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Flow-dependent epigenetic DNA methylation regulates endothelial gene expression and atherosclerosis

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In atherosclerosis, plaques preferentially develop in arterial regions of disturbed blood flow (d-flow), which alters endothelial gene expression and function. Here, we determined that d-flow regulates genome-wide DNA methylation patterns in a DNA methyltransferase–dependent (DNMT-dependent) manner. Induction of d-flow by partial carotid ligation surgery in a murine model induced DNMT1 in arterial endothelium. In cultured endothelial cells, DNMT1 was enhanced by oscillatory shear stress (OS), and reduction of DNMT with either the inhibitor 5-aza-2′-deoxycytidine (5Aza) or siRNA markedly reduced OS-induced endothelial inflammation. Moreover, administration of 5Aza reduced lesion formation in 2 mouse models of atherosclerosis. Using both reduced representation bisulfite sequencing (RRBS) and microarray, we determined that d-flow in the carotid artery resulted in hypermethylation within the promoters of 11 mechanosensitive genes and that 5Aza treatment restored normal methylation patterns. Of the identified genes, HoxA5 and Klf3 encode transcription factors that contain cAMP response elements, suggesting that the methylation status of these loci could serve as a mechanosensitive master switch in gene expression. Together, our results demonstrate that d-flow controls epigenomic DNA methylation patterns in a DNMT-dependent manner, which in turn alters endothelial gene expression and induces atherosclerosis.

Introduction

Endothelial cells undergo dramatic gene expression changes when exposed to disturbed blood flow (d-flow) as compared with unidirectional, stable blood flow (s-flow) (1–4). Atherosclerosis preferentially develops in areas of d-flow, where the dysfunctional endothelial cell phenotype initiates and perpetuates plaque development (5–7). S-flow upregulates “atheroprotective” genes and downregulates “proatherogenic” genes, while d-flow enhances proatherogenic genes and suppresses atheroprotective genes. However, the mechanisms by which d-flow causes changes in endothelial cell gene expression are still unclear.

Gene expression can be regulated epigenetically by histone modifications, DNA methylation, and microRNAs (miRNAs) (8). Of these, flow has been shown to regulate gene expression by histone modifications (9, 10) and miRNAs (11–18). It is not known, however, whether flow regulates DNA methylation patterns and whether this plays a critical role in mechanosensitive gene expression.

DNA methylation is the most stable epigenetic modification and involves the addition of a methyl group to the 5′ carbon of a cytosine base pair that occurs most often in a CG dinucleotide (CG site) (20, 21). CpG islands are dense regions of CG sites that are normally unmethylated and are associated with approximatively 40% of human genes (22–26). However, repetitive elements such as Alu, Line1, and B1 are generally highly methylated (24). DNA methylation in the promoter region of a gene, near the transcription start site (TSS), is associated with repression of gene expression (27–29).

DNA methyltransferases (DNMTs) catalyze the addition of the methyl group to cytosine. DNMT1 is classically referred to as a maintenance methylase (it preferentially methylates hemimethylated DNA), although it also has de novo methylation capabilities (30, 31). DNMT3a and DNMT3b are referred to as de novo methyltransferases that preferentially add methyl groups to fully unmethylated DNA during development (32). DNA methylation is a gene-regulatory mechanism known to play a key role in various diseases, particularly in cancer, by silencing tumor suppressor genes via aberrant hypermethylation, and drugs that inhibit DNA methyltransferases have proven to be promising treatment options. 5-Aza-2′-deoxycytidine (5Aza, also known as decitabine) is a nucleoside analog that traps DNMT1 in a covalent complex with DNA, and also preferentially targets DNMT1 via ubiquitin-dependent proteasomal degradation, resulting in DNMT1 inhibition (33). 5Aza is an FDA-approved drug and is currently used to treat myelodysplastic syndromes including leukemia (33–38), but its specific mechanism of action and gene targets need to be further determined.

Recently, DNA methylation has been implicated as a novel risk factor for atherosclerosis in smooth muscle cells (39–43). Hypermethylation of the estrogen receptor α (ER-α) promoter, and high plasma homocysteine levels, a source of the methyl group used for DNA methylation, were found in atherosclerosis patients (44, 45). 5-Methylcytosine (5mC) is elevated in the intima of plasma homocysteine-fed a Western diet, and high 5mC levels are linked to LDL receptor and p53 mutation in vascular cells (43, 46). 15-Lipoxygenase,
a gene implicated in oxidative modification of LDL, is also regulated by DNA methylation (47). It is unknown, however, whether the proatherogenic mechanical force caused by d-flow regulates epigenetic DNA methylation.

Here, we show that DNMT1 is induced by d-flow in endothelial cells both in vivo and in vitro. The DNMT inhibitor 5Aza prevented endothelial inflammation in vitro and atherosclerosis development in vivo. The genome-scale studies demonstrated that d-flow induces genome-wide DNA methylation pattern changes and revealed novel mechanosensitive genes, such as HoxA5, that are epigenetically regulated and contain cAMP response elements (CREs).

