Enhancer methylation dynamics contribute to cancer plasticity and patient mortality

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During development, enhancers play pivotal roles in regulating gene expression programs; however, their involvement in cancer progression has not been fully characterized. We performed an integrative analysis of DNA methylation, RNA-seq, and small RNA-seq profiles from thousands of patients, including 25 diverse primary malignances and seven body sites of metastatic melanoma. We found that enhancers are consistently the most differentially methylated regions (DMR) as cancer progresses from normal to primary tumors and then to metastases, compared to other genomic features. Remarkably, identification of enhancer DMRs (eDMRs) enabled classification of primary tumors according to physiological organ systems, and in metastasis eDMRs are the most correlated with patient outcome. To further understand the eDMR role in cancer progression, we developed a model to predict genes and microRNAs that are regulated by enhancer and not promoter methylation, which shows high accuracy with chromatin architecture methods and was experimentally validated. Interestingly, among all metastatic melanoma eDMRs, the most correlated with patient survival were eDMRs that “switched” their methylation patterns back and forth between normal, primary, and metastases and target cancer drivers, e.g., KIT. We further demonstrated that eDMR target genes were modulated in melanoma by the bone metastasis microenvironment, suggesting that eDMRs respond to microenvironmental cues in metastatic niches. Our findings that aberrant methylation in cancer cells mostly affects enhancers, which contribute to tumor progression and cancer cell plasticity, will facilitate development of epigenetic anticancer approaches.

[Supplemental material is available for this article.]
cancer types (overall 25 data sets). We used the genome-wide scale Illumina HumanMethylation450 arrays (Dedeurwaerder et al. 2011) to identify differentially methylated regions (DMRs) by analyzing over 6200 DNA methylation profiles of patients’ tumors and normal tissues (Supplemental Table S1). We identified 123,649 DMRs (minimum region-wise mean methylation difference >0.3, false discovery rate, q <0.2). Interestingly, we found that enhancers had more DMRs than any other genomic region; promoters and CpG islands exhibited the least variation (Fig. 1A). This result suggests that alterations of enhancer methylation have a significant role in cancer progression, in support of data reported previously (Aran and Hellman 2013, 2014; Ziller et al. 2013; Taberlay et al. 2014). To test whether the selected threshold of 0.3 for differential methylation represented an optimal measure, we calculated the fraction of DMRs in each genomic feature using higher and lower thresholds (from 0.2 to 0.5); enhancers were consistently the most variable regions (Supplemental Fig. S1A). As the frequency of CpG dinucleotides varies with genomic features (Supplemental Fig. S1B), we repeated the differential methylation analysis using methylation levels of individual CpG sites in the array (Supplemental Fig. S1C) and observed consistent results, namely that CpG methylation within enhancers is the most variable compared to all other genomic features.

Given the parallels between embryonic development and oncogenic transformation (Hon et al. 2013), we examined the binding patterns of central pluripotent transcription factors (TFs) POU5F1, SOX2, and NANOG in various DMRs. We observed that multiple binding occurred more frequently at differentially methylated enhancers than at promoters or CpG islands (Fig. 1B). These results support previous findings (Göke et al. 2011; Whyte et al. 2013) suggesting that eDMRs bound by pluripotent TFs may mediate central expression programs. To qualitatively examine the altered enhancers, we analyzed their ChromHMM states (Ernst and Kellis 2012). We found that, compared to enhancers that were not differentially methylated (static enhancers), eDMRs were significantly enriched for the ChromHMM state of “strong enhancers” (P = 2 × 10^{-16}) (Fig. 1C), similar to previous reports (Aran et al. 2013). These results suggest that enhancers altered in cancer may regulate their target genes more strongly than do unaltered enhancers. It is known that promoter and CpG island hypermethylation are associated with gene silencing in cancers (Bergman and Cedar 2013), and, indeed, we found that these regions were frequently hypermethylated (63% and 94%, respectively). In comparison, enhancers were mostly differentially hypomethylated (67%) (Fig. 1D). Next, we used chromatin marks to qualitatively determine the chromatin features of the hypermethylated and hypomethylated enhancers. Using data from the ENCODE Project (Rosenbloom et al. 2013), we found that differentially hypomethylated enhancers exhibited significantly higher levels of chromatin marks of active DNA (increased DNase I, H3K4me1, H3K4me2, H3K27ac, H2A.Z, EP300, POL2, and decreased CTCF). Differentially hypermethylated enhancers showed the footprints of closed and inactive DNA (Fig. 1E; Supplemental Fig. S1D,E), consistent with previous results (Zhou et al. 2011; Aran et al. 2013; Lam et al. 2014). Taken together, the analysis of multiple tumor types strongly suggests that alteration of the enhancer methylome is a frequent feature of transformed cells and that it...
generally leads to chromatin activation, likely involved in cancer progression.

