Genetic variant at coronary artery disease and ischemic stroke locus 1p32.2 regulates endothelial responses to hemodynamics

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Biomechanical cues dynamically control major cellular processes, but whether genetic variants actively participate in mechanosensing mechanisms remains unexplored. Vascular homeostasis is tightly regulated by hemodynamics. Exposure to disturbed blood flow at arterial sites of branching and bifurcation causes constitutive activation of vascular endothelium contributing to atherosclerosis, the major cause of coronary artery disease (CAD) and ischemic stroke (IS). Conversely, unidirectional flow promotes quiescent endothelium. Genome-wide association studies (GWAS) have identified chromosome 1p32.2 as strongly associated with ischemic stroke and CAD/IS; however, the causal mechanism related to this locus remains unknown. Using statistical analyses, assay of transposase accessible chromatin with whole-genome sequencing (ATAC-seq), H3K27ac/H3K4me2 ChIP with whole-genome sequencing (ChIP-seq), and CRISPR interference in human aortic endothelial cells (HAECs), our results demonstrate that rs17114036, a common noncoding polymorphism at 1p32.2, is located in an endothelial enhancer dynamically regulated by hemodynamics. CRISPR-Cas9-based genome editing shows that rs17114036-containing region promotes endothelial quiescence under unidirectional shear stress by regulating phospholipid phosphatase 3 (PLPP3). Chromatin accessibility quantitative trait locus (caQTL) mapping using HAECs from 56 donors, allelic imbalance assay from 7 donors, and luciferase assays demonstrate that CAD/IS-protective allele at rs17114036 in PLPP3 intron 5 confers increased endothelial enhancer activity. ChIP-PCR and luciferase assays show that CAD/IS-protective allele at rs17114036 creates a binding site for transcription factor Krüppel-like factor 2 (KLF2), which increases the enhancer activity under unidirectional flow. These results demonstrate that a human SNP contributes to critical endothelial mechanosensing mechanisms and suggest that human haplotypes and related cis-regulatory elements provide a previously unappreciated layer of regulatory control in cellular mechanosensing mechanisms.

Significance

Biomechanical stimuli control major cellular functions and play critical roles in human diseases. Although studies have implicated genetic variation in regulating key biological functions, whether human genetic variants participate in the processes by which cells sense and respond to biomechanical cues remains unclear. This study provides a line of evidence supporting an underappreciated role of genetic predisposition in cellular mechanotransduction. Using genetics approaches and genome editing, our data demonstrate that rs17114036, a common noncoding polymorphism implicated in coronary artery disease and ischemic stroke by genome-wide association studies, dynamically regulates endothelial responses to atherosclerosis-related blood flow (hemodynamics) via a noncoding DNA region important for transcription activation (enhancer). These results provide molecular insights linking disease-associated genetic variants to cellular mechanobiology.

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Mechanical stimuli regulate major cellular functions and play critical roles in the pathogenesis of diverse human diseases (1). This is especially important in the vasculature, where endothelial cells are activated by local disturbed flow in arterial regions prone to atherosclerosis (2–5), the major cause of coronary artery disease (CAD) and ischemic stroke (IS). The role of biomechanical forces on the noncoding and regulatory regions of the human genome is unexplored. Recent studies demonstrated that the noncoding, nontranscribed human genome is enriched in cis-regulatory elements (6). In particular, enhancers are distinct genomic regions that contain binding sites for sequence-specific transcription factors (7). Enhancers spatially and temporally control gene expression with cell type- and cell state-specific patterns (8). Notably, top-associated human disease-associated SNPs are frequently located within enhancers that explicitly activate genes in disease-relevant cell types (9). The nature of mechanosensitive enhancers and their biological roles in vascular functions have not been identified.

Atherosclerotic disease is the leading cause of morbidity and mortality worldwide. Genome-wide association studies (GWAS) identified chromosome 1p32.2 as one of the most strongly associated loci with susceptibility to CAD and IS (10–12). Six candidate genes in this locus are phospholipid phosphatase 3 (PLPP3; also known as phosphatidic acid phosphatase-type 2B), which inhibits endothelial inflammation and promotes monolayer integrity by hydrolyzing lysophosphatic acid (LPA) that activates endothelium (13, 14). Our recent study demonstrated that PLPP3 expression is significantly increased in vascular endothelium by unidirectional flow in vitro and in vivo (13). Moreover, expression quantitative trait locus (eQTL) mapping showed that CAD-protective allele at 1p32.2 is associated with increased PLPP3 expression in an endothelium-specific manner.
(13). However, whether genetic variants and mechanosensing mechanisms converge on PLPP3 expression is unclear. In addition, causal SNPs at locus 1p32.2 remain unknown.