Results

D-flow stimulates DNMT1 expression in endothelial cells in vivo and in vitro. We initially identified DNMT1 as a potential mechanosensitive gene in endothelial cells from our previous in vivo mRNA array study (48). To identify flow-sensitive genes in arterial endothelium in vivo, endothelial-enriched RNA was obtained directly from mouse carotid arteries 2 days after partial carotid ligation surgery in C57BL/6 mice (Figure 1A). The mRNA array showed that DNMT1 expression was approximately 2.4-fold higher in the left carotid artery (LCA), which was partially ligated and exposed to d-flow for 48 hours, than in the contralateral right common carotid artery (RCA), exposed to s-flow (48). Here, we validated this endothelial microarray data by quantitative PCR (qPCR), immunostaining, and Western blotting both in vivo and in vitro.

DNMT1 expression showed a significant increase, more than 2-fold, in the LCA endothelium as compared with that of the RCA at 48 hours after partial carotid ligation (Figure 1B). Similarly, robust DNMT1 protein expression was observed in the flow-disturbed LCA (48 hours after ligation) compared with the contralateral RCA and sham-controlled LCA and RCA (Figure 1C). Importantly, a robust DNMT1 protein expression was also observed in endothelial cells in the lesser curvature of the aortic arch, which is naturally and chronically exposed to d-flow without the ligation surgery, but not in the greater curvature, an atherosclerosis-resistant region, which is naturally exposed to s-flow (Figure 1, A and D). Interestingly, DNMT1 expression was increased in the intimal and medial layers of the LCA (Figure 1C), suggesting that d-flow affected the protein expression in both endothelial and smooth muscle cells. These in
...hypothesis that d-flow induces hypermethylation of antiatherogenic endothelial cell genes at the promoter regions, thereby silencing their expression, and that 5Aza treatment prevents this mechanism by which 5Aza inhibited atherosclerosis in vivo, we tested the hypothesis that d-flow induces genome-wide DNA methylation changes and regulates global gene expression in a 5Aza-dependent manner. To determine the mechanisms by which 5Aza inhibited atherosclerosis in vivo, we used 2 different ApoE−/− mouse models to study the role of DNMT in atherosclerosis: the acute partial carotid ligation model with Western diet, which rapidly develops atherosclerosis within 3 weeks (49), and the conventional Western diet-fed model that chronically develops atherosclerosis in 3 months (19, 50). In the carotid ligation model, the LCA rapidly developed atherosclerosis within 3 weeks (Figure 3A), and this was inhibited by 5Aza treatment in a dose-dependent manner with 0.2 mg/kg/d being the lowest effective dose (Figure 3, A and B). Immunostaining for the leucocyte marker CD45 showed a remarkable inhibitory effect of 5Aza on inflammatory cell infiltration at the 0.2-mg/kg/d dose (Supplemental Figure 3A), which was consistent with the atherosclerosis data. In the conventional Western diet model, 5Aza treatment (0.2 mg/kg/d) also significantly inhibited atherosclerosis development in the aortic arch and brachiocephalic arterial branch (Figure 3, C and D). In both models, 5Aza treatment did not show a significant effect on the serum lipid profile (Supplemental Figure 4, A and C) and apparent health, but induced a smaller body-weight gain than was seen in the control groups (acute study: 25 g in control vs. 23 g in 5Aza mice at 3 weeks; chronic study: 34 g in control vs. 31 g in 5Aza group at 3 months) (Supplemental Figure 4, B and D). As additional controls, we examined the effect of d-flow, atherosclerosis, and 5Aza treatment on 5mC content and DNMT1 expression by immunostaining. The dramatically increased 5mC and DNMT1 levels in the partially ligated LCA at the 3-week time point were significantly reduced by 5Aza treatment in a dose-dependent manner (Supplemental Figure 3, B and C). Taken together, these results provide strong evidence that 5Aza treatment prevented atherosclerosis by reducing DNMT expression and activity in the arterial wall.

D-flow induces genome-wide DNA methylation changes and regulates global gene expression in a 5Aza-dependent manner. To determine the mechanisms by which 5Aza inhibited atherosclerosis in vivo, we tested the hypothesis that d-flow induces hypermethylation of antiatherogenic endothelial cell genes at the promoter regions, thereby silencing their expression, and that 5Aza treatment prevents this hypermethylation, leading to prevention of atherosclerosis. Since it was not known whether d-flow regulates genome-wide DNA meth-
ylation patterns in endothelial cells, we first carried out a reduced representation bisulfite sequencing (RRBS) study using endothelial-enriched genomic DNA (gDNA) obtained from the RCAs and LCAs of partially ligated mice. To further determine whether the flow-dependent DNA methylation changes were regulated in a DNMT-dependent manner, mice treated with 5Aza were compared with a saline vehicle control. Moreover, to determine which mechanosensitive genes were downregulated, potentially by DNA hypermethylation at their promoter, we carried out a concomitant gene transcript microarray study using endothelial-enriched RNA obtained from the LCAs and RCAs of partially ligated mice treated with either saline (RCA vs. LCA) or 5Aza (Aza-RCA vs. Aza-LCA). To rule out the effect of confounding factors such as the hypercholesterolemic condition in ApoE−/− mice, C57BL/6 mice were used for these studies. Mice pretreated with 5Aza (0.2 mg/kg/d) or saline for 2 weeks were partially ligated, and the 5Aza treatment was continued for 1 week after ligation. For the RRBS with next-generation sequencing and gene transcript array studies, gDNA from 20 RCAs versus LCAs and total RNA from 3 RCAs versus LCAs were pooled, respectively. As a control for 5Aza treatment, blood was collected 1 week after the start of the treatment, and global DNA methylation was measured by high-resolution melt curve (HRM) analysis. This showed that global methylation decreased as early as 1 week after 5Aza treatment, demonstrating the efficacy of this treatment strategy (Supplemental Figure 5, A and B).

