# Whole genome sequence analysis of blood lipid levels in >66,000 individuals

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### **Abstract:**

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Plasma lipids are heritable modifiable causal factors for coronary artery disease, the leading cause of death globally. Despite the well-described monogenic and polygenic bases of dyslipidemia, limitations remain in discovery of lipidassociated alleles using whole genome sequencing, partly due to limited sample sizes, ancestral diversity, and interpretation of potential clinical significance. Increasingly larger whole genome sequence datasets with plasma lipids coupled with methodologic advances enable us to more fully catalog the allelic spectrum for lipids. Here, among 66,329 ancestrally diverse (56% non-European ancestry) participants, we associate 428M variants from deep-coverage whole genome sequences with plasma lipids. Approximately 400M of these variants were not studied in prior lipids genetic analyses. We find multiple lipid-related genes strongly associated with plasma lipids through analysis of common and rare coding variants. We additionally discover several significantly associated rare non-coding variants largely at Mendelian lipid genes. Notably, we detect rare LDLR intronic variants associated with markedly increased LDL-C, similar to rare LDLR exonic variants. In conclusion, we conducted a systematic whole genome scan for plasma lipids expanding the alleles linked to lipids for multiple ancestries and characterize a clinically-relevant rare non-coding variant model for lipids.

### Introduction

Discovery of rare alleles linked to plasma lipids (i.e., low-density lipoprotein cholesterol [LDL-C], high-density lipoprotein cholesterol [HDL-C], total cholesterol [TC], and triglycerides [TG]) continue to yield important translational insights toward coronary artery disease (CAD), including PCSK9 and ANGPTL3 inhibitors now available in clinical practice<sup>1,2,3,4,5</sup>. The monogenic and polygenic bases of plasma lipids are well-suited to population-based discovery analyses and confer broader insights for genetic analyses of complex traits. We now evaluate numerous newly catalogued, largely rare, alleles never previously systematically analyzed with lipids.

Analyses of imputed array-derived genome-wide genotypes and whole exome sequences in hundreds of thousands of increasingly diverse individuals continue to uncover low-frequency protein-coding variants linked to lipids. Due to purifying selection, causal variants conferring large effects tend to occur relatively more recently, and are thus rare and often specific to families or communities<sup>6</sup>. Most discovery analyses for large-effect rare alleles have focused on the analysis of disruptive protein-coding variants given (1) well-recognized constraint in coding regions, (2) incomplete genotyping of rare non-coding sequence given relative sparsity of deep-coverage (i.e., >30X) whole genome sequencing (WGS), and (3) better prediction of coding versus non-coding sequence variation consequence<sup>1,7,8,9,10,11,12</sup>. We recently described a statistical framework incorporating multi-dimensional reference datasets paired with genomic data to improve rare coding and non-coding variant analyses for WGS analysis of lipids and other complex traits<sup>13,14</sup>. Furthermore,

including individuals of non-European ancestry facilitates the discovery of both novel alleles at established loci as well as novel loci<sup>14,15,16</sup>.

Here, we examine the full allelic spectrum with plasma lipids using whole genome sequences and harmonized lipids from the National Heart, Lung, and Blood Institute (NHLBI) Trans-Omics for Precision Medicine (TOPMed) program<sup>17,18</sup>. We studied 66,329 participants and 428 million variants across multiple ancestry groups – 44.48% European, 25.60% Black, 21.02% Hispanic, 7.11% Asian and 1.78% Samoan. We identified robust allelic heterogeneity at known loci with several novel variants at these loci; we additionally identified novel loci and pursued replication in independent cohorts (31.50% non-European samples). We then explored the association of genome-wide rare variants with lipids, with detailed explorations of rare coding and non-coding variant models at known Mendelian dyslipidemia genes. Our systemic effort yields new insights for plasma lipids provides a framework for population based WGS analysis of complex traits.

#### Results

### Overview

We studied the TOPMed Freeze8 dataset of 66,329 samples from 21 studies and performed genome-wide association studies (GWAS) separately for the four plasma lipid phenotypes (i.e., LDL-C, HDL-C, TC and TG) using 28M individual autosomal variants (minor allele count [MAC] > 20) and aggregated rare autosomal variant (minor allele

frequency [MAF] < 1%) association testing for 417M variants (**Fig. 1, Supplementary Fig. 1**). Secondarily, we associated individual variants with minor allele frequencies (MAF) > 0.01% within each ancestry group to detect ancestry-specific lipid-associated alleles. We intersected our results with currently published array-based GWAS results<sup>15</sup> to identify novel associations with lipids. We performed replication analyses for the putative novel associations identified, in up to approximately 45,000 independent samples with array-based genotyping imputed to TOPMed. Finally, we conducted rare variant association studies as multiple aggregate tests across the genome to identify gene-specific functional categories and non-coding genomic regions influencing plasma lipid concentrations.

