Genetics of blood lipids among ~300,000 multi-ethnic participants of the Million Veteran Program

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The Million Veteran Program (MVP) was established in 2011 as a national research initiative to determine how genetic variation influences the health of US military veterans. Here we genotyped 312,571 MVP participants using a custom biobank array and linked the genetic data to laboratory and clinical phenotypes extracted from electronic health records covering a median of 10.0 years of follow-up. Among 297,626 veterans with at least one blood lipid measurement, including 57,332 black and 24,743 Hispanic participants, we tested up to around 32 million variants for association with lipid levels and identified 118 novel genome-wide significant loci after meta-analysis with data from the Global Lipids Genetics Consortium (total n > 600,000). Through a focus on mutations predicted to result in a loss of gene function and a phenome-wide association study, we propose novel indications for pharmaceutical inhibitors targeting PCSK9 (abdominal aortic aneurysm), ANGPTL4 (type 2 diabetes) and PDE3B (triglycerides and coronary disease).

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cardiovascular risk. Lastly, we performed a PheWAS for a set of DNA sequence variants within genes that have already emerged as therapeutic targets for lipid modulation, leveraging the full catalog of International Classification of Disease, ninth edition (ICD-9) diagnosis codes in the Veteran Affairs EHR to better understand the potential consequences of pharmacological modulation of these genes or their products. We followed up significant findings from our PheWAS with multivariate Mendelian randomization analyses.

**Results**

**Demographics of genotyped MVP participants.** A total of 353,323 veterans had genetic data available in MVP, with clinical phenotypes recorded in the Veteran Affairs EHR for over 3,088,030 patient-years prior to enrollment (median of 10.0 years per participant) and 61,747,974 distinct clinical encounters (median of 99 per participant). We categorized veterans into three mutually exclusive ancestral groups for association analysis: (1) non-Hispanic white, (2) non-Hispanic black and (3) Hispanic participants. Admixture plots depicting the genetic background of the black and Hispanic groups are shown in Supplementary Figs. 1 and 2. Demographics and participant counts for a number of cardiometabolic traits for the 312,571 white, black and Hispanic MVP participants that passed our quality control are depicted in Table 1.

A subset of 297,626 participants passing quality control had at least one laboratory measurement of blood lipids in their EHR. These individuals collectively had a total of 15,456,328 laboratory entries for blood lipids, or a median of 12 measurements per lipid fraction per participant. To minimize potential confounding from the use of lipid-altering agents with variable adherence, we selected a participant’s maximum LDL-C, triglycerides and total cholesterol as well as his or her minimum HDL-C for genetic association analysis. Table 2 summarizes characteristics at enrollment and the distribution lipid levels for MVP participants included in our analysis. As expected, most of the participants were male and 28% were of non-European ancestry. While approximately 45% of participants had evidence of a statin prescription at the time of enrollment, only 8–9% participants had such evidence at the time of their maximum LDL-C or total cholesterol measurement used for our GWAS analysis.

**Lipid genetic association and conditional analyses.** We successfully imputed (INFO > 0.3, minor allele frequency (MAF) > 0.0003) 19.3, 31.4, and 30.4 million variants for white, black and Hispanic veterans, respectively, using the 1000 Genomes Project reference panel (Table 2). Black and Hispanic participants had substantially more variants available for analysis, reflecting the known greater genetic diversity within these populations. We also identified 6,657 pLOF variants in 4,294 genes across the three ethnicities (Supplementary Fig. 3).

We compared the Z scores and effect estimates from the published literature with those observed in MVP for 444 previously reported exome-wide significant variants for lipids. We found a strong correlation of genetic associations across all four traits, validating the lipid data obtained from the EHR (Supplementary Figs. 4, 5).

We performed association testing separately among individuals of each of three ancestries (white, black, and Hispanic) in our initial discovery analysis and then meta-analyzed results across ancestry groups using an inverse-variance-weighted fixed-effects method (Fig. 1a, Supplementary Fig. 6). Following trans-ethnic meta-analysis in the discovery phase of our study (stage 1), a total of 46,526 variants at 188 of the 268 known loci for lipids met the genome-wide significance threshold (P < 5 x 10^-8) (Supplementary Tables 1–4). We performed pairwise comparisons of the allele frequencies and effect estimates between white and black participants as well as between white and Hispanic participants for 354 of the 444 previously established independent variants for lipids that were well-imputed in all three ancestral groups in MVP (Fig. 2). We observed a much stronger correlation for effect allele frequencies between white and Hispanic participants (Pearson’s correlation coefficient R = 0.96) than between white and black participants (R = 0.72), likely reflecting the greater European admixture in the MVP Hispanic participants. The effect estimates among the three ethnicities varied by lipid trait (Fig. 2, Supplementary Fig. 7).

We sought replication for variants within MVP with suggestive associations (P < 1 x 10^-4) in either stages 2a or 2b (Fig. 1b). We first attempted replication of these variants using summary statistics from the 2017 GLGC exome array meta-analysis (stage 2a). If association statistics for promising DNA sequence variants from stage 1 were not available for replication in the 2017 exome array-focused study, we sought replication of these variants in publicly available datasets (stage 2b).

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**Fig. 1** | GWAS study design. **a.** DNA sequence variants across three separate ancestry groups in the MVP were meta-analyzed using an inverse-variance-weighted fixed-effects method in the discovery phase (stage 1). Variants with suggestive association were then brought forward for independent replication. **b.** DNA sequence variants with suggestive associations (two-sided linear regression P < 1 x 10^-4) in the discovery analysis (stage 1) were brought forward for independent replication and tested using summary statistics from the 2017 exome-array-focused GLGC meta-analysis (stage 2a). Only variants with suggestive associations in stage 1 that were not present in the GLGC 2017 exome-array study (stage 2a) were alternatively replicated in the 2013 GLGC joint meta-analysis (stage 2b).
available summary statistics from the 2013 GLGC ‘joint meta-analysis’ (stage 2b). We did not attempt replication of any variant in both studies given the substantial overlap of participants in these two studies. A total of 170,925 variants demonstrated suggestive association (P < 10^{-8}) in the MVP discovery analysis. Among these variants, 39,663 were also available for in silico replication in either stage 2a (GLGC 2017) or stage 2b (GLGC 2013). We defined significant novel associations as those that were at least nominally significant in replication (P < 0.05) with consistent direction of effect and had an overall P < 5 × 10^{-8} (genome-wide significance) in the discovery and replication cohorts combined. Following replication, 118 novel loci (from 141 lead variants) exceeded genome-wide significance (P < 5 × 10^{-8}, Supplementary Tables 5–8). MAFs of lead variants ranged from 0.08% to 49.9%, with effect sizes ranging from 0.01 to 0.243 standard deviations (σ). For example, carriers of a rare missense mutation in the gene encoding nixin 8 (SNX8 Ile414Thr, rs144787122, NC_000007.13: 2296552A>G) MAF = 0.35% in MVP demonstrated a 0.10σ (3.8 mg dl^{-1}) higher plasma LDL-C after testing in 587,481 individuals.

