Bivalent and Broad Chromatin Domains Regulate Pro-metastatic Drivers in Melanoma

- 3 Christopher Terranova^{#,1}, Ming Tang^{#,1,10}, Mayinuer Maitituoheti^{#,1}, Ayush T. Raman^{1,7,8},
- 4 Jonathan Schulz¹, Samir B. Amin^{1,6}, Elias Orouji^{1,11}, Katarzyna Tomczak¹, Sharmistha
- 5 Sarkar¹, Junna Oba³, Caitlin Creasy³, Chang-Jiun Wu¹, Dongyu Zhao⁹, Kaifu Chen⁹,
- 6 Lauren E. Haydu⁵, Wei-Lien Wang², Alexander J. Lazar², Scott E. Woodman^{1,3,4},
- 7 Chantale Bernatchez³, and Kunal Rai^{1,*}
- ⁸ ¹ Department of Genomic Medicine, University of Texas MD Anderson Cancer Center,
- 9 Houston, TX.
- ² Department of Pathology, University of Texas MD Anderson Cancer Center, Houston,
 TX.
- ¹² ³ Department of Melanoma Medical Oncology, University of Texas MD Anderson Cancer
- 13 Center, Houston, TX.
- ⁴ Department of Systems Biology, University of Texas MD Anderson Cancer Center,

15 Houston, TX.

- ¹⁶ ⁵ Surgical Oncology, University of Texas MD Anderson Cancer Center, Houston, TX.
- ⁶ The Jackson Laboratory for Genomic Medicine, Farmington, CT.
- ¹⁸ ⁷ Epigenomics Program, Broad Institute of MIT and Harvard, Cambridge, MA.
- ¹⁹ ⁸ Graduate Program in Quantitative Sciences, Baylor College of Medicine, Houston, TX.
- ⁹ Institute for Academic Medicine, Methodist Hospital Research Institute, Houston, TX.
- ¹⁰ FAS informatics, Department of Molecular Biology, Harvard, Cambridge, MA
- ¹¹ Epigenetics Initiative, Princess Margaret Genomics Centre, Toronto, ON Canada
- 23 *Corresponding Author: krai@mdanderson.org
- 24

25 ABSTRACT

Chromatin deregulation is an emerging hallmark of cancer. However, the extent of 26 epigenetic aberrations during tumorigenesis and their relationship with genetic 27 aberrations are poorly understood. Using ChIP-sequencing for enhancers (H3K27ac and 28 29 H3K4me1), promoters (H3K4me3), active transcription (H3K79me2) and polycomb 30 (H3K27me3) or heterochromatin (H3K9me3) repression we generated chromatin state 31 profiles in metastatic melanoma using 46 tumor samples and cell lines. We identified a 32 strong association of NRAS, but not BRAF mutations, with bivalent states harboring 33 H3K4me3 and H3K27me3 marks. Importantly, the loss and gain of bivalent states 34 occurred on important pro-metastasis regulators including master transcription factor 35 drivers of mesenchymal phenotype including ZEB1, TWIST1, SNAI1 and CDH1. 36 Unexpectedly, a subset of these and additional pro-metastatic drivers (e.g. POU3F2, SOX9 and PDGFRA) as well as melanocyte-specific master regulators (e.g. MITF, ZEB2, 37 38 and TFAP2A) were regulated by exceptionally wide H3K4me3 domains that can span tens of thousands of kilobases suggesting roles of this new epigenetic element in 39 40 melanoma metastasis. Overall, we find that BRAF, NRAS and WT melanomas may use 41 bivalent states and broad H3K4me3 domains in a specific manner to regulate pro-42 metastatic drivers. We propose that specific epigenetic traits – such as bivalent and broad 43 domains - get assimilated in the epigenome of pro-metastatic clones to drive evolution of 44 cancer cells to metastasis.

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48 Introduction

Melanoma is a deadly disease with an estimated 96,000 new cases each year¹⁻³. While targeted therapy and immunotherapy have become the standard of care with significant improvement in clinical response, more than 7,000 patients still succumb to this disease per year due to primary or acquired resistance^{4,5}. Therefore, it is critical to gain a deeper understanding of the disease biology to design more effective therapies.

Large scale efforts from consortiums such as The Cancer Genome Atlas (TCGA) 54 have provided deeper understanding of molecular aberrations in the disease⁶⁻⁸. These 55 56 studies identified critical somatic mutations in this disease that likely occur due to UV 57 exposure. Among these, somatic mutations in important bona fide oncogenes and tumor suppressors, such as BRAF, NRAS, NF1, INK/ARF, PTEN and TP53, have been 58 59 demonstrated to be well-chronicled drivers of this malignancy⁶⁻⁸. One of the important 60 findings from these studies were genetic aberrations in several key epigenetic regulators 61 such as EZH2, IDH1/2, ARID2, KMT2C and KMT2D⁶⁻⁸. Many of these proteins are enzymes that regulate covalent modifications of histones^{7,9-13}. Although recent studies 62 63 provide insight into the correlation of isolated histone marks, there are a myriad of 64 possible combinatorial patterns of histone modifications, and it is these combinatorial 65 states - not individual modifications - that dictate epigenetic status of associated genomic loci¹⁴⁻¹⁶. These observations suggest that epigenetic alterations, including those in 66 histone modifications, may play important roles in melanomagenesis. Indeed, specific 67 68 functional roles have been assigned to some of these players as well as histone variants such as macroH2A and H2A.z in melanomas^{17,18}. In addition, our previous study showed 69 70 alterations in specific chromatin states during transition from pre-malignant to malignant phenotype in melanoma¹⁹. 71

72 These studies provide strong rationale for a systematic mapping of epigenome to 73 obtain comprehensive understanding of epigenetic elements that may act as driver events 74 in specific melanoma tumors. This concept has been epitomized by the DNA methylation 75 profiles in large number of tumors by the TCGA study which has provided key concepts 76 about roles of this important epigenomic mark in cancer progression¹¹. For example, a 77 subset of cancer types are demonstrated to harbor hypermethylation phenotype, termed CpG Island Methylation Phenotype, which associated with mutations in IDH1/2 genes¹¹. 78 79 Chromatin state mapping in large number of tumors has the potential to identify similar 80 concepts²⁰. Furthermore, several projects such as ENCODE and Roadmap Epigenomics 81 have cataloged extensive histone modification in normal human tissues and cell lines which allows for identification of cancer specific alternations in chromatin states^{21,22}. Since 82 83 epigenetic aberrations are reversible by targeting of their enzyme regulators, the 84 chromatin mapping efforts are likely to identify potentially novel therapeutic strategy in 85 specific genetic context. For example, our previous study suggested that HDAC inhibitors could be a good strategy to block pre-malignant to malignant transition in melanoma¹⁹. 86

87 In this study, we present a comprehensive chromatin state analysis of metastatic 88 melanoma in 46 tumor samples (profiled by TCGA) and cell lines (Profiled by Cancer Cell 89 Line Encyclopedia (CCLE) or internal efforts at MD Anderson) by performing ChIP-90 sequencing of 6 histone modification marks. Overall, investigation revealed a mechanism 91 in which alterations of distinct chromatin states, including bivalent and exceptionally wide 92 H3K4me3 and H3K27me3 domains, regulate key drivers of a mesenchymal/invasive 93 phenotype, a network of genes that includes master transcription factors and melanocyte-94 specific regulators. Together, this study encompasses the most complete description

- 95 regarding the epigenetic circuitry governing melanoma metastasis that can serve as an
- 96 important resource for advancing the understanding of the melanoma epigenome.

103 Results

104 Bivalent polycomb repressive chromatin domains distinguish metastatic 105 melanoma tumors based on mutational subtypes.