### **TOPMed baseline characteristics**

The TOPMed Informatics Research Center (IRC) and TOPMed Data Coordinating Center (DCC) performed quality control, variant calling, and calculated the relatedness of population structures of Freeze 8 data<sup>17</sup>. We studied 66,329 samples across 21 cohorts and 41,182 (62%) were female. The ancestry distribution was 29,502 (44.46%) White, 16,983 (25.60%) Black, 13,943 (21.02%) Hispanic, 4,719 (7.11%) Asian, and 1,182 (1.78%) Samoan (**Supplementary Table 1**). The mean (standard deviation [SD]) age of the full cohort was 53 (15.00) years which varied by cohort from 25 (3.56) years for Coronary Artery Risk Development in Young Adults (CARDIA) to 73 (5.38) years for Cardiovascular Health Study (CHS). The Amish cohort had a higher-than-average concentration of LDL-C (140 [SD 43] mg/dL) and HDL-C (56 [SD 16] mg/dL) as well as lower TG (median 63 [IQR 50] mg/dL) consistent with the known founder mutations in *APOB* 

and *APOC3*<sup>7,8,14</sup>. In the Women's Health Initiative (WHI) cohort, the TC (230 [SD 41] mg/dL) and TG (median 129 [IQR 87] mg/dL) concentrations were higher than for other cohorts as previously described<sup>12</sup>. We accounted for lipid-lowering medications and fasting status and inverse rank normalized the phenotypes as before<sup>12,14</sup> which are further detailed in the **Methods**. The adjusted normalized lipid concentrations for the four lipids were similar across the cohorts.

A total of 428M variants passed the quality criteria with an average depth >30X in 22 autosomes. 202M variants were singletons, 417M were rare variants (MAF<1%), and 11M were common or low frequency variants (MAF>1%) with differences by cohort (**Supplementary Table 2**).

# Individual variant associations with lipids

Approximately 28M variants with MAC > 20 were individually associated with LDL-C, HDL-C, TC and TG. We used p-value < 5x10<sup>-9</sup> to claim significance as previously recommended for whole genome sequencing common variant association studies<sup>14,19</sup>. The total numbers of variants that met our significance threshold were 2,214, 2,314, 2,697 and 2,442 for LDL-C, HDL-C, TC and TG, respectively, and after clumping<sup>20</sup> the numbers of variants were 357, 338, 324, and 289, respectively. Of these variants, most were previously demonstrated to be associated with plasma lipids either at the variant- or locus-level<sup>15</sup> (**Supplementary Table 3**, **Supplementary Fig. 2**).

To identify putative novel variant associations, we compared our results to a recent multi-ethnic lipid GWAS among 312,571 participants of the Million Veteran Program (MVP)<sup>15</sup> as well as the GWAS Catalog (All associations(v1.0) file

dated 06/04/2020) (**Fig. 2**). We clumped (window 250 kb, r<sup>2</sup> 0.5) significant variants using Plink<sup>20</sup> and queried these in the GWAS Catalog and MVP. Among genome-wide significant variants, we tabulated 'known-position' (variant previously associated), 'known-loci' (variants not previously significantly associated with the corresponding lipid phenotype but within 500 kb of a known locus, thereby representing additional allelic heterogeneity), and 'novel' variants (variants not in a known lipid locus) (**Supplementary Table 3**).