More than one variant may independently affect plasma lipid levels at any given genetic locus. We performed a conditional analysis using combined summary statistics from MVP and publicly available data from GLGC for each lipid trait (Supplementary Fig. 8) and identified a total of 826 independently associated lipid variants across 118 novel and 268 previously identified loci (Supplementary Table 9).

Variance explained obtained from multiple lipid measures. The previously mapped 444 lipid variants explain about 7.5–10.5% of the phenotypic variance in lipid levels in the MVP population. The 118 novel loci in our study explain an additional 0.38–0.74% in phenotypic variance, and the 826 independent variants identified in our conditional analysis increase the overall explained phenotypic variance to 8.8–12.3% (Supplementary Table 10).

We subsequently explored the impact of multiple lipid measurements in an analysis restricted to 171,314 European MVP participants with ≥5 lipid measurements in their EHR. We constructed a weighted genetic risk score (GRS) of 223 variants across 268 of the previously mapped loci with effect estimates available in the 2017 GLGC exome array analysis summary statistics (Supplementary Table 11). Generally across the four lipid traits, the GRS explained a larger proportion of the phenotypic variance with an increasing number of lipid measurements included in the analysis (Supplementary Table 12). In addition, when the maximum/minimum lipid values were used as in our discovery GWAS, the GRS explained more total variance than when using up to five lipid measurements for the LDL-C, triglycerides and total cholesterol phenotypes.

Transcriptome-wide association study. We next performed a TWAS using: (1) pre-computed weights from expression array data measured in peripheral blood from 1,245 unrelated control

### Table 1 | Demographic and clinical characteristics of black, white and Hispanic individuals passing quality control in the MVP

| Basic demographics | Genotyped veterans |
|---------------------|---------------------|
| n                   | 312,571             |
| Age at enrollment in years (mean ± σ) | 62.4 ± 13.5 |
| Male, n (%)         | 287,441 (92.0%)     |
| Body mass index in kg m^{-2} (mean ± σ) | 30.3 ± 6.0 |
| Current smoker, n (%) | 59,385 (19.0%)     |
| Former smoker, n (%) | 159,459 (51.0%)    |
| n with ≥1 measurement of plasma lipids (%) | 297,626 (95.2%) |
| Number of lipid measurements (median per lipid fraction) | 15,456,328 (12) |

**Race/ethnicity**
- Black, n (%) | 59,007 (18.9%)
- White, n (%) | 227,817 (72.8%)
- Hispanic, n (%) | 25,747 (8.1%)

**Cardiometabolic disease at enrollment**
- Coronary artery disease, n (%) | 67,912 (21.7%)
- Type 2 diabetes, n (%) | 92,079 (29.5%)
- Peripheral artery disease, n (%) | 21,418 (6.9%)
- Abdominal aortic aneurysm, n (%) | 5,618 (1.8%)
- Deep venous thrombosis or pulmonary embolism, n (%) | 7,009 (2.2%)

* Diseases are defined by ICD-9 diagnosis codes.

### Table 2 | Demographic and clinical characteristics for 297,626 veterans in the Million Veteran Program lipids analysis

|                         | White          | Black          | Hispanic        |
|-------------------------|----------------|----------------|-----------------|
| Veterans, n (%)         | 215,551 (72.4%) | 57,332 (19.3%) | 24,743 (8.3%)   |
| Age at enrollment in years (mean ± σ) | 64.2 ± 13    | 57.7 ± 11.8    | 56.3 ± 15.0     |
| Male, n (%)             | 200,900 (93.2%) | 50,059 (87.3%) | 22,601 (91.3%)  |
| Body mass index in kg m^{-2} (mean ± σ) | 30.1 ± 5.9   | 30.4 ± 6.3     | 30.7 ± 5.8      |
| Statin therapy prescription at enrollment, n (%) | 100,024 (46.4%) | 23,302 (40.6%) | 9,646 (39.0%)   |
| Statin therapy prescription at time of maximum LDL-C blood draw, n (%) | 18,818 (8.7%) | 5,024 (8.8%) | 2,262 (9.1%)    |
| Statin therapy prescription at time of maximum total cholesterol blood draw, n (%) | 18,433 (8.6%) | 5,027 (8.8%) | 2,162 (8.7%)    |
| Minimum HDL-C in mg dl^{-1} (mean ± σ) | 36.2 ± 11.4   | 38.9 ± 12.8    | 36.4 ± 11.0     |
| Maximum LDL-C in mg dl^{-1} (mean ± σ) | 139 ± 38.4     | 142.2 ± 40.7   | 141.3 ± 38.1    |
| Median maximum triglycerides IQR in mg dl^{-1} | 211 ± 174      | 179 ± 149      | 221 ± 184       |
| Maximum total cholesterol in mg dl^{-1} (mean ± σ) | 218.6 ± 46.7  | 220.8 ± 47.2   | 221.9 ± 48.0    |
| Variants included in analysis | 19,342,852   | 31,448,849     | 30,455,745      |

IQR, interquartile range.
Articles

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RNA-sequencing data measured in adipose tissue from 563 control individuals from The Netherlands Twin Registry24, RNA-sequencing data measured in post-mortem liver (97 individuals) and tibial artery (285 individuals) tissue from the Genotype-Tissue Expression project25 (GTEx V6); and (2) combined MVP and GLGC summary statistics for each of the four lipid traits. In brief, this approach integrates information from expression reference panels (variant–expression correlation), GWAS summary statistics (variant–trait correlation), and linkage disequilibrium (LD) reference panels (variant–variant correlation) to assess the association between the cis-genetic component of expression and phenotype23. The results yield candidate causal genes from the GWAS results under the assumption that the causal mechanism of the tested genes involves changes in cis-expression.

