Targeted Deletion of the 9p21 Noncoding Coronary Artery Disease Risk Interval in Mice

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Sequence polymorphisms in a 58kb interval on chromosome 9p21 confer a markedly increased risk for coronary artery disease (CAD), the leading cause of death worldwide \(^1\). The variants have a substantial impact on the epidemiology of CAD and other life-threatening vascular conditions since nearly a quarter of Caucasians are homozygous for risk alleles. However, the risk interval is devoid of protein-coding genes and the mechanism linking the region to CAD risk has remained enigmatic. Here we show that deletion of the orthologous 70kb noncoding interval on mouse chromosome 4 affects cardiac expression of neighboring genes, as well as proliferation properties of vascular cells. Chr4\(^{Δ70\text{kb}/Δ70\text{kb}}\) mice are viable, but show increased mortality both during development and as adults. Cardiac expression of two genes near the noncoding interval, \(Cdkn2\text{a}\) and \(Cdkn2\text{b}\), is severely reduced in chr4\(^{Δ70\text{kb}/Δ70\text{kb}}\) mice, indicating that distant-acting gene regulatory functions are located in the noncoding CAD risk interval. Allele-specific expression of \(Cdkn2\text{b}\) transcripts in heterozygous mice revealed that the deletion affects expression through a \textit{cis}-acting mechanism. Primary cultures of chr4\(^{Δ70\text{kb}/Δ70\text{kb}}\) aortic smooth muscle cells exhibited excessive proliferation and diminished senescence, a cellular phenotype consistent with accelerated CAD pathogenesis. Taken together, our results provide direct evidence that the CAD risk interval plays a pivotal role in regulation of cardiac \(Cdkn2\text{a/b}\) expression and suggest that this region affects CAD progression by altering the dynamics of vascular cell proliferation.

Each day, cardiovascular disease causes 2,400 deaths in the United States alone, more than cancer, accidents and diabetes combined \(^3\). The largest proportion of this mortality is due to coronary artery disease (CAD), which causes approximately 1 of every 5 deaths in the United States. CAD has a complex etiology and there is strong evidence that both environmental and genetic factors
are major determinants of disease risk. However, identifying the genomic loci associated with increased CAD susceptibility has been a challenge, and most of the known risk loci explain only small proportions of CAD cases (e.g., ref. 4). Genome-wide association studies have recently identified common sequence variants on human chromosome 9p21 that confer an increased risk for CAD and myocardial infarction. These associations have been confirmed in multiple additional cohorts and were extended to other severe arterial diseases. Even in homozygous individuals, the variants increase the relative risk for CAD only moderately by a factor of 1.3 to 2. However, since the risk alleles are very common, they contribute substantially to the epidemiology of CAD. Between 20% and 25% of Caucasians are homozygous for risk alleles, resulting in estimates of 10% to 31% population attributable risk, depending on cohort and cases considered.

Despite compelling genetic evidence for association, the mechanism by which 9p21 sequence polymorphisms confer an increased CAD risk is unknown, preventing the development of pharmacological or behavioral intervention strategies. The variants are not associated with established CAD risk factors such as plasma lipoprotein levels, hypertension, or diabetes, suggesting that they influence CAD pathogenesis through a previously unappreciated pathway. The CAD-associated SNPs are located within a 58kb linkage disequilibrium block on chromosome 9p21.3 that does not contain any known protein-coding genes. Several expressed sequence tags of apparently noncoding transcripts, including a proposed long noncoding RNA, have been mapped to the risk interval, but their functional relevance remains elusive. Gene expression studies in human peripheral blood cells and in vitro reporter assays have provided support for the notion that gene regulatory elements might be located within the risk interval, but there is conflicting evidence whether the CAD risk variants are associated with increased or decreased regulatory activity. More importantly, it is unclear whether altered gene regulation would result in cellular or physiological
phenotypes that are relevant to CAD pathogenesis, highlighting the need to study the function of this noncoding interval in a suitable *in vivo* system.

To create a mouse model for investigating the function of the human 58kb noncoding CAD risk interval, we sought to generate a severe (null) allele for this locus by its targeted removal from the mouse genome. Human-mouse orthology could be unambiguously established since 50% of the basepairs in the human region are alignable to mouse and synteny with flanking genes is preserved (Fig. 1a-c). The mouse interval is 70kb in size and thus 20% longer than the human orthologous region, partially due to increased repetitive sequence content. Due to the large size of the interval of interest, its targeted deletion was accomplished through a sequential double-targeting strategy followed by Cre-mediated recombination (Fig 1d; Suppl. Figures 1-3). Chr4Δ70kb/Δ70kb mice are viable and fertile. While the majority of live-born homozygous animals survive to weaning and beyond without obvious morphological or behavioral phenotypes, we did observe reduced embryonic, post-natal, and adult survival due to the deletion (for details see Suppl. Material and Suppl. Fig. 4). Clinical analysis of adult animals at seven months of age showed no significant general aberrations in urine and blood chemistry markers, differential blood cell counts, or histopathological appearance of internal organs including heart, liver, lung, kidney, spleen, and gastrointestinal tract. However, in a larger cohort of chr4Δ70kb/Δ70kb mice on which gross necropsy was performed between 7 and 14 months of age (or at time of premature death), 9 of 20 (45%) animals were found to have internal neoplasms or tumors of various types (see Suppl. Material), compared to none in a cohort of age-matched wild-type controls (*P* = 0.0012, FET). In addition to this increased tumor incidence, both male and female chr4Δ70kb/Δ70kb mice that were fed standard mouse chow *ad libitum* gained weight significantly faster than wild-type controls, resulting in a 17% increased body mass by 30 weeks of age (Suppl. Fig. 5).
To investigate the effects of the deletion in more detail, we examined the possibility that the CAD risk interval is required for distant-acting regulation of gene expression. To compare mRNA expression levels of surrounding genes between wild-type and chr4Δ70kb/Δ70kb mice, we isolated mRNA from hearts and other adult mouse tissues and performed reverse transcription followed by quantitative PCR. In heart tissue, chr4Δ70kb/Δ70kb mice had substantially depressed expression levels of the neighboring Cdkn2a and Cdkn2b genes, but no significant alteration of expression levels of two other neighboring genes, Mtap and Dmrt1a (Fig. 2a). Cardiac expression of Cdkn2a and Cdkn2b was more than ten-fold decreased compared to wild-type controls. These results indicate that the CAD risk interval is required for appropriate expression of Cdkn2a and Cdkn2b in the heart. Due to the known roles of these genes in several disease-related pathways 16-19, these results support the possibility of a regulation-mediated mechanism by which the chr4Δ70kb deletion might impact on cellular, physiological and pathological processes.

