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Global loss of DNA methylation uncovers intronic enhancers in genes showing expression changes

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Abstract

Background: Gene expression is epigenetically regulated by a combination of histone modifications and methylation of CpG dinucleotides in promoters. In normal cells, CpG-rich promoters are typically unmethylated, marked with histone modifications such as H3K4me3, and are highly active. During neoplastic transformation, CpG dinucleotides of CG-rich promoters become aberrantly methylated, corresponding with the removal of active histone modifications and transcriptional silencing. Outside of promoter regions, distal enhancers play a major role in the cell type-specific regulation of gene expression. Enhancers, which function by bringing activating complexes to promoters through chromosomal looping, are also modulated by a combination of DNA methylation and histone modifications.

Results: Here we use HCT116 colorectal cancer cells with and without mutations in DNA methyltransferases, the latter of which results in a 95% reduction in global DNA methylation levels. These cells are used to study the relationship between DNA methylation, histone modifications, and gene expression. We find that the loss of DNA methylation is not sufficient to reactivate most of the silenced promoters. In contrast, the removal of DNA methylation results in the activation of a large number of enhancer regions as determined by the acquisition of active histone marks.

Conclusions: Although the transcriptome is largely unaffected by the loss of DNA methylation, we identify two distinct mechanisms resulting in the upregulation of distinct sets of genes. One is a direct result of DNA methylation loss at a set of promoter regions and the other is due to the presence of new intragenic enhancers.

Background

Genes are regulated by epigenetic modifications and transcription factor binding at their promoters and at distally located regulatory regions. Studies over the past two decades have shown that promoters having high levels of DNA methylation are not transcriptionally active [1-3]. Recent genome-wide epigenetic profiling efforts demonstrate that promoter regions with high levels of DNA methylation have low levels of active marks such as H3K4me3 and that methylated distal regulatory regions lack the active mark H3K27ac [4-8]. During neoplastic transformation, DNA methylation is reduced genome-

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wide, but accumulates at certain promoters. Because some of the promoters that become highly methylated are tumor suppressor genes [9-11], DNA de-methylating agents are being used in the clinic to reactivate silenced promoters. However, it has yet to be determined whether the global eradication of DNA methylation is advantageous for the cell or the patient. One could imagine that global loss of DNA methylation would have major effects on the transcriptome and epigenome of the cell. The DNA de-methylating drug 5-azacytidine (5-Aza-CR) has been approved for use as an epigenetic chemotherapeutic agent [12,13]. 5-Aza-CR functions by incorporating into DNA in place of cytosine and trapping DNA methyltransferases (DNMTs), which leads to their degradation and a subsequent passive loss of DNA methylation via replication. Previously, we treated HEK293 cells with 5-Aza-CR and analyzed the effects on histone modifications and RNA expression [12]. We found that 5-Aza-CR treatment



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caused changes in gene expression in approximately 1,500 genes (out of the 24,000 genes analyzed) but less than 800 of the genes were up-regulated as a result, and most genes that showed increased expression were not regulated by promoters that displayed DNA methylation prior to treatment. In addition to affecting DNA methylation, 5-Aza-CR can also incorporate into RNA and interrupt normal cellular processes such as ribosomal assembly and translation [14,15]. Therefore, it was not clear if the observed changes in transcript levels were due to changes in transcription rate from de-methylated promoters or to changes in RNA stability caused by intercalation of the 5-Aza-CR into the transcripts, affecting cellular signaling pathways due to translational defects. In addition, treatment with 5-Aza-CR does not completely abolish DNA methylation. Even with high doses, the overall levels of DNA methylation are reduced only 50 to 60% [12]. Therefore, it was also possible that de-repression of genes was incomplete after treatment with the drug (due to the remaining DNA methylation) and that many more transcripts whose promoters are normally silenced by DNA methylation would be identified if a more dramatic reduction in DNA methylation could be achieved. Here we explore the relationship between DNA methylation and the epigenome using both HCT116 colorectal cancer cells and DKO1 cells, a derivative of HCT116 cells that have a bi-allelic knockout of DNMT1 and bi-allelic deletion of exons 2 to 21 of DNMT3b [16]. Surprisingly, we found only a modest effect on the transcriptome and very limited increases in active marks on promoter regions. In order to fully understand the effects of global DNA methylation loss on the transcriptome and the epigenome at promoters and distal regulatory regions, we employed genome-wide methods for examining DNA methylation, RNA expression changes, histone modification patterns, and RNA polymerase II (RNAPII) occupancy. We found that the most robust epigenomic changes occurring after loss of DNA methylation were due to the acquisition of thousands of new enhancers. Interestingly, many of the genes that were up-regulated in DKO1 cells via mechanisms distinct from de-methylation of promoter regions had multiple newly acquired intragenic enhancers.