Differential methylation patterns of eDMRs clusters tumors according to their organ system

Next, we compared the patterns of methylation change in enhancers between the various types of cancer. Since enhancers show tissue-specific patterns of histone modifications and TF binding in normal cells (Bulger and Groudine 2011), we expected that alteration of methylation patterns would be cancer-/tissue-specific. Indeed, most eDMRs (54%) were unique to a single cancer type (Fig. 2A). Strikingly, principal components analysis (PCA) of eDMRs showed a higher order clustering of cancers into groups related to the same organ system (Fig. 2B). For example, the central nervous system cluster, which includes cancers arising from glia and astrocytes (Fig. 2B, green circle); similarly, the reproductive system cluster consists of breast, uterine, and prostate tumors (Fig. 2B, pink circle). Importantly, PCA of differentially methylated CpG islands, promoters, and intergenic regions showed no such

Figure 2. Variation in enhancer methylation classifies primary tumors according to their organ systems. (A) Pie chart shows that most eDMRs are unique (77% of the eDMRs appear in only one or two cancers, excluding the same cancer type, e.g., GBM1 and GBM2). (B) Principal components analysis (PCA) shows that differential methylation patterns of eDMRs are similar for cancers belonging to the same organ systems (circled and colored similarly). First and second principal components (x-axis: PC1; y-axis: PC2) account for 72% of variation (see Supplemental Fig. S2A–C for PCA analysis of methylation changes within CpG islands, promoters, and intergenic regions). (C) Schematic of method for identification of eDMR-gene and eDMR-miRNA pairs (see Supplemental Fig. S2H for flow chart). (D) (Top) Precision of eDMR-gene pair predictions is compared to predictions from CAGE (Andersson et al. 2014), IM-PET (Teng et al. 2015), ChIA-PET, and Hi-C (Teng et al. 2015) methods (see text). (Bottom) Percent of predicted eDMR-gene pairs. Results in both plots are divided into groups based on the distances between eDMR and transcription start site (TSS). Most predictions (>56%) lie within a distance of <500 kb (abbreviations: 100K, 0–100,000 bp, 200K: 100,001–200,000 bp, and so forth). (E) eDMR-associated genes are distinctly enriched for genes involved in diseases related to a particular organ system. Disease-related genes were derived from the DISEASES database (see Methods). Scores represent −log₁₀(FDR-corrected P-values). (F) Scatter plots (a point for each patient) show examples of tissue-specific genes linked to cancer risk and their eDMRs (the same six cancer types shown in E). Spearman’s correlations between eDMRs methylation and associated-gene expression are shown. (G) Landscape of two differentially hypomethylated eDMRs in uterine (UCEC) and head and neck (HNSC) cancers, associated with a single miRNA, miR-9-1, which is up-regulated in both cancers. (H) Landscape of two differentially hypomethylated eDMRs in breast (BRCA) and colon (COAD) cancers associated with two miRNAs of the same family, miR-200a and miR-200b; each is up-regulated in the respective tumors. In G,H, yellow arrows mark hypomethylated eDMRs, green arrows mark up-regulated miRNAs; gray boxes mark eDMR and miRNA locations in the hg19/GRCh37 genome; distances between miRNAs and eDMRs are indicated on arrows. All parts of the figure refer to eDMRs determined between normal samples and primary tumors.
clustering (Supplemental Fig. S2A–C). No library preparation batch affect was found either (Supplemental Fig. S2D,E). This evidence provides confidence that our findings were not affected by batch effect and strengthen our hypothesis that enhancer methylation changes may have meaningful biological roles.