106 Chromatin state profiling remains a powerful tool for determining the regulatory status of 107 annotated genes and identifying novel elements in non-coding genomic regions^{3,4}. Using 108 ChIP-sequencing for enhancers (H3K27ac and H3K4me1), promoters (H3K4me3), active 109 transcription (H3K79me2) and polycomb (H3K27me3) or heterochromatin (H3K9me3) 110 repression coupled with tissue-matched mutation, transcriptomic and methylation data, 111 we describe the cis-regulatory landscape across 46 melanoma samples. These 112 constituted 20 metastatic melanoma tumors (profiled by the TCGA study⁶), 10 patient-113 derived melanoma short term cultures (passage n < 10; profiled by internal effort at MD 114 Anderson; manuscript in preparation) and 16 established melanoma lines profiled by the Cancer Cell Line Encyclopedia/Sanger (CCLE)²³ (Supplementary Table 1). Using our 115 116 primary cohort of 20 metastatic melanoma tumor samples we computed multiple 117 chromatin state models (8-states through 30-states) with the ChromHMM algorithm (Fig. 118 **1a** and Supplementary Fig. **1a**). We chose an 18-state model because it is large enough 119 to identify important functional elements while still being small enough to interpret easily. 120 This model includes active (E1) and transcribed promoters (E2), harboring high levels of 121 H3K4me3, H3K27ac, H3K4me1 without (E1) and with H3K79me2 (E2) within the TSS or 122 TSS flanking regions (Supplementary Fig. 1b); transcribed genes (E3 and E4); genic (E5, 123 E6) and active enhancers (E7, E8) harboring high levels of H3K27ac and H3K4me1 with 124 concomitant enrichment of H3K79me2 within (E5-E6) or outside (E7-E8) the TSS 125 flanking regions; and heterochromatic (E10) or polycomb (E14) based repression

harboring high levels of either H3K9me3 (E10) or H3K27me3 (E14) respectively. In 126 127 addition, we also observed two prominent bivalent/poised states: first, harboring both 128 H3K4me3 and high levels of H3K9me3 (E12, annotated as "bivalent H3K9me3"), and 129 second, H3K4me3 and high levels of H3K27me3 (E13, annotated as "bivalent 130 H3K27me3"). Overall, the chromatin state profiles in metastatic melanoma tumors are 131 associated with both gene expression patterns and DNA methylation levels (Fig. 1b, c). 132 As expected, active promoters (E1 and E2) are associated with high levels of gene 133 expression and low levels of methylation, whereas repressed states (E12 and E13) are 134 associated with low levels of gene expression and high levels of methylation (Fig. 1b, c). Consistent with the previous Roadmap epigenome analysis^{21,22} in normal samples, DNA 135 136 methylation patterns also showed some dynamic patterns. For example. 137 hypermethylation in active chromatin states E3 through E8 are also associated with high 138 levels of transcriptional activation (Fig. 1b, c).

139 This TCGA tumor cohort is inclusive of multiple melanoma subgroups⁶, including 140 mutation subtypes (BRAF, NRAS, WT), transcriptomic subtypes (Immune, Keratin, MITF-141 low) and DNA-methylation subtypes (CpG, Hypermethlyated, Hypomethylated, Normal). 142 Before comparing chromatin state data and molecular subtypes, we first ensured the 143 generated ChIP-seq profiles could be uniquely mapped to the specific donor through CNV 144 analysis. In 18 of 20 tumors, the CNV analysis from ChIP-seq data correlated with the 145 CNV classification from TCGA (Supplementary Fig. 1c, d). Projection of chromatin state 146 data using Multidimensional Scaling (MDS) analysis revealed chromatin state E13 147 (bivalent state containing high levels of H3K27me3 and H3K4me3 and modest 148 enrichment of H3K9me3) was able to separate NRAS-mutant melanoma tumors from 149 BRAF-mutants and WT samples in the first dimension (Fig. 1d, e and Supplementary Fig. 150 1e). Moreover, differential analysis of bivalent chromatin states (E12 and E13) between 151 mutational subtypes (BRAF vs NRAS vs WT) demonstrated NRAS-enrichment specific 152 to bivalent H3K27me3 high (Fig. 1f-h and Supplementary Fig. 2a-c). Together, this data 153 suggested that a substantial number of genomic loci in melanoma tumors harbor bivalent 154 chromatin states. To determine whether this could be a reflection of tumor heterogeneity, 155 we assessed the presence of these bivalent chromatin states in 10 melanoma short-term 156 cultures (MSTC) and 16 commercially available cell lines from CCLE (Supplemental 157 Table 1), allowing us to eliminate the possibility of signals emerging from other cell types 158 in the tumor microenvironment. Using Model Based Analysis of ChIP-seg (MACS), we 159 identified all potential bivalent combinations by directly overlapping H3K4me3 peaks with 160 either H3K27me3 peaks (bivalent H3K27me3), H3K9me3 peaks (bivalent H3K9me3) or 161 H3K27me3 peaks + H3K9me3 peaks (bivalent H3K4/H3K9/H3K27me3) in each 162 individual sample. Bivalent loci were further identified as "common" in tumors and cell 163 lines if they were present in \sim 50% of the samples in each subgroup (BRAF = 7/13, NRAS) 164 = 2/4, WT = 2/3, MSTC = 5/10 and CCLE = 8/16). In accordance with our chromatin state 165 analysis, NRAS-mutants displayed the greatest number of bivalent H3K27me3 loci out of 166 all the subgroups whereas WT samples displayed the least number of bivalent loci (Fig. 167 **1i** and Supplementary Fig. **2d**). Importantly, both the MSTC and CCLE subgroups also 168 displayed a large number of bivalent H3K27me3 loci that were shared with melanoma 169 tumors (Fig. 1i, j and Supplementary Fig. 2e-g), further suggesting these domains are 170 enriched in cancer cells and are not a product of tumor heterogeneity.

Bivalent domains are lost and gained on key mesenchymal genes in metastaticmelanoma.

174 In embryonic stem cells (ESCs), bivalent promoters mark critical lineage-specific genes 175 which gain or lose these modifications as cells differentiate towards a particular 176 phenotype^{24,25}. Previous studies have demonstrated various cancer-related genes (i.e. 177 CDKN2A) maintain or regain bivalent promoters in normal tissues, however their role in cancer progression has yet to be described²⁶. To this end, we first computed unique 178 179 bivalent loci (H3K4me3 and H3K27me3) in BRAF-mutant, NRAS- mutant and WT 180 samples by overlapping common peaks in each tumor subtype which further suggested 181 NRAS-mutants contained the highest number of bivalent loci (Fig. 2a and Supplementary 182 Fig. **3a-c**). To determine how subtype-specific bivalent polycomb losses and gains 183 influence melanoma progression, we calculated the overlaps of bivalent sites from NRAS-184 and BRAF- mutants to those in primary melanocytes (Fig. 2b, c and Supplementary Fig. 185 **3d**, e). We focused primarily on the comparisons between NRAS- and BRAF-mutants as 186 most significant differences in bivalency were observed to occur between these 187 subgroups (Fig. 1f-i). We posited that removal of H3K27me3 mark from bivalent loci in 188 melanocytes would lead to transcriptionally 'active' loci in melanoma tumors and such loci 189 were termed under "bivalent losses" (Fig. 2d). Similarly, gain of H3K27me3 mark on loci 190 (bivalent in tumors) that harbor only H3K4me3 in melanocytes will lead to transcriptional 191 repression which were termed as "bivalent gains" (Fig. 2d). Determination of the gene 192 targets (within +/- 10KB of each locus) and subsequent pathway enrichment analyses by 193 GSEA MSigDB tool identified critical melanoma-associated "hallmark pathways" within 194 each genetic subgroup (Fig. 2e). For example, in NRAS-mutants, losses of melanocyte-

195 specific bivalency included genes associated with the "epithelial-mesenchymal transition" 196 and "KRAS-signaling up", while gains of tumor-specific bivalency included genes 197 associated with "KRAS-signaling down" and "apical junction" (Fig. 2e and Supplementary 198 Table 2). Furthermore, RNA-seq analysis between NRAS-mutant (n = 81) and BRAF-199 mutant (n = 118) TCGA SKCM tumors confirmed "epithelial-mesenchymal transition" and 200 "KRAS-signaling up" signatures to be upregulated specifically in NRAS-mutant samples 201 (Fig. 2f and Supplementary Table 3), demonstrating an association between shifts in 202 bivalent domains and gene expression patterns within critical melanoma pathways. 203 Indeed, apart from obvious activation of RAS pathway genes, previous studies have 204 demonstrated importance of activation of mesenchymal drivers and phenotypes in 205 invasive behavior of metastatic melanoma and other malignancies²⁷⁻²⁹.

206 In order to identify important metastasis driver genes that are subjected to 207 epigenetic regulation, we focused on genes that do not harbor genetic changes in 208 cancers, which included key EMT transcription factors (EMT-TF) ZEB1, TWIST1, SNAI1 209 and CDH1 (Fig. 2e and Supplementary Fig. 3f). In response to activation of NRAS or 210 BRAF, the EMT-TF network undergoes a reorganization that includes activation of ZEB1 211 and TWIST1 along with the loss of CDH1. This phenomenon is accompanied with 212 increased invasion and correlates with poor prognosis in metastatic melanoma patients²⁸. 213 We observed that ZEB1, TWIST1, and additionally SNAI1 and TGFBI are held in a 214 bivalent state in melanocytes and transition to an active state in metastatic melanoma 215 tumors, which occurs in a subtype specific manner (Fig. 2g and Supplementary Fig. 3g). 216 Importantly, these events are significantly correlated with their mRNA expression levels 217 (Fig. **2h** and Supplementary Fig. **3h**). In addition, genes such and *CDH1* and *CDH3* 218 harbor H3K4me3 in melanocytes and gain repressive bivalency in both NRAS-mutant and 219 BRAF-mutant tumors subtypes (Fig. 2i). This transition to repressive bivalency is 220 associated with their downregulation (Fig. 2). Interestingly, ZEB1 and TWIST1 harbored 221 bivalent chromatin states in embryonic stem cells and germ-layer stem cells, but not in 222 mesenchymal stem cells where they are active (Supplementary Fig. 4a-d). Various other tissues such as breast (TWIST1 and ZEB1^{30,31}), brain (TWIST1 and ZEB1), colon 223 (TWIST1), lung (TWIST1) and ovarian (TWIST1) normal tissues or cell lines 224 225 (Supplementary Fig. 4a-d) showed varying degree of bivalency which correlated well with 226 gene expression patterns demonstrating this "bivalent loss" mechanism can be expanded 227 to various other tissues. In addition to EMT-TF, we observed bivalent shifts on several well-characterized drivers of melanoma metastasis, such as BMI1³², RNF2³³, and CDK6¹⁷ 228 229 (Supplementary Fig. **3i-k**), further strengthening the hypothesis that shifts in bivalent 230 domain can be a key epigenetic event during metastasis.