Our TWAS identified a total of 655 genome-wide significant ($P < 5 \times 10^{-8}$) gene–lipid associations (summed across expression reference panels) in 333 distinct genes, including 194 that were significant in more than one tissue or lipid trait (Supplementary Tables 13–16, Supplementary Figs. 9–10). The 333 distinct genes fell within 122 genomic loci, 117 of which were within a lipid GWAS region ($\pm$1 Mb around a mapped sentinel GWAS variant) identified in either a prior analysis or in the current study. However, five genes identified by TWAS fell outside of previously mapped GWAS regions, representing potentially novel genomic loci for lipids (Supplementary Table 17). Previous work has suggested that future lipid GWAS with larger sample sizes will likely confirm the novel lipid loci identified by our TWAS26. Results from additional competitive gene-set pathway and tissue expression analyses are available in the Supplementary Note.

Fig. 2 | Comparison of 354 independent lipid-associated variants across ethnicities. a, b. Allele frequencies of lipid-associated variants observed in white individuals ($n=215,196$; x axes) compared to black (a; $n=57,280$; Pearson’s $R=0.72$), or Hispanic (b; $n=24,742$; Pearson’s $R=0.96$) individuals. c, d. Linear regression effect estimates for LDL-C associations in white individuals ($n=215,196$; x axes) compared to black (c; $n=57,280$; $\beta=1.07$) or Hispanic (d; $n=24,742$; $\beta=1.06$) individuals.

Non-European low-frequency missense variant associations. We next focused on ancestry specific low-frequency (MAF < 5%) missense variants, as these variants have been suggested to have a higher likelihood of causality27,28. We identified several novel low-frequency missense variants associated with one or more lipid levels at genome-wide significance that were specific to black or Hispanic participants. We found a total of five variants associated with LDL-C and/or total cholesterol among black individuals (Supplementary Table 18) and two associated with HDL-C and/or total cholesterol among Hispanic individuals (Supplementary Table 19) in PCSK9, LDLR, APOB and ABCA1. All ten associations were directionally consistent with the 2017 GLGC exome chip meta-analysis with nine reaching nominal significance ($P < 0.05$) among 17,009 black and 5,084 Hispanic individuals included in the GLGC study. In addition, the seven variants that we identified were either monomorphic or had a MAF < 0.0005 in the approximately 215,000 white veterans in MVP. Of note, we observed the low-frequency 443Thr allele in

| Effect allele frequency (white participants) | Effect allele frequency (black participants) | Effect allele frequency (Hispanic participants) |
|---------------------------------------------|---------------------------------------------|-----------------------------------------------|
| 0.00 0.25 0.50 0.75 1.00 | 0.00 0.25 0.50 0.75 1.00 | 0.00 0.25 0.50 0.75 1.00 |

| Effect estimate for white participants (a) | Effect estimate for black participants (b) | Effect estimate for Hispanic participants (c) |
|---------------------------------------------|---------------------------------------------|-----------------------------------------------|
| -0.25 0.00 0.25 0.50 0.75 | -0.25 0.00 0.25 0.50 0.75 | -0.25 0.00 0.25 0.50 0.75 |
PCSK9 within Hispanic individuals to be eightfold more common in black individuals (MAF = 0.011 in Hispanic versus 0.092 in black individuals). We also found that this variant was associated with total cholesterol in black individuals at genome-wide significance.

**Predicted loss of gene function lipid associations.** We focused next on the subset of genotyped or imputed pLOF variants (variants that were annotated as premature stop (nonsense), canonical splice sites (splice-donor or splice-acceptor) or insertion/deletion variants that shifted frame (frameshift) by the Variant Effect Predictor software). A total of 15 distinct pLOF variants demonstrated genome-wide significant lipid associations across individuals of all three ethnic groups (Supplementary Table 20). We replicated known pLOF associations at PCSK9, APOC3, ANGPTL8, LPL, CD36 and HBB, and we observed genome-wide significant associations of comparable magnitude of effect in each of the three ethnic groups for two pLOF variants: a base substitution in APOC3 55+1G>A and a mutation in LPL encoding Ser747Ter.

We identified one novel pLOF association. Among white MVP participants, carriers of a rare stop-gain mutation in PDE3B (encoding Arg783Ter; carrier frequency of 1 in 625), exhibited 4.72 mg dl⁻¹ higher blood HDL-C levels (P = 2.8 × 10⁻¹⁶) and 43.3 mg dl⁻¹ lower blood triglyceride levels (P = 7.5 × 10⁻¹⁶). We found this signal to be independent of a previously reported genome-wide significant association in the region involving a common polymorphism, rs10373781 (Arg783Ter; conditional analysis P = 6.3 × 10⁻¹⁶ for HDL-C and P = 8.91 × 10⁻⁸ for triglycerides). We also identified one individual who was homozygous for Arg783Ter. This PDE3B ‘human knockout’ was in his sixth decade of life and had HDL-C and triglycerides levels of 73 and 56 mg dl⁻¹, respectively. He was not on lipid-lowering medication and was free of coronary artery disease (CAD). We replicated the triglyceride and HDL-C associations for this pLOF variant in an independent sample of approximately 45,000 participants of the DiscovEHR study (Fig. 3a,b).

**Loss of PDE3B function and risk of coronary artery disease.** Hypothesizing that mutations that were damaging or causing loss of function in PDE3B could protect against the development of CAD based on their association with lifelong lower levels of triglycerides in blood, we conducted a case–control study of CAD involving five cohorts: MVP, UK Biobank, Myocardial Infarction Genetics Consortium (MIGen), Penn Medicine Biobank (PMBB) and DiscovEHR. For three studies that underwent exome sequencing (MIGen, PMBB and DiscovEHR), we combined pLOF variants with missense variants that were predicted to be damaging or possibly damaging by each of five computer prediction algorithms (LRT score, MutationTaster, PolyPhen-2, HumDiv, PolyPhen-2) and we observed genome-wide significant associations of comparable magnitude of effect in each of the three ethnic groups for two pLOF variants: a base substitution in APOC3 55+1G>A and a mutation in LPL encoding Ser747Ter. We leveraged a median of 65 unique ICD-9 diagnosis codes per participant prior to enrollment in MVP to explore the spectrum of phenotypic consequences of genetic variation within genes targeted by lipid-lowering medicines. We selected five lipid-associated genes currently being targeted by pharmaceutical agents and identified functional variants in these genes: two nonsense variants (ANGPTLA Glu40Lys, APOA5 Ser19Thr, PCSK9 Arg46Leu). We considered phenotypes to be significantly associated with a variant if they met a Bonferroni corrected P < 4.98 × 10⁻⁵ (0.05/1,004
traits), a conservative threshold given the correlation structure present among PheWAS phenotypes.