Results

Loss of DNA methylation does not result in an increase in active histone marks at promoters

To determine the relationship between a reduction of DNA methylation and global epigenetic marks, we performed functional genomic analyses using DNMT-deficient HCT116 DKO1 cells. The DKO1 cell line has a bi-allelic knockout of DNMT1 and bi-allelic deletion of exons 2 to 21 of DNMT3b and is reported to have 5% of the overall DNA methylation levels relative to the parental HCT116 cell line [16]. However, these results were obtained using a liquid chromatography approach which monitored overall 5-methylcytosine content genome-wide and thus did not examine DNA methylation reduction in specific genomic compartments such as promoters or gene bodies. We therefore performed whole genome bisulfite sequencing (WGBS) on HCT116 parental and DKO1 cells (Figure 1A); to achieve adequate coverage of GC-rich promoters, WGBS library preparation was performed as discussed in the Materials and methods section. We found that promoters, gene bodies, and randomly selected regions of the genome showed extensive losses of DNA methylation, with the median level of DNA methylation in DKO1 cells being <1%, 13%, and 9% of the parental cell line, respectively (Figure 1B). We note that randomly selected regions of the genome showed an overall reduction of 89% of their original methylation levels, which is slightly different than the value determined previously (95% reduction in methylation). However, Rhee et al. [16] used a method that monitors percentage methylation of all cytosines in the genome whereas we measured methylated cytosines in the context of CpG dinucleotides that are located in the uniquely mappable, non-repetitive regions of the human genome. As shown in Figure 1B, the promoters in parental HCT116 cells display a wide range of methylation levels. These promoters generally fall into two major groups, those having very high methylation or very low methylation levels (Figure 1C); essentially all promoters have greatly reduced DNA methylation in DKO1 cells.

Because approximately 30,000 promoters can be classified as highly methylated in HCT116 cells, having an average DNA methylation level greater than 50% at the CpGs within -100 to +700 bp of the transcription start site (TSS), we anticipated that a loss in DNA methylation in DKO1 cells at these promoters may reveal previously inaccessible transcription factor binding sites, resulting in a new set of active promoters marked by increased levels of active histones. To determine the effect of losses in DNA methylation on active histone marks, we compared ChIPseq datasets for H3K4me3 and H3K27ac in HCT116 and DKO1 cells. The H3K4me3 data for HCT116 cells was available as part of the ENCODE project [17]. To obtain the other datasets, two biological replicates for each of the H3K4me3 (DKO1), H3K27ac (DKO1), and H3K27ac (HCT116) ChIP-seq samples were produced. The sequencing and peak metrics for all ChIP-seq datasets used in this study are provided in Additional file 1; see Additional file 2 for ChIP-seq peaks in HCT116 cells and Additional file 3 for ChIP-seq peaks in DKO1 cells. Surprisingly, we found that the levels of H3K4me3 and H3K27ac signals were greatly reduced in both replicates of the DKO1 ChIP-seq datasets compared with the HCT116 ChIP-seq replicates. However, a small number of genomic locations do have increases in the levels of H3K4me3 or H3K27ac in DKO1 cells (Figure 2). To quantify these differences, we first identified approximately 12,000 promoter-proximal



Figure 1 Whole genome bisulfite sequencing comparative analysis of HCT116 and DKO1 cells. (A) Light blue and pink tracks represent the sequencing coverage along a segment of human chromosome 19 in HCT116 and DKO1 cells, respectively. Dark blue and red tracks illustrate the percentage of methyl-C/C in HCT116 and DKO1, respectively. Light-colored lines within the percentage methylation (%mCpG) tracks represent the average percentage methylation in the immediate region. CpG islands are shown above the RefSeq genes as green bars. (B) Box plot illustrating the percentage methylation in promoters, gene bodies, and random regions of the genome; the horizontal line in each bar indicates the median value. For HCT116 cells the median values are 30% (promoters), 84% (gene bodies), and 84% (random regions) and for DKO1 cells the median values are <1% (promoters), 13% (gene bodies), and 9% (random regions). (C) The number of promoters containing varying levels of methylation in HCT116 and DKO1 are shown; the minimum and maximum DNA methylation values for the region between -100 and +700 relative to the start site at the promoters in each group is indicated by the color key.

H3K4me3 peaks in both HCT116 and in DKO1 cells. A comparison of the two datasets revealed that very few new promoter-proximal H3K4me3 peaks were identified (Additional file 4). In fact, the overall level of H3K4me3 was greatly reduced at promoters originally unmethylated in HCT116, and promoters losing methylation in DKO1 did not gain H3K4me3 (Figure 3). To determine if other active epigenetic marks were also reduced at these promoter regions, we examined H3K27ac levels. Again, we found that relatively few promoters gained H3K27ac (Additional file 5) and that the active H3K27ac mark was reduced in DKO1 cells at promoters having H3K27ac in HCT116 (Figure 3). It was possible that the loss of DNA methylation did result in more active promoter regions, but somehow also interfered with recruitment of histone modifying enzymes, thus

altering the histone ChIP-seq profiles. Therefore, we next monitored the binding of RNAPII and found that essentially no promoters gained RNAPII in DKO1 cells (Additional file 6) and that, similar to the active histones, the levels of RNAPII also decreased at the vast majority of promoters in the DKO1 cells (Figure 3).