To test whether the observed regulatory effect on gene expression occurs through a cis- or trans-acting mechanism, we performed allele-specific expression analysis. We used for this purpose crosses of mice with the deletion linked to the Cdkn2b allele of the 129Sv strain (in which the deletion was originally created) and wild-type C57BL/6 strain mice. Strains C57BL/6 and 129Sv are distinguished by several transcribed SNPs in the Cdkn2b gene that can reveal quantitative differences in expression from the two alleles. Direct sequencing of PCR product from tail genomic DNA confirmed that the mice were heterozygous for the expected SNPs and the two alleles were detected at the expected 1:1 ratio (Fig. 3a,b). As a control, in cDNA derived from tissues of C57BL/6 (wt) x 129Sv (wt), the two alleles were also expressed at indistinguishable levels, confirming the absence of general strain-specific differences in Cdkn2b expression levels (Fig. 3b). In contrast, direct sequencing of RT-PCR product derived from RNA isolated from
chr4+(C57BL/6)/Δ70kb(129Sv) hearts and other tissues revealed that Cdkn2b was predominantly expressed from the wild-type C57BL/6 allele and expression from the chr4Δ70kb 129Sv allele was strongly diminished (Fig. 3b). Among five organs and cell types examined, the most severe allele-specific down-regulation was observed for the heart and aorta (Fig. 3c). These results support that the CAD risk interval controls gene expression in cardiac and other tissues through a distant-acting cis-regulatory mechanism.

The proteins encoded by Cdkn2a, Cdkn2b and other cyclin-dependent kinase inhibitor genes have been implicated in cellular phenotypes including regulation of proliferation and cellular senescence. Given the severe impact of the chr4Δ70kb deletion on cardiac expression of Cdkn2a and Cdkn2b, we tested if cell proliferation and senescence are affected in chr4Δ70kb/Δ70kb mice. To test for an effect on cell proliferation, we measured the proliferation rates of primary cultures of aortic smooth muscle cells (aSMCs) and mouse embryonic fibroblasts (MEFs) during early passages. In both cell types, cells derived from chr4Δ70kb/Δ70kb mice proliferated excessively compared to wild-type controls, with daily proliferation rates nearly 2-fold increased in aSMCs and nearly 3-fold increased in MEFs (Fig. 4a,b). During later passages of these primary cultures, wild-type aSMCs and MEFs became senescent, whereas chr4Δ70kb/Δ70kb-derived cells that had been isolated and cultured under identical conditions continued to proliferate and did not show signs of senescence (Fig. 4c,d). These cellular phenotypes are consistent with known and proposed functions of Cdkn2a, Cdkn2b, and other cyclin-dependent kinase inhibitors.

The risk interval affects human CAD through a mechanism that appears to be independent of plasma lipid levels and other known risk factors. To study possible in vivo effects of the chr4Δ70kb allele on plasma lipids and early stages of atherogenesis, we placed 40 chr4Δ70kb/Δ70kb mice and 40 wild-type controls in an isogenic 129Sv background on a high-fat, high-cholesterol (“Western”) diet for 20
weeks 21. As expected, this diet caused substantial alterations in plasma lipid levels, however, no significant differences in this physiological response were observed between wild-type and chr4Δ70kb/Δ70kb mice (Suppl. Figure 6; see Suppl. Material for details). In addition, we did not observe significant differences in fatty lesion formation (Suppl. Figure 7). Nevertheless, the high-fat, high-cholesterol nutrition caused substantially increased mortality among chr4Δ70kb/Δ70kb mice compared to isogenic wild-type controls, indicating an overall increased susceptibility to detrimental effects of this noxious diet (Suppl. Fig. 8). Studies in complementary background strains and additional genetic manipulations of the lipid metabolism in chr4Δ70kb/Δ70kb mice may be required to mimic the full course of human atherosclerosis 21-23. Such data could also help to distinguish if diet-induced increase in mortality is due to cardiovascular phenotypes other than the aortic fatty lesions examined here, or to extra-cardiovascular phenotypes that may be present in chr4Δ70kb/Δ70kb mice. Irrespective of the underlying etiology, these observations indicate that the CAD risk interval is not required for maintenance of normal plasma lipid levels in mice, consistent with the observation that variation in the human interval influences CAD risk independent of altered lipid levels.