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The novel variants, tabulated in **Table 1**, are divided into two subsets – 'novel variants' or variants at established lipid loci for another lipid phenotype, and 'novel loci,' representing new loci associations for any lipid phenotype. For example, the CETP locus is well-known for its link to HDL-C, but we now found that rs183130 (16:56957451:C:T, MAF 28.3%) at the locus is associated with LDL-C. Similarly, the variants rs7140110 (13:113841051:T:C, MAF 27.8%) GAS6 and rs73729083 (7:137875053:T:C, MAF 4.5%) CREB3L2 are newly associated with TC, while previous studies showed that rs73729083 associates with LDL-C<sup>21</sup> and rs7140110 associates with LDL-C<sup>22</sup> and TG<sup>23</sup>. Index variants at novel loci were typically low frequency variants often observed in non-European ancestries, so we also conducted ancestry-specific association analyses for these alleles (**Supplementary Table 4**). For example, 12g23.1 (12:97352354:T:C, MAF 0.3%) and 4g34.2 (4:176382171:C:T, MAF 0.2%) associations with LDL-C are specific to Hispanic (MAF 1.3%) and Black (MAF 0.6%) populations, respectively and among Asians (MAF 1.5%) alone, 11q13.3 (11:69219641:C:T, MAF 0.2%) was associated with TG. One variant initially passing the novel locus filter for HDL-C (RNF111 - rs112147665, beta = 8.664, pvalue = 6.51x10<sup>-10</sup>), was in LD (r=0.7) with LIPC p.Thr405Met (rs113298164) which is known to be associated with HDL-

C. The lead variant from MVP was 604 kb away from the *RNF111* variant but the rare *LIPC* missense variant p.Thr405Met was 421 kb away. Conditional analysis accounting for *LIPC* p.Thr405Met rendered the non-coding variant near *RNF111* variant non-significant (beta = 4.351, p-value = 2.47x10<sup>-02</sup>), therefore we reclassified *RNF111* variant as a known-position variant. Ancestry-specific GWAS did not yield additional novel loci beyond our larger trans-ancestry GWAS. Majority of genome significant single variants were captured by previous lipid GWAS<sup>15</sup>, but ancestry specific novel-hits are unique to WGS TOPMed data.

Due to the paucity of available diverse WGS datasets with lipids of comparable size, we pursued replication with two genome-wide array-based genotyped datasets imputed to TOPMed WGS<sup>17,24</sup>: Mass General Brigham (MGB)

Biobank (N=25,137) and Penn Medicine Biobank (N=20,079)<sup>25,26</sup>, the replication cohorts had diverse ancestry distribution, where non-European samples accounted for 15.77% in MGB Biobank and 51.20% in Penn Medicine Biobank

(Supplementary Table 5). We brought seven putative novel variants with p-values < 5x10<sup>-9</sup> forward for replication. The three common variants, rs183130 (*CETP*), rs7140110 (*GAS6*) and rs73729083 (*CREB3L2*), that were associated with both LDL-C and TC in TOPMed also replicated in MGB and two (rs183130, rs73729083) replicated in Penn Biobank at an alpha level of 0.05 and consistent direction of effect (Table 1). The two variants that were associated in both replication studies were most significantly associated among African Americans in TOPMed (rs183130: beta = -2.762 mg/dL, p-value = 5.71x10<sup>-07</sup>; rs73729083: beta = -3.725 mg/dL, p-value = 5.25x10<sup>-07</sup>). Low-frequency variants from specific ancestry groups associated with lipids in TOPMed were not replicated but we cannot rule out the possibility of reduced power due

to general underrepresentation of non-white ancestry groups in the replication data. In exploratory analyses, we extended the same approach for variants discovered to have  $5x10^{-9} < p$ -value  $< 5x10^{-7}$  but did not observe replication (**Supplementary Table 6**).

## CETP locus, HDL-C, and LDL-C

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CETP is a well-recognized Mendelian HDL-C gene and the locus was previously known to be significantly associated with HDL-C, TC and TG at genome-wide significance<sup>15</sup>. Pharmacologic CETP inhibitors have shown strong associations with increased HDL-C but mixed effects for LDL-C reduction in clinical trials 27,28,29,30. We found that the CETP locus variant rs183130 (chr16:56957451:C:T, MAF 28.3%, intergenic variant) was associated with reduced LDL-C concentration (beta = -1.568 mg/dL, SE = 0.264, p-value = 2.88x10<sup>-09</sup>). The lead HDL-C-associated variant at the locus. rs3764261 (chr16:56959412:C:A, MAF 30.3%), was associated with 3.5 mg/dL increased HDL-C (p-value = 8.03x10<sup>-283</sup>), and rs183130 was associated with 3.9 mg/dL increased HDL-C (p-value < 1x10<sup>-284</sup>) as well. Among the ancestry groups analyzed, rs183130 was most significantly associated with LDL-C among those of African ancestry (beta = -2.762 mg/dL, p-value = 5.71x10<sup>-07</sup>) (**Supplementary Table 7**). We next investigated variants by their HDL-C and LDL-C effects within this locus (+/- 500kb of rs183130 and rs3764261) (Fig. 3). We identified five variants showing at least suggestive (p-value < 5x10<sup>-07</sup>) association with both HDL-C and LDL-C. Though variants with strong LD (linkage disequilibrium) existed, ancestry-specific analyses showed that the stronger LDL-C effects were among those of African ancestry.