The 4 RRBS data sets (RCA, LCA, Aza-RCA, and Aza-LCA) were mapped to the mm9 mouse genome assembly (Build 37, NCBI, July 2007), which contains a total of 21,342,492 CG sites. Our 4 RRBS data sets showed a methylome coverage of approximately 23% (4,967,716 unique CG sites on average) of all CG sites in the mouse genome, including promoters, gene bodies, and intergenic regions (Supplemental Table 1). Interestingly, approximately 53% (1,125,997 CG sites on average) of all CG sites in the promoter regions (defined as transcription start site ± 1 kb) were covered by our RRBS data sets (Supplemental Table 1), which shows a higher coverage in the promoter regions as compared with the other genomic sites.

To measure the DNA methylation changes at each CG site in the RRBS data sets, only those covered in all 4 data sets (RCA, LCA, Aza-RCA, and Aza-LCA) were selected for further analysis (3,232,969 unique CG sites). The methylation ratio at each CG site was compared between the RCA and LCA as well as the Aza-RCA and Aza-LCA using heat-map correlation scatter plots (Figure 4, A and B). Of the 3.2 million CG sites examined, 4.1% (131,176 CG sites) showed hypermethylation by more than 40% in the LCA as compared with the RCA, whereas 93% showed no significant difference in methylation, defined as less than a 40% difference between the RCA and LCA (Figure 4A). 5Aza treatment reduced the number of hypermethylated CG sites in the LCA by more than 50% as compared with the saline group (from 131,176 to 55,224 CG sites), while 97% of the CG sites showed no significant difference between the Aza-RCA and Aza-LCA (Figure 4B). This result showed that the small number (4.1%) of CG sites that were hypermethylated in the RCA by d-flow was substantially reduced by the 5Aza treatment (to 1.7%), suggesting that d-flow regulates genome-wide DNA methylation patterns in a DNMT-dependent manner. This result was further corroborated by an a priori methylation pattern analysis that
was performed by ranking the methylation ratio of each gene promoter among the 4 groups. This revealed a pronounced emergent methylation pattern, wherein gene promoters in the LCA (2,086 out of 16,493 genes) were hypermethylated as compared with those in the RCA, Aza-LCA, and Aza-RCA (listed in decreasing order of methylation ratios; Supplemental Figure 6A). This supports our hypothesis that d-flow induces promoter hypermethylation in a DNMT-dependent manner.

To determine how many of these mechanosensitive DNA methylation patterns at the promoters correlate to gene expression patterns, we compared the promoter DNA methylation data with the gene transcript array data. First, the gene array data analysis showed that expression of 1,319 genes (569 down- and 750 upregulated) changed by more than 33% in the LCA compared with the RCA (Figure 4C and Figure 5A). Treatment with 5Aza decreased the number of differentially expressed genes to 385 (152 down- and 233 upregulated in the LCA) between the Aza-RCA and Aza-LCA (Figure 4D). Importantly, 5Aza treatment rescued the downregulation of 540 of the 569 mechanosensitive genes in the LCA (Figure 4D and Figure 5A), demonstrating the dominant effect of 5Aza treatment on mechanosensitive gene expression, either directly or indirectly (Figure 4, C and D, Figure 5A, and Supplemental Figure 6, D and E).

Further analysis revealed that our RRBS and gene array data sets showed the expected pattern, wherein percentage promoter methylation inversely correlated with gene expression levels on a global scale (Supplemental Figure 7A), whereas the same trends were not observed for the gene body and 3′ gene regions surrounding the transcription termination sites (TTSs). This finding is consistent with previous reports showing a close correlation between promoter hypermethylation and gene silencing (51), validating that our RRBS and transcriptome data sets behave in an expected and reliable manner. In addition, as expected, we found that overall promoter CGs were far less methylated (~39%) in comparison with genome-wide CGs (~78%) (Supplemental Figure 7B). Contrary to our hypothesis, however, the genome-wide and promoter-wide CG methylation status was similar regardless of flow conditions and 5Aza treatment in the RCA, LCA, Aza-RCA, and Aza-LCA (Supplemental Figure 7B), although this result is consistent with the small difference in the number of hypermethylated CG sites on the genome-wide level in LCA and Aza-LCA groups, shown in Figure 4, A and B. These results suggest a possibility that the 5Aza effect may be specific and limited to a relatively small number of genes.