Identification of genes affected by loss of DNA methylation We had expected that the large increase in the number of accessible promoters in DKO1 cells due to the loss of DNA methylation (approximately 30,000 additional unmethylated promoters) would result in the transcriptional activation of a similar number of genes. However, we identified very few promoters that gained RNAPII in DKO1, suggesting that we would not find increased levels of transcripts for many



and light colors represent HCT116 and DKO1 cells, for RNAP II (blues), H3K4me3 (reds), H3K27ac (greens), and H3K36me3 (oranges). Dark and light colors represent HCT116 and DKO1 cells, respectively. Although peaks are reduced in height at most sites, some genomic locations do have increases in H3K4me3 (blue boxes) or H3K27ac (red boxes) in DKO1 cells.



the transcription start sites of promoters that had less than 50% methylation (left) and promoters that had more than 50% methylation in HCT116 cells (right). Light-colored dashed lines represent the marks in DKO1 cells.

genes. Rather, because of the reduction in overall levels of active marks at promoters, we thought it was possible that lack of DNA methylation was causing a global decrease in transcription. We also note that DNA methylation is found in the body of genes and that studies have shown a correlation between gene body methylation and transcript levels [18-20]. Therefore, it was possible that the loss of methylation was causing decreased transcription throughout the genome. To examine the consequences of the loss of DNA methylation on the transcriptome, we performed RNA-seq in HCT116 and DKO1 cells; see Additional file 7 for replicate comparisons of the RNA-seq samples and Additional file 8 for all gene expression values. Even though the relative levels of DNA methylation across gene



bodies is much lower in DKO1 cells (Figure 4A), we found relatively similar transcriptome profiles in the two cell lines (Figure 4B), suggesting that gene body methylation does not play a significant role in regulating gene expression in HCT116 and DKO1 cells.

We did, however, identify three sets of transcripts that were affected by loss of DNA methylation (Additional file 8). One set of transcripts affected by a loss of methylation at promoters, shown in green in Figure 4B, corresponds to 1,089 transcripts that are lowly expressed (log2 normalized expression <1.5) in HCT116 and highly expressed (log2 normalized expression >2.5) in DKO1 cells. The promoters of these genes, which we have termed de-repressed, were highly methylated in HCT116 and unmethylated in DKO1 (Figure 5A). Because there are similar numbers of de-repressed genes as DKO1-specific H3K4me3 peaks (Additional file 4), we thought that perhaps the promoters of the de-repressed genes would show an increase in levels of H3K4me3. Indeed, tag density plots across the promoters of de-repressed genes show that, on average, H3K4me3 is increased at these promoters. However, an overlap analysis showed that only approximately 40% of the de-repressed genes gained a new H3K4me3 peak (Figure 5B). It was possible that the promoters of the 599 de-repressed genes that did not have a new H3K4me3 site were bound by H3K4me3 in HCT116, even though they were silenced. However, we found that only 78 of these 599 promoters were in the set of common H3K4me3 peaks. To investigate the possibility that the remaining 521 de-repressed genes might be utilizing a new (alternative) promoter region, we determined the distance from the TSS of each gene to the nearest DKO1specific H3K4me3 site. Only 48 of these genes had a newly acquired H3K4me3 site within 20 kb, with the median distance from the known TSS to the nearest new H3K4me3 site in DKO1 cells being over 250 kb, suggesting that if



this mechanism is used, the alternative promoters are quite far from the rest of the gene. One hallmark of cancer cell gene expression is the repression of tumor suppressor genes via promoter methylation. Therefore, one would expect that some of the de-repressed genes would be tumor suppressor genes. Of the 1,089 de-repressed genes in DKO1, 38 were identified as tumor suppressor genes present in the current TSGene database [21] (Additional file 9). A Gene Ontology analysis of the entire site of derepressed genes revealed enrichment for zinc finger and Krueppel-associated genes (Figure 5C), many of which are found in large clusters on chromosome 19. Zinc finger genes were particularly enriched in the set of de-repressed genes overlapping new H3K4me3 peaks in DKO1.