To better understand the mechanisms for HDL-C and LDL-C effects at the CETP locus, we pursued colocalization with eQTLs from 3 tissues (Liver, Adipose Subcutaneous and Adipose Visceral [Omentum]) from GTEx<sup>31</sup>. We analyzed 5 LDL-C and 441 HDL-C associated (p-values < 5x10<sup>-07</sup>) variants. We correlated eQTL effect estimates for genes at the locus with lipid outcome effect estimates. Indeed, CETP gene expression effects were strongly negatively correlated with HDL-C effects (Liver: ρ -0.933, p-value 4.01x10<sup>-17</sup>; Adipose Subcutaneous: ρ -0.762, p-value 8.87x10<sup>-12</sup>; Adipose Visceral: ρ -0.739, p-value 5.52x10<sup>-10</sup>) (**Supplementary Fig. 3**). However, *CETP* expression effects were not significantly correlated with LDL-C (Liver: ρ 0.007, p-value 0.99; Adipose Subcutaneous: ρ 0.344, p-value 0.57; Adipose Visceral: ρ -0.59, p-value 0.29). Given the possibility that the observed lack of correlation for LDL-C could be due to reduced power from a limited number of variants attaining a suggestive p-value (< 5x10<sup>-07</sup>), we repeated the analysis with a subset of 122 nominally significant (p-value < 0.05) LDL-C associated variants in this locus. Indeed, CETP gene expression effects were strongly positively correlated with LDL-C effects (Liver: p 0.957, p-value 2.28x10<sup>-08</sup>; Adipose Subcutaneous: p 0.922, pvalue 1.34 x10<sup>-15</sup>; Adipose Visceral:  $\rho$  0.868, p-value 6.09x10<sup>-11</sup>).

# Rare variant aggregates associated with lipids

## I) Gene-Centric associations

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We next evaluated the association of aggregated rare (MAF<1%) variants, linked to protein-coding genes ('genecentric'). We employed a Bonferroni-corrected significance threshold of 0.05/20,000=2.5x10<sup>-06</sup> for coding and non-coding

gene-centric rare variant analyses (**Supplementary Fig. 4**). We identified 102 coding and 160 non-coding gene-centric rare variant aggregates significantly associated with at least one of the four plasma lipid phenotypes in nonconditional analysis (**Supplementary Table 8-9**). We secondarily conditioned our significant aggregate sets on variants individually associated with lipid levels from the GWAS catalog, MVP summary statistics and the TOPMed data. We identified 74 coding and 25 non-coding rare variants aggregates associated with at least one lipid level after conditional analyses (**Supplementary Table 10-11**).

Most of the coding gene-centric sets remained significant after secondary conditioning while a minority of non-coding gene-centric sets remained significant after conditioning. Significant genes identified from coding rare variant analyses included multiple known Mendelian lipid genes including *LCAT*, *LDLR*, and *APOB* (**Supplementary Table 10**). *RFC2* putative loss-of-function mutations (combined allele frequency < 0.002%) were significantly associated with triglycerides (p-value 2x10<sup>-06</sup>) representing a putative novel association for triglycerides. The *RFC2* aggregate set (plof) was associated with reduced TG (beta = -0.89 for log[TG]). The persistently significant regions identified from non-coding rare variant analyses linked to genes included the UTR (untranslated region) for *CETP* and promoter-CAGE (CAGE- Cap Analysis of Gene Expression sites) around *APOA1* for HDL-C, and *APOE* promoter-CAGE, *APOE* enhancer-DHS (DHS - DNase hypersensitivity sites), and *EHD3* promoter-DHS for total cholesterol (**Supplementary Table 11**). Most of the coding aggregates had larger effects compared to non-coding aggregates, and among the non-coding aggregates *SPC24* 

non-coding aggregate (enhancer-CAGE) at the *LDLR* locus had the strongest effect for LDL-C (beta = 2.320 mg/dL; p-value =  $1.75 \times 10^{-05}$ ).

