells (Figure 7, E–G). These results demonstrated that PRC2-loss tumors were resistant to ICB therapy.

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Intratumoral delivery of immunostimulatory heat-inactivated MVA sensitizes PRC2-loss tumors to ICB therapy. Immunogenic modified MVA infection of DCs can induce type I IFN by activating the cyclic GMP-AMP synthase (cGAS) and stimulator of IFN gene–mediated (STING-mediated) cytosolic DNA–sensing pathways (49). Intratumoral (i.t.) delivery of heat-inactivated MVA (heat-iMVA) can generate local and systemic antitumor immunity mediated by CD8+ T cells and Batf3-dependent CD103+CD8+ DCs (50). Thus, we tested whether i.t. delivery of heat-iMVA could enhance the IFN response and innate immunity, modulate the immune-desert TME, and sensitize PRC2-loss tumors to ICB. We observed that heat-iMVA in PRC2-loss murine tumor cells (AT3 [sgEed], SKP605 [sgEed]) not only triggered type I IFN production (e.g., Ifnb1 and Ifna4) (Figure 8A), but also

We observed that NK cell depletion did not significantly affect the ICB treatment response and that CD4+ T cell depletion abolished the ICB treatment response. In contrast, IFN-γ and CD8+ T cell depletion not only diminished the ICB treatment responses, but also led to accelerated tumor growth compared with untreated controls (Figure 7H). Consistently, expression levels of Ifng and its response genes (e.g., Cd274, Irf1, Ccl4) were diminished by depletion of IFN-γ+, CD4+, or CD8+ T cells compared with expression levels in the ICB treatment controls (Figure 7I and Supplemental Figure 7E). These data indicate that IFN-γ in the TME was critical for immune surveillance and control of tumor growth as well as for mediating the ICB treatment response, and that the source of IFN-γ was mainly CD8+ T cells and partially CD4+ T cells.

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Figure 8. Intratumoral injection of heat-iMVA sensitizes PRC2-loss tumors to anti–PD-1 and anti-CTLA4 therapies in murine cancer models. (A and B) Fold changes in mRNA expression levels of type I IFN (A) and IFN-γ–responsive (B) genes by qRT-PCR, 24 hours after heat-iMVA infection in vitro (n = 3). (C) Schematic of the treatment plan for heat-iMVA and ICB therapy in PRC2-loss AT3 (sgEed) or SKP605 (sgEed) tumors. (D and E) Tumor growth curves (D) and Kaplan-Meier survival curves (E) over time in mice with AT3 sgEed tumors under the indicated treatment conditions (n = 10 per cohort). (F) Percentages and representative FACS profiles of tumor-infiltrating CD45+ immune cells in AT3 (sgEed) tumors under the indicated treatment conditions (n = 5 per cohort). (G) Percentages of CD4+ and CD8+ T cells among all live cells and the relative Ki67+ subpopulation in AT3 (sgEed) tumors under the indicated treatment conditions. (H and I) Percentages of FoxP3+ Tregs among CD4+ T cells (H) and GzmB+ cells among CD8+ T cells (I) in AT3 (sgEed) tumors under the indicated treatment conditions (n = 5 per cohort). (J and K) Tumor growth curves for each tumor (J) and Kaplan-Meier survival curves (K) over time for mice with SKP605 (sgEed) tumors under the indicated treatment conditions. (L) Tumor growth curves for each tumor s.c. grafted onto the left flank of C57BL/6J mice (n = 8–10 tumors per cohort). *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001, by unpaired, 2-tailed t test (D and G–I) between 2 groups; by log-rank (Mantel-Cox) test (E and K); and by 1-way ANOVA (G–I) among 3 groups. Data indicate the mean ± SEM.
rescued the PRC2-loss-mediated IFN-γ signaling deficiency, resulting in increased chemokines and Cd274 (Pdcd4) expression (Figure 8B).