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232 Melanocyte-specific bivalent genes transition to transcriptionally active broad 233 H3K4me3 in melanoma tumors.

While surveying the transitions of bivalent domains to a transcriptionally active state through the genome, we observed two distinct types of H3K4me3 profiles present in many of the samples. For example, the *TWIST1* domain spanned well beyond the gene body whereas the *ZEB1* domain was localized within the promoter region (Fig. **2g** and Supplemental Fig. **4a**, **b**). In contrast to typical H3K4me3 domains that are usually 200-1000bp long, broad H3K4me3 domains can span thousands of kilobases and have been implicated in various cellular processes, including increased gene expression, enhancer activity and tumor-suppressor gene regulation³⁴⁻³⁶. Hence, we posited that some important driver genes that lose bivalency in melanocytes may retain or gain different types (broad or non-broad) of H3K4me3 domains in melanoma tumors.

244 Hence, we systematically identified broad H3K4me3 domains by computing the 245 overall width and density from MACS2 broad peaks in BRAF-mutant, NRAS-mutant and 246 WT tumors (Fig. **3a-c**). Globally, NRAS-mutant and BRAF-mutant subtypes harbored the largest number of broad H3K4me3 peaks, in some cases spanning >30kb in NRAS-247 248 tumors (Fig. 3b). Similarly, the top 1% broadest domains in these subtypes extended well 249 beyond the typical H3K4me3 domain (200-1000bp), with peaks reaching over 4kb in 250 10/17 the individual samples profiled (Fig. 3d). In contrast, WT samples harbored much 251 shorter H3K4me3 domains with the widest peaks spanning around 15kb (Fig. 3c). Here, 252 the top 1% broadest peaks did not markedly extend beyond the typical H3K4me3 domain, 253 with only 1/3 samples harboring peaks over 4kb at this percentage cutoff (Fig. 3d). Based 254 on these observations and previous reports, broad domains were defined as peaks that 255 extended at least 4x (> 4kb; merge size = 1kb) beyond that of typical H3K4me3 domain 256 (Fig. 3d and Supplementary Fig. 5a). Using this method, we identified two distinct types 257 of H3K4me3 in melanoma tumors including broad domains (> 4kb) spanning outside of 258 the TSS (mean length = 12.3kb) and non-broad domains (< 4kb) localized within the 259 promoter region (mean length = 2.2kb) (Fig. **3e-f** and Supplementary Fig. **5b**). GSEA 260 analysis suggested occurrence of broad domains on genes regulating important biological 261 processes implicated in melanoma metastasis and tumorigenesis, such as "TNFA signaling via NFKB" ³⁷, "WNT signaling" ^{38,39}, and Hedgehog signaling" ⁴⁰⁻⁴² (Fig. **3g-i** and 262 263 Supplemental Table 4), further suggesting a role for broad domains in melanoma

metastasis. Additionally, important melanoma drivers such as NEAT1, MALAT1, MYC 264 265 and EVX1 harbored broad H3K4me3 domains in and around their gene bodies (Fig. 3i). 266 To identify the subsets of melanocyte-specific bivalent genes that transition to 267 transcriptionally active broad domains (and lose H3K27me3 in tumors), we further 268 overlapped genes that harbor 1) bivalent domains uniquely in melanocytes (but not in 269 tumors), 2) tumor-specific broad or non-broad H3K4me3 domains, 3) active transcription 270 mark H3K79me2 and 4) gene expression [using RNA-seq data from NRAS (n=81), BRAF 271 (n=118) and WT (n=38) metastatic samples]. Integrative analysis revealed NRAS-specific 272 broad domains were associated with increased gene expression (Fig. 4a,b and 273 Supplemental Table 5) and enriched for melanoma pathways such as "UV response up", 274 "KRAS signaling up" and "Glycolysis" (Fig. 4c). Genes displaying the transition to 275 transcriptionally active broad H3K3me3 included additional metastatic drivers known to 276 function in the switch to an mesenchymal/invasive state, including SOX9, POU3F2, 277 PDGFRA and MYCN (3.7kb width) (Fig. 4d, f), While we also identified a broad 278 H3K4me3-associated increase of transcription in WT samples, this occurred on a 279 markedly lower number of genes many of which were shared with NRAS samples (Fig. 280 **4a**, **b**). In contrast to broad domains, we identified a relatively large and constant number 281 of genes transitioning from a bivalent state to non-broad H3K4me3 in melanoma tumors, 282 many of which were present within all mutational subtypes (Fig. 4a). On a global level, 283 we did not observe a change in the mean expression levels of melanocyte-specific 284 bivalent genes that transitioned to non-broad H3K4me3 domains in any melanoma 285 subtype (Fig. 4b), however, gene expression changes in specific genes were noted. 286 These included various genes known to be misregulated upon the aberrant activation of 287 RAS/RAF signaling, including *VCAN* as well as various RAS-effectors such as 288 *RASGEF1B*, *RASEF*, *RASGRP1*, *RAPGEF5*, and *ARHGAP27* (Fig. **4e**, **g** and 289 Supplementary Fig. **5c-e**), Overall, these results suggest that melanocyte-specific 290 bivalent losses that transition to broad H3K4me3 domains in NRAS-mutant tumors 291 regulate/increase the expression of mesenchymal/invasive drivers during metastatic 292 progression.

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Broad H3K4me3 domain spreading is associated with increased transcriptional activation.

296 Our results suggest that during the transition from a bivalent to an active state, genes that 297 retain broad H3K4me3 domains are associated with increased transcriptional activation 298 compared to non-broad domains in melanoma tumors (Fig. 4a-c). However, this was only 299 observed on a small subset of genes (NRAS = 97; Supplementary Table 5). We 300 considered that another mechanism of gene activation could be spreading of the 301 H3K4me3 signal, while gene repression may be associated with shortening of the broad 302 H3K4me3 domains. Indeed, a previous report demonstrated that H3K4me3 domain 303 shortening is associated with decreased gene expression of tumor suppressor genes in lung and liver cancers³⁶. Therefore, we investigated whether the size of H3K4me3 304 305 domains are altered in metastatic melanoma. First, on a genome-wide level, we observed 306 preferential shortening (< 2kb) of all H3K4me3 peaks in melanoma tumors relative to 307 melanocytes (Fig. 5a). Focusing on the promoter-associated broad H3K4me3 peaks (-308 /+10kb TSS), we further observed preferential shortening (< 2kb) of these peaks in 309 melanoma tumors which was associated with a decrease in gene expression levels (Fig.

5b-d and Supplemental Table **6**). Promoters harboring some of the broadest H3K4me3 310 311 domains displayed marked shortening on critical components of the melanocyte 312 regulatory network in melanoma tumors, including PMEL (aka GP100), MITF, ZEB2 and 313 TFAP2A⁴³ (Fig. 5e and Supplementary Fig. 6a, b). Because genes such as MITF and 314 ZEB2 are differentially expressed in alternate phenotypic states, displaying high 315 expression in melanocytes (or proliferative phenotype) and low expression in melanoma 316 tumors^{28,44} (Fig. **5f** and Supplementary Fig. **6c**), this finding illustrates a new epigenetic 317 mechanism for their misregulation in metastatic melanoma. Similar to that of bivalent 318 losses, we also identified a smaller subset of promoters that gained broad H3K4me3 in 319 melanoma tumors. The top promoters displaying a broad H3K4me3 transition (> 10kb, 320 29 genes) were enriched for critical developmental regulators including DLX1/2, TBX3, 321 HMX2/3 and LBX1, many of which were also upregulated in metastatic melanoma tumors 322 and hence suggest potential oncogenic roles for these proteins (Fig. 5g, h and 323 Supplementary Fig. 6d, e). Together, these results suggest broad H3K4me3 domains are 324 associated with higher expression that decreases upon shortening on key melanocyte-325 specific promoters during the transition to metastatic melanoma.

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Broad H3K27me3 domains display preferential lengthening in metastatic
 melanoma.