Data from a total of 176,913 white veterans were available for analysis after quality control. Among these individuals, we identified 33 statistically significant phenotypic associations across the five variants, all of which are correlated with lipids (Supplementary Table 22). We replicated known associations with CAD for \( \text{LPL} \), \( \text{ANGPTL4} \) and \( \text{PCSK9} \). Notably, carriers of triglyceride-lowering and/or HDL-C-raising mutations in \( \text{ANGPTL4} \) (Glu40Lys; 7,013 carriers) were also found to have a reduced risk of type 2 diabetes (Fig. 4). We replicated the type 2 diabetes association for the \( \text{ANGPTL4} \) E40K variant in an independent sample of approximately 452,000 participants in the recently published trans-ethnic diabetes GWAS (odds ratio, 0.89; 95% confidence interval, 0.86–0.93; \( P = 9.24 \times 10^{-10} \); Supplementary Fig. 11). In addition, carriers of LDL-C-lowering mutations in \( \text{PCSK9} \) (Arg46Leu; 5,537 carriers) also demonstrated a reduced risk of abdominal aortic aneurysm (AAA, Fig. 5).
Lipids and AAA Mendelian randomization analysis. To further explore the causal relationship of lipids on AAA development, we performed a multivariate Mendelian randomization analysis using a weighted GRS of 223 lipid-associated variants and summary data from a GWAS of 5,002 AAA cases and 139,968 controls in MVP. Consistent with our PheWAS results, a 1σ genetically elevated LDL-C was associated with an increased risk of AAA (odds ratio, 1.47; 95% confidence interval, 1.28–1.68; \(P = 4.4 \times 10^{-13}\)); a 1σ genetically elevated HDL-C level was associated with a decreased risk of AAA (odds ratio, 0.79; 95% confidence interval, 0.68–0.91; \(P = 0.001\)); and a 1σ genetically elevated triglyceride level was associated with an increased risk of AAA (odds ratio, 1.40; 95% confidence interval, 1.18–1.66; \(P = 8.5 \times 10^{-5}\); Fig. 6). An MR-Egger analysis indicated no pleiotropic bias of our lipid genetic instruments (MR-Egger intercept \(P > 0.05\) for all three lipid fractions (Supplementary Table 23)).

### Discussion
We leveraged clinical and genetic data from the MVP to investigate the inherited basis of blood lipids in nearly 300,000 US veterans. Our investigation resulted in several key findings. First, we robustly

| Disease                          | Cases   | Controls | Odds ratio | 95% CI       | \(P\)     |
|----------------------------------|---------|----------|------------|--------------|-----------|
| Cardiovascular                   |         |          |            |              |           |
| Coronary disease                 | 41,449  | 111,656  | 0.83       | 0.78–0.88    | \(1.38 \times 10^{-13}\) |
| Abdominal aortic aneurysm        | 5,001   | 139,925  | 0.72       | 0.62–0.83    | \(2.05 \times 10^{-6}\) |
| Peripheral vascular disease      | 13,023  | 139,925  | 0.85       | 0.76–0.95    | 0.004     |
| Dermatologic                     |         |          |            |              |           |
| Psoriasis                        | 5,783   | 129,186  | 0.91       | 0.77–1.06    | 0.218     |
| Actinic keratosis                | 31,866  | 113,003  | 1.01       | 0.94–1.08    | 0.811     |
| Atopic dermatitis                | 20,599  | 120,106  | 1.11       | 1.02–1.21    | 0.011     |
| Digestive                        |         |          |            |              |           |
| Ulcerative colitis               | 1,601   | 114,158  | 1.07       | 0.81–1.41    | 0.621     |
| Diverticulosis                   | 17,837  | 114,158  | 1.08       | 0.99–1.18    | 0.085     |
| Cholelithiasis                   | 5,372   | 164,226  | 1.09       | 0.94–1.27    | 0.249     |
| Endocrine/Metabolic              |         |          |            |              |           |
| Graves’ disease                  | 561     | 147,098  | 0.96       | 0.79–1.56    | 0.876     |
| Type 2 diabetes                  | 57,203  | 95,789   | 1.03       | 0.97–1.09    | 0.362     |
| Gout                             | 13,868  | 158,066  | 1.01       | 0.92–1.12    | 0.826     |
| Genitourinary                    |         |          |            |              |           |
| Chronic renal failure            | 14,978  | 139,056  | 0.99       | 0.90–1.10    | 0.911     |
| Pyelonephritis                   | 619     | 135,058  | 0.87       | 0.54–1.42    | 0.584     |
| Urinary calculus                 | 12,101  | 158,922  | 1.02       | 0.92–1.14    | 0.856     |
| Hematopoietic                    |         |          |            |              |           |
| Anemia of chronic disease        | 2,445   | 134,039  | 0.98       | 0.77–1.23    | 0.831     |
| Neutropenia                      | 1,136   | 153,819  | 0.81       | 0.56–1.17    | 0.262     |
| Lymphadenitis                    | 2,790   | 153,819  | 0.90       | 0.72–1.13    | 0.36      |
| Mental disorders                 |         |          |            |              |           |
| Schizophrenia                    | 4,013   | 81,587   | 0.95       | 0.79–1.14    | 0.588     |
| Bipolar                          | 10,923  | 81,587   | 1.10       | 0.98–1.22    | 0.108     |
| Depression                       | 29,881  | 81,587   | 1.03       | 0.95–1.11    | 0.461     |
| Musculoskeletal                  |         |          |            |              |           |
| Spinal stenosis                  | 11,473  | 120,150  | 1.05       | 0.95–1.17    | 0.346     |
| Osteoarthritis                   | 19,501  | 90,523   | 1.03       | 0.94–1.13    | 0.479     |
| Osteoporosis                     | 6,294   | 157,501  | 1.02       | 0.89–1.18    | 0.758     |
| Neoplasms                        |         |          |            |              |           |
| Colon cancer                     | 3,082   | 112,365  | 1.20       | 0.99–1.45    | 0.064     |
| Cancer of bronchus               | 2,357   | 173,116  | 0.99       | 0.78–1.25    | 0.914     |
| Squamous cell carcinoma          | 2,095   | 132,018  | 1.22       | 0.98–1.53    | 0.078     |
| Neurological                     |         |          |            |              |           |
| Sleep apnea                      | 41,784  | 102,716  | 1.07       | 1.00–1.14    | 0.045     |
| Migraine                         | 9,330   | 162,328  | 1.10       | 0.98–1.23    | 0.106     |
| Epilepsy                         | 1,486   | 145,349  | 0.89       | 0.66–1.22    | 0.475     |
| Respiratory                      |         |          |            |              |           |
| Pneumonia                        | 1,429   | 144,216  | 0.87       | 0.64–1.20    | 0.407     |
| Asthma                           | 12,415  | 146,753  | 1.11       | 1.00–1.22    | 0.052     |
| Chronic airway obstruction       | 8,894   | 146,753  | 1.09       | 0.97–1.23    | 0.139     |