Using statistical analyses, assay of transposable accessible chromatin with whole-genome sequencing (ATAC-seq), H3K27ac ChIP with whole-genome sequencing (ChIP-seq), H3K4me2 ChIP-seq, luciferase assays, and CRISPR-based approaches, we report that rs17114036-containing genomic region at 1p32.2 causatively promotes endothelial expression of PLPP3 and governs the atheroresistant endothelial phenotype under unidirectional shear stress by functioning as a mechanosensitive endothelial enhancer. Using human aortic endothelial cells (HAECs) isolated from a cohort of human subjects, we performed transcriptome analyses and chromatin accessibility quantitative trait locus (cQTL) mapping showing nucleotide-specific epigenetic and transcriptomic effects of rs17114036 in humans. Allelic imbalance (AI) assays, ChIP-PCR, and luciferase assays collectively demonstrate that, due to a single base pair change, the CAD/IS-protective allele at rs17114036 and rs17114036 contains 15 SNPs with high linkage disequilibrium (LD; $r^2 > 0.8$) with rs17114036, and any of these SNPs could conceivably be a causal variant. To predict possible causal SNPs at the 1p32.2 locus, we conducted two statistical analyses. First, we used a Bayesian statistical approach to assign posterior probabilities and credible sets of SNPs that refine the association signals of GWAS-detected loci (15). Second, we applied conditional and joint association analyses using summary-level statistics of GWAS data to predict causal variants (16). Using Bayes’ theorem in the cohort of 45 SNPs at 1p32.2, we identified 15 SNPs with >95% posterior probability to be causal (Fig. 1A and SI Appendix, Table S1). Using the approximate conditional and joint association analyses, we identified seven 1p32.2-associated SNPs to be possible causal (Fig. 1A and SI Appendix, Table S1). Only two SNPs, rs17114036 and rs2184104, were predicted to be causal by both methods.

Results
Bayesian Refinement and Conditional Joint Multiple-SNP Analyses Predict That rs17114036 and rs2184104 Are Possible Causal SNPs Located in CAD/IS Locus 1p32.2. rs17114036 is the tag SNP used in most CAD/IS GWAS (10–12) and in our eQTL mapping (13); however, there are 44 common SNPs in high linkage disequilibrium (LD; $r^2 > 0.8$) with rs17114036, and any of these SNPs could conceivably be a causal variant. To predict possible causal SNPs at the 1p32.2 locus, we conducted two statistical analyses. First, we used a Bayesian statistical approach to assign posterior probabilities and credible sets of SNPs that refine the association signals of GWAS-detected loci (15). Second, we applied conditional and joint association analyses using summary-level statistics of GWAS data to predict causal variants (16). Using Bayes’ theorem in the cohort of 45 SNPs at 1p32.2, we identified 15 SNPs with >95% posterior probability to be causal (Fig. 1A and SI Appendix, Table S1). Only two SNPs, rs17114036 and rs2184104, were predicted to be causal by both methods.

CRISPR-Based Approaches Identified rs17114036-Containing Region as a cis-Regulatory Element for Endothelial PLPP3 Expression. To determine the causal role of rs17114036-containing genomic locus in regulating endothelial PLPP3 expression, the bacterial CRISPR-Cas9 system was used to selectively delete an ~66-bp genomic region (chr1:56962783–56962849) enclosing rs17114036 in HAECs. A pair of guide RNAs (Fig. 2A) was designed according to the method described by Ran et al. (19). We pre-assembled the Cas9–guide RNA ribonucleoprotein (RNP) complex by incubating guide RNAs with recombinant Streptococcus pyogenes Cas9 followed by the delivery to cells using cationic liposome transfection reagents. To improve efficiency of the CRISPR-Cas9–mediated deletion, cells were reverse transfected with the RNP complex four times before flow cytometry to sort single cells (Fig. 2B). Immortalized HAECs (carrying major alleles at rs17114036) with high proliferating capacity were used for single-cell clonal isolation to select a genetically identical cell line. Among 459 HAEC colonies that we grew to confluency, PCR assays detected 17 lines with ~66-bp genetic deletion in PLPP3 intron 5 enclosing rs17114036 (SI Appendix, Fig. S6A). DNA deletion was further confirmed by cloning and Sanger sequencing (SI Appendix, Fig. S6B). Endothelial PLPP3 expression is significantly reduced in the genome-edited cells compared with teloHAECs that underwent CRISPR-Cas9 treatment and single-cell clonal isolation but showed no sign of deletion at chr1:56962783–56962849 (Fig. 2C). In addition, deletion of this putative enhancer in PLPP3 intron 5 resulted in an increase of LPA-induced E-selectin expression (Fig. 2D) and leukocyte adhesion (Fig. 2E), in agreement with the antiinflammatory/ adhesion role of endothelial PLPP3 (13, 14). Moreover, transendothelial electrical resistance decreased detected increased monolayer

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**Table S1.** SNPs at 1p32.2 in 459 human subjects.