The chromosome 9p21 common haplotype linked to CAD represents an important but particularly puzzling risk interval and the present study provides key insights regarding the in vivo function of this noncoding region. We have demonstrated that the precise orthologous mouse interval, despite its large distance from any protein-coding genes, is critically required for normal cardiac expression of two cell cycle inhibitor genes, Cdkn2a and Cdkn2b. These observations raise the question as to what type of underlying molecular mechanism mediates these regulatory effects. Subregions of the 58kb noncoding risk interval increase transcriptional activity in cell-based in vitro transfection assays 13, but the location and function of distinct small enhancer sequences with relevant in vivo activities remains to be established (see Supplementary Material).
Alternatively, our results are also consistent with models in which the noncoding interval shields the \textit{Cdkn2a/b} genes from the influence of very distal negative regulatory elements, either by spacing effects or due to presence of insulator elements. In contrast, any mechanism mediated by freely diffusible RNA molecules is not expected to result in the allele-selective regulation observed in chr4\textsuperscript{$\Delta$70kb/$\Delta$70kb} mice (Fig. 3). Support for an RNA-mediated mechanism\textsuperscript{11,12,24} is therefore restricted to scenarios in which the transcriptional activity itself affects local chromatin state or in which RNA molecules remain tethered to the chromosome from which they are transcribed. We have also shown that the aberrations of \textit{in vivo} expression of \textit{Cdkn2a} and \textit{Cdkn2b} coincide with abnormal regulation of vascular cell proliferation and senescence. These phenotypes are reminiscent of mouse models in which the \textit{Cdkn2a/b} genes themselves have been deleted. Specifically, primary cultures of \textit{Cdkn2a-} or \textit{Cdkn2b-deficient fibroblasts}\textsuperscript{16,19} and \textit{Cdkn2a-deficient} aSMCs\textsuperscript{25} exhibit elevated proliferation rates. Hence, a parsimonious explanation for the cellular phenotypes observed in chr4\textsuperscript{$\Delta$70kb/$\Delta$70kb} mice is that the noncoding CAD risk interval affects vascular cell proliferation and senescence by modulating the expression levels of \textit{Cdkn2a} and \textit{Cdkn2b}. Altered proliferation rates of vascular cells resulting from genetic manipulation of other \textit{cyclin-dependent kinase inhibitors} have been closely linked to the dynamics of CAD pathogenesis\textsuperscript{26} and \textit{Cdkn2a} deficiency causes altered vascular injury responses in a mouse model of CAD\textsuperscript{25}. Moreover, sequence polymorphisms in the promoters of at least two other human \textit{cyclin-dependent kinase inhibitor} genes have been implicated in increased cardiovascular disease risk\textsuperscript{27,28}. Thus, variation in distant-acting regulatory sequences required for cardiovascular expression of \textit{CDKN2A} and \textit{CDKN2B} provides a plausible mechanistic model for the increased CAD risk associated with the 9p21 region independently of lipid levels and other known risk factors.
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**Author Contributions**

A.V. and L.A.P. wrote the manuscript. All authors contributed to data collection and analysis and provided comments on the manuscript.

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Methods Summary

Targeted deletion of the 70kb noncoding interval was performed by two sequential targeting steps, followed by Cre-mediated recombination (Suppl. Figures 1-3). See Suppl. Table 1 for primer sequences used for vector construction and genotyping. For quantitative real-time reverse transcription PCR (RT-PCR), total RNA was extracted from wild-type and knockout mouse tissues using Trizol Reagent (Invitrogen) following the manufacturer's instruction. Total RNA was treated with RNase-free DNase and first-strand cDNA was synthesized by standard methods. RT-PCR was performed by standard methods; primer sequences are provided in Suppl. Table 2. For allele-specific expression profiling, RNA extraction from tissues, DNase treatment and first-strand cDNA synthesis were done as described above. Genomic DNA was extracted from the tails of the same mice used for RNA extraction by standard methods. PCR products were generated using gene-specific primers containing M13 primer tags (M13 -40 forward or M13 -20 reverse) and Platinum Taq DNA Polymerase (Invitrogen). Primer sequences are provided in Suppl. Table 2. PCR products were gel purified using QiaGen MinElute Gel Extraction Kit (Qiagen) and sequenced using M13 primers. For proliferation and senescence assays, primary mouse embryonic fibroblasts (MEFs) were isolated from embryonic day 12.5 to 14.5-day-old chr4^Δ70kb/Δ70kb, wild-type and heterozygous littermates. Each embryo was disaggregated in 0.25% trypsin, and fragments were cultured in Dulbecco’s modified Eagle's medium (DMEM) with 10% fetal bovine serum. Aortic smooth muscle cells were isolated from thoracic aorta of 4-week old chr4^Δ70kb/Δ70kb mice and wild-type littermates as previously described 29. Cells were counted at each passage and re-cultured in constant concentrations (2x10^4/well). For senescence assays, cells were grown to senescence (passage 12), trypsinized and plated at 2x10^4/well on day 0. Cell counts were determined after 4 days. Senescence staining by X-Gal was done as previously described 30.
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Figure Legends

Figure 1: Deletion of the noncoding region orthologous to the 58kb CAD risk interval on human chromosome 9p21. a) Noncoding CAD risk interval with SNPs found to be most significantly associated with CAD in genome-wide association studies 1,2 and SNPs defining the boundaries of the linkage disequilibrium block. b) Overview of the human locus including neighboring genes (blue, intron/exon structure not shown). A noncoding RNA of unknown function transcribed from this locus 12 is shown in green (CDKN2BAS, also known as ANRIL, GenBank NR_003529.3). c) Orthologous region on mouse chromosome 4. The exon structure of a noncoding transcript of unknown function, AK148321, is shown in green. d) Chr4Δ70kb after successful targeting and deletion of the 70kb CAD risk interval.

Figure 2: Deletion of the CAD risk interval affects expression of neighboring genes Cdkn2a and Cdkn2b. a) Expression levels of Cdkn2a (INK splice variant) and Cdkn2b are significantly reduced in heart tissue of chr4Δ70kb/Δ70kb mice. In contrast, expression of the neighboring genes Mtap and Dmrtai is not significantly altered. b) Overview of locus with locations of genes. Error bars are standard error of the mean; asterisks indicate P ≤ 0.005 (t-test, one-tailed).

Figure 3: Deletion of the CAD risk interval affects gene expression through a cis-regulatory mechanism. a) Two of seven transcribed single nucleotide polymorphisms (SNPs) that were used to distinguish expression of the C57BL/6 and 129Sv strain alleles of the Cdkn2b gene. Representative electropherograms from direct Sanger sequencing of PCR product are shown. b) No differences between alleles are observed in genomic DNA or tissue-derived cDNA in chr4+(C57BL/6)/+(129Sv) mice or in genomic DNA of chr4+(C57BL/6)/Δ70kb(129Sv) mice. In contrast, in tissues of chr4+(C57BL/6)/Δ70kb(129Sv) mice, the C57BL/6 allele is expressed
four-fold higher than the 129Sv allele (all tissues combined). c) Allelic expression differences shown by individual tissues. Error bars indicate standard deviation (b) and standard error of means (c). P-values: t-test, two-tailed.