Fig. 5 | PCSK9 Arg46Leu carrier disease associations. Forest plot for a representative 33 of the 1,004 disorders tested in the PCSK9 Arg46Leu PheWAS.
confirmed 188 previously identified loci while concurrently uncovering an additional 118 novel genome-wide significant loci. Next, we identified a total of 826 independent lipid-associated variants increasing the phenotypic variance explained by nearly 2%. We performed a TWAS in four tissues identifying five additional novel lipid loci at a genome-wide level of significance and performed a pathway analysis highlighting lipid transport mechanisms in our GWAS results. We identified ancestry-specific effects of rare coding variation on lipids among white, black, and Hispanic participants, and observed 15 pLOF mutations associated with lipids at a genome-wide level of significance, including a protein-truncating variant in \textit{PDE3B} that lowers triglycerides, raises HDL-C and protects against CAD. Finally, we examined the full spectrum of phenotypic consequences for mutations in lipid genes emerging as therapeutic targets, identifying protective effects of functional mutations in PCSK9 for abdominal aortic aneurysm and in \textit{ANGPTL4} for type 2 diabetes.

We obtained four main insights through our findings. First, we confirm the enormous potential of a large-scale multi-ethnic biobank built within an integrated health-care system in the discovery of the genetic basis of human traits. Specifically, we leveraged the Veteran Affairs’ mature nationwide EHR to efficiently extract existing repeated laboratory measures of lipids collected during the course of clinical care in nearly 300,000 veterans over a median of 10 years for GWAS analysis. Our results highlight the expected increase in variance explained by known loci when repeated lipid measurements are considered but also demonstrate the efficiency of examining the single most extreme lipid value least likely influenced by the use of lipid-altering medications. Subsequent meta-analysis (combined \(n > 600,000\)) with existing datasets increased the number of known independent genetic lipid loci to nearly 400, including several lipid pathways with links to human disease. For example, common variants near genes such as \textit{COL4A2} and \textit{ITGA1} identified for LDL-C and/or total cholesterol suggest links to extracellular matrix and cell adhesion biology, two pathways recently implicated by GWAS of CAD\(^{46,47}\).

We also demonstrated that carriers of a rare missense mutation in the gene encoding perilipin 1 (\textit{PLIN1} Leu90Pro) possess a markedly higher plasma HDL-C (0.243 vs 0.68–0.91 0.001) and also has well-documented, substantial effects on triglycerides and HDL-C levels—likely through antagonism of \textit{PDE3B}. We demonstrate that a \textit{PDE3B} pLOF variant recapitulates the known lipid effects of cilostazol and extend these findings to show that damaging \textit{PDE3B} mutations are also associated with reduced risk of CAD. Randomized control trials to date have demonstrated the efficacy of cilostazol in intermittent claudication\(^{70}\) and prevention of restenosis following percutaneous coronary intervention\(^{71}\). The drug is also currently used off-label for the prevention of stroke recurrence through a presumed anti-platelet effect\(^{49}\). We note that mice genetically deficient in \textit{Pde3b} display reduced atherosclerosis\(^{72}\) as well as decreased infarct size and improved cardiac function following experimental coronary artery ligation\(^{73}\). In light of our findings, use of cilostazol, or one of its derivatives, for the primary or secondary prevention of CAD deserves further consideration.

Our final insight highlights the potential benefit of PhewAS across a large-scale EHR-based biobank to predict both potentially adverse and beneficial consequences of artificially inhibiting gene function. Here, we provide evidence that pharmacologic PCSK9 inhibition may reduce abdominal aortic aneurysm risk in addition to its known effects on atherosclerotic cardiovascular disease\(^{41}\).

This finding is further supported by: our Mendelian randomization results; a recently published analysis using an independent AAA dataset\(^{52}\); and a recent report demonstrating that a \textit{Pcsk9} gain-of-function mutation augments AAA development in a mouse model\(^{53}\). However, we also recognize the possibility that these results may be a consequence of pleiotropic effects induced by a high phenotypic correlation between AAA and the presence of advanced atherosclerotic disease. Thus, additional studies are necessary before definitive conclusions can be made on causality. Similarly, we expand on the potential indications for \textit{ANGPTL4} inhibition to include type 2 diabetes. Future PhewAS efforts may identify associations that facilitate prioritization of drugs currently in development, repurposing of therapies already in clinical use, or prediction of adverse or off-target effects prior to investigation through expensive and time-consuming clinical trials.

Several limitations deserve to be mentioned. First, our MVP lipid phenotype definitions are based entirely on EHR data with a high prevalence of use of lipid-lowering therapy at enrollment. We used maximum or minimum values to capture untreated lipid levels, but the possibility of misclassification of lipid levels remains for participants entering the Veteran Affairs healthcare system on therapy. Such misclassification, however, would be expected to generally reduce our power to detect genetic associations. Second, participants in MVP are overwhelmingly male. Although almost 25,000 women were included in our discovery analysis, we did not attempt to detect genetic associations specific to females or of the largest single-cohort GWAS to date for these ethnic groups for any trait. Among these individuals, we compared the effect estimates and allele frequencies of lipid-associated variants across ancestral groups and identified seven novel low-frequency coding variants associated with lipids only in non-European populations. Conversely, we also confirmed a shared genetic architecture across all three ethnic groups for pLOF variation at the \textit{LPL} and \textit{APOC3} loci. Previous work identifying low-frequency missense and pLOF variation in lipid genes have led to the development of the next generation of pharmaceutical agents for cardiovascular disease\(^{41,44,45}\).