Next, in order to uncover the functional effect of altered enhancer methylation, we first compared the genomic neighborhood of eDMRs with static enhancers (those showing no differential methylation). We found that eDMRs were flanked by more genes and microRNAs (miRNAs) than were static enhancers (Supplemental Fig. S2F,G). To identify which genes the eDMRs regulate, we developed an integrated model combining multi-omics data for associating enhancers with their target genes (eDMR-associated genes) (Fig. 2C; Supplemental Fig. S2H for pipeline). Since it is known that there is an inverse correlation between DNA methylation and chromatin activity (Zhou et al. 2011), we retained only inversely correlated eDMR-gene pairs, which represented the majority of our data (Supplemental Fig. S2I). In order to assess the validity of our eDMR-gene pair predictions, we compared our results to other methods that identify enhancer-promoter associations based on physical interactions IM-PET (He et al. 2014), ChIA-PET, Hi-C (Teng et al. 2015), and transcriptional activities of interacting enhancer-promoters (cap analysis gene expression; CAGE) (Andersson et al. 2014). Our model predicted eDMR-gene pairs separated by 400 kb or less at a precision rate of 75% or better (Fig. 2D), indicating that our model can reliably predict interacting enhancer gene pairs.

Similar to their enhancers (Fig. 2B), the genes predicted to be regulated by the eDMRs also showed organ system patterns of enrichment (Fig. 2E; Supplemental Table S3). Using our model, we were able to identify known tissue-specific and ubiquitous oncogenic genes and microRNAs (Fig. 2F–H), such as: ESR1 in breast cancer (Holst et al. 2012; Aran and Hellman 2014), ECT2 in lung cancer (Murata et al. 2014), and WNT3A in prostate cancer (Verras et al. 2004), hsa-miR-9-1 (Ma et al. 2010) and the clustered miRNAs hsa-miR-200a and miR-200b (Korpal et al. 2008; see Supplemental Table S4 for full list of predicted eDMR-gene pairs). Taken together, our model defines cancer-related eDMRs as organ-specific regulators of genes and miRNAs that are central to malignant transformation.

**Accumulation of eDMR hypomethylation correlates with likelihood of metastasis**

Our analysis showed that changes in methylation patterns of enhancers in primary tumor tissues are influenced by their relation to the organ system (Fig. 2B). We were curious what would occur to DNA methylation patterns of cells that disseminate from the primary location and colonize in metastatic sites. To evaluate this, we focused on melanoma, a highly metastatic cancer (Braeuer et al. 2014). We compared methylation data from metastatic melanoma patients (seven distinct locations) to data from patients with primary in situ melanoma (Supplemental Table S1). Consistent with our observation of differential methylation patterns in primary tumors (Fig. 1A), the majority of “metastatic DMRs” occurred within enhancers (Fig. 3A), suggesting that enhancers play important roles, not only in malignant transformation but also in metastatic progression. Additionally, metastatic eDMRs differentiate much better between patient outcomes than do DMRs from any other genomic feature (Fig. 3B; Supplemental Fig. S3A). The majority of eDMRs were specific to a single metastatic site (Fig. 3C; Supplemental Fig. S3B) and were mostly (74%) hypomethylated (Fig. 3D), similar to our observations in primary cancers (see Figs. 1D, 2A). Strikingly, Figure 3E shows that accumulation of hypomethylated enhancers highly correlates with the likelihood of forming metastases at distant organs derived from Meyers and Balch (1998). For example, melanoma is more likely to metastasize to lymph nodes than to the brain; correspondingly, a greater number of enhancers are differentially hypomethylated in brain metastases than in lymph node metastases (2.24% and 0.33%, respectively) (Fig. 3E). In contrast, we found no correlation between the fraction of differentially hypermethylated metastatic eDMRs and the frequency of spreading to organs (Supplemental Fig. S3C). These results are in agreement with studies suggesting that global hypomethylation is a common feature of diseased states (Pogribny and Beland 2009). Since widespread DNA methylating changes are associated with aging (Richardson 2003), we calculated the correlations between patient ages and accumulation of enhancer methylation changes. Encouragingly, we found no significant correlation between eDMR methylation changes in metastatic tissues and the patient age (Supplemental Fig. S3D), suggesting that metastatic progression corresponds to cancer-related changes and not age-related changes.