- **SNP**: rs17114036, rs2184104
- **Frequency**: >95%
- **Allelic Imbalance (AI)**: 95% of subjects with the CAD/IS-protective allele at rs17114036 had an allelic imbalance (AI) of >95%
- **ChIP-PCR**: rs17114036 in human umbilical vein endothelial cells (HUVECs) (6) (SI Appendix, Fig. S3). A 1,200-bp DNA sequence corresponding to human chr1:5696213–56963412 was cloned upstream of firefly luciferase that was driven by a minimal promoter. Reporter assays demonstrated that insertion of this putative enhancer region with major allele T at rs17114036 significantly increased the luciferase activity in HAECs (Fig. 1C). No significant enhancer activity was implicated when rs2184104-containing region (chr1:56911623–56912023) was cloned into the same reporter vector compared with the vector with only PLPP3 promoter (Fig. 1C). Minimal enhancer activities chr1:56962213–56963412 were detected when the constructs were expressed in the nonendothelium cell line HEK 295 (SI Appendix, Fig. S5). ATAC-seq, H3K27ac/H3K4me2 ChIP-PCR, and luciferase assays altogether demonstrate that chr1:56962213–56963412 functions as an enhancer in HAECs.

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**SI Appendix.** Fig. S2 shows the ATAC-seq and H3K27ac/H3K4me2 tracks in HAECs at 1p32.2 locus. The enhancer activity of chr1:56962213–56963412 was experimentally demonstrated by a luciferase reporter assay (Fig. 1C). Plasmid transfection was first validated in HAECs using electroporation of pmax green fluorescent protein (pmaxGFP)-expressing constructs (SI Appendix, Fig. S3). A 1,200-bp DNA sequence corresponding to human chr1:56962213–56963412 was cloned upstream of firefly luciferase that was driven by a minimal promoter. Reporter assays demonstrated that insertion of this putative enhancer region with major allele T at rs17114036 significantly increased the luciferase activity in HAECs (Fig. 1C). No significant enhancer activity was implicated when rs2184104-containing region (chr1:56911623–56912023) was cloned into the same reporter vector compared with the vector with only PLPP3 promoter (Fig. 1C). Minimal enhancer activities chr1:56962213–56963412 were detected when the constructs were expressed in the nonendothelium cell line HEK 295 (SI Appendix, Fig. S5). ATAC-seq, H3K27ac/H3K4me2 ChIP-PCR, and luciferase assays altogether demonstrate that chr1:56962213–56963412 functions as an enhancer in HAECs.

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**SI Appendix.** Fig. S6 shows the ATAC-seq and H3K27ac/H3K4me2 tracks in HAECs at 1p32.2 locus. The enhancer activity of chr1:56962213–56963412 was experimentally demonstrated by a luciferase reporter assay (Fig. 1C). Plasmid transfection was first validated in HAECs using electroporation of pmax green fluorescent protein (pmaxGFP)-expressing constructs (SI Appendix, Fig. S3). A 1,200-bp DNA sequence corresponding to human chr1:56962213–56963412 was cloned upstream of firefly luciferase that was driven by a minimal promoter. Reporter assays demonstrated that insertion of this putative enhancer region with major allele T at rs17114036 significantly increased the luciferase activity in HAECs (Fig. 1C). No significant enhancer activity was implicated when rs2184104-containing region (chr1:56911623–56912023) was cloned into the same reporter vector compared with the vector with only PLPP3 promoter (Fig. 1C). Minimal enhancer activities chr1:56962213–56963412 were detected when the constructs were expressed in the nonendothelium cell line HEK 295 (SI Appendix, Fig. S5). ATAC-seq, H3K27ac/H3K4me2 ChIP-PCR, and luciferase assays altogether demonstrate that chr1:56962213–56963412 functions as an enhancer in HAECs.
permeability in rs17114036-deleted HAECs (Fig. 2F), consistent with PLPP3’s role in maintaining endothelial monolayer integrity (13, 14). CRISPR interference was recently developed to suppress the activity of cis-regulatory elements (20). Here, we showed that rs17114036-targeted (Fig. 2G) but not rs2184104-targeted (SI Appendix, Fig. S7) guide RNAs significantly reduce PLPP3 mRNA expression in HAECs. These results demonstrate that rs17114036-containing region causatively regulates PLPP3 expression and endothelial functions.