Figure 4: Deletion of the CAD risk interval disrupts normal dynamics of cellular proliferation and senescence. a) Increased proliferation of primary aSMC cultures. b) Increased proliferation of primary MEF cultures. c) Failure of normal cellular senescence of primary aSMC cultures in late passages. Cells derived from wild-type and chr4Δ70kb/Δ70kb mice were grown to senescence under identical conditions, seeded at equal densities and cell counts were determined after 4 days. d) Chr4Δ70kb/Δ70kb-derived MEFs fail to enter cellular senescence, as evident from absence of senescence staining by X-Gal in comparison to wild-type MEFs30. Mean daily proliferation rates are shown in a) and b), viable cell count at day 4 is shown in c). Error bars indicate SEM. *, P < 0.05; **, P<0.01; t-test.
**Online Methods**

**Targeted Deletion of 70kb Noncoding Interval**

Two targeting vectors were generated for the deletion. See Suppl. Table 1 for primer sequences used for vector construction and genotyping. One targeting vector, containing a homologous region at the 5’ end of the region of interest (proximal to Cdkn2a/b), was in the ploxPneoTK-2 vector. The second targeting vector, containing a homologous region at the 3’ end of the region (distal from Cdkn2a/b), was in a ploxPhygTK vector backbone. Both ploxPneoTK-2 and ploxPhygTK were generated in this laboratory.

Homologous arms were generated by PCR from W4/129S6 ES cell genomic DNA. The PCR product of the 5’ homologous arm (with BamHI and EcoRI tags on primers) was cloned into ploxPneoTK-2 BamHI/EcoRI sites next to a loxP site, generating vector pCHD5’neo for 5’ targeting. The PCR product of the 3’-homologous arm (with NotI and BglII tags on primers) was cloned into ploxPhygTK NotI/BamHI sites at the 3’ side of the PGK terminator of the PGKtk cassette, generating targeting vector pCHD3’hyg for 3’ targeting.

Targeting vector pCHD5’neo was electroporated into W4/129S6 ES cells (Taconic Farms). Neomycin-resistant clones were picked, screened by PCR and confirmed by Southern hybridization (Suppl. Fig. 1). Successfully 5’-targeted clones were pooled and electroporated with 3’ targeting vector pCHD3’hyg. Hygromycin-resistant clones were picked, screened by PCR and confirmed by Southern hybridization (Suppl. Fig. 2).

Double-targeted clones (Hyg’/Neo’, note that the two loxP sites could be in cis or in trans) were pooled together, expanded and electroporated with about 20 ug of Cre-recombinase-expressing plasmid pTURBO-Cre. When loxP sites were present in cis, the loxP bracketed sequence included: 1) the region chr4:89,054,800-89,126,878 (mm9) to be deleted, 2) the PGKhyg, PGKneo and HSV-tk cassettes, which can be deleted by Cre-recombinase-mediated loxP recombination (Suppl. Fig. 3). When loxP sites were present in trans, a translocation could be generated that resulted in one deleted and one duplicated allele of the region of interest. ES cells that underwent cis-recombination and
deletion were identified by selecting for neomycin or hygromycin sensitivity and negative selection of HSV-tk for 1-(2-deoxy-2-fluoro-b-D-arabinofuranosyl)-5-iodouracil (FIAU) resistance. Cells surviving this selection were screened by PCR for the predicted deletion using a primer outside the deleted region and T7 primer within the vector backbone left on the chromosome after deletion. The predicted deletion was further confirmed by PCR using primers outside the deleted region, negative PCR of Neo or Hyg primers, and by Southern blot analysis using a probe outside the deletion.

Assessment of Survival

To determine embryonic survival, embryos from timed pregnancies were dissected between E9.5 and E15.5. Embryos whose size and appearance was normal for the respective stage were considered as surviving and genotyped by PCR.

To determine the survival from birth to weaning, live-born pups resulting from \( \text{wt} \times \text{wt}, \text{chr}^4_{+/\Delta70kb} \times \text{chr}^4_{+/\Delta70kb} \) and \( \text{chr}^4_{\Delta70kb/\Delta70kb} \times \text{chr}^4_{\Delta70kb/\Delta70kb} \) crosses were counted at day P0 or P1 and the same litters were counted again at weaning at day P20 or P21. Any dead or missing pups were considered as having died between birth and weaning. Pups from heterozygous crosses were counted regardless of their genotype since missing pups could generally not be recovered for genotyping.

For adult survival analysis on standard chow and on high-fat diet, animals that were found dead, met euthanasia criteria, or had an expected remaining survival time of less than 7 days (based on progression of pre-mortal symptoms in previously died animals, which included severe weight loss and general inactivity) when sacrificed for histological analysis were considered as died. Tick marks in survival plots indicate “censored” animals, which includes animals that were healthy when removed from the study for histological analysis. In order to exclude the possibility of bias in assessing euthanasia criteria, we also performed survival analysis for animals on the high-fat diet up to 120 days. All animals that died during this initial study phase were found dead and no animals were euthanized or removed for histological analysis prior to day 120. Consistent with the full study duration, a significantly increased mortality among \( \text{chr}^4_{\Delta70kb/\Delta70kb} \) animals on high-fat diet.
compared to wild-type controls was observed ($P = 0.006$, Kaplan-Meier log-ranked survival test).

**High-fat Diet, Plasma Lipid and Aortic Fatty Lesion Analysis**

Deletion and wild-type mice at 6 weeks of age were fed with high-fat diet containing about 15.8% fat, 1.25% cholesterol and 0.5% sodium cholate (Harlan, TD.88051). Mice were under the diet for 18-22 weeks. The control group was fed with chow diet containing about 6.5% fat (Labdiet, Formulab Diet 5008). For plasma lipid analysis, mice were fasted overnight (approximately 15-17 hours). Whole blood was collected into EDTA Capillary Blood Collection tube (Fisher) by tail bleeding. Blood cells were removed by centrifugation at 4 °C. Clear plasma was transferred to a new tube and frozen at -80 °C until analysis. Plasma lipids were measured by standard assays at the Cincinnati Mouse Metabolic Phenotyping Center. Aortic fatty lesion analysis was done as previously described. Briefly, mouse hearts were excised and the upper third, including the proximal aorta, was embedded in optimal cutting temperature (OCT) compound. Serial 10 µm thick cryosections were cut in the region extending from the appearance to the disappearance of the aortic valves. Sections were mounted on Superfrost (VWR) slides, fixed in 10% neutral buffered formalin vapor and stained with Oil-Red O in PEG and Gill's III hematoxylin. Lesion areas were determined using a calibrated eyepiece at 200 X magnification.