In order to specifically explore whether melanoma eDMRs could promote metastatic growth, we identified genes differentially expressed between primary and metastatic melanoma and compared the correlation of these genes with methylation patterns of differentially methylated enhancers and promoters (eDMRs and pDMRs, respectively). We found that eDMRs were significantly more correlated with differentially expressed genes than were pDMRs (Supplemental Fig. S3E). Next, we divided the genes into two groups: eDMR-associated genes, and all other genes not identified by our model to be associated with eDMRs (control genes) (Fig. 3F), and examined the differential expression patterns of these two groups. The percentage of genes that were differentially expressed between normal and primary melanoma was similar for both groups (60% and 53% for eDMR-associated genes and control genes, respectively) (Fig. 3F); however, between primary and metastatic melanoma, the eDMR-associated genes were significantly more variable than the control genes (binomial distribution, \( P < 2 \times 10^{-16}; 57\% \) and \( 3\% \), respectively) (Fig. 3F, marked by gray areas). This observation supports our notion that a functional interaction exists between the eDMRs and their predicted associated genes. Additionally, we note that a fraction of the control genes were down-regulated in both primary and metastatic melanoma (Fig. 3F, left panel, marked with dashed box). This group was enriched (\( q < 0.01, \) FDR corrected) with Gene Ontology (GO) terms associated with tissue development (GO:0008544, GO:0007398) and differentiation (GO:0030216, GO:0009913, GO:0030855), likely a feature of tumor de-differentiation (Brabletz 2012).

**Methylation plasticity of melanoma eDMRs is associated with increased patient mortality**

The transition of cancer from the in situ primary stage to the metastatic stage involves phenotypic plasticity (Craene and Berx 2013), which facilitates migration through tissues and adaptation to changing microenvironments. Underlying tumor plasticity are genetic and epigenetic regulatory layers that are reprogrammed in the context of cancer (Friedl and Alexander 2011). Interestingly, we found that 18% (\( N = 277 \)) of melanoma metastatic eDMRs (\( N = 1539 \)) switched the direction of differential methylation as melanoma progressed (denoted “switched” eDMRs) (Fig. 4A, bottom circle). For example, regions that were hypermethylated...
between normal melanocytes and primary melanoma were hypomethylated between primary and metastatic melanoma. We also defined two other eDMR groups: “consistent” and “de novo.” The “consistent” eDMRs exhibited differential methylation changes in the same direction (e.g., hypomethylated between normal melanocytes and primary melanoma, and hypomethylated between primary and metastatic melanoma) (Fig. 4A, right circle). The “de novo” group exhibited differential methylation only between primary and metastatic melanoma samples and not between normal melanocytes and primary tumors (Fig. 4A, left circle). We examined whether these three groups of eDMRs differed with respect to metastatic progression by comparing their ability to correlate with survival of patients (see Methods). To our knowledge, this is one of the first examples (Stone et al. 2015) of a survival analysis applied to DNA methylation patterns of enhancers. We identified 30 eDMRs that were associated with patient survival rates (marked “survival” eDMRs) (Fig. 4A, middle circle). Remarkably, these survival eDMRs were enriched with the switched eDMRs ($P = 9.4 \times 10^{-5}$) (Fig. 4A, bottom circle), whereas de novo eDMRs were significantly depleted ($P = 1.6 \times 10^{-2}$) (Fig. 4A, left circle). In addition, the switched eDMRs exhibited the highest conservation scores (Fig. 4B) and the fewest copy number variations (CNVs) in melanoma patients (Fig. 4C). These results suggest that eDMRs, in particular the switched eDMRs that exhibit methylation plasticity, are functionally important in cancer.

Given these results, we hypothesized that methylation plasticity may play important roles in melanoma metastatic progression. To examine this, we performed survival analyses also on the eDMR-associated genes, using their expression patterns across patients. Remarkably, we found that 40% were able to significantly differentiate between survival outcomes ($q^2, q < 0.1$, FDR corrected); these genes include: ATP2B1 (Lee et al. 2002), FN1L2 (Zhu et al. 2008), KIT (Tian et al. 1999), PRKCE (Sharif and Sharif 1999), and VGF (Mitra et al. 2008). Figure 4D represents such an example in which the oncogene KIT and the eDMR (located at Chr4:55708295–55709294, hg19/GRCh37) not only exhibit switched patterns of methylation and expression, but both independently distinguish between patient survival rates based on their expression andmethylation patterns, respectively (Fig. 4D, left and right panels). The KIT promoter was not differentially methylated, nor did the methylation pattern differentiate between survival times (Fig. 4D, middle panel).