Unidirectional Flow Increases the Enhancer Activity at chr1:56962213–56963412 in Vascular Endothelium. Given the critical role of hemodynamics in controlling endothelial PLPP3 transcription (13), we tested whether shear stresses regulate the enhancer activity of chr1:56962213–56963412. ATAC-seq and H3K27ac ChIP-seq collectively identified an enhancer-like region in chr1:56962213–56963412 in HAECs. ATAC-seq and H3K27ac/H3K4me2 ChIP-seq collectively identified an enhancer-like region in chr1:56962213–56963412 in HAECs. All sequencing experiments were performed in duplicate, and the merged tracks are shown. (C) Enhancer activity of chr1:56962213–56963412 in vascular endothelium. DNA sequences of chr1:56962213–56963412 were cloned into luciferase reporters (firefly luciferase construct pGL4) that contain a minimal promoter or human PLPP3 promoter. The red asterisks denote the relative position of rs17114036 in the luciferase construct. Dual luciferase reporter assays were conducted in HAECs 24 h after the electroporation-based transfection (using pRL-TK plasmid carrying Renilla luciferase as transfection controls) in HAECs, detecting increased firefly luciferase as the result of insertion of chr1:56962213–56963412. Data represent mean ± SEM. ***P < 0.0005 as determined by Student’s t test.

Fig. 1. CAD-associated SNP rs17114036 is located in an enhancer element (chr1:56962213–56963412) in HAECs. (A) In silico prediction of causal SNPs in the CAD locus 1p32.2. Diagrams demonstrate the association (−log10P) and LD pattern of a total of 45 common SNPs in the 1p32.2 locus. Green circles indicate possible causal SNPs predicted by the Bayesian Statistical Approach. Red circles indicate possible causal SNPs predicted by the Approximate Conditional and Joint Association Analysis. Purple circles indicate putative causal SNPs predicted by both statistical analyses. (B) Chromatin accessibility and canonical enhancer marks in chr1:56962213–56963412 region enclosing rs17114036 in HAEC. ATAC-seq and H3K27ac/H3K4me2 ChIP-seq collectively identified an enhancer-like region in chr1:56962213–56963412 in HAECs. All sequencing experiments were performed in duplicate, and the merged tracks are shown. (C) Enhancer activity of chr1:56962213–56963412 in vascular endothelium. DNA sequences of chr1:56962213–56963412 were cloned into luciferase reporters (firefly luciferase construct pGL4) that contain a minimal promoter or human PLPP3 promoter. The red asterisks denote the relative position of rs17114036 in the luciferase construct. Dual luciferase reporter assays were conducted in HAECs 24 h after the electroporation-based transfection (using pRL-TK plasmid carrying Renilla luciferase as transfection controls) in HAECs, detecting increased firefly luciferase as the result of insertion of chr1:56962213–56963412. Data represent mean ± SEM. ***P < 0.0005 as determined by Student’s t test.
impaired unidirectional flow-induced PLPP3 expression in HAECs (Fig. 3B). Moreover, enhancer activity at chr1:56962213–56963412, measured by H3K27ac ChIP-PCR, was increased in control HAEC but not in rs17114036-(biallelic) deleted cells when subjected to 24 h of unidirectional flow (Fig. 3C). These results collectively demonstrate that enhanced activity of chr1:56962213–56963412 is dynamically activated by the atheroprotective unidirectional flow to regulate endothelial PLPP3.

**CAD/IS-Protective Allele C at rs1711403 Confers a Higher Enhancer Activity of chr1:56962213–56963412.** GWAS have linked the minor allele C at rs17114036 at 1p32.2 to reduced CAD/IS susceptibility (10–12), and our eQTL mapping described increased PLPP3 expression in HAECs with minor allele C (13). Here, we investigated the genotype-dependent effect of rs17114036 on the enhancer activity of chr1:56962213–56963412 by ATAC-seq and luciferase assays. In addition to HAEC lines carrying major (risk) allele T at rs17114036, we conducted ATAC-seq in HAECs isolated from donors who are heterozygous (T/C; ~20% of Europeans) at rs17114036, allowing us to perform caQTL mapping. caQTL was recently developed to detect between-individual signaling in cis-regulatory elements as a function of genetic variants (22). ATAC-seq detected significantly increased numbers of reads corresponding to rs17114036-containing region in HAEC lines that contain one CAD-protective allele (T/C) compared with HAECs from donors homozygous of CAD risk allele (T/T) (Fig. 4A), supporting increased chromatin accessibility associated with C allele at rs17114036. In addition, we conducted RNA-seq analysis in these cells, demonstrating that there