Intratumoral heat-iMVA alone in syngeneically transplanted PRC2-loss AT3 (sgEed) tumors only mildly retarded the tumor growth; however, i.t. heat-iMVA, when combined with anti–PD-1 and anti–CTLA4 ICB treatment, significantly reduced tumor growth (Figure 8, C and D, and Supplemental Figure 8A). This was accompanied by significant prolongation of survival of mice in the heat-iMVA and ICBCombination treatment group compared with mice that received ICBC alone or vehicle treatment (Figure 8E). Consistently, we observed significantly more cell death in virus-treated PRC2-loss tumors (Supplemental Figure 8B), suggesting an enhanced antitumor effect. Consistently, i.t. heat-iMVA significantly increased CD45+ immune cell infiltration in PRC2-loss AT3 (sgEed) tumors compared with vehicle treatment, which was further augmented when combined with ICBC treatment (Figure 8F), including TCRβ+ T cells and MHC-II+CD11c+ DCs (Supplemental Figure 8C). We did not observe significant changes in F4/80+CD11b+ macrophages or B220+ B cells in heat-iMVA-injected tumors (Supplemental Figure 8D). We observed significant enrichment of CD4+ and CD8+ T cells accompanied by increased proliferation (Ki67+) in the heat-iMVA treatment group, with and without ICBC treatment, compared with the vehicle-treated tumors (Figure 8G). Moreover, heat-iMVA treatment led to a significant decrease in immune-suppressive FoxP3+ Tregs (Figure 8H and Supplemental Figure 8D) as well as a significant increase in GzmB+CD8+ cytotoxic T cells compared with vehicle-treated tumors (Figure 8I and Supplemental Figure 8E), especially when combined with ICBC therapy.

We further investigated i.t. heat-iMVA in a newly established PRC2-isogenic murine MPNST model (SKP605) amenable for syngeneic transplantation that resembles human MPNSTs (Figure 5 and Supplemental Figure 4). In the PRC2-wt context, SKP605 (sgCon) tumors already had low levels of tumor immune infiltrates, which were further diminished with PRC2 loss (sgEed) (Figure 5E); both PRC2-wt and PRC2-loss SKP605 tumors were resistant to ICBC treatment (Supplemental Figure 8F). Using the murine MPNST model, we observed that i.t. heat-iMVA sensitized the PRC2-loss SKP605 tumors to ICBC treatment and significantly prolonged survival (Figure 8, J and K). Notably, 7 of 10 mice treated with the combined i.t. Heat-iMVA and ICBC had a complete response (CR) (Figure 8J). Moreover, we did not observe any tumor regrowth when the 7 mice with a CR were rechallenged with SKP605 (sgEed) tumor cells on the opposite side of the previously treated tumor grafts (Figure 8L), suggesting an adaptive immune response. These results demonstrate that i.t. delivery of immunogenic MVA therapy combined with ICBC was an effective initial strategy to modify the cold TME and elicit an antitumor effect in PRC2-loss tumors.

Discussion

Unlike previously described mechanisms of tumor cell–intrinsic immune evasion that primarily affect selective subpopulations of tumor immune infiltrates by, for example, decreasing T lymphocyte infiltration and function or recruiting myeloid-suppressive cells (e.g., LKB1 mutation in lung cancer) (47, 51–59), our study showed that PRC2 loss in tumors affected a broad spectrum of subpopulations of immune infiltrates, including DCs, T cells, B cells, and/or macrophages, and therefore drives an immune-desert TME, consistent with a prior proteomics study of MPNST (60). Rather than influencing MHC-I and MHC-II gene expression directly (29, 60), we found that PRC2 genetic inactivation in tumors led to a reprogrammed H3K27ac enhancer and promoter landscape and pervasive shifts of multiple chromatin templated processes, particularly activation of WNT/β-catenin signaling, diminished IFN signaling responses, reduced chemokine production (e.g., CCL2), and, consequently, impaired antigen presentation. These effects were further amplified in vivo and ultimately resulted in an immune-desert TME with diminished recruitment of the major subclasses of immune cells and resistance to ICBC treatment in both MPNSTs and other cancer types. Importantly, our study demonstrated that immunotherapeutic viruses could overcome a PRC2-loss–mediated immune-desert TME and sensitize PRC2-loss tumors to ICBC treatment.