## II) Region-Based associations

We also performed unbiased region-based rare variant association analyses tiled across the genome with both static and dynamic window sizes. We first evaluated 2.6M regions statically at 2 kb size and 1 kb window overlap by the sliding window approach. Statistical significance was assigned at 0.05/(2.6x1<sup>-06</sup>)=1.88x10<sup>-08</sup>. We identified 28 significantly associated windows with at least one lipid phenotype. After conditioning on variants individually associated with the corresponding lipid phenotype, we identified two regions at *LDLR* still significantly associated with both total cholesterol and LDL-C although these regions included both intronic and exonic variants (**Supplementary Table 12**). *LDLR* intron 1, which encodes *LDLR-AS1* (LDLR antisense RNA 1) on the minus strand, had suggestive evidence for association with TC (p-value 3.17x10<sup>-6</sup>) with -2.76 mg/dL reduction in TC. A prior study identify that a common variant (rs6511720, MAF 0.11) in *LDLR* intron 1 is associated with increased *LDLR* expression in a luciferase assay and reduction in LDL-C<sup>32</sup>. When adjusting for rs6511720, the significance improved (p-value 1.43x10<sup>-8</sup>) with -3.35 mg/dL reduction in TC.

For dynamic window scanning of the genome, we implemented the SCANG method<sup>33</sup>. The SCANG procedure accounts for multiple testing by controlling the genome-wide error rate (GWER) at 0.1<sup>33</sup>. In the dynamic window-based workflow, STAAR-O detected 51 regions significantly associated with at least one lipid phenotype after conditioning on

known variants (Supplementary Table 13). Most of the regions mapped to known Mendelian lipid genes, including LCAT (8.7x10<sup>-13</sup>) for HDL-C, and LDLR (2.4x10<sup>-28</sup>, 7.3x10<sup>-26</sup>) and PCSK9 (2.9x10<sup>-12</sup>, 5.5x10<sup>-12</sup>) for LDL-C and TC, respectively. Exon 4 aggregates of LDLR were specifically associated with 20 mg/dL increase in LDL-C. PCSK9 Exon2-Intron2 region spanning chr1:55043782-55045960 had significantly reduced LDL-C by 6 mg/dL (p-value = 3x10<sup>-13</sup>), and the effect persisted even with only Intron 2 rare variants of PCSK9 (-5 mg/dL, p-value =  $2x10^{-8}$ ). Strikingly, in secondary analyses, we found evidence for very large effects for rare variants in LDLR Introns 2 and 3 (+21 mg/dL, p-value =  $7x10^{-4}$ ) and LDLR Introns 16 and 17 (+17 mg/dL, p-value = 0.02), similar to rare coding LDLR mutations. While 32 of the significant dynamic windows also included exonic regions, there were also several dynamic windows significantly independently associated with lipids not containing exonic regions. For example, four non-coding windows (two overlapping) at 2p24.1, which harbors the Mendelian APOB gene, were significantly associated with LDL-C. Intronic non-coding regions were associated with both LDL-C and TC -associated windows at LPAL2-LPA-SLC22A3; for example LPAL2 Intron 3 was associated with a 3.7 mg/dL increase in TC. Non-coding TC-associated significant dynamic windows were near TOMM40/APOE. One rare variant signal observed was at TOMM40 Intron 6, where the 'poly-T' variant in this region is on the APOE4 haplotype and influences expressivity for Alzheimer's disease age-of-onset<sup>34,35</sup>. For HDL-C, we identified significant non-coding windows at an intergenic region near LPL and CD36 Intron 4. In the generation of the spontaneously hypertensive rat model, the deletion of intron 4 in Cd36 with resultant Cd36 deficiency has been mapped to

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defective fatty acid metabolism in this model<sup>36</sup>. Several regions significant in SCANG were not even nominally significant in burden association analyses indicating the likelihood of causal variants with bidirectional effects.