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**Fig. 2.** CRISPR-based approaches identify rs17114036-containing genomic locus as a cis-regulatory element regulating endothelial PLPP3 expression. (A) RNA RNP complex that contains recombinant S. pyogenes Cas9 and two sgRNAs flanking rs17114036. (B) Experimental overview of CRISPR-Cas9–mediated genomic deletion in HAECs. TeloHAECs were treated with RNP complex four times before single-cell sorting and isogenic clone selection. (C) Reduced PLPP3 expression, (D) elevated E-selectin adhesion, and (E) higher monolayer permeability in teloHAECs with genomic deletion at rs17114036-containing region. (G) Reduced PLPP3 expression in HAECs treated with CRISPR interference targeting rs17114036-containing region. n = 3–8. Data represent mean ± SEM. *P < 0.05 as determined by Student’s t test; **P < 0.005 as determined by Student’s t test; ***P < 0.0005 as determined by Student’s t test.
Fig. 3. Unidirectional flow (UF) increases enhancer activity at chr1:56962213–56963412 that transcriptionally activates PLPP3 in human aortic endothelium. (A) Increased chromatin accessibility and H3K27ac mark at chr1:56962213–56963412 in HA ECs subjected to 24-h atheroprotective UF compared with cells under 24-h atherosusceptible disturbed flow. The PLPP3 locus is shown and zoomed in to demonstrate details around the enhancer region of interest, with rs17114036 highlighted with a vertical blue line. (B) Reduced UF-induced PLPP3 expression in HA ECs with genomic deletion at rs17114036-containing genomic locus. Control and genome-edited (∼66-bp deletion) HA ECs were subjected to 24-h UF. The y axis represents the fold change of PLPP3 mRNA quantities between the static conditions and UF for each individual clone. Nondeletion clones, n = 3; deletion clones, n = 4. (C) H3K27ac ChIP-PCR performed in two CRISPR clones; one nondeletion clone and one biallelic deletion clone harboring the 66-bp deletion near rs17114036. The cells were subjected to static or UF conditions before cross-linking and ChIP. PCR primers were designed to amplify a region within the enhancer but not overlapping the deleted region. Data are shown as fold change of the UF-treated samples compared with static conditions. Data represent mean ± SEM. *P < 0.05 as determined by Student’s t test; ***P < 0.0005 as determined by Student’s t test.
is a strong correlation between enhanced chromatin accessibility in rs17114036-containing region and increased mRNA levels of PLPP3 in HAECs (Fig. 4B) and further suggesting that chr1:56962213–56963412 functions as an enhancer in promoting endothelial PLPP3 transcription. Moreover, ATAC-seq experiments in HAEC lines heterozygous at rs17114036 further allow us to determine whether the chromosome with C at rs17114036 exhibits higher chromatin accessibility at chr1:56962213–56963412 compared with the chromosome with T allele. This is achieved by the AI analysis, which assigns next generation sequencing reads overlapping heterozygous sites to one chromosome or the other for allele-specific signals (23). ATAC-seq detected reads enriched from the C-containing chromosome compared with that from T allele in HAECs heterozygous at rs1711403 (Fig. 4C), further supporting the increased chromosome accessibility associated with C allele at rs17114036. Lastly, luciferase assays were conducted to support the genotype-dependent enhancer activity of chr1:56962213–56963412. Replacement of T allele with C allele led to a much higher luciferase activity (~5.2-fold, C vs. T) in endothelium (Fig. 4D). Polymorphisms (A or G) at rs2184104 had no effect of the chr1:56911623–56912823 in the luciferase assay (SI Appendix, Fig. S8). Taken together, these results demonstrate that CAD-protective allele C at rs17114036 confers a higher enhancer activity of chr1:56962213–56963412 to promote PLPP3 expression in vascular endothelium.

**CAD/IS-Protective C Allele at rs1711403 Promotes Flow-Induced, KLF2-Mediated Enhancer Activity of chr1:56962213–56963412.** We further examined whether the genetic variants at rs17114036 modulate