Next, we compared the list of mechanosensitive genes that were hypermethylated at their promoter and downregulated in the LCA in a 5Aza-dependent manner to determine the genes that exhibit...
the expected correlation between methylation and gene expression. This comparative analysis revealed 11 flow-sensitive genes that were hypermethylated at their promoter and silenced by d-flow that could be rescued by 5Aza (Figure 5, A and B, and red points in Figure 4, C and D). These included HoxA5, Tmem184b, Adamts15, Klf3, Cmklr1, Pkp4, Acrvl1, Dok4, Spry2, Zfp46, and F2rl1. Notably, HoxA5 showed the most robust flow-dependent changes in promoter methylation and gene silencing that was significantly reversed by 5Aza treatment, indicating the potential importance of DNMT-dependent mechanosensitive gene regulation by promoter methylation. Detailed analysis of the RRBS DNA methylation data in the HoxA5 gene cluster is shown in Figure 6A using a University of California, Santa Cruz, genome browser track to display our RRBS data at the nucleotide resolution. The HoxA5 promoter region was clearly hypermethylated in the d-flow LCA as compared with the s-flow RCA, the Aza-LCA, and the Aza-RCA (Figure 6B). This RRBS result was further confirmed by manual bisulfite sequencing of additional endothelial gDNA obtained from the RCA, LCA, Aza-RCA, and Aza-LCA, validating the RRBS data (Figure 7A). Further, we validated the HoxA5 expression data obtained from the microarray study (Figure 7B) by qPCR (Figure 7C). The qPCR result showed that HoxA5 expression was downregulated in the LCA by more than 50% compared with that in the RCA and was reversed by the 5Aza treatment, validating the microarray result.

Promoter CREs are hypermethylated by d-flow in a 5Aza-dependent manner. To explore whether there are any functional regulatory elements that are common within the promoter regions among the 11 genes, we first searched for transcription factors that are known or predicted to regulate these genes. The systems biological analysis using MetaCore with these 11 genes as the input predicted CREB1 (cAMP response element binding protein) as the transcription factor with the highest potential connectivity. This led us to a computational sequence analysis designed to look for the full palindromic or half CRE sequences (TGACGTCA, TGACG, and CGTCA) within the differentially methylated regions (DMRs) of the 11 gene promoters. This search revealed that 5 of the 11 gene promoter DMRs contained CRE sites, including HoxA5, Klf3, Cmklr1, Acrvl1, and Spry2. We then examined the DNA methylation status of these gene promoter DMRs specifically at the CRE CG site and found that HoxA5, Klf3, Cmklr1, and Acrvl1 indeed showed significant hypermethylation at this CG in the LCA, which was prevented by 5Aza treatment (Figure 8, A–D). For example, percentage methylation of the HoxA5 promoter CRE CG site was dramatically higher in the LCA (55.6%) as compared with the RCA (11.5%), the Aza-LCA (17.2%), and the Aza-RCA (20.0%). The promoter CRE CG site in Spry2, however, was not fully covered by our RRBS data; therefore we could not determine its flow- and 5Aza-dependency.

We next examined whether the CREs across the genome display this property of d-flow–induced hypermethylation that can be prevented by 5Aza. We found that CRE sequences across the genome were nearly fully methylated (95%–100%) in all 4 groups (Figure 8E). However, CREs within just the promoter regions showed significantly less methylation compared with that of genome-wide CREs (Figure 8F). In the LCA, the average percentage methylation of promoter CREs was 8.7%, which was decreased by approximately 2% in the Aza-LCA (Figure 8F). In both untreated and 5Aza-treated RCAs,
promoter CRE methylation was nearly zero, indicating the dramatic effect of s-flow in preventing methylation of promoter CREs in the RCA (Figure 8F). It is interesting to note that 5Aza treatment showed only a small reduction in average promoter CG site methylation, while it dramatically decreased CG methylation at specific gene promoters such as HoxA5, Klf3, Cmklr1, and Acvrl1, indicating a targeted effect of 5Aza toward some gene promoters containing CREs.

Given the potential role of the transcription factors HoxA5 and Klf3 as master regulators of gene expression networks, we further examined their relevance to the endothelial flow response. We found that HoxA5 is strongly suppressed in response to d-flow in vivo by microarray and qPCR (Figure 7, B and C, respectively), and to OS in HUVECs in vitro at both the mRNA and the protein level (Figure 9, A and B). Similarly, Klf3 mRNA expression is also suppressed by d-flow in vitro (Figure 9A). We were not able to determine Klf3 protein because of a lack of adequate antibody for Western blot. To analyze the role of these transcription factors in endothelial inflammation, we performed monocyte adhesion assays using siRNAs to HoxA5 and Klf3 in HUVECs under shear. We found that in the LS condition, siRNA-mediated knockdown of HoxA5 significantly increased monocyte adhesion to HUVECs, partially simulating the OS condition of increased inflammation (Figure 9C and Supplemental Figure 8A). However, knockdown of Klf3 did not have a significant effect on endothelial inflammation, suggesting its potential importance in other flow-responsive pathways (Figure 9D and Supplemental Figure 8A).