The other sets of transcripts that were altered in DKO1 cells are those that were expressed in HCT116 (and thus did not have high levels of DNA methylation at the promoter region in HCT116 cells) but showed expression changes (either up or down) in DKO1 cells (Figure 6A). A Gene Ontology analysis of the 274 genes up-regulated in DKO1 did not reveal any significant gene categories, but 22 genes were identified as tumor suppressor genes present in the TSGene database. Thus, loss of DNA methylation resulted in the up-regulation of tumor suppressor genes by both promoter methylationdependent (38 genes) and promoter methylation-independent (22 genes) mechanisms. Analysis of the 1,366 down-regulated genes showed enrichment for chaperonins (Figure 6B). Chaperonins, which are involved in protein folding, are overexpressed in cancers [22]; our studies suggest reducing global DNA methylation levels may be a suitable option for reducing the levels of these proteins in cancer cells. However, we note that the expression differences of these genes in response to loss of DNA methylation are modest. Unlike the promoters of the de-repressed genes (Figure 5A), analysis of the promoter regions of the upregulated genes (Figure 6C, right panel) did not show an increase in levels of active marks. Strikingly, active histone marks and RNAPII have similar profiles for the promoters of genes up- or down-regulated in DKO1, indicating that perhaps some other mechanism is responsible for the differential regulation of these genes. However, we have also considered that there may be a modest global change in gene expression due to loss of DNA methylation that is difficult to observe using RNA-seq as a read-out. For example, if expression of all genes is modestly increased in DKO1 cells, then the genes identified as 'down-regulated' may simply be those that did not increase as much as the rest of the transcriptome. Because modest global changes are difficult to quantify using current methods, it remains possible that the 1,336 down-regulated genes do not show a loss of active histone marks because they in fact represent a set of genes that simply do not respond to a loss of DNA methylation (with all other genes showing increased expression). Finally, to further classify the promoters of the altered genes, we determined if they were located within a CpG island or if they had a TATA box. We found that for all three categories of genes that responded to loss of DNA methylation, the majority of promoters were categorized as CpG island promoters. However, a smaller percentage of the de-repressed genes were CpG islands (58%) compared with the up-regulated (75%) or down-regulated (83%) genes. For all cases, the CpG islands averaged approximately 1 kb in length. Allowing one mismatch to the TATAWAW motif, we found that few promoters in any class contained a TATA box within -20 to -40 of the TSS (de-repressed: 10%; up-regulated: 10%; down-regulated: 7%).

Loss of DNA methylation has major effects on distal regulatory regions

We were particularly interested in the set of genes whose expression was increased in DKO1 cells but whose promoters were not highly methylated in HCT116 cells. Because the promoters of these genes were not highly methylated in HCT116 cells and did not show large increases in active histone marks in DKO1 cells, we hypothesized that a loss of DNA methylation at enhancer regions may be responsible for the increased transcript levels. To test this hypothesis, we identified active enhancer regions, as defined by H3K27ac regions more than 2 kb from a TSS. Interestingly, although relatively few new active promoter regions were identified in DKO1 cells (Additional files 4, 5 and 6), the enhancer landscape was greatly altered between the two cell types, with many new enhancers in DKO1 cells (Figure 7A; Additional file 10), most of which were highly methylated in HCT116 cells (Figure 7B). Similar to previous studies of DNase I hypersensitive sites, the enhancers in HCT116 and DKO1 cells are evenly divided between intergenic or intragenic locations [23]. Although the overall level of H3K27ac on common enhancers is lower in DKO1 cells, the unique enhancers have higher levels of H3K27ac. While it is difficult to link an enhancer to a specific gene, we reasoned that, in general, enhancers regulate genes in the 'nearby' vicinity (studies from ENCODE have shown that an enhancer loops to the nearest active promoter approximately 50% of the time [17]). Therefore, for the sets of upregulated and down-regulated genes, we calculated the distance to the nearest enhancer that was unique to HCT116, unique to DKO1, or common to both cell lines (Figure 7C). We found that genes that are up-regulated in DKO1 cells have more DKO1-unique enhancers than HCT116-unique enhancers located within 20 kb of their TSS (right panel). In fact, 35% of the up-regulated genes in DKO1 have a new enhancer within 20 kb of the promoter (shown in red). By contrast, only 12% of these genes are within 20 kb of a lost enhancer region (shown in blue).

-log10(p-value) 0 -4 à 0 log2(Fold Change) Down in DKO1 Up in DKO1 (1,366) (274) -log10(Binomial p value) В 3 5 6 0 4 8 Chaperonin TCP-1, conserved site 8.14 Nucleic acid-binding, OB-fold-like Chaperone, tailless complex polypeptide 1 Nucleic acid-binding, OB-fold Chaperonin Cpn60/TCP-1 Proteasome component (PCI) domain 7.31 7.06 6.52 6.17 5.27 С Promoters of down-regulated genes Promoters of up-regulated genes 50 20 K4me3 HCT116 45 18 K4me3 DKO1 K27ac HCT116 peak Tags per bp per peak 16 K27ac DKO1 35 14 Pol2 HCT116 per 30 12 Pol2 DKO1 per bp | 25 10 Tags r HCT116 %mC DKO1 %mC -2500 -2000 -1500 -1000 -500 0 500 1000 1500 2000 2500 -2500 -2000 -1500 -1000 -500 0 500 1000 1500 2000 2500 Distance from transcription start site Distance from transcription start site %Methylation 100 ò 50 Figure 6 (See legend on next page.)

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Figure 6 Identification and characterization of de-regulated genes. (A) The 1,640 genes significantly differentially expressed between HCT116 and DKO1 cells are shown in red (*P*-value <0.05, fold-change >1.2). **(B)** Gene Ontology results for down-regulated genes; up-regulated genes were not enriched for any Gene Ontology terms. **(C)** The epigenetic profiles at the promoters of genes up-regulated (right panel) and down-regulated in DKO1 (left panel).

The median distance of a DKO1-unique enhancer from a gene up-regulated in DKO1 cells is approximately 42 kb, whereas the median distance to an HCT116-unique enhancer is over 200 kb (Figure 7C, right panel). For comparison, we show that the DKO1-unique enhancers have a median distance of approximately 132 kb to the down-regulated genes and are farther from the down-regulated genes than are the HCT116-unique enhancers (Figure 7C, left panel).