Expansion of these efforts to larger sample sizes and additional ancestries may help to explain differences in blood lipid levels and risk of atherosclerosis among select populations.

Our third insight centers around our findings for the deleterious exonic variants within \textit{PDE3B}. These findings lend human genetic support to \textit{PDE3B} inhibition as a therapeutic strategy for atherosclerosis. Cilostazol, an inhibitor of both the 3A and 3B isoforms of the phosphodiesterase enzyme, is known to have anti-platelet\(^{43}\), vasodilatory\(^{44}\) and inotropic\(^{45}\) effects through inhibition of \textit{PDE3A}, and also has well-documented, substantial effects on triglycerides and HDL-C levels—likely through antagonism of \textit{PDE3B}. We demonstrate that a \textit{PDE3B} pLOF variant recapitulates the known lipid effects of cilostazol and extend these findings to show that damaging \textit{PDE3B} mutations are also associated with reduced risk of CAD. Randomized control trials to date have demonstrated the efficacy of cilostazol in intermittent claudication\(^{70}\) and prevention of restenosis following percutaneous coronary intervention\(^{71}\). The drug is also currently used off-label for the prevention of stroke recurrence through a presumed anti-platelet effect\(^{49}\). We note that mice genetically deficient in \textit{Pde3b} display reduced atherosclerosis\(^{72}\) as well as decreased infarct size and improved cardiac function following experimental coronary artery ligation\(^{73}\). In light of our findings, use of cilostazol, or one of its derivatives, for the primary or secondary prevention of CAD deserves further consideration.

**Fig. 6 | Lipid associations with abdominal aortic aneurysm.** Logistic regression association results of the 223 variant lipid genetic risk score with abdominal aortic aneurysm in a multivariable Mendelian randomization analysis. Odds ratios are displayed per 1σ genetically increased lipid fraction. Two-sided \(P\) values are displayed.
heterogeneity of effects between sexes due to suspected limited power. Third, our TWAS identifies candidate causal genes under the assumption that the causal mechanism of the tested genes involves changes in cis-expression. However, we are unable to discriminate between instances of pleiotropy (when a given variant may alter gene expression and affect lipid levels independently) with TWAS alone and further functional analysis may be necessary. Fourth, our analysis demonstrating a lack of association between HDL-C raising alleles and CAD risk may be underpowered given the small number of examined alleles, although this finding has been demonstrated consistently in previous studies\cite{ASS1, ASS2, ASS3, ASS4}. Lastly, power to detect associations for less common diseases in our PheWAS may be also be limited despite the overall number of participants included in the analysis.

In conclusion, we identified more than 100 new genetic signals for blood lipid levels utilizing a biobank that exploits existing EHRs of US veterans. We demonstrate the potential of this approach in the discovery of novel genetic associations and the development of novel therapeutic agents.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, statements of data availability and associated accession codes are available at https://doi.org/10.1038/s41588-018-0222-9.

Received: 6 February 2018; Accepted: 3 August 2018; Published online: 1 October 2018

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Acknowledgements

Data on patients with coronary artery disease and myocardial infarctions have been contributed by the CARDIoGRAMplusC4D investigators and the Myocardial Infarction Genetics and CARDIoGRAM Exome investigators. Both datasets were obtained online (see URLs). This research is based on data from the MVE Office of Research and Development, Veterans Health Administration, and was supported by the Department of Veterans Affairs Cooperative Studies Program award G002. This research was also supported by three additional Department of Veterans Affairs awards (i01BX003340, i01BX003362, and i01CX001025) and the NIH (T32 HL007734, K01HL125751, R101HL127564). The content of this manuscript does not represent the views of the Department of Veterans Affairs or the United States Government.

Author contributions

Concept and design: D.K., T.L.A., S.M.D., K.K., K.-M.C., P.S.T., S.K., D.J.R., P.W.E.W., J.C. and J.M.G. Acquisition, analysis or interpretation of data: D.K., S.M.D., Y.V.S., K.C., T.M.T.; J.Ho., D.R.G., S.L.D., J.L., G.M.P., M.C., A.M.S., J.H.A., H.T., J.S.Y., Y.L.H., D.J.L., C.A.E., A.H.L., J.A.L., R.C., P.N., D.S., M.V., R.S.P., E.D.A., R.M.N., A.v.K., J.D., K.-M.C., G.A., C.W., F.E.D., J.E.H. and D.I.C. Drafting of the manuscript: D.K. and T.L.A. Critical revision of the manuscript for important intellectual content: S.M.D., Y.V.S., K.C., K.-M.C., P.S.T., S.K., D.J.R. and P.W.W. Administrative, technical or material support: D.K., Y.V.S., K.C., J.Ho., D.R.G., S.L.D., J.A.L., Y.H., J.C., J.M.G., C.W., J.E.H., and P.W.W.

Competing interests

S.K. reports grant support from Regeneron and Bayer, grant support and personal fees from Aegerion, personal fees from Regeneron Genetics Center, Merek, Celera, Novartis, Bristol-Myers Squibb, Sanofi, AstraZeneca, Ablynym, Eli Lilly and Leerink Partners, personal fees and other support from Catabasis, and other support from San Therapeutics outside the submitted work. He is also the chair of the scientific advisory board at Genomics Plc. T.M.T., A.H.L., A.B., F.E.D. and D.J.C. are employees of Regeneron Pharmaceuticals. G.A. has received consulting income from Regeneron Genetics Center, 23andMe and Helix. S.L.D. has received research grant support from the following for-profit companies through the University of Utah or the Western Institute for Biomedical Research (VA Salt Lake City’s affiliated non-profit): AbbVie Inc., Anolix LLC, Astellas Pharma Inc., AstraZeneca Pharmaceuticals LP, Boehringer Ingelheim International GmbH, Celgene Corporation, Eli Lilly and Company, Genentech Inc., Genomic Health Inc., Gilead Sciences Inc., GlaxoSmithKline PLC, Innocrin Pharmaceuticals Inc., Janssen Pharmaceuticals Inc., Kantar Health, Myriad Genetic Laboratories Inc., Novartis International AG and PAREXEL International Corporation.