**Discussion**

Here, we demonstrate, for the first time to our knowledge, that blood flow epigenetically controls endothelial gene expression by regulating genome-wide DNA methylation patterns via a DNMT-dependent mechanism. This novel mechanism has broad implications for understanding the atherogenesis mechanisms and identifying potential therapeutic targets of atherosclerosis. DNMT1 is a mechanosensitive gene upregulated by d-flow, and DNMT inhibition prevents d-flow-induced endothelial inflammation in vitro and atherosclerosis development in vivo. Given the flow-sensitivity of DNMT1, but not DNMT3a or DNMT3b, it is highly likely that DNMT1 plays a dominant role in endothelial inflammation and atherosclerosis.
We applied a two-pronged parallel approach, using the RRBS and transcriptome studies with in vivo–derived endothelial gDNA and RNA to determine the mechanism by which 5Aza inhibited atherosclerosis. These combined genome-wide studies showed that flow epigenetically regulates gene expression by controlling genomewide DNA methylation patterns in a DNMT-dependent manner. RRBS is widely used in genome-scale methylation studies, although incomplete CG coverage is an inherent limitation of RRBS due to the MspI restriction enzyme digest, PCR, and sequencing steps (52). These limitations produce partial genomic coverage, as evidenced by the 23% genomic and 53% promoter coverage of our RRBS assays. Additionally, we were only able to make determinations about genes and genomic regions that were covered in all samples by both the microarray and by RRBS. Therefore, genomic regions with partial coverage between samples or assays were not considered in this particular study, which could have resulted in the overlooking of potentially important CG sites. 5Aza treatment showed a dramatic effect in rescuing the mechanosensitive gene expression patterns, recovering nearly 540 of 569 genes that were suppressed by d-flow in the LCA (Figure 5A) while having a minimal effect on the RCA gene expression (Supplemental Figure 6E). The lack of any significant effect of 5Aza on the RCA gene expression was surprising, but it may indicate that the 5Aza effect is limited only to the d-flow–sensitive genes in the LCA. This preferential effect of 5Aza in the LCA may be due to the following reasons: (a) 5Aza in blood is in direct contact with the target cell types, such as arterial endothelium and circulating leukocytes. (b) Arterial regions exposed to d-flow have high endothelial permeability to blood-borne macromolecules, including anti-miRNAs as demonstrated by our group previously (19), albumin (53), and LDL (54). For a similar reason, 5Aza may be preferentially taken up by the endothelial cells in d-flow regions. (c) The endothelial cells in d-flow regions are hyperproliferative, and this facilitates 5Aza incorporation into these dividing cells’ DNA (55, 56), while the RCA endothelial cells remain quiescent. This notion is further corroborated by the lack of a widespread effect of 5Aza on the genome-wide DNA methylation patterns in the RCA, although 5Aza did reduce the number of hypermethylated CG sites significantly for a limited subset of the methylome (from ~130,000 to ~55,000 sites in the LCA), and those CG sites may have a functional importance in gene expression (Figure 4, A and B).

Using these 2 data sets, our goal was to identify those genes that are hypermethylated in the promoter regions and silenced by d-flow, but that could be rescued by 5Aza treatment. We would like to emphasize the importance of the user-defined threshold values in our method of gene discovery, which, by using more strict criteria, enabled us to filter out a manageable number of genes for further study. This unbiased genome-wide comparative analysis revealed 11 genes that fit these criteria. This result was surprising at first, since most of the genes downregulated by d-flow in the LCA were rescued by 5Aza treatment (540 of 569 genes), and because the promoter DNA methylation patterns were rescued by 5Aza in the majority of genes (335 of 421 genes) (Figure 5A). This discrepancy between the 5Aza effect on gene expression and methylation and the number of genes that fit our hypothesis could be due to the following possibilities: (a) 5Aza may target a few master regulators such as transcription factors that could in turn regulate the rest of the mechanosensitive genes; (b) there may be only a few key CG sites on promoters that determine the binding of master transcription factors, which control the expression of most of the mechanosensitive genes; and (c) the majority of other CG sites in the promoters do not necessarily lead to functional consequences. To test these possibilities, we first searched for potential master transcription factors that could regulate the 11 mechanosensitive and 5Aza-sensitive genes using the MetaCore analysis. This suggested CREB as a potential master transcription factor that could bind to 5 of the 11 genes. The subsequent computational sequence-based analysis showed that 4 of these genes (HoxA5, Klf3, Cmklr1, and Acr11) contained a CG site in the promoter CRE that was hypermethylated by d-flow in a 5Aza-dependent manner (Figure 5B and Figure 8, A–D).

It is interesting that our study finds HoxA5 and Klf3 as novel mechanosensitive transcription factors regulated by DNA methylation that may serve as master regulators of global gene expression.

Figure 7

D-flow induces DNA hypermethylation of the HoxA5 gene promoter and downregulates its expression in a 5Aza-dependent manner. (A) DNA methylation at the HoxA5 promoter region in endothelial-enriched genomic DNA from the LCA and RCA at 1 week after partial ligation from mice treated with saline or 5Aza was further examined in independent samples by bisulfite sequencing. Black and white circles represent methylated and unmethylated cytosines, respectively. Eight CG sites (denoted by the columns) were probed in this assay, and 8 to 10 colonies (denoted by the rows) were chosen for analysis. Percentages below the figures denote the average percent methylation for this region of the HoxA5 promoter. (B and C) HoxA5 gene expression was examined in endothelial-enriched RNA from the LCA and RCA obtained at 1 week after partial ligation from saline- or 5Aza-treated mice by microarray analysis. Data for B are shown as the mean ± SEM. **P < 0.01, n = 3 each and were validated by qPCR. Data for C are shown as the mean ± SEM. **P < 0.01, n = 5 each.