Another possible transition of bivalent domains is the loss of H3K4me3 and retention or extension of H3K27me3 domains. Therefore, we identified H3K27me3 domains by computing overall width and density from MACS2 broad peaks in BRAF-mutant, NRASmutant and WT samples (Supplementary Fig. **7a-c**). In all subtypes, we observed

333 exceptionally wide H3K27me3 domains spanning tens of thousands of kilobases with 334 multiple peaks extending well over 100kb in BRAF- and NRAS-mutant samples 335 (Supplementary Fig. 7a-c). Globally, the top 1% broadest H3K27me3 domains in 14/20 336 tumors extended well over 10kb (Fig. **6a** and Supplementary Fig. **7e**; merge size = 1kb). 337 In contrast to the results for H3K4me3, we observed preferential lengthening of 338 H3K27me3 domains on a global level (> 2kb) in BRAF-, NRAS- mutants and WT samples 339 (Fig. 6b). However, since H3K27me3 is already known as a broad mark, we focused on 340 promoter regions that displayed large extensions or constrictions (> 10kb) of H3K27me3 341 domains. Although we observed relatively consistent number of broad (\geq 4kb) 342 H3K27me3-associated promoters across mutational subtypes (Fig. 6c), each subtype 343 displayed a different pattern of H3K27me3 in comparison to melanocytes, with NRAS-344 and BRAF- mutant samples displaying preferential lengthening and WT preferential 345 shortening (Fig. 6d). Importantly, melanocytes harbored much shorter broad H3K27me3 346 domains (~50kb) compared to all tumors (Supplemental Fig. 7d), While the largest 347 constriction of H3K27me3 was just over 50kb, the largest extension spanned over 100kb 348 in BRAF- and 85kb in NRAS- mutant samples (Supplemental Table 7), further indicating 349 polycomb domains are exceptionally broad in melanoma tumors. Overall, the widest 350 H3K27me3 domain lengthening observed was on the HOXC locus spanning 102.8kb in 351 BRAF samples (Fig. 6e). Interestingly, as 3' HOX genes are known to be expressed first 352 during normal development, we observed H3K27me3 spreading specifically towards the 353 3' end of the HOXC cluster in melanoma tumors. This 5' -> 3' H3K27me3 spreading was 354 also observed on other HOX cluster genes (HOXB/HOXD) and further associated with 355 the downregulation of various 3' members relative to melanocytes (Fig. 6f-h and

356	Supplementary Fig. 7f, g). Together, this exceptionally wide H3K27me3 spreading
357	illustrates a new epigenetic signature for gene silencing of critical regulatory genes in
358	metastatic melanoma.
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364 **Discussion**

365 Our study describes an epigenetic program for reorganization of an interconnected 366 regulatory network encompassing EMT-TF, melanocyte master regulators and 367 mesenchymal/invasive genes as melanocytes progress to metastatic melanoma. Our 368 work provides three major advances regarding the role of the epigenome in melanoma 369 metastasis: 1) Specific genetic events such as BRAF or NRAS mutations may utilize 370 specific chromatin states to bring about transcriptional changes unique that genotype. 2) 371 Chromatin state switches involving bivalent domains mark master regulatory genes 372 associated with a metastatic phenotype, including the epithelial-to-mesenchymal 373 transition. 3) In addition to bivalent domain switches, contraction or extension of broad 374 H3K4me3 and H3K27me3 domains may be a key driver in the regulation of pro-metastatic 375 genes. In addition to these findings, the current study encompasses the most 376 comprehensive dataset from a large number of well-annotated samples which can help 377 the community better understand the epigenetic circuitry governing melanoma 378 metastasis.

379 Our results suggest that assimilation of epigenetic traits may be critical to evolution of metastatic clones, consistent with some prior studies^{17,45}. It has been argued that 380 381 evolution of cancer cells through metastasis entails constant acquisition and switching of 382 cellular phenotypes during its journey from the primary site to colonization in a distant organ⁴⁶. Hence, the molecular processes governing metastasis may mirror those driving 383 384 tissue differentiation during normal development. At the same time, those molecular traits 385 that allow establishment/evolution of neo-phenotypes, such as the ones in play during 386 selection of *crossweinless* phenotype observed by Waddington many decades ago⁴⁷,

387 may also play an important driver role in evolution of cells undergoing metastasis. 388 Waddington's "genetic assimilation" model suggested that specific traits (genetic or 389 epigenetic) driving a cell towards a specific differentiation state in response to certain 390 environmental pressure are assimilated in the genome (or epigenome) during evolution. 391 Similarly, we propose that during evolution of cancer cells to metastasis, under selective 392 environmental pressure specific cell clones assimilate epigenetic traits – such as shifts in 393 bivalent and broad domains – in their genome which eventually help them disseminate 394 and colonize distant organs.

395 In ESCs, bivalent promoters mark critical lineage-specific genes which gain or lose these modifications as cells differentiate towards a particular phenotype^{24,25}. Moreover, 396 397 various cancer-related genes maintain or regain bivalent promoters in normal tissues²⁶. 398 Here, we expand on those studies by demonstrating that bivalent domains are lost and 399 gained on critical EMT-TF, melanoma-specific drivers and mesenchymal genes as 400 melanocytes progress to metastatic melanoma. Shifts from bivalent states to active or 401 repressive marks are likely the least energy-intensive route to gene activation or 402 repression respectively (https://doi.org/10.1101/456525). Since metastasis involves 403 several cell fate transitions, which likely employ transcriptional circuitry consisting of 404 multiple genes, energetically the use of shifts involving bivalent chromatin states would 405 be favored as the preferred mode of gene regulation. It can be suggested that this specific 406 epigenetic process could constitute one of the primary 'forces' that can help with the 407 'canalization' (per Waddington model) of the pro-metastatic clones.

408 Interestingly, our study for the first time suggests that melanomas harbor 409 exceptionally wide H3K4me3 domains on key drivers of invasion and metastasis (POU3F2^{48,43}, SOX9⁴⁹, PDGFRA⁵⁰ and PDGFA) and this transition was associated with increased transcriptional activation. It is likely, barring physical and chemical barrier, that addition of H3K4me3 marks on adjacent nucleosome may require less energy due to already recruited methylation machinery, and hence may be a preferred mode for relative increase in gene expression to meet the needs of metastatic cells. The observations of broad domains on critical components of the melanocyte regulatory network, including MITF^{51,52}, ZEB2^{28,44}, TFAP2A⁴³ and PMEL (aka GP100), as well as other genes not known to be previously associated with melanoma, including MYCN, KLF6, NR2F2 and CPEB2, may allow for identification of pro-metastatic drivers whose expression is mediated by epigenetic events. Overall, our work identifies new principles of epigenome regulation in melanoma metastasis and strengthens the hypothesis that metastatic dissemination is likely driven by specific epigenetic events.

431 Author contributions

432 CT and KR conceptualized and designed the study; CT and MM generated the ChIP-seq 433 data; CT, MT, AR, JS and SA processed and performed data analysis for ChIP-seq; CT, 434 MT and AR processed and performed data analysis for RNA-seq; JO, CC, CJW, CB and 435 SW generated and characterized the Short Term Culture datasets; EO, SS, KT, DZ and 436 KC provided intellectual input; CT and KR wrote and prepared the manuscript. All authors 437 read and approved the manuscript.

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454 Methods

455 **Collection of melanoma tumor samples**

456 Melanoma tumors were obtained from the Melcore tumor bank at MD Anderson Cancer

457 Center.

458 **Characterization of melanoma short term cultures**

459 Melanoma short term cultures were generated from metastatic tumor specimens as part 460 of the Adoptive T-cell Therapy Clinical Program at the University of Texas, MD Anderson 461 Cancer Center (LAB06-0755 and 2004-0069), as previously described^{53,54,55}). Briefly a 462 tumor specimen from metastatic tumor was collected and incubated with an enzymatic 463 digestion cocktail (0.375% collagenase type I, 75 µg/ml hyaluronidase and 250 U/mL DNAse I) in tumor digestion medium (RPMI, 10 mM of HEPES, 1x Penicillin-Streptomycin 464 and 20 ug/mL of Gentamicin; Gibco/Invitrogen) in a humidified incubator at 37°C with 5% 465 CO₂ and with a gentle rotation for 2-3h to obtain a single cell suspension. The tumor 466 467 digest was filtered through a 70-µm filter, washed, and re-suspended in a serum free 468 media, which was then plated in one well of a 6-well culture plate and incubated at 37 °C. 469 After 24h, the media was replaced with fresh tumor media, comprised of RPMI with 470 GlutaMAX, 10% FBS, Penicillin/Streptomycin, Gentamicin, β -mercaptoethanol (50 uM, Gibco/Invitrogen), HEPES (10 mM), 471 and insulin-selenium-transferin (5 ug/ml, 472 Gibco/Invitrogen). Cells were grown in enriched DMEM/F12 culture media 473 (Gibco/Invitrogen) supplemented with all growth factors including 10%FBS, sodium 474 pyruvate (1mM), insulin-selenium-transferin, MycoZap-PR (Lonza), HEPES (10mM) and 475 β-Mercaptoethanol. Once enough cells were grown, the purity of the tumor was tested 476 using a melanoma tumor surface marker (MCSP-1) by flow cytometry. Cultures were

477 deemed established when the cells stained positive for a melanoma tumor marker 478 (MCSP-1) and negative for a fibroblast marker (CD90). Appropriate serum starving was 479 performed to eliminate fibroblasts. All cell lines were tested for mycoplasma using 480 MycoAlert detection kit (Lonza), and fingerprinted by STR fingerprinting, and validated by 481 comparing with matched blood samples. A few passages after a pure tumor line was 482 established, the cells were cryopreserved and kept in stocks in liquid nitrogen until use.