To experimentally test the role of enhancer methylation on the transcriptional regulation of KIT, we first examined KIT expression upon treatment with 5-aza-2′-deoxycytidine (5-aza-dC), a DNA methyltransferase inhibitor. There was a significant increase in KIT levels in treated compared with untreated cells (Fig. 4E). Next, we cloned the KIT enhancer upstream of a luciferase reporter and demonstrated that its expression was reduced upon enhancer methylation (Fig. 4F; Supplemental Fig. S4A). Finally, we examined the role of KIT up-regulation on the invasive potential of
melanoma cells (Fig. 4D). Remarkably, noninvasive melanoma cells (Golan et al. 2015) acquired significant invasion ability upon KIT overexpression (Fig. 4G; Supplemental Fig. 4B). Taken together, our data demonstrate that enhancer methylation contributes to cancer progression by directly regulating oncogene expression.

Next, we asked whether the dynamics of eDMR methylation and eDMR-associated genes expression are a result of tumor heterogeneity, evolution of the disseminated cancer cells, or are induced by the metastatic niche. To assess this, we selected two eDMR-associated genes, CTYL1 and KIF14, which were differentially expressed only between primary and metastatic melanoma (not between normal and primary melanoma tissues). CTYL1 is a cytokine-like protein implicated in lung cancer and neuroblastoma (Kwon et al. 2012; Wen et al. 2012), and KIF14 is an oncogene...
essential for cytokinesis (Corson et al. 2005); both are up-regulated in melanoma bone metastases ($P < 0.05$) (Fig. 5A, top and bottom panels). High expression levels of these genes is correlated with poor patient survival rates ($\chi^2 q < 0.1$, FDR corrected) (Fig. 5A, green stepwise curves). Enhancers of these genes were differentially methylated, yet their promoters were not (Fig. 5A, right and middle, panels, respectively). To experimentally analyze the role of enhancer methylation on the transcriptional regulation of CYTL1 and KIF14, we examined whether their expression is sensitive to DNA methylation. CYTL1 and KIF14 expression was significantly up-regulated upon treatment of cells with 5-aza-dC (Fig. 5B). Next, we cloned the identified enhancers of CYTL1 and KIF14 upstream of a luciferase reporter and observed a decrease in the reporter expression upon enhancer methylation (Fig. 5C; Supplemental Fig. S4A). These results strengthen the validity of our model by demonstrating that the identified eDMRs elicit a response to changes in their methylation and directly regulate the expression of their associated genes: KIT, CYTL1, and KIF14.

Finally, to assess changes in gene expression induced by metastatic melanoma cells colonization in the bone tissue, we first generated a melanoma cell line stably expressing the GFP gene to enable tracking of the melanoma cells. Next, we established a coculture of melanoma cells with human primary osteoblasts (Fig. 5D; Dillon et al. 2012) and found that levels of CYTL1 and KIF14 were significantly increased compared to their levels in melanoma cells cultured alone (Fig. 5E). Our data support the notion that homing into a new microenvironment induces expression of pro-cancer genes, which are regulated by eDMRs in melanoma metastatic cells. Taken together, our data suggest that methylation changes at enhancers contribute to melanoma phenotypic plasticity and ultimately to the patient’s chance of survival.

**Discussion**

Here, we performed an analysis of DNA methylation alterations in over 6200 cancer patients from 31 cancer data sets, including...
23 diverse primary malignant tumors, two benign tumors, and melanoma metastases to seven distinct organs. This extensive analysis revealed that most methylation variation occurs at enhancers (Figs. 1A, 3A; Supplemental Fig. S1C). Changes in methylation patterns could result from competition between methylation and de-methylation processes or from errors in replication (Jones 2012). Our findings support the latter hypothesis, for two reasons. First, replication-related methylation errors occur more frequently in methylated regions, such as enhancers, where replication errors lead to loss of methylation; this is what we observe (Fig. 1D). Second, expression of DNMT3B, which encodes a methylase enzyme, was up-regulated in many tumors, and hence, we would expect increased methylation since we observed no significant differences in expression of TET genes, which encode demethylase enzymes (Supplemental Fig. S2J). However, since the majority of changes in enhancers’ methylation involves de-methylation, we hypothesize that loss of enhancer methylation could have occurred during replication.