We have established and characterized a PRC2-isogenic SKP605 murine MPNST model amenable for syngeneic transplantation into C57BL/6J mice. This model harbors the characteristic genetic alterations of and exhibits the histopathological and TME features of human MPNSTs. We noted some differences in tumor immune infiltrates between the murine and human MPNSTs, e.g., B220+ tumor-infiltrating B cells that appeared more enriched in murine MPNSTs compared with human tumors (Figure 1, D and E, and Figure 5F), probably due to host differences between mice and humans, syngeneic transplant systems, a lack of full recapitulation of the entire genetic alteration landscape, and the differential timing of tumor development. Nevertheless, we believe this model represents a valuable resource for the NF1 and sarcoma community for future mechanistic and therapeutic investigations of MPNSTs in immunocompetent hosts.

Mechanistically, PRC2 loss in tumor cells led to genome-wide redistribution of chromatin accessibility and chromatin modifications (e.g., H3K27ac), and hence reprogramming of the chromatin landscape. Although with PRC2 loss, global H3K27me2/3 histone modifications were lost, with a reciprocal global gain of H3K27ac on the chromatin, the genome-wide change in H3K27ac was not uniform, with distinct genomic regions of increased and decreased H3K27ac enrichment, as well as an increase in diffuse H3K27ac signals over large genomic territories. The global increase in H3K27ac was not due to increased enzymatic activity of CBP/P300, the canonical histone acetyltransferase (HAT) for H3K27ac, as other classical CBP/P300 substrates, such as H3K9ac and H3K18ac, had remained stable with PRC2 loss (data not shown). It is conceivable that the global increase in H3K27ac levels was due to an increased availability of unmodified H3K27 substrate because of H3K27me1/2/3 loss in PRC2-loss tumors. Moreover, global H3K27me1/2/3 loss led to a redistribution and spreading of H3K27ac from adjacent H3K27ac-enriched regions to previously H3K27me3-enriched regions and resulted in dampened enrichment at adjacent preexisting H3K27ac regions (Supplemental Figure 2, H and N). Consistent with chromatin accessibility changes mediated by PRC2 loss, there were many genes with markedly increased H3K27ac enrichment at distal regulatory enhancers and intergenic regions with a corresponding increase in transcriptional activity of master regulators, developmental and lineage specification pathways, and, importantly, genes involved in the WNT signaling pathway that had previously been shown to actively suppress T cell recruitment and
affect the local antitumor response through defects in T cell priming (47, 57, 58, 61). The decreased H3K27ac enrichment mediated by PRC2 loss mainly occurred at active promoters and distal regulatory enhancers including existing SEs in PRC2-wt tumors and correlated with the decrease in transcriptomes. This negative impact of H3K27ac enrichment preferentially affected the immune signaling pathways, including those for chemokatic cytokines (e.g., CCL2) and type I and type II IFN response genes, leading to significantly ($P < 0.05$) diminished expression of chemokines and dampened IFN-γ signaling responses in vitro. The diminished antitumor immune response was further amplified in vivo. Thus, in the relevant cell context, PRC2 loss reprogrammed the chromatin landscape and shifted a baseline primed immune signaling–dependent cellular response to the PRC2-regulated development and cellular differentiation transcriptional programs. Through these mechanisms, PRC2 loss reprogrammed the tumor cell and the TME to decrease antigen presentation and reduce chemokatic cytokine secretion, leading to diminished tumor immune infiltrates and ICB primary resistance. Our observations are consistent with recent reports demonstrating that inactivating mutations of the core components, e.g., PBRM1 of the PBAF complex, that are antagonistic to the PRC2 complex in tumor cells enhanced IFN-γ signaling and sensitized the PBAF-deficient tumors to ICB therapies (62, 63).