483

484 **Chromatin immunoprecipitation assays**

485 ChIP assays were performed as described previously⁵⁶. Briefly, $\sim 2 \times 107$ cells were 486 harvested via cross-linking with 1% (wt/ vol) formaldehyde for 10 min at 37 °C with shaking. After guenching with 150 mM glycine for 10 min at 37 °C with shaking, cells were 487 washed twice with ice-cold PBS and frozen at -80 °C for further processing. Cross-linked 488 489 pellets were thawed and lysed on ice for 30 min in ChIP harvest buffer (12 mM Tris-Cl, 1 490 × PBS, 6 mM EDTA, 0.5% SDS) with protease inhibitors (Sigma). Lysed cells were 491 sonicated with a Bioruptor (Diagenode) to obtain chromatin fragments (~200–500 bp) and 492 centrifuged at 15,000 × g for 15 min to obtain a soluble chromatin fraction. In parallel with 493 cellular lysis and sonication, antibodies (5 μ g/3 × 106 cells) were coupled with 30 μ l of magnetic protein G beads in binding/blocking buffer (PBS + 0.1% Tween + 0.2% BSA) 494 495 for 2 h at 4 °C with rotation. Soluble chromatin was diluted five times using ChIP dilution 496 buffer (10 mM Tris-Cl, 140 mM NaCl, 0.1% DOC, 1% Triton X, 1 mM EDTA) with protease 497 inhibitors and added to the antibody-coupled beads with rotation at 4 °C overnight. After 498 washing, samples were treated with elution buffer (10 mM Tris-Cl, pH 8.0, 5 mM EDTA, 499 300 mM NaCl, 0.5% SDS), RNase A, and Proteinase K, and cross-links were reversed

500 overnight at 37. Immune complexes were then washed five times with cold RIPA buffer 501 (10mM Tris-HCl, pH 8.0, 1mM EDTA, pH 8.0, 140mM NaCl, 1% Triton X-100, 0.1% SDS, 502 0.1% DOC), twice with cold high-salt RIPA buffer (10mM Tris-HCl, pH 8.0, 1mM EDTA, pH 8.0, 500mM NaCl, 1% Triton X-100, 0.1% SDS, 0.1% DOC), and twice with cold LiCl 503 504 buffer (10mM Tris-HCl, pH 8.0, 1mM EDTA, pH 8.0, 250mM LiCl, 0.5% NP-40, 0.5% 505 DOC). ChIP DNA was purified using AMPure XP beads (Agencourt) and quantified using 506 the Qubit 2000 (Invitrogen) and Bioanalyzer 1000 (Agilent). Libraries for Illumina 507 sequencing were generated following the New England BioLabs (NEB) Next Ultra DNA 508 Library Prep Kit protocol. A total of 10 cycles were used during PCR amplification for the 509 generation of all ChIP-seg libraries. Amplified ChIP DNA was purified using double-sided 510 AMPure XP to retain fragments (~200–500 bp) and guantified using the Qubit 2000 and 511 Bioanalyzer 1000 before multiplexing.

512

513 ChIP-seq data processing

514 Raw fastg reads for all ChIP-seg experiments were processed using a snakemake based 515 pipeline⁵⁷. Briefly, raw reads were first processed using FastQC 516 (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and uniquely mapped reads were aligned to the hg19 reference genome using Bowtie version 1.1.2⁵⁸. Duplicate reads 517 were removed using SAMBLASTER ⁵⁹ before compression to bam files. To directly 518 519 compare ChIP-seg samples uniquely mapped reads for each mark were downsampled 520 per condition to 18 million, sorted and indexed using samtools version 1.5⁶⁰. To visualize ChIP-seq libraries on the IGV genome browser, we used deepTools version 2.4.0⁶¹ to 521 522 generate bigWig files by scaling the bam files to reads per kilobase per million (RPKM).

523 Super ChIP-seq tracks were generated by merging bam files from each cancer type,

524 sorting and indexing using samtools and scaling to RPKM using deepTools.

525

526 Chromatin state calls

ChromHMM ⁶² was used to identify combinatorial chromatin state patterns based on the 527 528 histone modifications studied. Normalized bam files were converted to bed files and 529 binarized at a 1000bp resolution using the BinarizeBed command. We specified that 530 ChromHMM should learn a model based on 18 chromatin states. As we considered 531 models between 8 and 30 chromatin states, we chose an 18-state model because it is 532 large enough to identify important functional elements while still being small enough to 533 interpret easily. To determine chromatin state differences between different groups we 534 used a two-step process. First, using the segmentation calls from the ChromHMM output 535 the entire genome is divided into non-overlapping windows of 10 Kb. We next count the number of times a chromatin state is observed in each of the 10 Kb windows and obtain 536 537 a frequency matrix for each state in the ChromHMM model (E1-E18). In the second step, 538 low variable genomic loci are removed from the frequency matrix and significant 539 differences between two groups of samples types are calculated by using a 540 nonparametric Mann Whitney Wilcoxon test with a P-value < 0.05 for each state 541 separately.

542

543 **Correlation of copy number from ChIPseq and SNP array.**

544 SKCM TCGA copynumber data were downloaded by TCGAbiolinks⁶³. Copy number 545 analysis for ChIPseq was carried out using copywriteR⁶⁴, which uses off-target reads for accurate copy number detection. The ChIP-seq input files were used, which represent low-pass whole genome sequencing data sets. Bin size of 200kb was used for analysis. The copy number of each gene was determined by overlapping the genes with the segmentation files from ChIPseq and SNP array, respectively. The Pearson correlation of the copy number of all genes among samples was calculated and plotted in heatmap by ComplexHeatmap.

552

553 Correlation of RNA expression and chromatin state data

554 SKCM TCGA RNAseq data were downloaded by TCGAbiolinks⁶³. TPM (transcript per 555 million) value was calculated from raw counts by scaling to the gene length first and then 556 the library size. For each gene in a sample, the transcription start site is overlapped with 557 the chromatin state segmentation file to determine the state of that gene. The expression 558 values (TPM) for all genes and all samples are then combined and spilt by categories 559 (18) of chromatin states. A box plot is plotted for each chromatin state.

560

561 **Correlation of DNA methylation and chromatin state data**

562 SKCM TCGA 450k DNA methylation data were downloaded by TCGAbiolinks⁶³. For each 563 sample and each chromatin state segmentation bin, the average of beta values from the 564 DNA methylation data were calculated for each bin. The average values of all bins from 565 all samples are combined and then split by categories (18) of chromatin states. A box plot 566 is plotted for each chromatin state.

567

568 Identification and visualization of ChIP-seq binding sites

We used Model-based analysis of ChIP-seq (MACS) version 1.4.2⁶⁵ peak calling 569 570 algorithm to identify H3K4me3 (p-value of 1e-5) and MACS version 2.1.0 to identify 571 H3K27me3 (p-value of 1e-5) enrichment over "input" background. Bivalent sites were 572 identified by overlapping H3K4me3 with H3K27me3 or H3K9me3 by a minimum of 1bp using intersectBed⁶⁶. Bivalent polycomb-heterochromatin regions were identified by 573 574 overlapping the H3K4me3+H3K27me3 output with H3K9me3 by a minimum of 1bp. 575 Common sites (NRAS = 2/4, BRAF = 7/13, WT = 2/3, MSTC = 5/10, CCLE = 8/16 and Melanocytes = 2/2) were identified using bedops⁶⁷ with the following command; cat *bed 576 577 | sort-bed - | bedmap --count --echo --delim '\t' - | unig | awk '\$1 >=x' | cut -f2- > 578 samplename common.bed. Final peaksets used for downstream analysis were 579 generated using mergeBed. Unique BRAF, NRAS, WT and melanocyte peaks for bivalent 580 polycomb, bivalent heterochromatin and bivalent polycomb-heterochromain were 581 identified using the concatenate, cluster and subtract tools from the Galaxy/Cistrome web 582 based platform ⁶⁸. To identify sites that were bivalent in melanocytes but active in 583 melanoma tumors, and visa-versa, common H3K27me3 sites were subtracted from the 584 opposing bivalent site in each comparison.