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**Fig. 4.** CAD-protective allele C at rs17114036 confers higher enhancer activity of chr1:56962213–56963412. (A) Increased ATAC-seq reads in chr1:56962213–56963412 region from HAECs isolated from people heterogeneous (T/C) at rs17114036 compared with HAECs from people heterozygous (T/T) at rs17114036. (B) A positive correlation ($R = 0.6$, $P$ value = 6.23e-06) between ATAC-seq reads at chr1:56962213–56963412 and PLPP3 mRNA detected by RNA-seq in 56 HAEC lines. (C) Increased ATAC-seq reads at rs17114036-containing genomic locus from C (rs17114036)-containing chromosome compared with T-containing chromosome in HAEC lines heterozygous (T/C) at rs17114036. (D) Increased enhancer activity of chr1:56962213–56963412 with C allele at rs17114036 compared with T allele. Dual luciferase reporter assays were conducted in teloHAEC. The red and green asterisks denote the relative position of rs17114036 polymorphisms in the luciferase construct. $n = 4$–6. Data represent mean ± SEM. ***$P < 0.0005$ as determined by Student’s $t$ test.
the flow-induced enhancer activity of chr1:56962213–56963412. Luciferase assays detected an increase of the enhancer activity of chr1:56962213–56963412 (with protective C allele at rs17114036) in cells under 18-h unidirectional flow compared with disturbed flow (Fig. 5A), while no significant increase of the enhancer activity by unidirectional flow was detected with the risk T allele (SI Appendix, Fig. S9). In addition to the ATAC-seq experiments in HAECs homozygous at rs17114036 under flow (Fig. 3), we performed ATAC-seq analysis in four HAEC lines heterozygous at rs17114036 under 24-h unidirectional flow to perform open chromatin AI analysis. SI Appendix, Fig. S10 demonstrates that, in all four selected HAEC lines heterozygous at rs17114036, unidirectional flow increases ATAC-seq peaks in the proposed enhancer region in PLPP3 intron 5, in agreement with increased ATAC-seq reads in rs17114036-containing region (Fig. 5B). Moreover, AI analysis showed an enrichment of ATAC from the genome harboring the protective C allele (Fig. 5B). In contrast, ATAC-seq detected no AI at rs6421497, a common SNP in high LD with rs17114036 (SI Appendix, Fig. S11). Indeed, the protective allele C at rs17114036 creates a CACC box that is a binding site for KLF2, which mediates the flow sensitivity of a cohort of endothelial genes, including PLPP3 (13, 24–27). We then tested whether KLF2 dynamically activates this rs17114036-containing enhancer and if rs17114036 alleles impact KLF2-mediated enhancer activity. The affinity of KLF2 to the rs17114036-containing locus was determined by KLF2 ChIP-PCR assays in HAECs carrying a protective allele at rs17114036 (T/C), showing a physical binding of KLF2 to the rs17114036-containing region and the CACC site in the PLPP3 promoter (Fig. 5C). Enhancer activities of chr1:56962213–56963412 were further determined in HAECs as a function of KLF2 expression. Constructs of enhancer (chr1:56962213–56963412) and PLPP3 promoter were cotransfected with KLF2-overexpressing plasmids. Luciferase assays detected a 2.9-fold increase of luciferase activity in the T allele-containing construct as the result of KLF2 overexpression (Fig. 5D). Moreover, KLF2 overexpression led to a 4.7-fold increase of luciferase activity when T allele was substituted by the protective allele C at rs17114036 (Fig. 5D). Collectively, KLF2 ChIP-PCR and luciferase assays demonstrate that CAD/IS-protective allele C at rs17114036 confers a higher KLF2-dependent enhancer activity of chr1:56962213–56963412 in vascular endothelium.

Discussion

Although it is proposed that genetic and environmental factors jointly influence the risk of most common human diseases, the interplay between genetic predisposition and biomechanical cues at the molecular level is poorly understood. The biology underlying the majority of CAD and IS GWAS loci remains to be elucidated (28). Most of the CAD and IS SNPs reside in the noncoding genome. Gupta et al. (29) recently reported that the noncoding common variant at rs943979, implicated in CAD by GWAS, regulates endothelin 1 expression in endothelium. Atherosclerotic lesions preferentially develop in elastic arteries where vascular endothelial cells are activated by local disturbed flow (2–5). As of now, it remains unknown whether disease-associated genetic variants contribute to mechanosensing mechanisms by which cells sense and convert biomechanical stimuli to biological signaling. Our results here elucidate the convergence of CAD/IS genetic predisposition and mechanotransduction mechanisms in endothelial PLPP3 expression. Statistical analyses, whole-genome chromatin accessibility/enhancer marks, CRISPR interference (CRISPRi), genome editing, enhancer assays, caQTL mapping, and AI assay collectively demonstrate that CAD/IS locus 1p32.2 harbors a mechanosensitive endothelial enhancer that regulates PLPP3 expression. Moreover, CAD/IS-protective allele at rs17114036 confers an increased enhancer activity that is dynamically regulated by unidirectional flow and transcription factor KLF2 (Fig. 5E).

Dysregulation of mechanosensing mechanisms contributes to the etiology of a wide range of human diseases in cardiovascular, pulmonary, orthopedic, muscular, and reproductive systems (1). The genetic basis of these complex human diseases has been strongly suggested by GWAS, but the interplay between genetic variants and mechanosensing mechanisms has not been investigated. Our data provide a line of evidence supporting the genetic regulation of mechanotransduction mechanisms in complex human diseases and suggest an underappreciated role of genetic predisposition in cellular mechanosensing processes.