**Real-time RT-PCR**

Total RNA was extracted from wild-type and knockout mouse tissues using Trizol Reagent (Invitrogen) following the manufacturer’s instruction. Total RNA was treated with Promega RQ1 RNase-Free DNase. First-strand cDNA was synthesized using SuperScript™ First-Strand Synthesis System for RT-PCR (Invitrogen). Real-time RT-PCR was performed using the Applied Biosystems SYBR Green PCR Master Mix and run on a 7500 Fast Real-Time PCR System (Applied Biosystems). Primer Sequences are provided in Suppl. Table 2.
Allele-specific Expression Profiling

RNA extraction from tissues, DNase treatment and first-strand cDNA synthesis were done as described above. Genomic DNA was extracted from the tails of the same mice used for RNA extraction. Tails were digested overnight at 50 °C in a buffer containing 1% SDS and 100-200 µg/ml Proteinase K. The lysate was heated at 95-100 °C for 5 minutes, and diluted for PCR. PCR products were generated using gene-specific primers containing M13 primer tags (M13 -40 forward or M13 -20 reverse) and Platinum Taq DNA Polymerase (Invitrogen). Primer sequences are provided in Suppl. Table 2. PCR products were gel purified using QiaGen MinElute Gel Extraction Kit (Qiagen) and sequenced using M13 primers.

Histopathology, Blood and Urine Analysis

Complete clinical blood chemistry profiles, hematological analysis, urine analysis, general histopathology, and histopathological analysis of neoplasms and tumors were performed at Charles River Research Animal Diagnostic Services (Wilmington, MA).

Proliferation and Senescence Assays

Primary mouse embryonic fibroblasts (MEFs) were isolated from embryonic day 12.5 to 14.5-day-old chr4\(^{Δ70kb/Δ70kb}\), wild-type and heterozygous littermates. Each embryo was disaggregated in 0.25% trypsin, and fragments were cultured in Dulbecco’s modified Eagle’s medium (DMEM) with 10% fetal bovine serum. Aortic smooth muscle cells were isolated from thoracic aorta of 4-week old chr4\(^{Δ70kb/Δ70kb}\) mice and wild-type littermates as previously described \(^{29}\). Cell counts were determined at each passage using a hemocytometer and re-cultured in constant concentrations (2x10\(^4\)/well). Fig. 4a shows mean daily proliferation rates over seven early passages in aSMC cultures derived from three animals per genotype. Fig. 4b shows mean daily proliferation rates over four early passages of MEFs derived from four (heterozygous) to six (wild-type and homozygous) animals per genotype. For senescence assays, cells were grown to senescence, trypsinized and plated at 2x10\(^4\)/well on day 0. Fig. 4c shows senescence data from primary aSMC cultures in late passages (passage 12), derived from five animals per genotype. Cells were grown to senescence under identical conditions, seeded at equal densities and cell counts
were determined after 4 days. Senescence staining by X-Gal in MEFs (Fig. 4d) was done as previously described \(^{39}\) after 8 passages.

**Reference (Online Methods)**

\(^{31}\) Ueda, Y. et al., Relationship between expression levels and atherogenesis in scavenger receptor class B, type I transgenics. *J Biol Chem* 275 (27), 20368-20373 (2000).
Figure 1
Figure 2

(a) Adult heart RNA expression levels normalized to β-actin (n=6) for wild-type littermates and chr4Δ70kb/Δ70kb (n=6).

(b) Genomic map showing the Mtap, Cdkn2a, Cdkn2b, and Dmrt1a regions, with the CAD risk interval (deleted in chr4Δ70kb) indicated.
Figure 3

a) Diagram showing genetic variation in the Cdkn2b gene. The figure illustrates genomic DNA and cDNA expression in different tissues: tail, heart, liver, and aorta. The genotypes include C57BL/6 and 129Sv with and without a 70kb non-coding sequence.

b) Bar graph showing genomic expression (all tissues) for C57BL/6 and 129Sv alleles. The expression is quantified for different allele ratios, and the statistical significance is indicated (P=6e-52, P=3e-6).

c) Bar graph indicating expressed Cdkn2b allele ratio by tissue for C57BL/6 and 129Sv alleles. The tissues include aorta, heart, leukocyte, liver, and spleen, with allele ratios ranging from 0% to 100%.
Figure 4
Targeted Deletion of the 9p21 Noncoding Coronary Artery Disease Risk Interval in Mice

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Reduced Survival in chr4^Δ70kb/Δ70kb Mice