585

586 Identification of broad H3K4me3 and H3K27me3 domains

Broad H3K4me3 domains were identified using MACS2.1.0 with the broad setting (pvalue of 1e-5) followed by merging adjacent peaks within 1kb using mergeBed^{66,69}. We determined the optimal distance to merge adjacent peaks based on the number of broad H3K4me3 and H3K27me3 domains at distance thresholds between 1kb through 10kb in each mutational subtype. Broad H3K4me3 domains were further classified within the top 592 1% of the broadest peaks that extended at least $4x (\geq 4kb)$ beyond that of a typical 593 H3K4me3 domain (0.2kb-1kb). Final peaksets for common sites broad and non-broad 594 H3K4me3 and H3K27me3 domains were defined as peaks present in ~50% of each 595 mutational subtype and melanocytes (NRAS = 2/4, BRAF = 7/13, WT = 2/3 and 596 Melanocytes = 2/2) as described in identification and visualization of ChIP-seq binding 597 sites.

598

599 Assigning binding sites to genes

A list of Ensembl genes was obtained from the UCSC Table browser (<u>http://genome.ucsc.edu/</u>). Proximal promoters were defined as ± 10 kb from the transcription start site (TSS) and genic regions were identified as ± 10 kbTSS to the transcription end site (TES). Peaks were assigned to genes if they overlapped the promoter or genic region by a minimum of 1bp using intersectBed. Gene body heatmaps and average density plots were generated using ngs.plot⁷⁰.

606

607 Gene set enrichment analysis

608 Gene Set Enrichment Analysis (GSEA) was performed using GSEA/MSigDB⁶⁹ 609 HALLMARK and KEGG pathways based on ensemble gene lists from peaks within -10kb-610 TES for bivalent domains and ±10kbTSS for broad H3K4me3 domains. All pathways are 611 significant based on FDR q-value.

612

613 RNA-sequencing data processing

614 obtained from TCGAbiolinks⁶³ For TCGA data. raw counts were R in 615 (http://www.rstudio.com/).and transformed to TPM. For Roadmap data, raw counts were 616 obtained from http://www.roadmapepigenomics.org/ and transformed to TPM. For all 617 RNA-seq boxplots, gene expression values from both datasets were normalized using quantile normalization function in R. For identification of differentially expressed genes 618 619 and GSEA analysis between NRAS and BRAF melanoma tumors, raw counts were obtained from TCGAbiolinks. DESeq2⁷¹ was employed for normalization of differential 620 gene expression analysis. GSEA⁶⁹ was run on a ranked gene list using HALLMARK gene 621 622 sets with default settings. 623 Data and software availability 624 The ChIP and mRNA sequencing data have been deposited in the NCBI GEO BioProject 625 database with the following accession number GSE134043. 626 627 628 629 630 631 632 633 634 635 636

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Figure 1

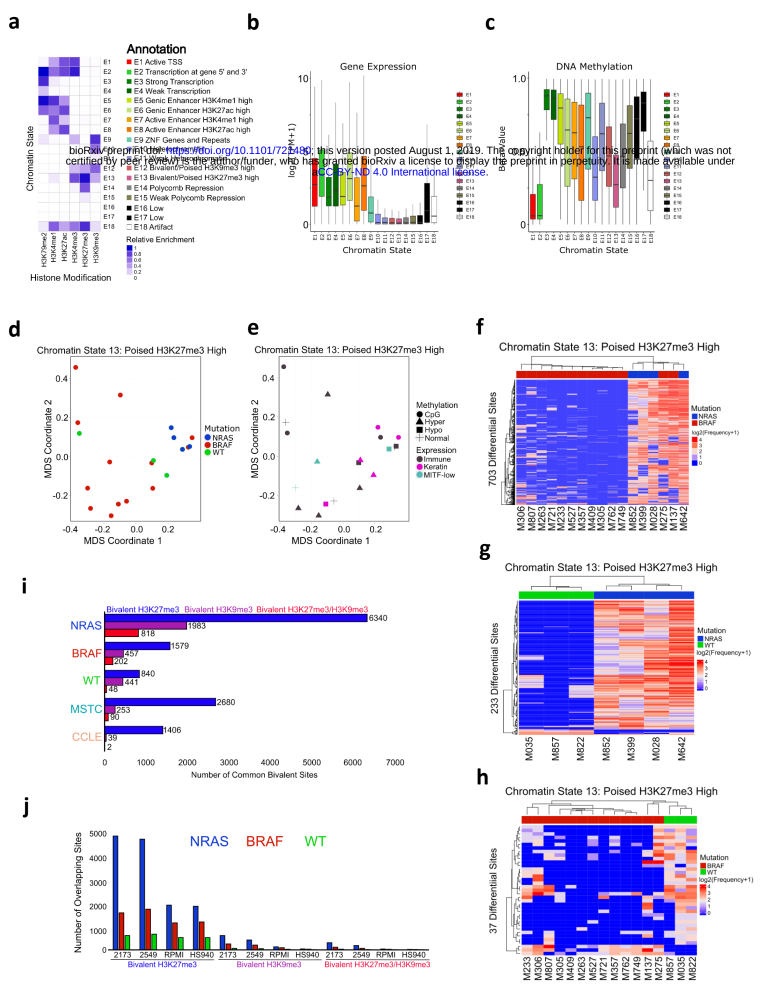


Figure 1: Bivalent chromatin domains distinguish metastatic melanoma tumors based on mutational subtypes

a) Combinatorial chromatin state definitions and histone mark probabilities identified in 20 metastatic melanoma tumor samples using the ChromHMM algorithm.

b) Boxplot illustrating mean expression of genes from RNA-seq based on genomic regions overlapping with each chromatin state.

c) Boxplot illustrating mean DNA methylation levels based on genomic regions overlapping with each chromatin state.

d) MDS analysis of chromatin state E13 (poised H3K27me3 high) annotated by mutation (NRAS, BRAF, WT), (e) RNA-expression (Immune, Keratin, MITF-low) and DNA methylation (Normal, CpG, Hyper, Hypo) classifications from The Cancer Genome Atlas.

f) Heatmap displaying differentially regulated regions (FDR < 0.05) of chromatin state E13 (poised H3K27me3 high) between NRAS and BRAF, (g) NRAS and WT and (h) BRAF and WT tumor subtypes.

i) Common bivalent H3K27me3 (H3K4me3+H3K27me3), bivalent H3K9me3 (H3K4me3+H3K9me3) and bivalent H3K27/H3K9me3

(H3K4me3+H3K27me3+H3K9me3) binding sites in melanoma tumor subtypes, MSTC and CCLE lines. Binding sites were identified as common if they were present in at ~50% of the samples (NRAS = 2/4, BRAF = 7/13, WT = 2/3, MSTC = 5/10 and CCLE = 8/16).

j) Co-occupancy analysis of common bivalent binding sites in melanoma tumor subtypes directly overlapping bivalent binding sites in representative MSTC or CCLE lines.



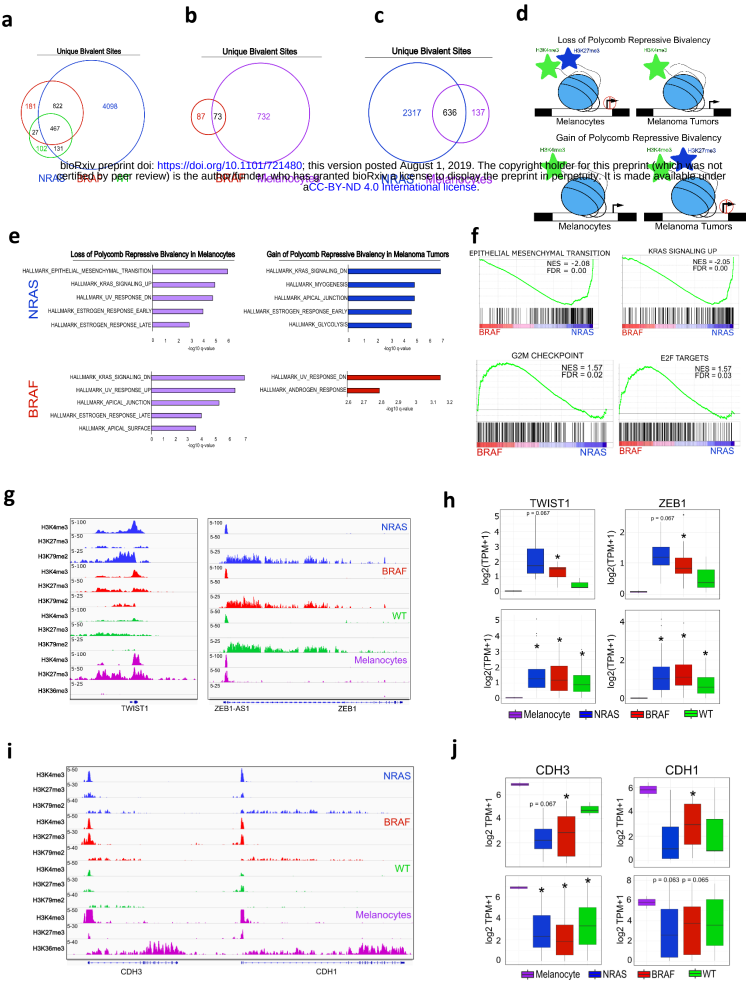


Figure 2: Bivalent domains are lost and gained on key mesenchymal genes in metastatic melanoma.

a) Venn diagram analysis of bivalent H3K27me3 sites in NRAS, BRAF and WT tumor subtypes.