Transcriptional enhancers orchestrate the majority of cell type-specific patterns of gene expression and play key roles in development, evolution, and disease (7), which are tightly regulated by mechanical cues (1). Our data provide molecular evidence that the noncoding genome actively participates in cellular mechanotransduction mechanisms that are influenced by human genetic variances. In addition to the flow regulation of the specific locus 1p32.2, our results provide a dataset to systematically determine the mechanosensitive chromatin accessibility and putative enhancer regions at the whole-genome scale in vascular endothelium. It is important to note that most of the epigenome studies including ENCODE were conducted in cells without physiological or pathophysiological mechanical stimuli, such as HUVECs under static (no flow) conditions (6). Since major endothelial functions are tightly and dynamically regulated by hemodynamics flow, our whole-genome epigenome profiling in HAECs under atherorelevant flows may benefit future studies to investigate mechanical regulation of the noncoding genome in vascular cells. Indeed, we have applied model-based analysis of ChIP-seq (30) and HOMER differential analysis (31), which unbiasedly identified rs17114036-containing locus as 1 of 36,965 open chromatin sites that are activated by unidirectional flow (SI Appendix, Fig. S12).

Mechanosensitive transcription factors have been proposed as major regulators to determine endothelial functions relevant to atherogenesis. For instance, nuclear factor-xB and HIF-1α mediate gene sets associated with proinflammatory, procoagulant, and glycolytic endothelial phenotypes under disturbed flow (32–35), while KLFs and nuclear factor erythroid 2-like 2 regulate gene networks promoting quiescent endothelial phenotype under unidirectional flow (25–27, 36–39). However, the interaction between flow-sensitive transcription factors and disease-associated genetic predisposition in vascular functions has not been suggested. Our results demonstrate that a genetic variant can influence important endothelial functions via a noncoding enhancer region recognized by the mechanosensitive transcription factor KLF2. These results are consistent with emerging evidence showing that top-scoring disease-associated SNPs are frequently located within enhancers explicitly active in disease-relevant cell types (9, 40). Moreover, the data suggest that disease-associated genetic variants, via modulation of transcription factor binding, may regulate the enhancer activities dynamically responding to biomechanical cues that are instrumental to key cellular processes.

GWAS related to atherosclerotic diseases have suggested previously unsuspected loci, genes, and biology involved in lipoprotein metabolism (28), resulting in the development of new cholesterol-lowering therapies (41). Despite that dyslipidemia is a major systemic risk factor of CAD and IS, atherosclerotic lesions largely initiate and develop at arterial regions of atypical vascular geometry associated with disturbed flow. Previous studies demonstrated that cellular mechanotransduction mechanisms, particularly endothelial responses to hemodynamics, causatively contribute to the focal nature of atherosclerotic lesions (2–5, 42). Our studies here demonstrate that genetic variants contribute to not only interindividual variation in plasma
Fig. 5. CAD-protective C allele at rs17114036 promotes flow-induced, KLF2-mediated enhancer activity of chr1:56962213–56963412. (A) Increased enhancer activity of chr1:56962213–56963412 (with C allele at rs17114036) under unidirectional flow (UF) but not disturbed flow (DF). Experiment was performed in biological triplicate and technical triplicate. Data represent mean ± SEM. * P < 0.05 as determined by two-way ANOVA; ** P < 0.005 as determined by two-way ANOVA. The green asterisk denotes the relative position of rs17114036 in the luciferase construct. (B) Increased ATAC-seq reads in rs17114036-containing region in HAECs under UF compared with DF. ATAC-seq was conducted in four HAEC lines heterozygous at rs17114036 under 24-h atherorelevant flows, detecting increased ATAC reads in cells under UF and higher reads from the C allele-containing chromosome compared with T allele. (C) KLF2 affinity to CACC sites in human PLPP3 promoter and intron 5. ChIP-qPCR was performed with either a control IgG antibody or the antibody against HA followed by qPCR using primers detecting CACC sites in PLPP3 promoter or rs17114036-enclosing region from control HAECs (Ctrl) or HAECs transfected with KLF2 transcripts with HA tag. Primers that detect a site ~200 bp from the CACC at rs17114036 were used as a negative control. n = 4. (D) Increased enhancer activity of chr1:56962213–56963412 by KLF2 overexpression. Dual luciferase reporter assays were conducted in HAECx transfected with luciferase constructs containing the PLPP3 promoter and enhancer with either the major (T) or minor (C) allele at rs17114036 along with KLF2-overexpressing or control plasmids. KLF2 overexpression resulted in a 2.9-fold increased luciferase activity in HAECs transfected with T allele-containing construct and a 4.7-fold increase in cells transfected with C allele-containing construct. n = 3. Data represent mean ± SEM. * P < 0.05 as determined by Student’s t test. (E) The interplay between hemodynamic forces, chromatin landscapes at PLPP3 intron 5, and rs17114036 at the molecular level in regulating endothelial PLPP3 expression and vascular functions.
lipid concentrations (43) but also, endothelial responses to blood flow. Indeed, genetic variants at rs17114036 predict CAD susceptibility independent of traditional systemic risk factors, such as cholesterol and diabetes mellitus (10, 11). Recent GWAS identified 15 new CAD risk loci near genes of key functions in endothelial, smooth muscle, and white blood cells (44), further highlighting the potential importance of genetic contribution to the arterial wall-specific mechanisms in atherogenesis. Our results indicate that CAD genetic predisposition and disturbed flow converge to inhibit endothelial PLPP3 expression and that restoration of endothelial PLPP3 in atherosusceptible regions may provide an attractive approach for future arterial wall-based atherosclerosis therapy complementary to current pharmacological treatments targeting systemic risk factors.