As noted above, the median distance between a DKO1unique enhancer and the start site of a gene up-regulated in DKO1 cells was approximately 42 kb, which is similar to the median size of the up-regulated genes. Because enhancers can function in either direction, this suggested that perhaps genes up-regulated in DKO1 cells contained DKO1-specific intragenic enhancers. We determined the number of enhancers within the genes up-regulated (Figure 8A) and down-regulated (Figure 8B) in DKO1 cells. We found that in the genes up-regulated in DKO1 cells, the number of DKO1-specific intragenic enhancers was much higher than the number of HCT-unique enhancers (40% of the genes had a DKO1-specific intragenic enhancer but only 12% of the genes had a HCTspecific intragenic enhancer). In contrast, the number of HCT116-specific versus DKO1-specific enhancers was more similar for the genes down-regulated in DKO1 cells. The relative levels of H3K27ac at the promoter regions did not change for the 111 up-regulated genes that had new intragenic enhancers. However, the H3K27ac marks at the intragenic enhancers was higher in DKO1 than in HCT116 (Figure 9A). A motif analysis of these new intragenic enhancers did not reveal an enrichment for any specific transcription factor binding sites. Of the 310 intergenic enhancers identified within the gene bodies of 111 genes, 285 were identified as intronic and 25 were exonic (Figure 9A). We observed that 65 of the 111 genes that had a DKO1-unique enhancer had more than one new intragenic enhancer. This suggested that perhaps the enhancers were spreading throughout the gene. This is in fact what we observed, and an example is shown in Figure 9B. The SASH1 gene now has marks of active enhancers throughout the transcribed region and is expressed 3.7-fold higher in DKO1 cells.

Discussion

Using DKO1 cells that have a severe reduction in global DNA methylation levels due to genetic deletion of the DNMT1 and DNMT3b genes, we have investigated the Page 9 of 16

global relationship of DNA methylation with histone modifications, RNAPII binding, and gene expression. Although other groups have previously analyzed DNA methylation and gene expression in DKO1 cells [24-26], this had not been done on a genome-wide scale. Because methylated promoters are in condensed chromatin which cannot be accessed by transcription factors and because DNA methylation of recognition motifs has been shown to inhibit transcription factor binding [27,28], we had anticipated that the de-methylation of promoters in DKO1 cells would expose thousands of previously inaccessible binding motifs, many of which would be recognized by transcription factors that are ubiquitously expressed. Therefore, we hypothesized that a reduction in DNA methylation would create thousands of new binding sites for transcription complexes, which would recruit RNAPII, leading to reactivation of these promoters. In contrast, active enhancers are highly cell type-specific and thus it seemed likely that even upon removal of methylation at the distal regulatory regions, few new enhancers would be created because the cell type-specific factors would not be present in DKO1 cells to bind to the motifs and recruit histone acetylation complexes. In contrast, we found that relatively few promoters were activated in DKO1 cells but thousands of newly active enhancers (which were highly methylated in parental HCT116 cells) were created. Interestingly, 3,008 (47%) of the 6,376 new enhancers that appeared in DKO1 cells had the H3K4me1 mark of a poised enhancer in the parental HCT116 cell line. Our studies indicate that 1) in general, loss of DNA methylation does not lead to the acquisition of newly active promoters, suggesting that DNA methylation is not the primary driver of promoter repression, and 2) loss of DNA methylation has a major effect on promoter-distal regulatory regions, uncovering intragenic enhancers within genes whose expression increases upon loss of DNA methylation.

We investigated the effect of loss of DNA methylation on histone modifications at promoters, enhancers, and gene bodies. We found that overall levels of all active histone marks were reduced at most promoters in DKO1 cells, regardless of their methylation status in HCT116 cells. The reason for this overall decrease in active promoter marks in DKO1 cells is not clear. However, DKO1 cells do grow slightly more slowly than HCT116 cells [16] and it is possible that small differences in the percentage of cells in S phase may influence



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Figure 7 Loss of DNA methylation uncovers thousands of distal H3K27ac sites. (A) Venn diagram comparing distal H3K27ac peaks in HCT116 and DKO1 cells. (B) DNA methylation status of common and unique enhancers. Pie charts show the breakdown of genomic locations for peaks residing within intergenic, intronic, and exonic regions of the genome. Tag density plots show the density of H3K27ac tags relative to the centers of these peaks. (C) Distances from the TSS of genes up-regulated (right panel) or down-regulated (left panel) in DKO1 to the nearest categorized enhancer. Also indicated is the median distance from each category of enhancer to the nearest up-regulated or down-regulated gene.