Enhancers play central roles in normal development and differentiation by responding to complex environmental cues. Cancer cells are exposed to changing environmental conditions that require their adaptation; this mainly occurs through epigenetic reprogramming (Friedl and Alexander 2011; Godding et al. 2014). We hypothesize that, in the context of cancer, enhancer methylation may be primed to respond to microenvironmental signals. To test this, we mimicked the microenvironment of melanoma bone metastases and recapitulated the perturbation of expression of eDMR-associated genes that were altered in the bone metastases. These results suggest that the metastatic niche can alter expression of eDMR-associated genes.

Interestingly, we show that, in comparison to DMRs in other genomic features (CpG islands, promoter, exons, introns, etc.), eDMRs can differentiate best between patient outcomes (Supplemental Fig. S4D). These results demonstrate the important role of alteration of enhancer methylation in cancer progression. However, we do not exclude that other factors may drive cancer progression, nor are we suggesting a direct causality between enhancer methylation and cancer progression; it may well be that alteration of enhancer methylation is a secondary event of the malignancy.

Nevertheless, we did observe that methylation patterns of eDMRs may be informative of patient survival rates (Fig. 3B). Within this group was a subset (18%) of highly conserved eDMRs that displayed methylation plasticity (“switched” eDMRs) (Fig. 4A–C) and provided insight into alteration of their associated genes (Fig. 4D). This is one of the first studies (Stone et al. 2015) to suggest that methylation patterns of enhancers can be used to predict patient outcome. Moreover, it has been shown that, in many diseases, including cancer, methylation changes are accumulative as the cancer progresses (Pogribny and Beland 2009); here, we show that methylation plasticity may also play important roles in cancer progression (see Supplemental Fig. S4C; for model of methylation plasticity and cancer progression). A prime example of the relationships between metastatic progression, methylation plasticity, and patient mortality is that of the eDMR-associated oncogene, KIT (Fig. 4D–F). Both the methylation of the eDMR and the expression of the KIT display plasticity (Fig. 4D) and significantly distinguish between patient survival rates. Our results suggest that enhancer methylation patterns may be informative of patient outcomes and that they may influence malignant progression via methylome plasticity.

Methods

TCGA and GEO data sources: DNA methylation, RNA-seq, smRNA-seq

Publicly available data of DNA methylation, RNA-seq gene expression, and small RNA-seq miRNA expression from cancer patient tissues were obtained from The Cancer Genome Atlas (TCGA, https://tcga-data.ncl.nih.gov/tcga) and from the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) repository (http://www.ncbi.nlm.nih.gov/geo). See Supplemental Table S1 data set information.

Chromatin marks, transcription factors, sequence conservation, and copy-number variation data

For analysis of chromatin marks (Fig. 1E; Supplemental Fig. 1D,E), we downloaded RNA polymerase II (POL2), histone modifications H2AZ, H3K4me1, H3K4me2, and H3K27ac, histone acetyltransferase (EP300), CCCTC-binding factor (CTCF), DNase I hypersensitivity, and Chromatin State Segmentation by HMM (ChromHMM) from the ENCODE Project. Replicate experiments were averaged. See Supplemental Materials for URL and description and Supplemental Table S2 for ENCODE Project cell types.

For the transcription factor analysis (Fig. 1B), we downloaded processed ChIP-seq data of POUSF1, SOX2, and NANOG, downloaded from NCBI/GEO (GSE46130). Replicate experiments were averaged. The TF peaks were overlapped with the coordinates of the differentially methylated enhancers, promoters, and CpG islands.

For the conservation analysis of enhancer (Fig. 4B), we used 100-way PhastCons conservation data.

Copy number variations for skin cutaneous melanoma (SKCM) were obtained from TCGA database as genomic segments after removal of germline CNV (TCGA_SKCM_GSNP660CNV_gSeg).