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b, c) Venn diagram analysis of unique bivalent H3K27me3 sites in NRAS and BRAF tumor subtypes overlapping bivalent polycomb sites in primary melanocytes from Roadmap.

d) Schematic of bivalent H3K27me3 losses and gains in melanocytes vs melanoma tumors.

e) Top 5 significant MSigDB/GSEA HALLMARK pathways based on bivalent H3K27me3 sites that are lost and gained within -/+10kbTSS-TES of a gene in NRAS and BRAF tumor subtypes.

f) Significant GSEA HALLMARK pathways based on differentially expressed genes between NRAS (n = 81) vs BRAF (n = 118) metastatic melanoma tumor samples from TCGA.

g) Genome browser view of ChIP-seq tracks for H3K4me3, H3K27me3 and active transcription (H3K79me2/H3K36me3) on the TWIST1 and ZEB1 genes in melanocytes and melanoma tumor subtypes.

h) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the TWIST1 and ZEB1 genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom). Asterisk denotes p-value < 0.05. i) Genome browser view of ChIP-seq tracks for H3K4me3, H3K27me3 and active transcription (H3K79me2/H3K36me3) on the CDH3 and CDH1 genes in melanocytes and melanoma tumor subtypes.

j) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the CDH3 and CDH1 genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom). Asterisk denotes p-value < 0.05.

Figure 3

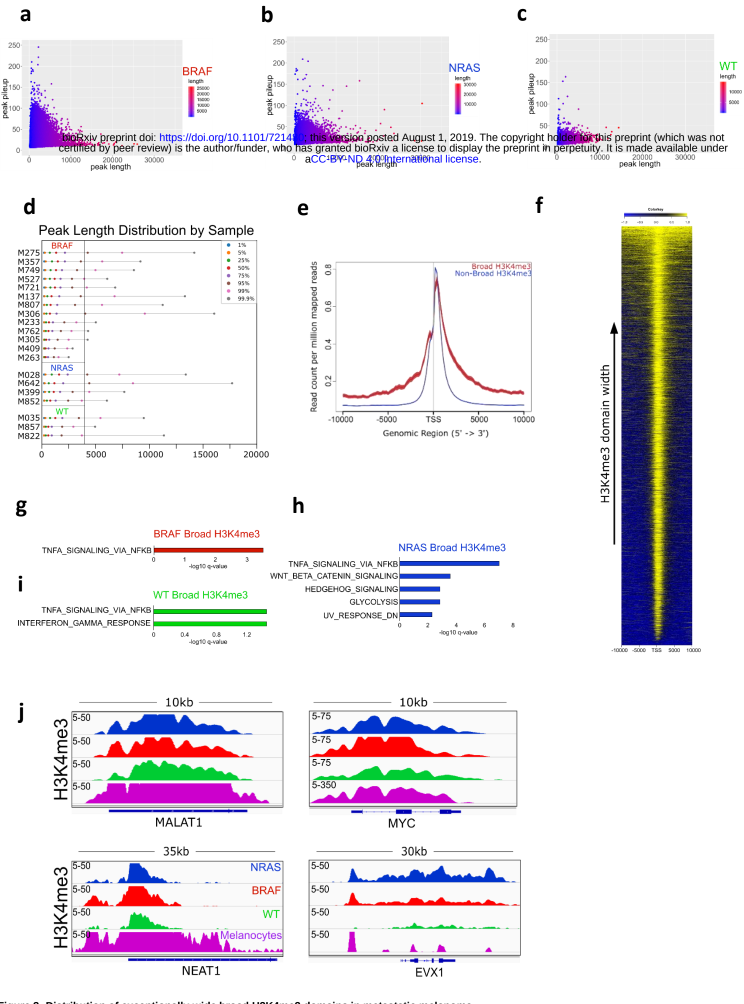


Figure 3: Distribution of exceptionally wide broad H3K4me3 domains in metastatic melanoma

a) Scatterplots of peak width (x-axis) and height (y-axis) from MACS2 broad peak calls (p-value 1e-5) for H3K4me3 in BRAF mutant tumors.

b) Scatterplots of peak width (x-axis) and height (y-axis) from MACS2 broad peak calls (p-value 1e-5) for H3K4me3 in NRAS mutant tumors.

c) Scatterplots of peak width (x-axis) and height (y-axis) from MACS2 broad peak calls (p-value 1e-5) for H3K4me3 in WT tumors.

d) Quartile plot displaying peak length distribution based on percentage of broadest H3K4me3 domains in melanoma tumors. Black line denotes 4kb peak length.

e) Average density profile displaying broad (>4kb) and non-broad (<4kb) H3K4me3 domains at 10kb to +10kb around TSS.

f) Heatmap of H3K4me3 signal sorted by width at -10kb to +10kb around transcription start sites (TSS).

g) Top significant MSigDB/GSEA HALLMARK pathways based on broad H3K4me3 domains within -/+10kbTSS of a gene in BRAF-mutant tumors.

h) Top significant MSigDB/GSEA HALLMARK pathways based on broad H3K4me3 domains within -/+10kbTSS of a gene in NRAS-mutant tumors.

i) Top significant MSigDB/GSEA HALLMARK pathways based on broad H3K4me3 domains within -/+10kbTSS of a gene in WT tumors.

j) Genome browser view of ChIP-seq tracks for H3K4me3 on the MALAT1, NEAT1, MYC and EVX1 genes in melanocytes and melanoma mutational subtypes.

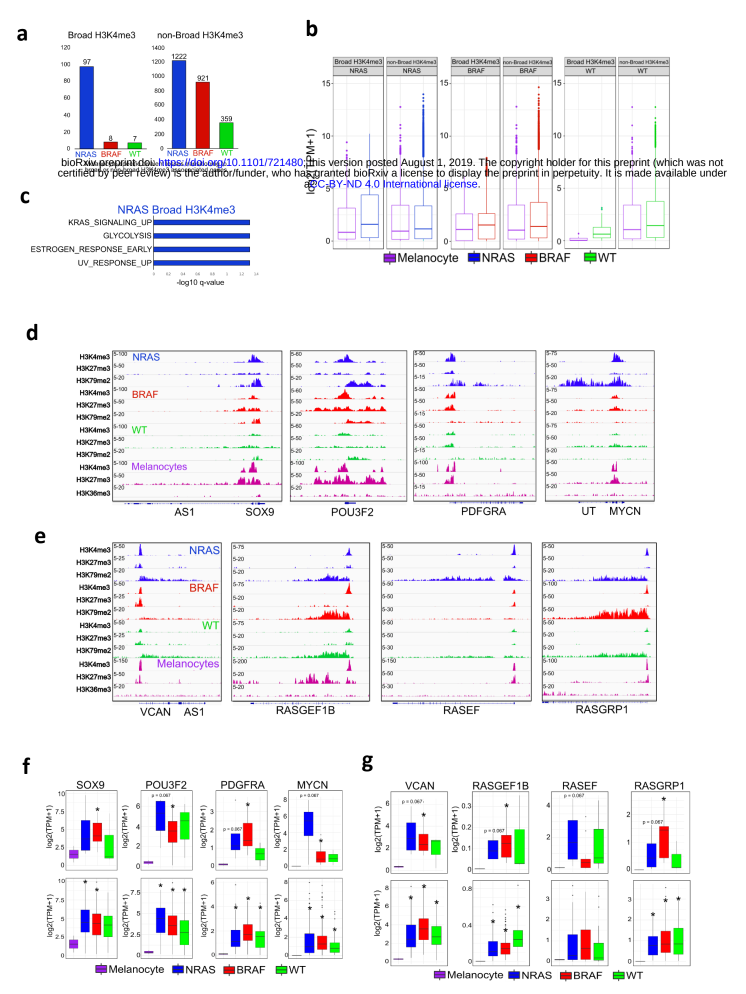


Figure 4: Melanocyte-specific bivalent genes transition to transcriptionally active broad H3K4me3 in NRAS mutant subtypes

a) Barplot of bivalent H3K27me3 genes (-/+10kbTSS-TES) that are lost in melanocytes and associated with broad or non-broad H3K4me3 in NRAS, BRAF and WT tumor subtypes.

b) Barplot displaying quantile normalized mean RNA-expression profiles of NRAS (n=81), BRAF (n=118) and WT (n = 38) tumor subtypes based on broad or non-broad H3K4me3 genes identified in (a).
 c) Top significant GSEA HALLMARK pathways based on bivalent H3K27me3 genes (-/+10kbTSS-TES) that are lost in melanocytes and associated with

d) Genome browser views of ChIP-seq tracks for H3K4me3, H3K27me3 and active transcription (H3K79me2/H3K36me3) on the SOX9, POU3F2,

d) Genome browser views of ChIP-seq tracks for H3K4me3, H3K27me3 and active transcription (H3K79me2/H3K36me3) on the SOX9, POU3F2, PDGFRA and MYCN genes in melanocytes and melanoma tumor subtypes.

e) Genome browser views of ChIP-seq tracks for H3K4me3, H3K27me3 and active transcription (H3K79me2/H3K36me3) on the VCAN, RASGEF1B, RASEF and RASGRP1 genes in melanocytes and melanoma tumor subtypes.

f) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the SOX9, POU3F2, PDGFRA and MYCN genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom). Asterisk denotes p-value < 0.05.
 g) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the VCAN, RASGEF1B, RASEF and RASGRP1 genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom).