Several gene-centric non-coding aggregates associated with lipids near known monogenic lipid genes but mapped to another gene at the locus via annotations. Therefore, we performed downstream conditional analyses adjusting the gene-centric non-coding results for rare coding variants (MAF<1%) within known lipid monogenic genes (**Supplementary Table 14**). When accounting for both common and rare coding variants at the nearby familial hypercholesterolemia *LDLR* gene, *SPC24*-enhancer DHS was significantly associated with total cholesterol (p-value= 3.01x10<sup>-11</sup>) and with suggestive evidence for LDL-C (p-value= 1.57x10<sup>-06</sup>). In a similarly adjusted model, *LDLR*-enhancer-DHS showed a strong association with TC (p-value 5.18x10<sup>-12</sup>). When adjusting for known common variants as well as rare coding variants in *PCSK9*, both *PCSK9*-enhancer DHS and *PCSK9*-promoter DHS were significantly associated with total cholesterol. (**Fig. 4, Supplementary Fig. 5**). Through this procedure, *CETP* UTR retained significance for its independent association with HDL-C as well as the putatively novel gene *EHD3*-promoter DHS association with TC. However, the non-coding genecentric *APOC3* and *APOE* associations were rendered non-significant for HDL-C and TC, respectively.

Since we cannot rule out the possibility of reduced power for genome-wide rare variant analyses, we leveraged current knowledge of 22 Mendelian lipid genes for more focused exploratory analyses<sup>14</sup>. We validated most genes in rare variant coding analyses. The genes with the strongest coding signals typically had at least nominal evidence of genecentric non-coding rare variant associations (**Supplementary Table 15, Supplementary Fig. 6**). When rare coding

variants were introduced into the model, the evidence for non-coding rare variant associations were largely unchanged.

Our findings expanding the currently described genetic basis for hypercholesterolemia to include rare non-coding variation at LDLR and PCSK9 (Fig. 5).

### **Discussion**

Conducting one of the largest population-based WGS association analyses, we now simultaneously interrogate and establish a common, rare coding, and rare non-coding variant model for a complex trait. Utilizing 66,329 diverse individuals with deep-coverage WGS, we interrogated 428M variants with plasma lipids expanding the allelic series to rare non-coding variants, often within introns, of Mendelian lipid genes with prior robust rare coding variant support. Our observations have important implications for plasma lipids as well as the genetic basis of complex traits more broadly.

WGS of diverse ancestries enables both allelic and locus heterogeneity for complex traits. Population genetic analyses have largely been enriched for individuals of European descent<sup>37</sup>. Genetic association of plasma lipids using arrays or whole exome sequencing among Europeans have yielded several important insights regarding plasma lipids and the causal determinants of CAD<sup>5,4,38,39,40</sup>. Similar increasingly larger studies among non-Europeans have often yielded new genetic loci and sometimes new genes, such as *PCSK9*<sup>1,15,41,42,16</sup>. Such differences have also led to concerns about the use of polygenic risk scores gleaned from much larger European GWAS of complex traits for non-Europeans<sup>43</sup>. Aided by the availability of WGS data, we identify new putative loci associated with lipids in non-Europeans. Furthermore, our

study enabled the discovery of several novel alleles at known loci, with richly distinct allelic heterogeneity across ancestry groups. For example, HDL-C-raising *CETP* locus variants linked to *CETP* gene expression were only associated with LDL-C reduction among those of African ancestry. While all pharmacologic CETP inhibitors increase HDL-C, only those that decrease LDL-C also reduce cardiovascular disease risk<sup>27,28,30,29</sup>. Given the contribution of genetic differences, clinical trials with more diverse samples would show insights.

Our study now provides increasingly robust evidence for a rare non-coding variant model for complex traits. Our rare non-coding variant associations in both gene-centric and sliding window models were largely restricted to the introns of Mendelian lipid genes with prior robust rare coding variant support consistent with biologic plausibility<sup>44</sup>. Rare intronic variants, often impacting splicing, have been previously implicated in afflicted Mendelian families or small exceptional case series, often through candidate gene approaches<sup>45,46,47,48</sup>. We discovered one example of a rare non-coding signal without prior rare coding support – i.e., *EHD3*. We obtained estimates of phenotypic effect using burden tests. For most regions, even nominal significance was not detected using burden testing indicating the likelihood of variants with bidirectional effects further complicating clinical interpretation. When burden signals were detected, observed effects were typically larger than common non-coding variants and less than rare coding variants, with the exception of *LDLR*, consistent with whole genome mutational constraint models<sup>49,50,51</sup>.

The detection of independent rare non-coding variant signals has remained elusive largely due to limited sample sizes with requisite WGS and limitations in the interpretation of rare non-coding variation functional consequence.