Defining differentially methylated regions

First, we matched the CpG codes in the Illumina HumanMethylation450 microarray to their genomic coordinates using the microarray platform (NCBI/GEO record GPL13534; human genome release hg19/GRCh37). Second, we annotated the CpGs using the information in this record, identifying CpG sites belonging to CpG islands, shores, shelves, enhancers, and UTRs. Given that this record does not have information of CpG sites in promoters, exons, or introns, we used the UCSC table “knownGene” to identify these sites. Third, some CpG sites had several annotations; thus, we divided the CpGs into unique (non-overlapping) genomic features using the following prioritization: (1) promoters, (2) CpG islands, (3) enhancers, (4) introns, (5) exons, (6) UTRs, (7) shores, (8) shelves, and (9) intergenic regions. Fourth, for each separate genomic feature, we constructed intervals using a window of 500 bp directly upstream of and downstream from the CpG coordinate. Overlapping intervals (same genomic feature) were joined, and extended into a larger interval. Sixty-nine percent of the regions had a length of 1000 bp, 28% a length >1000 bp and <2000 bp, and no region was greater in length than 7500 bp. Fifth, we used these genomic intervals to calculate region-wise methylation levels based on the average methylation of all CpG sites within the interval; we performed this for all normal and all tumor samples in each cancer data set. Sixth, we used the two-sample Wilcoxon test to identify differentially methylated regions between normal and primary samples (methylation threshold >0.3, q < 0.2, FDR corrected); metastatic DMRs were identified by comparing primary and metastatic melanoma (methylation threshold >0.2 and q < 0.2). The fractions of DMRs in each
Predictions for enhancer and transcription start site (for genes) or promoter, methylation variability (performed only for predicting genes with expression change due to enhancer, rather than by promoter). Spearman’s correlation between eDMR methylation patterns and gene/miRNA expression patterns across the same (matched) patients. Highly correlated eDMR-genes or eDMR-miRNAs pairs were retained (correlation below ~0.4). Third, we filtered out genes whose promoters had a differential methylation greater than 0.2, enriching for genes with expression change due to enhancer, rather than by promoter, methylation variability (performed only for predicting eDMR-associated genes and not associated-miRNAs, since annotation for miRNA promoters is incomplete). Fourth, we selected eDMR-associated genes and not associated-miRNAs, since annotations for miRNA promoters are incomplete. We developed an integrated model to predict genes or miRNAs regulated by eDMRs (Fig. 2C; flowchart in Supplemental Fig. S2H).

We used the DISEASES website (http://diseases.jensenlab.org) (Pletscher-Frankild et al. 2015) to evaluate enrichment for diseases in particular organ systems (Fig. 2E). Diseases were identified using keywords: digestive (digest-, gastro-, gastric, stomach), endocrine (endocrine-, gland disease, gland cancer, gland neoplasm, thyroid, pancreas), renal (kidney, renal, nephron-, nephri-), reproductive (reproductive-, breast, prostate, uterine, cervic-, cervix, uterus), respiratory (lung, respiratory; duplicate genes were removed to ensure unique values for hypergeometric distribution significance testing and FDR corrected).

Survival analysis
Survival time was derived from the “overall survival” column of the clinical data files obtained for TCGA samples. First, patients were divided into two groups (high and low) by comparing the eDMR methylation level to the median methylation of the eDMRs across all patients. Similarly, for gene expression survival analysis, patients were divided into two groups by comparing the gene expression of each patient to the median expression of all patients. Significant differences between the two groups were determined using the $\chi^2$ distribution ($q<0.1$, FDR corrected).

Enrichment for association of DMRs with patient survival (Fig. 3B) was determined by comparing the amount of DMRs that can significantly differentiate between patient outcomes with the amount of static regions that can do the same (determined separately for each genomic feature).

Multiple testing
All significance tests were corrected for false discovery rates (FDR) using the Benjamini and Hochberg adjustment (Benjamini and Hochberg 1995).

Computational data analysis
Data analyses were performed using R statistical language (R Core Team 2015). We used the following packages for R in the analysis “GenomicRanges” (version 1.16.4) (Lawrence et al. 2013), “TxDb.Hsapiens.UCSF.hg19.knownGene” (version 3.1.2) (http://bioconductor.riken.jp/packages/3.1/data/annotation/html/TxDB.Hsapiens.UCSF.hg19.knownGene.html), “survival” (version 2.37–7) (http://cran.r-project.org/package=survival) and “reshape2” (version 1.4.1) (Wickham 2007). Custom R scripts for determining differentially methylated regions and for predicting and ranking eDMR-gene pairs were provided, together with sample data sets derived from the TCGA, available in the Supplemental Data (see “INFO_README.txt” file in Supplemental Scripts).