Our studies demonstrate that the latest human genetics approaches, such as caQTL mapping (22), AI assay (23), CRISPRi (20), and CRISPR-based assays (19), are powerful tools to investigate possible genetic contributions to cellular mechanotransduction. Miao et al. (45) recently applied CRISPR-Cas9 to achieve high efficiency of a 10-kb deletion of an enhancer region in bulk HUVECs. Here, we expanded the applications of CRISPR-based techniques to investigate key vascular functions. Isogenic adult aortic endothelial lines subjected to CRISPR-based deletion were successfully selected to determine the causal role of a 66-bp genomic region in regulating endothelial PLPP3. Nevertheless, one limitation here is that we are still unable to replace this human SNP at rs17114036 in adult aortic endothelium, although we have tried various methods to promote homology-directed repair. This will be the subject of a future study. Nevertheless, caQTL mapping (22) and AI assay (23) provide complementary approaches detecting at the single-nucleotide resolution that CAD/IS-protective allele at rs17114036 provides a higher enhancer activity at the PLPP3 intron 5.

Cellular mechanotransduction is required for physiological control of tissue homeostasis, while abnormal cell response to mechanical forces promotes pathologies of numerous human diseases. Although investigations, such as GWAS, have suggested the genetic basis of complex human diseases, the interplay between genetic predispositions of mechanosensing mechanisms remains virtually unknown. Our results identified that CAD-associated genetic variant at rs17114036 interacts with hemodynamics in concert to regulate endothelial PLPP3 expression and consequently, key vascular functions. Moreover, our experiments provide evidence supporting the regulatory role of the non-coding, nontranscribed genome in mechanotransduction mechanisms. In summary, this study demonstrates that human haplotypes and related cis-regulatory elements provide an important layer of molecular control by which cells convert physical stimuli into biological signaling.

Methods

ATAC-seq. ATAC-seq was performed as previously described (17) using Tn5 transposase (Illunima). Libraries were sequenced on an Illumina HiSeq 4000 according to the manufacturer's specifications by the Genomics Core Facility at the University of Chicago. The reads were aligned to the UCSC hg19 genome using Bowtie2 (46). ATAC-seq was conducted in HAECs under static conditions or subjected to 24-h unidirectional flow or disturbed flow.

cQTL Mapping and AI. caQTL mapping was performed to test for association between genotype at rs17114036 and chromatin accessibility measured by ATAC-seq. We pulled genotypes for HAE donors from our previous study (47) and imputed linked alleles using IMPUTE2 and SHAPEIT as we published previously (40). Association testing between ATAC-seq tags at the rs17114036 enhancer and genotype was performed using the Combined Haplotype Score test in WASP version 2.1.3.

To perform AI analysis that assigns next generation sequencing reads overlapping heterozygous sites to one chromosome or the other, we quantified ATAC-seq tags at the rs17114036 enhancer using HOMER’s annotatePeaks function to express the log(2)-normalized tags in this region.

CRISPR-Cas9–Mediated Deletion of Enhancer in TeloHAECs. The CRISPR reagents were adapted from the Alt-R system from IDT. The guide RNAs were designed using an online tool at crispr.mit.edu/ to minimize off-targeting effects using two guide RNAs to create a 66-bp deletion. The guide RNAs were made by annealing the transactivating CRISPR RNA (tracrRNA) to the single guide RNA (sgRNA). The RNP complex of S. Pyogenes Cas9 and sgRNA was formed by placing the components together at room temperature and then immediately transfected into cells using Lipofectamine RNAiMAX. For each successive treatment, the reagent amounts were scaled relative to the size of the destination vessel to compensate for the number of cells in the reaction. The volumes for each part of the reaction were increased 4x when treating cells from the 96-well to the 6-well and 16x when moving from the 6-well to the T-75 flask.

Detailed methods are available in SI Appendix.

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