Figure 5

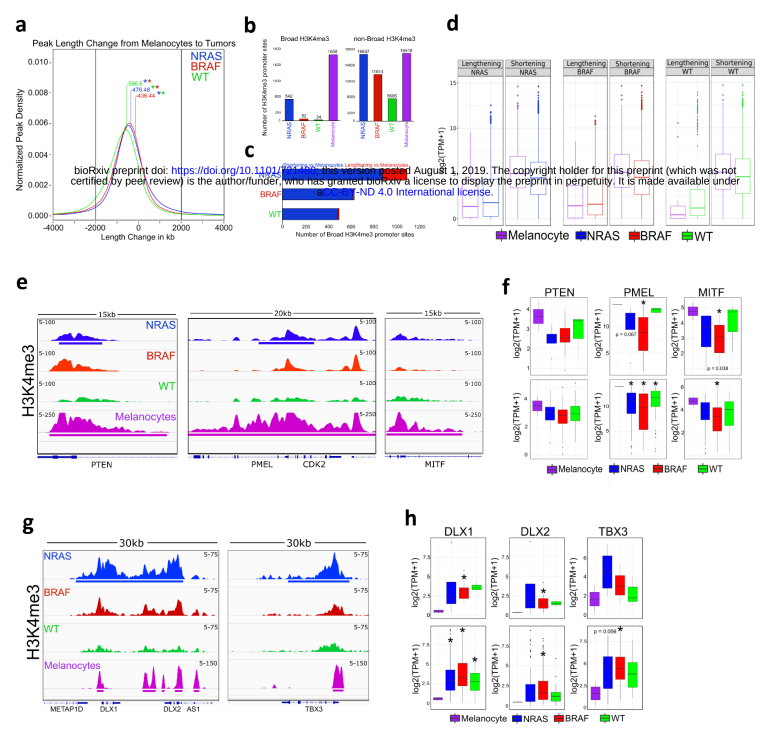


Figure 5: Broad H3K4me3 domain spreading is associated with increased transcriptional activation.

a) Kernel density estimation plot displaying H3K4me3 peak length change (-/+2kb) from melanocytes to mutational subtype. Number denotes mean length change in kilobases between melanocytes and melanoma subtype. Asterisk denotes p-value < 1e-50 between melanoma subtype. b) Barplot of broad and non-broad H3K4me3 promoter associated sites in each mutational subtype.

c) Barplot of broad H3K4me3 promoter associated sites displaying shortening or lengthening (-/+2kb) in melanoma tumors relative to melanocytes. d) Boxplot displaying quantile normalized mean RNA-seq expression profiles from NRAS (n=81), BRAF (n=118) and WT (n=38) metastatic samples

based on genes displaying shortening or lengthening (-/+2kb) in melanoma tumors relative to melanocytes identified in (c). e) Genome browser view of ChIP-seq tracks displaying H3K4me3 shortening on the PTEN, PMEL and MITF genes in melanoma tumors relative to melanocytes.

f) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the PTEN, PMEL and MITF genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom). Asterisk denotes p-value < 0.05.

g) Genome browser view of ChIP-seq tracks displaying H3K4me3 lengthening on DLX1/2 and TBX3 genes in melanoma tumors relative to melanocytes. h) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of the DLX1/2 and TBX3 genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=4, BRAF=13, WT=3 (top), NRAS=81, BRAF=118, WT=38 (bottom). Asterisk denotes p-value < 0.05.

Figure 6

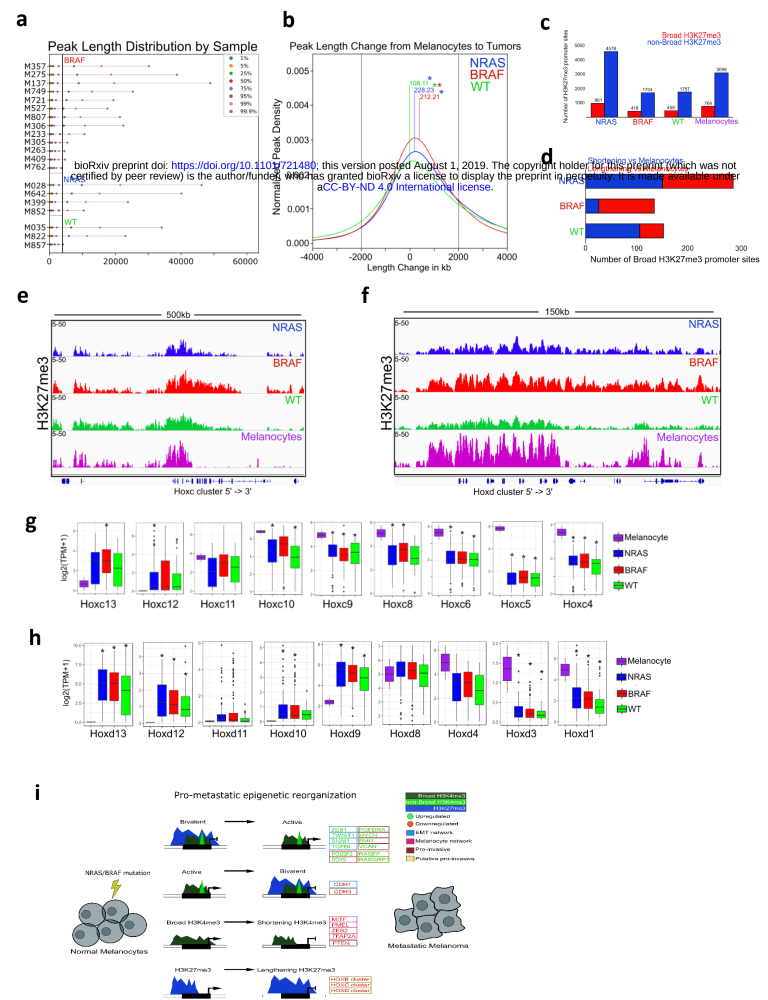


Figure 6: Broad H3K27me3 domains display preferential lengthening in metastatic melanoma.

a) Quartile plot displaying peak length distribution based on percentage of broadest H3K27me3 domains in melanoma tumors. Black line denotes 4kb peak length.

b) Kernel density estimation plot displaying H3K27me3 peak length change (-/+2kb) from melanocytes to mutational subtype. Number denotes mean length change in kilobases between melanocytes and melanoma subtype. Asterisk denotes p-value < 1e-15 between melanoma subtypec) Barplot of broad (>4kb) and non-broad (<4kb) H3K27me3 promoter associated sites in each mutational subtype.

d) Barplot of broad H3K4me3 promoter associated sites displaying shortening or lengthening (>10kb) in melanoma tumors relative to melanocytes. e) Genome browser views of ChIP-seq tracks displaying exceptionally wide H3K27me3 lengthening from the 5'->3' end on HOXC cluster genes in BRAF mutational subtypes.

f) Genome browser views of ChIP-seq tracks displaying exceptionally wide H3K27me3 lengthening from the 5'->3' end on HOXB cluster genes in NRAS mutational subtypes.

g) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of all HOXC cluster genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=81, BRAF=118, WT=38). Asterisk denotes p-value < 0.05.

h) Boxplot displaying quantile normalized mean RNA-expression profiles (log2 TPM) of all HOXD cluster genes in melanocytes (n=2) and melanoma tumor subtypes (NRAS=81, BRAF=118, WT=38). Asterisk denotes p-value < 0.05.

i) Model for potential mechanism in which bivalent switches, the shortening of H3K4me3 domains and lengthening of H3K27me3 domains facilitate the reorganization of an interconnected regulatory network encompassing EMT-TF, melanocyte master regulators and mesenchymal/invasive genes to promote melanoma metastasis