Previously, we used annotated functional non-coding sequence in 16,324 TOPMed participants, and found that rare non-coding gene regions associated with lipid levels, but they were not independent of individually associated single variants<sup>14</sup>. Using STAAR, we observed putative rare non-coding variant associations for lipids independent of individual variants associated with lipids in TOPMed.

WGS can improve diagnostic yield beyond the current standard of next-generation gene panel sequencing for dyslipidemias. A very small fraction with severe hypercholesterolemia and features consistent with strong genetic predisposition have a familial hypercholesterolemia variant in *LDLR*, *APOB*, or *PCSK9*<sup>52,53</sup>. The presence of familial hypercholesterolemia variants is independently prognostic for CAD, beyond lipids, and merits the consideration of more costly lipid-lowering medications<sup>52,53,54,55</sup>. We now observe that rare *LDLR* variants in Introns 2, 3, 16, and 17 lead to approximately 0.5 standard deviation increase in LDL-C, approximating effects observed with clinically reported exonic familial hypercholesterolemia variants in *LDLR*<sup>55</sup>. Small studies have indicated the possibility of rare intronic *LDLR* variants causing familial hypercholesterolemia due to altered splicing, which we now observe in our unbiased population-based WGS study<sup>56,57</sup>. A WGS approach to lipid disorders, particularly for familial hypercholesterolemia, will markedly improve the diagnostic yield beyond existing limited approaches.

Our dynamic window approach may also improve the clinical curation of exonic variants. Among the data used to curate exonic variants is the use of *in silico* functional prediction tools<sup>58</sup>. Although evolutionary constraint measures are typically employed, such tools are largely agnostic to functional domain. As it relates to lipids, disruptive *APOB* and

*PCSK9* exonic variants can lead to strikingly opposing directions with large effects for LDL-C depending on locations<sup>1,8,59,60</sup>. Using SCANG<sup>33</sup>, we detect a significant association with large effect for *LDLR* Exon 4 itself. This observation supports the pathogenicity of *LDLR* Exon 4 disruptive variants among patients with severe hypercholesterolemia. The majority of familial hypercholesterolemia variants worldwide occur in Exon 4 of *LDLR*<sup>61,62,63,64</sup>. Conventional rare coding variant analyses aggregate all exonic variants for a transcript. Here, we demonstrate an opportunity for exon-level rare variant association testing.

Our study has important limitations. First, while our study is large for a WGS study by contemporary standards, it is dwarfed by existing GWAS datasets limiting power for novel discovery. Nevertheless, by using WGS in diverse ancestries, we can study hundreds of millions new variants. Second, prediction of rare non-coding variation consequence to prioritize causal variants remains a challenge thereby limiting power<sup>65</sup>. The striking difference for most STAAR and burden results also highlights bidirectional effects for rare non-coding variants within the same region and further challenges for clinical utility. Third, given the paucity of multi-ancestral WGS datasets with lipids, our analyses are largely restricted to TOPMed. For single variant associations, we pursued TOPMed-imputed GWAS datasets but were limited by the lack of ancestral diversity. As TOPMed is a consortium of multiple different cohorts, we demonstrate consistencies by cohort. Furthermore, rare variant non-coding signals were largely restricted to regions with rare variant coding signals supporting biological plausibility.

In conclusion, using WGS and lipids among 66,329 ancestrally diverse individuals we expand the catalog of alleles associated with lipids, including allelic heterogeneity at known loci and locus heterogeneity by ancestry. We characterize the common, rare coding, and rare non-coding variant model for lipids. Lastly, we now demonstrate a monogenic-equivalent model for rare *LDLR* intronic variants predisposing to marked alterations in LDL-C, currently not recognized in current population or clinical models for LDL-C.

#### **Online Methods**

#### Dataset

# i) Contributing studies

The discovery cohort includes whole genome sequenced (WGS) data of 66,329 samples from 21 studies of the Trans-Omics for Precision Medicine (TOPMed) program with blood lipids available <sup>17</sup>. The overall goal of TOPMed is to generate and use trans-omics, including whole genome sequencing, of large numbers of individuals from diverse ancestral backgrounds with rich phenotypic data to gain novel insights into heart, lung, blood, and sleep disorders. The Freeze 8 data includes 140,306 samples out of which 66,329 samples qualified with lipid phenotype. Freeze 8 dataset passed the central quality control protocol implemented by the TOPMed Informatics Research Core (described below) and was deposited in the dbGaP TOPMed Exchange Area.