ChIP-seq analysis of promoters. Conversely, we identified thousands of new enhancers that have increased levels of H3K27Ac in DKO1 cells, indicating that the ChIP-seq assay is able to detect high levels of modified histones in DKO1 cells. These results suggest that DNA methylation is in fact a primary regulator of the activity of enhancers. In addition, we found only modest changes in H3K36me3 or H3K9me3 levels in HCT116 versus DKO1 cells (data not shown). Our previous studies using ChIP-chip had suggested that DNA methylation





may be required for H3K9me3 deposition [29]. In fact, follow-up studies using ChIP-seq showed a major reduction in H3K9me3 across the entire chromosome 19 (PJ Farnham and S Iyengar, unpublished data). However, in those studies DNA methylation levels were reduced by treatment of cells with the DNMT inhibitor 5-Aza-CR. It is likely that the effect on H3K9me3 may have been due to redistribution of the KAP1/SETDB1 histone methyltransferase complex due to the activation of the DNA damage response and not directly due to loss of DNA methylation [30]. We now show that the H3K9me3 patterns are essentially the same in HCT116 and DKO1 cells. An analysis of chromosome 19 in DKO1 cells using a ChIP-chip assay [26] is in agreement with our ChIP-seq data showing that H3K9me3 and H3K36me3 are not dramatically affected in DKO1 cells. Thus, interpretation of the mechanisms leading to changes in histone marks caused by 5-Aza-CR must be made with caution.

Unexpectedly, we found that very few genes increased in expression in DKO1 cells. However, we did identify two sets of genes whose expression increased by different mechanisms in these cells (Figure 10). One set of genes was up-regulated due to the removal of DNA methylation from their promoters (de-repressed genes). The other set of up-regulated genes had unmethylated, active promoters in HCT116 cells, but showed increases in overall transcript levels in the DKO1 cells. Interestingly, the later set of genes contained multiple active enhancers within their gene bodies in DKO1 cells, and these same enhancers were methylated in HCT116 cells. Intronic enhancers have been previously described to regulate the genes they reside within [23,31-33]. Here, we have identified several hundred genes that may be controlled by increases in active histones at intragenic enhancers. However, further functional studies in which these



enhancer regions are knocked out or repressed will be required to truly determine whether these intronic enhancers are responsible for the activation of these genes in DKO1 cells. Both the de-repressed and the up-regulated genes included tumor suppressor genes, and the resulting mutant cells display characteristics of normal cells (slightly slower doubling times relative to HCT116) as was previously shown by Rhee *et al.* [16]. Thus, although reducing DNA methylation has modest effects on gene expression, perhaps it will prove suitable as a therapeutic target.

Conclusions

While DNA methylation can play a role in the repression of gene expression, our studies indicate that it is likely not the key determinant in the regulation of most promoters in HCT116 cells. Our finding that loss of DNA methylation does not result in the acquisition of active histone marks or an increase in gene expression from most de-methylated promoters suggests that these promoters may remain in a closed or condensed confirmation. This interpretation is supported by other studies showing that promoters that lose DNA methylation in DKO1 cells do not gain accessibility to a specific restriction enzymes [34] and do not gain nucleosomedepleted regions (FD Lay, Y Liu, and BP Berman, personal communication). Thus, DNA methylation may be a consequence of, not causation for, promoter silencing. However, DNA methylation plays a much greater role in the silencing of distal regulatory elements, with many of the enhancers activated by loss of DNA methylation falling within the bodies of their probable target genes.

Materials and methods

Cell growth conditions

The human cell lines HCT116 (ATCC #CCL-247) and DKO1 [16] were grown in McCoy's 5A Medium supplemented with 10% fetal bovine serum and 1% penicillin/ streptomycin and were harvested for downstream experiments at 80% confluence.

Whole genome bisulfite sequencing

Genomic DNA was collected from HCT116 and DKO1 cells using a Qiagen (Valencia, CA, USA) QIAeasy DNA mini kit. Genomic DNA (2 µg) was sonicated using a Covaris to an average molecular weight of 150 bp. Achievement of the desired size range was verified by Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA) analysis. Fragmented DNA was repaired to generate blunt ends using the END-It kit (Epicentre Biotechnologies, Madison, WI, USA) according to the manufacturer's instructions. Following incubation, the treated DNA was purified using AmpureX beads (Beckman Coulter, Brea, CA, USA). In general, magnetic beads were employed for all nucleic acid purifications in the following protocol. Following end repair, A-tailing was performed using the NEB dA-tailing module according to the manufacturer's instructions (New England Biolabs, Ipswich, MA, USA). Adapters with a 3' 'T' overhang were then ligated to the end-modified DNA. For whole genome bisulfite