Primary human osteoblasts and melanoma cells coculture
Primary human osteoblasts were isolated using a protocol described by Dillon et al. (2012). In short, trabecular bone was obtained from healthy donors undergoing total knee arthroplasty. Written and informed consent was obtained from all subjects. The protocol was approved by the institutional Ethics Committee at Tel-Aviv Sourasky Medical Center, in accordance with the Helsinki Declaration on the use of human subjects in research. The trabecular bone fragments were diced into small pieces and washed with sterile PBS. The diced bone extracts were then placed on a tissue culture plate with Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 20% fetal bovine serum, 100 U/mL penicillin, and 100 µg/mL streptomycin (all from Gibco, Life Technologies) and incubated at 37°C and 5% CO_2_. Five days later, the medium was replaced; medium was then replaced twice a week until the cells reached confluency. Osteoblasts were then seeded 24 h prior to addition of melanoma cells in a ratio of 1:5 melanoma cells to osteoblasts. In control plates, only melanoma cells were seeded.

Cell culture and FACS sorting
WM3682 melanoma cells were generously provided by Dr. Levi A. Garraway (Department of Medical Oncology and Center for Cancer Genome Discovery, Dana-Farber Cancer Institute, Boston, MA). Cells were cultured in DMEM medium supplemented with 10% fetal bovine serum (Sigma-Aldrich) and 1% penicillin/streptomycin/glutamine (Invitrogen). For establishment of
stable cell lines, cells were transfected with GFP-expressing plasmid and selected with 1 μg/mL puromycin (Sigma-Aldrich). For coculture, WM3682-GFP cells were cultured with bone cells for 5 d. Cells were collected by flow cytometry using BD FACSAria cell sorter.

RNA purification and qRT-PCR
Total RNA was purified from sorted melanoma cells using TRIzol (Invitrogen) according to the manufacturer’s instructions, followed by treatment with RNase-free DNase (Qiagen). RNA was quantified based on OD260/280. For qRT-PCR analysis, RNA was subjected to one-step qRT-PCR using a MultiScribe RT-PCR kit (Applied Biosystems) and FastStart Universal SYBR Green Master Mix (Roche).

Plasmids and cloning
EDMRs of the human KIT, KIF14, and CYTL1 loci (Chr4:55708294–55709294, Chr1:201198480–201199526, Chr4:5019645–5020678, respectively) were amplified from human genomic DNA (see Supplemental Table S5 for primer sequences used in cloning). The CYTL1 eDMR (1033 bp) and KIF14 eDMR (1046 bp) fragments were digested with NheI-XhoI restriction enzymes and inserted into the pGL3-promoter vector (Promega) upstream of a minimal promoter and firefly luciferase reporter gene (kindly provided by Professor Eran Barachar, Department of Cell Research and Immunology, Faculty of Life Sciences, Tel-Aviv University). The KIT eDMR (1000 bp) fragment was cloned into PGL3-promoter reporter plasmid digested with Smal-Xhol restriction enzymes. The pCDNA3-KIT expression vector was kindly provided from Prof. Lars Rönnstrand (Division of Translational Cancer Research and Lund Stem Cell Center, Lund University).

In vitro methylation, transfections, and dual luciferase assay
The KIT, CYTL1, and KIF14 eDMR firefly luciferase reporter vectors were in vitro methylated using the methylase SssI (New England Biolabs), according to the manufacturer’s recommendations, followed by purification using the Wizard SV PCR clean-up system (Promega). Successful methylation was verified by restriction enzyme digestion with the methylation-sensitive (HpaII) and methylation-insensitive (MspI) enzymes (New England Biolabs). The digestion patterns were analyzed by agarose gel electrophoresis. WM3682 melanoma cell lines were cotransfected using jetPEI, digestion patterns were analyzed by agarose gel electrophoresis. WM3682 melanoma cell lines were cotransfected using jetPEI, digestion patterns were analyzed by agarose gel electrophoresis.

Invasion assay
WM3682 melanoma cell lines were transfected with the KIT expression vector or empty vector (as control) using jetPEI. Forty-eight hours post-transfection, an invasion assay was performed as previously described (Golan et al. 2015).

5-aza-deoxycytidine treatment
WM3682 melanoma cell lines were treated with 10 μM 5-aza-dC (Sigma-Aldrich) for 48 h, following by RNA purification and qRT-PCR as described. Baseline expression was established by mock treatment of cells with DMSO.

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