Additional information

Supplementary information is available for this paper at https://doi.org/10.1038/s41588-018-0222-9.
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Methods
The design of the MVP has been previously described. In brief, individuals aged 19–104 years have been recruited from more than 50 Veterans Affairs Medical Centers nationwide since 2011. Each veteran’s EHR data are being integrated into the MVP biorepository, including incident ICD-9 diagnosis codes, Current Procedural Terminology procedure codes, clinical laboratory measurements and reports of diagnostic imaging modalities. The MVP received ethical and study protocol approval from the Veteran Affairs Central Institutional Review Board in accordance with the principles outlined in the Declaration of Helsinki. Informed consent was obtained from all participants of the MVP study.

Genetic data. DNA extracted from whole blood was genotyped using a customized Affymetrix Axiom biobank array, the MVP 1.0 Genotyping Array. With 723,305 total DNA sequence variants, the array is enriched for both common and rare variants of clinical importance in different ethnic backgrounds. Veterans were assigned one of three exclusive ethnicities: (1) non-Hispanic white veterans (European ancestry), (2) non-Hispanic black veterans (African ancestry) and (3) Hispanic veterans. Further details on the methods used to assign ancestry and perform sample quality control are described in the Supplementary Note.

Variant quality control. Prior to imputation, variants that were poorly called (genotype missingness >5%) or that deviated from their expected allele frequency based on reference data from the 1000 Genomes Project were excluded. After pre-phasing using Eagle v.2, genotypes from the 1000 Genomes Project phase 3, v.3 reference panel were imputed into MVP participants via Minimac3 software. Ethnicity-specific principal component analysis was performed using the EIGENSOFT software.

Following imputation, variant-level quality control was performed using the EasyQC R package (see URLs), and the following exclusion metrics were used: ancestry-specific Hardy–Weinberg equilibrium $P < 1 × 10^{-8}$, posterior call probability $<0.9$, imputation quality INFO $<0.3$, MAF $<0.0003$, call rate $<97.5\%$ for common variants (MAF > 1%) and call rate $<99\%$ for rare variants (MAF < 1%). Variants were also excluded if they deviated by >10% from their expected allele frequency based on reference data from the 1000 Genomes Project.

EHR-based lipid phenotypes. EHR clinical laboratory data were available for MVP participants from as early as 2003. We extracted the maximum LDL-C, triglycerides and total cholesterol values and minimum HDL-C values for each participant for analysis. These extreme values were selected to approximate plasma lipid concentrations in the absence of lipid-lowering therapy as described previously. For each phenotype (LDL-C, natural log-transformed triglycerides, HDL-C and total cholesterol), residuals were obtained after regressing on age, age$^2$, sex and 10 principal components of ancestry. Residuals were subsequently inverse-normal transformed for association analysis. Statin therapy prescription at enrollment was defined as the presence of a statin prescription in the EHR within 90 days of enrollment or after enrollment in the MVP. Statin therapy prescription at the maximum lipid measurement was defined as the presence of a statin prescription in the EHR within 90 days prior to the maximum lipid laboratory measurement used in our GWAS analysis. Further details on lipid-phenotype quality control are described in the Supplementary Note.

MVP association analysis. Genotyped and imputed DNA sequence variants with a MAF > 0.0003 were tested for association with the inverse-normal-transformed residuals of lipid values through linear regression assuming an additive genetic model. In our initial discovery analysis (stage 1), we performed association testing separately among individuals of each of three genetic ancestries (whites, blacks and Hispanics) and then meta-analyzed results across ethnic groups using an inverse-variance-weighted fixed-effects method. For variants with suggestive associations (association $P < 10^{-5}$), we sought replication of our findings in one of two independent studies: the 2017 GLGC exome array meta-analysis (stage 2a) or the 2013 GLGC joint meta-analysis (stage 2b). Replication was first attempted using summary statistics from the 2017 GLGC exome array study (stage 2a). A total of 242,289 variants in up to 319,677 individuals were analyzed after quality control and were available for replication. If a DNA sequence variant was not available for replication in the above exome array-focused study, we sought replication from publicly available summary statistics from the 2013 GLGC joint meta-analysis (stage 2b). An additional 2,044,165 variants in up to 188,587 individuals were available for replication in this study. In total, 2,286,454 DNA sequence variants in up to 319,677 individuals were available for independent replication in either stage 2a or stage 2b. We emphasize that if a variant was available for replication in both studies, replication was performed only using summary statistics from the 2017 GLGC exome array study given its larger sample size. We defined significant novel associations as those that were at least nominally significant in replication (peak $P < 0.05$) and had an overall $P < 5 × 10^{-8}$ (genome-wide significance) in the discovery and replication cohorts combined. Novel loci were defined as being greater than 1 Mb away from a known genome-wide-associated lead variant for lipids. Additionally, LD information from the 1000 Genomes Project was used to determine independent variants for which a locus extended beyond 1 Mb. All association $P$ values were two-sided. Further details on the association analysis are described in the Supplementary Note.

Conditional analysis. We used the COJO-GCTA software (see URLs) to perform an approximate, stepwise conditional analysis to identify independent variants within lipid-associated loci given that individual level data for the prior GLGC lipid analyses are not publicly available. We used summary statistics of ~1.9 million overlapping variants that we meta-analyzed across the two GLGC datasets (predominantly European) and the European MVP dataset to conduct this analysis (Supplementary Fig. 8) combined with an LD matrix obtained from 10,000 unrelated European individuals randomly sampled from the UK Biobank interim release.

Lipid TWAS. We performed a TWAS using summary statistics after a meta-analysis of ~1.9 million overlapping variants among GLGC (predominantly European) and European MVP datasets (Supplementary Fig. 8) and four gene-expression reference panels (whole blood from The Netherlands Twin Registry, adipose tissue from the Metabolic Syndrome in Men study, and tibial artery and liver from GTEx) in independent samples as previously described.

Identification of low frequency coding variant lipid association specific to blacks and hispanics. We estimated the proportion of the variance explained by the 444 previously described GWAS lipid variants weighted by their previously reported effect sizes (Supplementary Table 11) as a function of the number of lipid measurements in MVP in order to determine the approximate number of lipid measurements used in discovery. We performed this analysis using the mean of one, two, three, and five lipid measurements for each individual starting with their measurement closest to enrollment and moving toward the past. To account for the use of statin therapy, individuals with evidence of a statin prescription in their EHR at the time of enrollment had their LDL-C and total cholesterol values adjusted by dividing by 0.7 and 0.8, respectively, as previously described.