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in response to flow. Hox genes are highly conserved, are known to be dysregulated in cancer, and are controlled in part by DNA methylation (57–62). Hox family members have been implicated in vascular remodeling, angiogenesis, and disease as orchestrators of gene expression, extracellular matrix, and integrin changes (63). HoxA5 is known to regulate various endothelial functions such as migration, angiogenesis, and inflammation by controlling specific genes (thrombospondin-2, vascular endothelial growth factor receptor-2, ephrin-A1, hypoxia-inducible factor-1α, prostaglandin-endoperoxide synthase-2, and PTEN) (64–68). HoxA5 was shown to regulate expression of approximately 300 genes in breast cancer cell lines (69).

Our study shows, for the first time to our knowledge, that HoxA5 is regulated by flow in a DNA methylation–dependent manner and that HoxA5 regulates endothelial inflammation. Interestingly, Klf3 is suppressed in acute myeloid leukemia (AML), and 5Aza is used to treat this disease, although it has not yet been shown whether 5Aza directly targets Klf3. AML is also treated with all-trans retinoic acid, and, also interestingly, this drug was shown to rescue Klf3 expression by an unknown mechanism (19). Since all-trans retinoic acid is known to influence DNA methylation of a small number of specific genes (70), it would be interesting to test whether Klf3 is regulated by this mechanism in AML. Although our current study focused on examining the role of DNA methylation in vascular biology and disease, these results indicate that our work could have far-reaching implications across multiple diseases.

Our in silico analysis suggests CREB as another key transcription factor whose DNA binding is known to be regulated by methylation of the CRE sites of its target genes, such as HoxA5 and Klf3 (71, 72). It was shown previously that protein kinase A downregulation in d-flow leads to decreased CREB phosphorylation and decreased CREB binding to CREs (73). Here, we identified a novel mechanism by which CREB binding to its gene targets may be decreased by d-flow via DNA methylation at their promoter CREs.

One of the most striking findings from our RRBS data is that CRE-localized promoter CG sites were not methylated at all in the s-flow RCA, whereas CG sites across all regions of the promoter were approximately 30% methylated (Figure 8F and Supplemental Figure 7B). This suggests that there is an active hypomethylation mechanism that prevents these promoter CRE CG sites from being methylated. Our results also suggest that d-flow induces hypermethylation in these promoter CRE CG sites by a DNMT-dependent mechanism (Figure 8), which in turn has a major impact on gene expression.
In summary, we describe, for the first time to our knowledge, that d-flow–induced changes in global gene expression in endothelial cells are regulated at the epigenomic level in a DNMT- and DNA methylation–dependent manner. We propose that d-flow induces DNMT1 expression in endothelial cells, which in turn stimulates DNA methylation in specific promoter CRE sites of mechanosensitive, master transcription factors including HoxA5 and Klf3. These master genes may then regulate expression of the majority of other mechanosensitive genes, resulting in endothelial dysfunction and atherosclerosis. We further propose that the DNMT inhibitor 5AzA targets these mechanosensitive master regulators to restore antiatherogenic gene expression profiles. In conclusion, our study provides a novel insight into the mechanism by which flow regulates gene expression in a DNMT-dependent manner and uncovers novel antiatherogenic therapeutic targets.

Methods

Partial carotid ligation model. Eight-week-old C57BL/6 or apoE–/– mice (The Jackson Laboratory) were used for all animal studies according to the approved protocol of the IACUC at Emory University. Mice were partially ligated under isoflurane anesthesia and the resultant d-flow conditions in the LCA were determined by ultrasound measurements as described previously (49). Briefly, 3 of 4 caudal branches of the LCA (left external carotid, internal carotid, and occipital artery) were ligated with 6-0 silk suture, while the superior thyroid artery was left intact. Mice were sacrificed by CO2 inhalation and were subsequently perfused with saline containing heparin for 5 minutes. Total carotid intimal RNA or gDNA was isolated by flushing of the RCA and LCA with QIAzol (QIAGEN) or Buffer AL (QIAGEN), respectively, by our method that we have demonstrated to be endothelial-enriched without any significant contamination of smooth muscle cells and leukocytes (19, 48, 49).

Cell culture and in vitro shear stress system. HUVECs (Lonza cc-2519) were maintained in M199 (Fisher MT10060CV) base culture medium containing 20% FBS (SH30071), 10% penicillin/streptomycin/fungizone (Gibco 15240-062), 10% GlutaMAX (Mediatech 25-005-cl), 10% endothelial cell growth serum, and 0.4% heparin (McKesson Medical 404867). In vitro shear stress was applied to passage 8 HUVECs in 10-cm culture plates for 24 hours using a cone and plate viscometer that exerts 15 dyn/cm2 of unidirectional flow (laminar shear stress, or LS, conditions) and ±5 dyn/cm2 of bidirectional flow (oscillatory shear stress, or OS). The cell culture plates were secured in place at the bottom by a vacuum pump, and the incubator was maintained at 5% CO2 and 37°C. Total RNA or gDNA was collected by scraping of cells in QIAzol (QIAGEN) or Buffer AL (DNeasy Blood and Tissue Kit; QIAGEN), respectively.

qPCR. Total RNA was collected from HUVEC cultures in 700 μl of QIAzol and was purified by the QIAGEN miRNeasy Mini Kit. Total RNA was reverse transcribed into cDNA using SuperScript III and random primers (Invitrogen), as described previously (49). qPCR for specific genes was performed using Brilliant II SYBR Green qPCR Master Mix (Stratagene) with custom-designed primers on the ABI StepOne Plus Real-Time PCR System. Results were normalized to 18S RNA, and the fold change between LCA and RCA was determined using the 2ΔΔct method (74). Primer sequences can be found in Supplemental Table 2.