To assess a possible quantitative effect of the chr4^Δ70kb allele on general viability, we compared the embryonic, postnatal and adult survival of chr4^+/Δ70kb and chr4^Δ70kb/Δ70kb mice to wild-type controls. We genotyped a total of 78 litters at embryonic stages between E9.5 and E15.5 to determine a possible effect of the deletion on prenatal survival. At stages E9.5 and E10.5, no significant deviation from expected Mendelian ratios was observed among surviving morphologically normal embryos (n = 191, P = 0.29, G-test). In contrast, among embryos collected between E11.5 and E15.5, chr4^+/Δ70kb and chr4^Δ70kb/Δ70kb genotypes were markedly depleted (113 wild-type : 200 heterozygous : 70 null embryos, P = 0.004), indicating that the chr4^Δ70kb deletion negatively impacts on embryonic survival up to E15.5 (Suppl. Fig. 4a). Next, we tested a possible effect of the chr4^Δ70kb deletion on postnatal viability. Since the genotype of pups from chr4^+/Δ70kb × chr4^+/Δ70kb crosses was generally not determined before weaning age (~3 weeks), we compared the survival from birth to weaning among 1,964 pups from wt × wt, chr4^+/Δ70kb × chr4^+/Δ70kb and chr4^Δ70kb/Δ70kb × chr4^Δ70kb/Δ70kb crosses. None of 345 pups from wt × wt crosses died before weaning. In contrast, 28 of 1,154 pups from heterozygous crosses (2.4%) and 24 of 465 pups from null crosses (5.2%) died before weaning (Suppl. Fig. 4b), suggesting that the chr4^Δ70kb allele is associated with decreased survival of live-born pups to weaning age (P = 0.001 and P = 1.7×10^-6, respectively; two-tailed Fisher’s Exact test). To determine if this decrease in viability also extends into adulthood, we monitored the survival of 16 chr4^Δ70kb/Δ70kb mice that had survived to weaning and 16 wild-type controls for over one year under standard conditions with regular mouse chow fed ad libitum (Suppl. Fig. 4c). During this study period, 5 of the null mice (31%), but none of the wild-type controls died (P = 0.016, Kaplan-Meier test). The cause of death was usually not unambiguously determined and correlated only in some cases with the presence of tumors. Of note, in addition to expression changes in the heart and aorta, we also observed significant changes in expression of Cdkn2a/b in other tissues (Fig. 3c). Considering that these genes and the larger 9p21 locus have been implicated in several common forms of disease, the relative contribution of CAD-related phenotypes,
increased cancer incidence or other phenotypes remains to be determined. Taken together, these results indicate that the 70kb noncoding region containing the CAD risk interval is required for normal survival and suggest that molecular and physiological mechanisms negatively affected by the chr4Δ70kb deletion are not temporally restricted to a particular developmental stage, but rather persist throughout much of embryonic development and postnatal life in mice.

**Effects of High-Fat Diet in chr4Δ70kb/Δ70kb Mice**

Studying atherogenesis in mouse models represents a challenge since most mouse strains develop severe atherogenesis only upon genetic manipulation of their lipid metabolism. However, the use of a high-fat, high-cholesterol diet (“Western” diet) can provide a model for fatty streak formation, an early stage of the atherogenic process, in the genetic background strain for the chr4Δ70kb deletion, 129Sv. This strain is moderately susceptible to fatty lesion formation in this diet-induced model, allowing an initial assessment of genetic influences on early stages of atherogenic plaque formation. To determine if the chr4Δ70kb deletion affects fatty streak formation, 40 chr4Δ70kb null mice and 40 wild-type controls were fed Western diet ad libitum for approximately 20 weeks. The diet caused marked aberrations in plasma lipid levels, namely a 2.5-fold reduction in triglycerides and a 2-fold increase in plasma cholesterol (Suppl. Fig. 6). However, no significant difference in this physiological response to the diet was observed between chr4Δ70kb null mice and controls. Following the Western diet regimen, formation of aortic fatty lesions was quantitated. Both in wild-type controls and in chr4Δ70kb null mice, the occurrence of lesions varied widely, from virtually absent to moderate levels. While a mild increase in median lesion size was observed in chr4Δ70kb/Δ70kb mice, the difference was overall not significant (Suppl. Fig. 7). Despite this lack of significant differences in plasma lipid levels and fatty lesion formation, we observed a substantially lowered tolerance for the high-fat diet in chr4Δ70kb/Δ70kb mice, resulting in increased mortality during the course of the feeding study (Suppl. Fig. 8). This was in sharp contrast to wild-type control animals in which cases of apparently diet-related deaths were observed only after several months of high-fat diet and at a lower frequency. The cause of death in chr4Δ70kb/Δ70kb animals could not be unambiguously determined, but was in most cases preceded by general signs of malaise over several days, including severe weight loss and
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decreased activity. The etiological link between the high-fat diet and increased mortality in chr4Δ70kb/Δ70kb mice remains to be elucidated, but it is noteworthy that this noncoding interval appears to influence both CAD risk in humans 1,2 and increased mortality in mice on a high-fat diet through a mechanism that is independent of plasma lipid levels.

Tumors Observed in chr4Δ70kb/Δ70kb Mice

Seven solid masses that developed spontaneously in chr4Δ70kb/Δ70kb mice were resected following euthanasia and further examined histopathologically (Charles River Research Animal Diagnostic Services). Six of seven masses were confirmed to be tumors, of which five were different types of sarcomas (Suppl. Table 3). Of note, sarcomas occur spontaneously at increased frequency in Cdkn2aInK4a-/- mice 16, possibly indicating a phenotype that is mediated by transcriptional down-regulation of Cdkn2a in chr4Δ70kb/Δ70kb mice. In addition, remarkable hepatosplenomegaly was evident in several chr4Δ70kb/Δ70kb and chr4+/Δ70kb mice. Histological analysis of two chr4Δ70kb/Δ70kb cases revealed different degrees of extramedullary hematopoiesis, which is again consistent with a phenotype observed in Cdkn2a-/- mice 16. These observations are also interesting in light of the increased susceptibility for other types of cancer associated with the 9p21 locus, e.g. glioma 32,33.

Scanning the Noncoding Risk Interval for Conserved Enhancers

The down-regulation of Cdkn2a and Cdkn2b in heart and other tissues of chr4Δ70kb/Δ70kb mice (Fig. 2) in conjunction with the marked differences in expression between the wild-type and chr4Δ70kb alleles in heterozygous mice (Fig. 3) provides direct evidence for a cis-regulatory function of the noncoding 70kb interval. To further examine if this regulatory function can be attributed to a smaller single enhancer element, we performed comparative genomic analysis of the human 58kb CAD risk interval at chr9p21 and a series of transgenic mouse experiments. To identify enhancer candidate elements, we assessed multi-vertebrate genome alignment data at the UCSC browser 34. Taking conservation depth, as well as vertebrate evolutionary conservation scores 35 into account, we selected six highly conserved noncoding sequence elements from the larger noncoding interval for transgenic mouse experiments (Suppl. Fig. 9). These sequences were generally well-conserved within the mammalian clade.
and had high conservation scores compared to other regions of the risk interval. Based on previous studies using comparative vertebrate genomics for predicting distant-acting enhancers\textsuperscript{36}, these six sequences were considered to be the most likely subregions of the larger CAD risk interval to be associated with enhancer activity. We cloned all six candidate regions (see Suppl. Table 4 for primer sequences) and tested them using a previously described transgenic mouse reporter assay\textsuperscript{37}. In total, 1,243 pronuclear injections (average: 207 per construct) of single-cell stage mouse embryos were performed for these studies. For each construct, we obtained at least 5 (average: 7) LacZ-stained embryos resulting from independent genomic integration events. Embryos were isolated and stained for LacZ activity at embryonic day (e) 11.5 and reporter gene expression patterns were annotated using established reproducibility criteria for this type of assay\textsuperscript{37}. In this annotation scheme, elements are only considered to be an enhancer if LacZ staining is observed in the same anatomical structure in at least three embryos resulting from independent transgenic integration events.