Loss of gene function analysis. We used the Variant Effect Predictor software to identify pLOF DNA sequence variants defined as: premature stop (nonsense), canonical splice-sites (splice-donor or splice-acceptor) or insertion/deletion variants that shifted frame (frameshift). For the pLOF lipid analysis, we then merged these variants with data from the Exome Aggregation Consortium (v0.3.1. see URLs), a publicly available catalogue of exome-sequence data to confirm consistency in variant annotation. We required that pLOF DNA sequence
variants be observed in at least 50 individuals, and set a statistical significance threshold of $P < 5 \times 10^{-8}$ (genome-wide significance).

Loss of PDE3B gene function and CAD. We identified a novel lipid association for a pLOF mutation in the PDE3B gene (rs150090666, Arg783Ter). For carriers of damaging mutations in phosphodiesterase 3B, we examined the effects of the mutation on risk for CAD using logistic regression in five separate cohorts: MVP, UK Biobank and three cohorts with exome sequencing: the MIGen, the PMBB and DiscovEHRR. In studies with exome sequencing, we combined pLOF variants with missense variants predicted to be damaging or possibly damaging by each of five computer prediction algorithms (LRT score, MutationTaster, PolyPhen-2, HumDiv, PolyPhen-2 HumVar and SIFT) as performed previously30,33. Because any individual damaging mutation was rare, variants were aggregated together for subsequent phenotypic analysis. We performed logistic regression on disease status, adjusting for age, sex and principal components of ancestry as appropriate. Effects of PDE3B damaging mutations were pooled across studies using an inverse-variance-weighted fixed-effects method. Further details on participating cohorts and CAD case definitions are described in the Supplementary Note. We set a two-sided $P < 0.05$ threshold for statistical significance.

PheWAS of variation in genes targeted by lipid-lowering therapies. For a set of DNA sequence variants within genes targeted by lipid-lowering medicines, we performed a PheWAS leveraging the full catalog of EHR ICD-9 diagnosis codes. We selected five lipid genes currently being targeted by pharmaceutical agents and identified functional variants in these genes: two nonsense variants (LPL, Ser474Ter and ANGPTL4 Glu121Ter) and three missense variants (ANGPTL4 Glu40Lys, APOA5 Ser19Trp, PCSK9 Arg46Leu). Details on PheWAS quality control, case definitions and association analysis are described in the Supplementary Note. We considered phenotypes to be significantly associated with a variant if they met a Bonferroni-corrected two-sided $P < 4.98 \times 10^{-5}$ (0.05/1,004 traits). For replication of our ANGPTL4 Glu40Lys type 2 diabetes finding, we combined the PheWAS results with publicly available data from the recently published trans-ancestry type 2 diabetes GWAS30 using an inverse-variance-weighted fixed-effects method.

Lipids and AAA Mendelian randomization analysis. Summary-level data for 223 genome-wide lipid-associated variants were obtained from publicly available data from the Global Lipids Genetics Consortium11. We then utilized results from a GWAS of 5,002 AAA cases and 139,968 controls performed in white MVP participants using the previously proposed definition17. The effect alleles were matched with all lipid and AAA summary data and three different Mendelian randomization analyses were performed: (1) inverse-variance-weighted; (2) multivariable; and (3) MR-Egger to account for pleiotropic bias. First, we performed inverse-variance-weighted Mendelian randomization using each set of variants for each lipid trait as instrumental variables. This method, however, does not account for possible pleiotropic bias. Therefore, we next performed inverse-variance-weighted multivariable Mendelian randomization. This method adjusts for possible pleiotropic effects across the included lipid traits in our analyses using effect estimates from the variant–AAA outcome and effect estimates from variant–LDL-C, variant–HDL-C and variant–triglycerides as predictors in one multivariable model. We additionally performed MR Egger as previously described53. This technique can be used to detect bias secondary to unbalanced pleiotropy in Mendelian randomization studies. In contrast to inverse-variance-weighted analysis, the regression line is unconstrained, and the intercept represents the average pleiotropic effects across all variants. Bonferroni-corrected two-sided $P$ values ($P < 0.016$ (0.05/3)) for three tests were used to declare statistical significance.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The full summary-level association data from the trans-ancestry meta-analysis for each lipid trait from this report are available through dbGaP, with accession number phs001672.v1.p1.

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Statistical parameters

When statistical analyses are reported, confirm that the following items are present in the relevant location (e.g. figure legend, table legend, main text, or Methods section).

- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- An indication of whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- The statistical test(s) used AND whether they are one- or two-sided
- Only common tests should be described solely by name; describe more complex techniques in the Methods section.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- A full description of the statistics including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted. Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated
- Clearly defined error bars
  - State explicitly what error bars represent (e.g. SD, SE, CI)

Our web collection on statistics for biologists may be useful.

Software and code

Policy information about availability of computer code

Data collection  
Phenotypic data was collected from the electronic health record and genetic data using the Million Veteran Program (MVP) Axiom array. All data was collated using R-3.2 as documented in the URLs section

Data analysis  
Data was collected using the EasyQC package (exemplar code link documented in the URLs section), and SNPTEST software program as outlined in the supplementary methods (exemplar code link documented in the URLs section)

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Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | All samples available of three ancestries (European, African, Hispanic) were used for analysis (after quality control). Sample size was determined based on genetic data available from MVP. |
|-------------|--------------------------------------------------------------------------------------------------|
| Data exclusions | Data were excluded if they did not pass our QC metrics, or if they did not fall within the three main ancestries used for analysis |
| Replication | Replication was performed using data from one of 2 sources: 1) GLGC 2017 exome chip GWAS summary statistics, or 2) GLGC 2013 joint-meta-analysis GWAS summary statistics |
| Randomization | N/A |
| Blinding | N/A |

Reporting for specific materials, systems and methods

| Materials & experimental systems |
|----------------------------------|
| n/a Involved in the study |
| ☑ Unique biological materials |
| ☑ Antibodies |
| ☑ Eukaryotic cell lines |
| ☑ Palaeontology |
| ☑ Animals and other organisms |
| ☑ Human research participants |

| Methods |
|---------|
| n/a Involved in the study |
| ☑ ChIP-seq |
| ☑ Flow cytometry |
| ☑ MRI-based neuroimaging |

Human research participants

Policy information about studies involving human research participants

Population characteristics Demographics and participant counts for a number of cardiometabolic traits for the 312,571 white, black, and Hispanic MVP participants that passed our quality control are depicted in Table 1.

Recruitment Individuals aged 19 to 104 years have been recruited voluntarily from more than 50 VA Medical Centers nationwide for participation in the Million Veteran Program biobank study.