Western blotting. Western blotting was performed by the standard method using antibodies at a 1:1,000 dilution for the following: DNMT1 (sc-20701; Santa Cruz Biotechnology Inc.), DNMT3a (ab23565; Abcam), DNMT3b (ab2851; Abcam), β-actin (A5316; Sigma-Aldrich), and HoxA5 (ab82645; Abcam).

Tissue preparation, oil red O staining, and immunohistochemistry. Frozen blocks containing the heart, aortic arch, and carotid arteries were prepared in TissueTek and stored at −80°C. Sections from the blocks were stained with oil red O by fixing of the RCA with 10% formalin for 2 minutes, followed by 2 rinses using the 2× saline for 5 minutes, orbital shaking in 100% propanediol for 10 minutes, and incubation in 0.2% oil red O solution for 5 minutes at 85°C. For hematoxylin staining, the previous method was followed by 2 rinses with distilled water, a hematoxylin dip for 30 seconds, 2 rinses with distilled water, 1 rinse with Scott’s Tap Water (Sigma-Aldrich), 2 rinses with distilled water, and then mounting. Plaque size was quantified using ImageJ software (NIH). Immunohistochemistry was performed on sections from the frozen blocks using the following antibodies at a 1:50 dilution: 5-methylcytidine (BI-MEcy; Eurogentec), DNMT1 (sc-20701; Santa Cruz Biotechnology Inc.), DNMT3b (ab2851; Abcam), DNMT3a (ab23565; Abcam), and CD45 (13-0454; eBioscience).

Figure 9

HoxA5 and Klf3 are suppressed by OS in vitro, and HoxA5 increases LS-suppressed endothelial inflammation. (A) HoxA5 expression and Klf3 expression are suppressed by OS compared with LS. HUVECs exposed to LS (15 dyn/cm2) or OS (±5 dyn/cm2, at 1 Hz) for 24 hours were used for qPCR using 18S as an internal control (n = 6 each, data are shown as the mean ± SEM. **P < 0.01). (B) Western blot analysis for HoxA5 was performed on the same samples from A using β-actin as the loading control. The bar graph shows the ImageJ Western blot quantification normalized to β-actin (n = 3 each, data are shown as the mean ± SEM; *P < 0.05). (C and D) HUVECs were pre-treated with siRNA to HoxA5 or Klf3 (100 nM for 24 hours), and subsequently exposed to OS or LS for an additional 24 hours. Following shear, endothelial inflammation was determined by quantification of the number of THP-1 monocytes adhered to shear-adherent endothelial cells (n = 3 each, data are shown as the mean SEM. *P < 0.05).
SAza and siRNA treatment of HUVECs in vitro. Optimal 5-aza-2′-deoxycytidine (SAza, A3656; Sigma-Aldrich) treatment conditions were determined by a dose curve using 0, 0.5, 1, 5, and 20 μM SAza for 5 days. DNMT enzymatic activity and DNMT1 protein expression assays indicated the optimal dose of 5 μM. For shear studies, 5 μM SAza was treated for 5 days before in vitro shear, and the LS or OS conditions were applied for 24 additional hours. siRNA was transfected using Oligofectamine (Invitrogen 12252-011) with either 100 nM siDNMT1, siHoxA5, or siKlf3 (Dharmacon RNAi ON-TARGET plus SMART pool) or 100 nM stealth RNAi negative control (medium GC content, 12935-112; Invitrogen) in OptiMEM (31985-088; Invitrogen) for 6 hours followed by normal HUVEC medium replacement. After 48 hours, shear stress (either LS or OS) was applied for an additional 24 hours.

Quantification of DNMT activity. The DNMT enzymatic activity assay was performed according to the manufacturer’s protocol (P-3010; Epigentek).

Restriction enzyme (MspI/HpaII) assay to determine global DNA methylation status in vitro. Genomic DNA was collected and purified by the DNeasy Blood and Tissue kit (Qiagen) from HUVECs treated with either 5 μM SAza or vehicle control for 5 days. Either MspI or HpaII (ER0541; Thermosci) was reacted with the gDNA according to the recommended protocol for the restriction enzymes. The reaction products were electrophoresed on a 2.5% agarose gel stained with ethidium bromide alongside a 100-bp ladder (N3231L; New England BioLabs) and imaged under UV light.

Monocyte adhesion assay. HUVECs were pretreated with either SAza or siRNA and were subjected to either LS or OS for 24 hours. Separately, human peripheral blood mononuclear leukocytes (THP-1 cells) were cultured in serum-containing RPMI medium (MT10040CM; Fisher) with 10% fetal bovine serum and were labeled in serum-free RPMI with 5 μM 5′-bis-(2-carboxyethyl)-5′-7′-7′-bis-(2-carboxyethyl)-5′-deoxycytidine (5Aza, A3656; Sigma-Aldrich) treatment conditions were determined by a dose curve using 0, 1, 0.2, or 0.4 mg/kg/d of 5-aza-2′-deoxycytidine (SAza; Sigma-Aldrich) daily for 4 weeks (the acute d-flow–induced atherosclerosis model). 0.2 mg/kg/d for 3 months (the chronic diet-induced atherosclerosis model), or 0.4 mg/kg/d for 3 weeks (the RRBS and array studies). Mouse weight was monitored weekly, and food and water intake and feces were monitored daily.