Virus-based cancer immunotherapy is a promising strategy to induce host antitumor immunity through multiple mechanisms, including virus-induced oncolysis and alteration of the immunosuppressive TME (64, 65). MVA is a highly attenuated vaccinia strain that has been approved by the FDA as a safe and effective vaccine against smallpox and monkeypox and has been investigated as a promising vaccine vector (66, 67). Infection of tumor and immune cells by i.t. delivery of heat-iMVA has been shown to lead to a robust induction of type I IFN and proinflammatory cytokines that inactivating mutations of the core components, e.g., PBRM1, of the PBAF complex, that are antagonistic to the PRC2 complex in tumor cells enhanced IFN-γ signaling and sensitized the PBAF-deficient tumors to ICB therapies (62, 63).

For tumor growth, tumors were measured twice weekly by Vernier calipers and female NSG (6–8 weeks of age) mice were purchased from the The Jackson Laboratory (stock no. 000664), and female NSG (6–8 weeks of age) mice were purchased from the Transplant mouse model. Female C57BL/6J (6–8 weeks of age) mice were purchased from mouse model. Female C57BL/6J (6–8 weeks of age) mice were purchased from the Transplant mouse model. Female C57BL/6J (6–8 weeks of age) mice were purchased from the The Jackson Laboratory (stock no. 000664), and female NSG (6–8 weeks of age) mice were purchased from the MSK animal core facility. PRC2-isogenic M3 cells (3 million cells) and SKP605 cells (5 million cells) in 100 μL 1:1 PBS/Matrigel (Corning, catalog 356237) were orthotopically transplanted into the sciatic nerve pockets of NSG and C57BL/6J mice, respectively. For heat-iMVA treatment, PRC2-isogenic SKP605 cells (1 million cells in 100 μL 1:1 PBS/Matrigel) were s.c. grafted onto the flanks of C57BL/6J mice. PRC2-isogenic AT3 cells (100,000–150,000 cells in 100 μL 1:1 PBS/Matrigel) were orthotopically transplanted into the mammary fat pads of C57BL/6J mice. For tumor growth, tumors were measured twice weekly by Vernier calipers and calculated as follows: tumor volume (TV) = 4/3 × width × height × thickness/2. For survival studies, mice were euthanized when the TV reached 1500 mm3 for AT3 murine mammary tumors and 300 mm3 for SKP605 s.c. tumors or when the mice exhibited signs of illness and discomfort that required euthanasia.

**Methods**

**Human tumor tissue collection.** Clinical samples were collected during surgical resection from patients with MPNSTs and other cancers according to MSK IRB protocols. Frozen and paraffin-embedded tissue samples were banked, and TMA s were generated. All MPNSTs and other PRC2-loss cancers were pathologically reviewed and confirmed by an MSK pathology expert. Sample annotations are shown in Supplemental Table 1.

**Cell lines.** HEK-293T and sNF96.2 cell lines were purchased from the American Type Culture Collection (ATCC). The human MPNST cell line M3 (NF1+/−, CDKN2A/B−/−) was a gift and derived from an NF1-associated MPNST by William Gerald (MSK). The murine MPNST cell line, SKP605 (NF1−/−, Cdkn2a−/−), was generated in-house from skin-derived precursors (SKPs) according to a previously published protocol (46) and confirmed by a pathologist. The murine breast cancer cell line AT3 was obtained in-house. All cell lines were confirmed as mycoplasma free.

All cell lines were cultured in DMEM supplemented with 1-glutamine (2 mM), penicillin (100 U/mL), streptomycin (100 μg/mL), and 10% heat-inactivated FBS in 5% CO2 at 37°C.