In none of the six cases, a reproducible staining pattern in any embryonic structure (including the heart and vascular system) was observed, indicating that none of the conserved noncoding sequences tested was a reproducible \textit{in vivo} enhancer in this assay at this embryonic stage. In light of the functional evidence from our deletion studies, there are several possibilities to explain this observation. These include: a) one of these sequences could be an enhancer at later time points than e11.5; b) the risk interval contains an enhancer that is not well-conserved in evolution and therefore not easily identified by comparative genomic methods; c) either the enhancer itself or the transcription factors binding to it are not sufficiently conserved between human and mouse to detect \textit{in vivo} activity in this mouse assay using the human candidate sequences; d) the CAD risk interval does not contain any single small subregion (tested elements were 2.1kb-3.2kb in size) that acts a classical enhancer detectable in this assay. This would, e.g., be the case if combinatorial binding of transcription factors to different subregions of the interval was required or if the \textit{in vivo} regulatory effects observed in our deletion studies were primarily due to spacing effects or deletion of non-enhancer types of regulatory element. It is expected that additional experimental studies will be required to pinpoint the exact location and \textit{in vivo} function of such smaller functional elements in the risk interval.
Supplementary Figures

Suppl. Figure 1 – Targeting the 5’-end of the 70kb risk interval

a) Schematic strategy for introduction of a LoxP site near the 5’-end of the region of interest.
b) Validation of successful recombination by Southern hybridization.  
c) Validation of successful recombination by PCR.  See Methods for details.
Suppl. Figure 2 – Targeting the 3'-end of the 70kb risk interval

a) Schematic strategy for introduction of a LoxP site near the 3'-end of the region of interest.
b) Validation of successful recombination by Southern hybridization.
c) Validation of successful recombination by PCR. See Methods for details.
Suppl. Figure 3 – Deletion of 70kb-interval by Cre-mediated recombination after double-targeting

a) Schematic strategy for recombination of double-targeted loci, resulting in deletion of the 70kb region of interest. 
b) Validation of successful deletion by Southern hybridization. 
c) PCR genotyping results of wild-type, heterozygous and homozygous chr4<sup>∆70kb</sup> mice from crosses of chimera-derived heterozygous founders.
Suppl. Figure 4 – Decreased embryonic, postnatal and adult survival

a) chr4+/Δ70kb and chr4Δ70kb/Δ70kb genotypes are underrepresented among morphologically normal, non-aborted embryos collected between embryonic days E11.5 and E15.5, indicating reduced survival compared to wild-type embryos. b) Pups from heterozygous and homozygous chr4Δ70kb crosses are more likely to die between birth and weaning. c) Chr4Δ70kb/Δ70kb mice are less likely to survive up to 55 weeks of age than wild-type controls. P-values: G-test (a), two-tailed Fisher’s Exact test (b), Kaplan-Meier test (c).
Suppl. Figure 5 – Chr4$^{\Delta70kb/\Delta70kb}$ mice have increased body weight

Growth curve on standard chow (ad libitum) up to 30 weeks of age. Plotted values are means ± SEM for 8 animals per gender and genotype. P=0.002 (t-test, two-tailed, paired, across males and females at 30 weeks of age). No significant difference from wild-types was observed for heterozygotes (not shown).
Suppl. Figure 6 – Chr4\(^{Δ70kb/Δ70kb}\) mice have normal plasma lipid levels
Deletion mice and wild-type controls were placed on standard chow up to 6 weeks of age. Control groups were kept on standard chow for an additional 9 weeks, the test groups were placed on a high-fat, high-cholesterol diet for 9 weeks. No significant differences between wild-type and deletion mice was observed in base lipid levels or in the response to the noxious diet. NEFA = non-esterified free fatty acids.
Supl. Figure 7 – Chr4^{Δ70kb/Δ70kb} mice show no increase in atherogenic lesions

Deletion mice and isogenic controls in a 129Sv background were placed on a high-fat, high-cholesterol diet for ~20 weeks and assayed for atherogenic lesions. Extent of atherogenic lesions varied over a wide range and no significant increase in atherogenesis was observed in deletion mice (P=0.28, Mann-Whitney U test, non-directional).
Suppl. Figure 8 – Increased mortality on high-fat, high-cholesterol diet
Survival curve of wild-type and chr4Δ70kb/Δ70kb mice on high-fat, high-cholesterol diet fed ad libitum for 150 days. See Methods for details about diet and euthanasia criteria. P-value: Kaplan-Meier test for survival up to 150 days on high-fat diet.
Suppl. Figure 9 – Conserved noncoding sequences within the CAD risk interval
a) Six conserved noncoding sequences (blue boxes) located within the human CAD risk interval were identified based on 44-way vertebrate genome sequence alignments at the UCSC genome browser. In a transgenic mouse enhancer assay, no reproducible enhancer activity in day 11.5 mouse embryos was observed in the heart or any other structure. b) 44-way vertebrate conservation across the 58kb noncoding CAD risk interval. c) Conservation in selected vertebrate species.
## Supplementary Tables