HRM analysis to quantify global DNA methylation. Global gDNA methylation as represented by genomic repeat element methylation was quantified by LINE1 and B1 high-resolution melt curve (HRM) assays for each mouse in the RRBS and array studies to determine the effect of SAza treatment. HRM analysis was done according to the protocol set by Newman et al. (75). Whole blood was collected by the submandibular technique 1 week after the start of SAza treatment. gDNA was extracted from blood using the DNeasy Blood and Tissue kit (Qiagen) and was bisulfite converted using the EpiTect Bisulfite Conversion Kit (Qiagen). Bisulfite-converted gDNA was subjected to HRM analysis using the F_unmeth_mLINE1 and R_unmeth_mLINE1, the F_mod_unbiased_B1_Mm, and the R_mod_unbiased_B1_Mm primers from Newman et al. and normal qPCR methods as described above.

RRBS sample collection. Genomic DNA was collected from mouse carotid arteries (as described previously) by flushing with 200 μl of a 1:1 solution of Buffer AL and nuclelease-free water. gDNA was purified using the DNeasy Blood and Tissue Kit (Qiagen), and 20 gDNA samples were pooled per RRBS sample. Pooled gDNA for the RCA, LCA, Aza-RCA, and Aza-LCA were sent to Zymo Research Corp., where reduced representation bisulfite sequencing (RRBS) was carried out by bisulfite conversion, genome reduction by restriction enzyme digest and size selection, library preparation, and next-generation sequencing.

Computational analysis of genome-wide methylome (RRBS) and transcriptome (microarray) data sets. Next-generation sequencing reads from the RRBS study were mapped to the mm9 genome assembly by Zymo Research Corp. The resulting mapped files were analyzed by our lab. Methylome coverage was determined by division of the number of unique CG sites covered by each individual RRBS data set by the number of total CG sites in the mm9 genome assembly (21,342,492 CG sites in chromosomes 1–19, X, and Y). Promoter coverage was determined by confining the RRBS and mm9 assemblies to gene transcription start sites (TSSs, as defined by the NCBI reference sequences database) ± 1 kb, and then again calculating coverage in these confined regions.

Heatmap correlation scatter plots were generated using the smoothScatter function in R. The methylation ratios of only those unique CG sites covered by all 4 data sets was plotted, and the R^2 values were determined by fitting of linear models with the R lm function (1, 76, 77).

For the a priori emergent methylation pattern analysis, we used only those CG sites confined to promoter regions (TSS ± 1 kb) and calculated the promoter methylation ratio by the total number of methylated reads divided by the total number of all reads in the region, normalized to the number of CG sites in that region. This was done for each RRBS data set, and then the methylation ratio for each gene promoter was compared between data sets and ranked from highest to lowest methylation ratio. There are 24 possible combinatorial patterns that can result from this ranking analysis by ordering of the 4 samples from highest to lowest, and we counted the number of genes that fell into each of the 24 possible pattern categories.

Gene transcript microarray (transcriptome). Total intrimal RNA was collected by flushing of the RCA, LCA, Aza-RCA, and Aza-LCA with 150 μl of QIAzol and was purified using the miRNeasy Mini Kit (Qiagen). Three carotid RNA samples were pooled and the combined sample was amplified and reverse transcribed, and the cDNA was applied to Illumina Mouse WG6 microarray chips. The microarray detected 45,281 probes corresponding to 30,854 genes, and the array results are available in the NCBI’s Gene Expression Omnibus (GEO) database (GEO GSE56143).

Filtering of RRBS and transcript microarray data sets to discover genes with the hypothesized methylation and expression characteristics. The differentially methylated and differentially expressed genes were parsed according to our hypothesis that d-flow–induced hypermethylation at the promoter suppresses gene expression, and that this could be reversed by SAza. We performed thresholding to filter out genes that are hypermethylated in the promoter and suppressed by d-flow in a SAza-preventable manner. Specifically, we looked for differentially methylated regions (DMRs) in gene promoters, defined by TSS ± 1 kb containing a minimum of 10 CG sites with a minimum difference in percent methylation (% methylation in the LCA – % methylation in the RCA) of 10. There were 421 such genes. We further parsed this list to genes whose promoter hypermethylation in LCA was prevented by SAza (% methylation in the Aza-LCA – % methylation in the Aza-RCA < 5), and this resulted in 335 genes. For the mRNA expression criteria, we looked for genes downregulated in RCA versus LCA (LCA/RCA < 0.67), which resulted in 569 genes, and then we further looked for genes whose expression in LCA was rescued by SAza (resulting in 540 genes). By combining these methylation and gene expression criteria, we extracted 11 genes that have d-flow–inducible hypermethylation and gene suppression that can be prevented by SAza.


42. Surgery approval. All animal procedures were approved by the IACUC at Emory University.

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