**IHC.** Human tissue processing, embedding, sectioning, and staining with H&E were performed at the MSK Department of Pathology. IHC of human TMA tumor samples was performed using a Ventana BenchMark ULTRA Automated Stainer at the MSK Human Oncology and Pathogenesis Program (HOPP) automatic staining facility. Mouse fresh tissues were fixed in 4% paraformaldehyde (PFA) overnight, washed 3 times with PBS for 5 minutes each time, and stored in 70% ethanol. Tissue paraffin embedding, sectioning, and H&E staining were performed by Histovis Inc. IHC of CD45 (Cell Signaling Technology [CST], catalog 70257, 1:100) was performed at the MSK Molecular Cytology Core Facility. IHC stainings for H3K27me3 (MilliporeSigma, catalog 07-449, 1:500, protocol no. 313), Ki67 (Abcam, catalog ab55580, 1:500, protocol no. 312), and S100B (Abcam, catalog ab52642, 1:2000, protocol no. 313) were performed using the Ventana BenchMark ULTRA Automated Stainer. Slides were scanned at the MSK Molecular Cytology Core Facility and analyzed using CaseViewer software.

**Gene knockout by CRISPR/Cas9.** The pLP2B plasmid was generated by removing Cas9-P2A-TRFP from pL-CRISPR-EFS-TRFP (Addgene, no. 57819) and replaced with Cas9-P2A-Blast. The lentiCRISPR-v2 vector with puromycin was purchased from Addgene (no. 52961). The sgRNA oligonucleotides described below were engineered into vectors using the standard CRISPR/Cas9 knockout protocol. M3 sgCon cells were pooled from 6 single-cell clones, and M3 sgsUZ12 cells were pooled from 9 single-cell clones. Both SKP605 sgCon cells and sgEd cells were pooled from 4 single-cell clones. See the Supplemental Methods for details on the methods used and sgRNA and sequences.
Viruses and intratumoral injection. The MVA was provided by G. Sutter (University of Munich, Munich, Germany). MVA was propagated in BHK-21 cells (baby hamster kidney cell, ATCC, catalog CCL-10) and purified through a 36% sucrose cushion. Heat-IMVA was generated by incubating purified MVA at 55°C for 1 hour. Seven to 11 days after implantation, tumors were measured and heat-IMVA (an equivalent of 4 × 10⁹ PFU) in 200 μL or PBS was delivered via i.t. injection twice weekly, and the mice were subsequently monitored daily.

**OVA model antigen system.** The pMSCV-EGFP-PGK-Luc2-2A-USA plasmid, which included OT-I and OT-II-binding antigens for mice on a C57BL/6j genetic background, was obtained in-house. To make the retrovirus, HEK293T cells were transfected with pMSCV-EGFP-PGK-Luc2-2A-USA and the packaging plasmids pSVSVG (Addgene, catalog 8454) and pEco (Takara, catalog PT3749-5). Cells transduced with the retrovirus were sorted for EGFP positivity by flow cytometry, and these cells stably overexpressed OT-I and OT-II-binding OVA model antigens. APC anti–mouse H-2Kb bound to the anti-IFN receptor antibody (BioLegend, catalog 141605) was used to detect the OVA model antigen on the cancer cell surface. To detect OVA-specific T cell priming, OVA- and OVA AT3 cells were orthotopically grafted into mammary fat pads of C57BL/6j mice. Eighteen days after grafting, cells isolated from TdLNs were incubated with 2 μg/mL OVA 257-264 in T cell medi-