| Primer name | Primer sequence | Product | Note |
|-------------|-----------------|---------|------|
| CHD 5' fvd  | ACTTCAGTAGGATCTGATTTTGAGGTACTTTTTAGACAGTTTAGAAATG   | 6223 bp | generate homology sequence for 5' knock-in |
| CHD 5' rev  | ACTTCAGTAGAATTCAGGAAGAACCAGGACTCCAATGCTTGC      |         |      |
| CHD 3' fvd  | ACTTCAGTAGGGGGCCGTCAGGGGCCCAGGCTTCAATAGGAAATG   | 6284 bp | generate homology sequence for 3' knock-in |
| CHD 3' rev  | ACTTCAGTAGAATTCAGGAAGAACCAGGACTCCAATGCTTGC      |         |      |
| T5          | GCAATAAACCCCTCAATAGGGAAC   | 6913 bp | PCR screening for 5' knock-in wild-type no band |
| CHD5'del    | same as above       | 6867 bp | PCR screening for 3' knock-in wild-type no band |
| T7          | GTAATAGACCTACTATAGGGCG  | 236 bp  | PCR screening deletion event wt: no band; deletion: 236 bp |
| T7          | GTAATAGACCTACTATAGGGCG  |         |      |
| CHDp1.fwd   | TGGAGTCCTCAAGAATCTTACATCTC | 361 bp | probe 1 for southern |
| CHDp1.rev   | CAGATCTATAGTAAATATATATGCT  |         |      |
| CHDp3.fwd   | AAGGTATGCTTAAATATGTCACTTGCGAG  | 180 bp | probe 3 for southern |
| CHDp3.rev   | CGAGTCCAAATTTTCTCAGGTGTTTTCTCAGA |         |      |
| T7          | GTAATAGACCTACTATAGGGCG  |         |      |
| CHD3'arm.fwd | GTAATAGACCTACTATAGGGCG  | 3136 bp | PCR confirming deletion event wt: no band; deletion: 3136 bp |
| CHD3'arm.rev| GTAATAGACCTACTATAGGGCG  |         |      |
| T7          | GTAATAGACCTACTATAGGGCG  |         |      |
| Suppl. Table 1 - Primer sequences used for generation of targeting constructs and genotyping |

| Primer name | Primer sequence | Product |
|-------------|-----------------|---------|
| Cdkn2b-F    | AGATCCCAACGCCCTGCAAC | 110 bp  |
| Cdkn2b-R    | GCGATGCTATGGCTC    |         |
| p16-INK4aF  | CCCAAGGCCCAGACT    | 79 bp   |
| p16-INK4aR  | GAGAAGATGTCTGCTGAA |         |
| Mtap E2f    | TGGTGAGACAGCTTGGATACOC | 164 bp |
| Mtap E4r    | AGGGCCAGATGTGTCTGCTGCC |         |
| Dmrt1 E1f   | TGCCCTCTAGACACCCCTGGAGGC | 146 bp |
| Dmrt1 E2r   | GGTAGATGTCTGCTGGAGACCTGCC |         |
| CDKN2B-rtSNP-F | GTTTCCCAGTCAGAGGTTGTAAMAGGCAACTCAAATGTAGGAAA | 1019 bp |
| CDKN2B-rtSNP-R | AGGAACAGCAATGACCCTAGATGCTCAGGCAACTCAAATGTAGGAAA |         |
| CDKN2B-Genomic-F | GTTTCCCAGTCAGAGGTTGTAAMAGGCAACTCAAATGTAGGAAA | 470 bp |
| CDKN2B-Genomic-R | AGGAACAGCAATGACCCTAGATGCTCAGGCAACTCAAATGTAGGAAA |         |

Suppl. Table 2 – Primer sequences used for quantitative RT-PCR and allele-specific expression profiling
### Genotype Age (weeks) Sex Site of tumor/mass Histopathological diagnosis

| Chr | Genotype | Age | Sex | Site of tumor/mass | Histopathological diagnosis |
|-----|----------|-----|-----|-------------------|----------------------------|
| chr4 | ∆70kb/∆70kb | 51 | M | skin | hemangiosarcoma |
| chr4 | ∆70kb/∆70kb | 54 | F | skin | anaplastic sarcoma with features of hemangiosarcoma (necropsy also revealed an irregular liver, histological evaluation confirmed myeloid leukemia) |
| chr4 | ∆70kb/∆70kb | 28 | F | leg | osteosarcoma |
| chr4 | ∆70kb/∆70kb | 28 | F | skin | sarcoma with moderate eosinophilic inflammation |
| chr4 | ∆70kb/∆70kb | 55 | M | lung | sarcoma with features of hemangiosarcoma |
| chr4 | ∆70kb/∆70kb | 61 | F | uterus | metritis |
| chr4 | ∆70kb/∆70kb | 15 | M | abdomen | teratoma |

**Suppl. Table 3 – Summary of histopathological analysis of tumors incidentally found in chr4∆70kb/∆70kb mice.**

| Candidate sequence | Coordinates (hg18) | Primer sequence | Fragment size |
|--------------------|-------------------|-----------------|---------------|
| hs1518             | chr9:22068885-22072105 | Forward: CTTTTGGGTTTCCCCATTGT<br>Reverse: AACAAGTGAACCTGGGGACCA | 3221bp |
| hs1519             | chr9:22077552-22079798 | Forward: GCCTAGTGGAAATTTCTATTGCTG<br>Reverse: GTCAATTGGCTCATTAAATACCAA | 2247bp |
| hs1520             | chr9:22092664-22094753 | Forward: CACCTACCTAAAAAGCCCAAAACA<br>Reverse: CAATGCCTGGCACCTAGAAT | 2090bp |
| hs1521             | chr9:22096994-22099800 | Forward: AGGATAGTCTGCATTTTCATGTT<br>Reverse: CCACCTTAGGTTCCACCAAA | 2807bp |
| hs1695             | chr9:22117677-22120448 | Forward: GAGACAGGAGGAGGTCCCCAAT<br>Reverse: AGAGGAATACACCTCTGGAA | 2772bp |
| hs1696             | chr9:22106987-22109574 | Forward: CAAATGGGAGCTGGGAGGAGTT<br>Reverse: GAAGGATGCTGATTTCTGICCA | 2588bp |

**Suppl. Table 4 – Primer sequences and coordinates of candidate enhancer sequences that were tested in transgenic mouse embryos.**
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