sequencing, modified Illumina paired-end adapters were used in which cytosine bases in the adapter are replaced with 5-methylcytosine bases. Depending on the specific application, we utilized either Early Access Methylation Adapter Oligos that do not contain barcodes, or the adapters present in later versions of the Illumina DNA Sample Preparation kits, which contain both indices and methylated cytosines. Ligation was carried out using ultrapure, rapid T4 ligase (Enzymatics, Beverly, MA, USA) according to the manufacturer's instructions. The final product was then purified with magnetic beads to yield an adapter-ligation mix. Prior to bisulfite conversion, bacteriophage lambda DNA that had been through the same library preparation protocol described above to generate adapter-ligation mixes was combined with the genomic sample adapter ligation mix at 0.5% w/w. Adapter-ligation mixes were then bisulfite converted using the Zymo DNA Methylation Gold kit (Zymo Research, Orange, CA, USA) according to the manufacturer's recommendations. Final modified product was purified by magnetic beads and eluted in a final volume of 20 µl. Amplification of one-half the adapter-ligated library was performed using HiFi-U Ready Mix (Kapa Biosystems, Wilmington, MA, USA) for the following protocol: 98° for 2 minutes; then six cycles of 98° for 30 s, 65° for 15 s, 72° for 60 s; with a final 72° 10-minute extension, in a 50 μ l total volume reaction. The final library product was examined on the Agilent Bioanalyzer, then quantified using the Kapa Biosystems Library Quantification kit according the to manufacturer's instructions. Optimal concentrations to get the right cluster density were determined empirically but tended to be higher than for non-bisulfite libraries. Libraries were plated using the Illumina cBot and run on the Hi-Seq 2000 according to the manufacturer's instructions using HSCS v.1.5.15.1. Image analysis and base calling were carried out using RTA 1.13.48.0, and deconvolution and fastq file generation were carried out using CASAVA_v1.7.1a5. Raw reads were mapped using Bis-SNP [35], and percentage methyl-C/C was calculated for every CpG dinucleotide in the human genome. All CpG dinucleotides with a minimum sequencing coverage of 3× were used for downstream analyses. To determine the average DNA methylation surrounding a set of promoters or enhancers, HOMER was used to calculate the percentage methylation across promoter or enhancer regions using the annotatePeaks.pl script and the '-ratio' option for 2,500 bp surrounding the regions of interest using a bin size of 100 bp. The resulting bins were plotted as a heatmap using heatmap.2 in R.

ChIP-seq

ChIP assays were was performed in replicate for H3K4me3 (Cell Signaling Technology, Danvers, MA, USA; 9751S, lot# 4), H3K27ac (Active Motif, Carlsbad, CA, USA; #39133, lot# 21311004), and RNAPII (Covance, Princeton,

NJ, USA; MMS-126R (8WG16), Lot# D12LF0314) in DKO1 cells and one replicate of RNAPII ChIP-seq was performed in HCT116 cells, as previously described [36]. The ChIP-seq libraries were sequenced on a HiSeq2000; the H3K4me3 datasets from HCT116 were available via ENCODE and were downloaded from the UCSC browser, accession number [UCSC: wgEncodeEH000949]. All ChIPseq data were mapped to hg19 using BWA (default parameters) and peaks were called using Sole-Search [37,38] with the following parameters: Permutation:5; Fragment:250; AlphaValue: 0.00010 = 1.0E-4; FDR: 0.00010 = 1.0E-4; Peak-MergeDistance:0; HistoneBlurLength:1200 for H3K4me3 and H3K27ac. Each replicate ChIP-seg dataset was analyzed separately and only peaks present in both replicates were used for the subsequent analyses; see Additional file 1 for ChIP-seq reproducibility measures. Peaks were separated by their proximity to promoters, with promoterproximal peaks defined as those found within ±2 kb of a TSS and promoter-distal as everything else; enhancers were defined as promoter-distal H3K27ac sites. To create TSS-centered tag density plots, mapped reads were used with the HOMER annotatePeaks script [39] and the -hist option to average ChIP-seq tags relative to all (or in some cases to a subset of) RefSeq TSSs. To create enhancercentered tag density plots, the center of the H3K27ac peaks were used. The mergePeaks option in HOMER was used to determine overlapping H3K27ac regions between HCT116 and DKO1 at proximal and distal sites. H3K27ac peak proximity to TSSs was determined using annotate-Peaks. Gene Ontology analysis was performed using Stanford's Genomic Regions Enrichment of Annotations Tool (GREAT) [40].

RNA-seq

RNA-seq was performed for HCT116 and DKO1 in replicate. RNA was collected from cells using Trizol (Life Technologies/ThermoFisher, Waltham, MA, USA; catalog #15596018) and paired-end libraries were prepared using the Illumina TruSeqV2 Sample Prep Kit (catalog #15596-026), starting with 1 µg total RNA. Libraries were sequenced on an Illumina Hi-Seq 2000. RNA-seq data were analyzed with Partek Flow version 3 (Partek Inc., St Louis, MO, USA). Raw reads were trimmed using the Quality Score method and mapped to hg19 (Ensembl 72) using Tophat2 [41]. Gencode V17 annotation was used to quantify the aligned reads using the Partek E/M method. Quantified reads were normalized using TMM in EdgeR [42] and then analyzed for differential expression using Partek's Gene Specific Analysis. Differentially expressed genes were determined as those with a P-value <0.05 and a fold change >1.2. Gene expression scatter and volcano plots were created using ggplot2 in R. To address concerns about RNA and/or libraries, we prepared two replicates of RNA from each cell type; a comparison of the replicates

for HCT116 and for DKO1 shows that the RNA-seq data sets from the two replicates are highly reproducible (Additional file 7). In addition, visual inspection of the RNA-seq reads on a genome browser (with no normalization) shows very similar RNA profiles in HCT116 and DKO1 cells, except for the set of genes that were identified to have altered expression.