Protein extraction and Western blotting. Sample preparation and Western blotting were performed as described previously (31) and in the Supplemental Methods. The following primary antibodies were used for immunoblotting: anti-NF1 (1:2000, Bethyl Laboratories, catalog A300-140A); anti-CDKN2A (1:1000, Delta BioLabs, catalog DB018); anti-CDKN2B (1:500, Abcam, catalog ab53034); anti-SUZ12 (1:1000, CST, catalog 3737); anti-H3K27me3 (1:2000, CST, catalog 9733); anti-H3K27me2 (1:5000, CST, catalog 9728); anti-H3K27me1 (1:1000, Takara, catalog MABIO321-1001); anti-H3K27ac (1:4000, Abcam, catalog ab4729); anti-histone H3 (1:2000, CST, catalog 12648); anti–β-actin (1:5000, Proteintech, catalog 66009-1-lg); anti-lamin B1 (1:2000, Proteintech, catalog 12987-1-AP); anti-HIRA (1:1000, Active Motif, catalog 39557); and anti-GAPDH (1:3000, CST, catalog 5174S). See complete unedited blots in the supplemental material.

**RNA-Seq and analysis.** Poly-A capture RNA-Seq of total RNA isolated from fresh tissues or cells using TRIzol was performed at the MSK Integrated Genomics Operation (IGO) facility on an Illumina HiSeq 2500 platform with 50 bp paired-end (human MPNST tumors) or single-end (AT3 tumors) reads to obtain a minimum yield of 40 million reads per sample. The sequence data were processed and mapped to the human reference genome (hg19) or mouse reference genome (mm9) using STAR, version 2.3 (68). Gene expression was quantified as transcripts per million (TPM) using STAR (69) and log-transformed. GSEA was performed using the JAVA GSEA 2.0 program (70).

**ATAC-Seq and analysis.** ATAC-Seq was performed at the Center for Epigenetics Research as previously described (71, 72). For each sample, nuclei from 50,000 cells were prepared and incubated with 2.5 μL transposase (Illimina) in a 50 μL reaction for 30 minutes at 37°C. After purification of transposase-fragmented DNA, the library was amplified by PCR and subjected to paired-end, 50 bp high-throughput sequencing on an Illumina HiSeq 2500 platform. For data analysis, ATAC-Seq reads were quality and adapter trimmed using Trim Galore before alignment to the human reference genome (hg19) with Bowtie2 using the default parameters. Motif signatures were obtained using HOMER, version 4.5 (http://homer.ucsd.edu). See the Supplemental Methods for details on the analysis.

**ChIP-Seq, CUT&RUN, and analysis.** Chromatin isolation from the indicated cells and immunoprecipitation were performed as previously described (73). The libraries were sequenced on an Illumina HiSeq 2500 platform with 50 bp paired-end reads. Reads were trimmed using Trim Galore software and then aligned to the human genome (hg19) using Bowtie2 alignment software, version 2.3.5 (74). Duplicated reads were eliminated for subsequent analysis. Peak calling for H3K27ac was performed using MACS2 software, version 2.1.1 (75) in paired mode and comparing ChIP samples to the input, using an FDR of q < 10⁻³. H3K27me3 peaks were called using a sliding window approach to find regions enriched with H3K27me3 compared with input reads. Spike-in was used in H3K27me3 ChIP-Seq for normalization because of the global loss of H3K27me3 after PRC2 loss. We discarded peaks mapped to blacklisted genomic regions identified by ENCODE (76, 77). The primary antibodies used for ChIP were anti-H3K27me3 (CST, catalog 9733) and anti-H3K27ac (Abcam, catalog ab4729).

The H3K27ac peaks were separated into promoters and distal and intergenic peaks as described previously (78). They were also used for SE analysis using the ROSE R package (option: −t 2500) (79, 80). The peaks from controls and sgSUZ12 samples were merged to generate a nonoverlapping list of union peaks. ChIP-Seq reads located to the merged peaks were calculated and used by DESeq2 software to identify peaks with differential modifications at an adjusted P value of less than 0.05 and a fold change of greater than 2. The markedly increased or decreased peaks in the sgSUZ12 samples at promoters and nonpromoter regions were subjected to independent transcription factor binding motif analysis with HOMER software (version 4.7, default parameter) (81), using all peaks as a background. The fold changes in these peaks were also compared with gene expression changes in the RNA-Seq analysis for each of the promoter and distal peaks assigned to genes (note that 1 gene could have multiple peaks).

The CUT&RUN assay was performed using the commercial kit and protocol (CST, catalog 86652). The primary antibodies (1 μg per assay) used for the CUT&RUN assay were as follows: anti-H3K27me3 (CST, catalog 9733); anti-H3K27ac (Abcam, catalog ab4729); anti-H3K36me2
(Thermo Fisher Scientific, catalog MA5-14867); and anti-H3K36me3 (Active Motif, catalog 61021).

**Data availability.** All sequencing data were deposited in the NCBI’s Gene Expression Omnibus (GEO) database: RNA-Seq of human MPNST tumor tissues (GEO GSE206527); RNA-Seq of AT3 tumors (GEO GSE179703); ATAC-Seq of M3 cells (GEO GSE179699); ChIP-Seq of M3 cells (GEO GSM5420909, GSM5420912, GSM5420919, GSM5420922); and CUT&RUN of M3 cells (GEO GSE202555).

**Statistics.** All statistical analyses were performed using GraphPad Prism 7 or 9 (GraphPad Software). Unless otherwise noted in the figure legends, all data are shown as the mean ± SEM combined with a 2-tailed, unpaired t-test for statistical comparisons between 2 groups, 1-way ANOVA for comparisons of more than 2 groups, and a log-rank (Mantel-Cox) test for survival analyses. A P value of less than 0.05 was considered statistically significant. All experiments shown were repeated at least twice.

**Study approval.** Clinical samples were collected according to MSK IRB protocols (IRB nos. 02-060 and 06-107). All procedures related to mouse handling, care, and treatments were performed according to guidelines approved by the IACUC of MSK.

**Author contributions**

JY, Yuedan Chen, PC, and Yu Chen conceived the project and designed the experiments. JY, Yuedan Chen, SW, CJL, JS, CRA, SS, PC, and Yu Chen collected clinical samples. JY and Yuedan Chen performed most of the in vitro and in vivo experiments with assistance from AJP, BGN, EWPW, and MARM and technical support from SW, CJL, MRP, EG, and JS for the in vivo experiments. JY, LD, NY, and YW performed the heat-iMVA-based viral therapy experiments. CRA and PC performed pathological and immunohistochemical review of the clinical and murine tumor samples. MOL, LD, and BGN provided additional conceptual advice, experimental design assistance, and technical support for the immunology experiments. DZ, RPK, JLVM, YL, PMG, FT, EK, Yu Chen, JY, and PC performed RNA-Seq, ChIP-Seq, CUT&RUN, ATAC-Seq, and integrative analyses. JY, PC, and Yu Chen wrote the manuscript. All authors reviewed and edited the manuscript.

**Acknowledgments**

We thank all members of the Chi and Chen laboratory for general support. We also thank William Gerald and Leon (Xiaoliang) Xu for the patient-derived MPNST M3 cell line. Next-generation sequencing, ChIP-Seq, CUT&RUN, RNA-Seq, and ATAC-Seq were performed at the IGO/Center for Molecular Oncology and Center for Epigenetics Research. IHC of FFPE and MPNST samples were procured from and performed at the Pathology core facility, and animal studies were performed at the Center for Comparative Medicine and Pathology core facility at MSK. This work was supported by grants from the NIH and the National Cancer Institute (NCI) (R01 CA228216, DP2 CA174949); Department of Defense (DOD) grant W81XWH-15-1-0124; a Francis Collins Scholar NTAP grant; a grant from the Cycle for Survival and Linn Family Discovery Fund (to PC); NCI, NIH grant P50 CA179649 (to PC, SS, and CRA); NCI, NIH grant R01 CA208100 (to Yu Chen); a grant from the Geoffrey Beene Cancer Research Fund (to PC, MOL, and LD); Translation- Onology Research in Oncology Training Program T32 grant ST32CA160001-09 (to AJP); and NIH grant P30 CA 008748 to the MSK Cancer Center (Core Grant).

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