Data access

WGBS and RNA-seq datasets for HCT116 and DKO1 cells were produced for this manuscript and are deposited in Gene Expression Omnibus (GEO) [GEO: GSE60106]. H3K27ac and H3K4me3 ChIP-seq datasets in DKO1 cells and RNAPII ChIP-seq data for both HCT116 and DKO1 cells were produced for this manuscript and are deposited in GEO [GEO: GSE60106]. H3K4me3 [UCSC: wgEnco-deEH000949], H3K27ac [UCSC: wgEncodeEH002873], H3K4me1 [UCSC: wgEncodeEH002874] and part of the RNAPII [UCSC: wgEncodeEH000651] HCT116 ChIP-seq data were produced as part of the ENCODE Consortium [14] and can be downloaded from the UCSC genome browser.

Additional files

Additional file 1: Sequencing and peak metrics for all ChIP-seq, RNA-seq, and WGBS datasets. ChIP-seq reads were mapped using BWA and peaks were called using Sole-Search [37,38]. All data was collected as part of this study except for H3K4me3 from HCT116, which was from ENCODE. However, for consistency, the reads were remapped and peaks were called for this dataset using Sole-Search. High-confidence (HC) peaks were determined as those present in two independent biological replicates. Promoter-proximal or promoter-distal peaks were selected by their proximity to a TSS, with proximal being those peaks within 2000 bp of a TSS and distal being everything else.

Additional file 2: ChIP-seq peaks in HCT116 cells. Worksheet containing all peaks called using Sole-Search for H3K27ac (Sheet 1), H3K4me3 (Sheet2) and RNAPII (Sheet 3) in HCT116 cells.

Additional file 3: ChIP-seq peaks in DKO1 cells. Worksheet containing all peaks called using Sole-Search for H3K27ac (Sheet 1), H3K4me3 (Sheet2) and RNAPII (Sheet 3) in DKO1 cells.

Additional file 4: Characterization of H3K4me3 promoter-proximal peaks in HCT116 and DKO1 cells. Venn diagram showing differences in binding sites for promoter-proximal H3K4me3 peaks (top), and the density of H3K4me3 ChIP-seq tags in HCT116 and DKO1 for all three peak categories (bottom).

Additional file 5: Characterization of H3K27ac promoter-proximal peaks in HCT116 and DKO1 cells. Venn diagram showing differences in binding sites for promoter-proximal H3K27ac peaks (top), and the density of H3K27ac ChIP-seq tags in HCT116 and DKO1 for all three peak categories (bottom).

Additional file 6: Characterization of RNAP2 promoter-proximal peaks in HCT116 and DKO1 cells. Venn diagram showing differences in binding sites for promoter-proximal RNA polymerase 2 peaks (top), and the density of RNA polymerase 2 ChIP-seq tags in HCT116 and DKO1 for all three peak categories (bottom).

Additional file 7: Replicate RNA-seq analysis. log2 expression values are plotted comparing replicates for of the same cell type, and across cell types for HCT116 and DKO1 cells.

Additional file 8: RNA-seq analysis of HCT116 and DKO1 cells.

Worksheet containing the output from RNA-seq analysis using Partek Flow version 3.0.14.0321. Sheet 1 contains Genes, statistics, and expression values for all transcripts from the gene-specific analysis. Sheet 2 lists all de-repressed genes, and their expression values in HCT116 and DKO1. Sheet 3 lists all up-regulated genes and their expression values in HCT116 and DKO1. Sheet 4 lists all down-regulated genes and their expression values in HCT116 and DKO1.

Additional file 9: Differentially expressed tumor suppressor and zinc finger genes. Sheet 1 lists all de-repressed tumor suppressor genes and their expression values in HCT116 and DKO1. Sheet 2 lists all up-regulated tumor suppressor genes and their expression values in HCT116 and DKO1. Sheet 3 lists all de-repressed zinc finger genes and their expression values in HCT116 and DKO1.

Additional file 10: ChIP-seq peaks of enhancer regions. Sheet 1 lists all H3K27ac peaks that are unique to HCT116 cells. Sheet 2 lists all H3K27ac peaks that are unique to DKO1. Sheet 3 lists all H3K27ac peaks that are common to HCT116 and DKO1. Sheet 3 lists all H3K27ac peaks present within intragenic regions of genes up-regulated in DKO1.

Abbreviations

5-Aza-CR: 5-azacytidine; bp: base pair; ChIP: chromatin immunoprecipitation; DNMT: DNA methyltransferase; GEO: Gene Expression Omnibus; RNAPII: RNA polymerase II; TSS: transcription start site; WGBS: whole genome bisulfite sequencing.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

AB prepared samples, performed analyses for figures, and assisted in drafting the manuscript, LY performed analyses for figures, HW and YG performed ChIP-seq experiments, CMN and BPB carried out the DNA methylation library methods development, PJF conceived of the project, advised on all experiments and drafted the manuscript. All authors read and approved the final manuscript.

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