The studies included in the current dataset, along with their abbreviations and sample sizes, contains the Old Order Amish (Amish, n=1,083), Atherosclerosis Risk in Communities study (ARIC, n=8,016), Mt Sinai BioMe Biobank (BioMe, n=9,848), Coronary Artery Risk Development in Young Adults (CARDIA, n=3,056), Cleveland Family Study (CFS, n=579), Cardiovascular Health Study (CHS, n=3,456), Diabetes Heart Study (DHS, n=365), Framingham Heart Study (FHS, n=3,992), Genetic Studies of Atherosclerosis Risk (GeneSTAR, n=1,757), Genetic Epidemiology Network of Arteriopathy (GENOA, n=1,046), Genetic Epidemiology Network of Salt Sensitivity (GenSalt, n=1,772), Genetics of Lipid-Lowering Drugs and Diet Network (GOLDN, n=926), Hispanic Community Health Study - Study of Latinos (HCHS\_SOL, n=7714), Hypertension Genetic Epidemiology Network and Genetic Epidemiology Network of Arteriopathy (HyperGEN, n=1,853), Jackson Heart Study (JHS, n=2,847), Multi-Ethnic Study of Atherosclerosis (MESA, n=5,290), Massachusetts General Hospital Atrial Fibrillation Study (MGH AF, n=683), San Antonio Family Study (SAFS, n=619), Samoan Adiposity Study (SAS, n=1,182), Taiwan Study of Hypertension using Rare Variants (THRV, n=1,982) and Women's Health Initiative (WHI, n=8,263) (Please see Supplementary Text for additional details). The multi-ancestral data set included individuals from White (44%), Black (26%), Hispanic (21%), Asian (7%), and Samoan (2%) ancestries. Study participants granted consent per each study's Institutional Review Board (IRB) approved protocol. Secondarily, these data were analyzed through a protocol approved by the Massachusetts General Hospital IRB. Supplementary Table 1 details the number of samples across different studies and ancestral group.

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The replication cohorts include TOPMed-imputed genome-wide array data from the Mass General Brigham (MGB) and Penn Medicine Biobanks which consist of 25,137 samples and 20,079 samples respectively<sup>26,25</sup>. We curated the MGB Biobank and Penn Medicine Biobank phenotype data from the corresponding electronic health record databases in accordance with corresponding institutional IRB approvals. Consent was previously obtained from each participant regarding storage of biological specimens, genetic sequencing, access to all available electronic health record (EHR) data, and permission to recontact for future studies. The MGB Biobank consists of 54% and Penn Medicine Biobank consist of 52% female samples and average ages were 55.89 years and 58.35 years, respectively (**Supplementary Table 5**).

## ii) Phenotypes

The primary outcomes in this study included LDL cholesterol (LDL-C), HDL cholesterol (HDL-C), total cholesterol (TC) and triglycerides (TG) phenotypes. LDL-C was either directly measured or calculated by the Friedewald equation when triglycerides were <400 mg/dL. Given the average effect of lipid lowering-medicines, when lipid-lowering medicines were present, we adjusted the total cholesterol by dividing by 0.8 and LDL-C by dividing by 0.7, as previously done<sup>14</sup>. Triglycerides remained natural log transformed for analysis. Fasting status was accounted for with an indicator variable. We harmonized the phenotypes across each cohort<sup>18</sup> and inverse rank normalization of the residuals of each race within each cohort scaled by the standard deviation of the trait and adjusted for covariates<sup>12</sup>. We included covariates such

as age, age<sup>2</sup>, sex, PC1-11, study-groups as well as Mendelian founder lipid variants *APOB* p.R3527Q and *APOC3* p.R19X for the Amish cohort<sup>7,66,8</sup>. **Supplementary Table 1** provides the distributions of each of the four lipid phenotypes by cohort, ancestral groups, and gender. We executed similar steps of phenotype harmonization and normalization for the replication cohorts. Additionally, we adjusted the MGB Biobank for study-center and array-type, and Penn Medicine Biobank for ancestry and BMI in addition to the other common covariates.

## iii) Genotypes

Whole genome sequencing of goal >30X coverage was performed at seven centers (Broad Institute of MIT and Harvard, Northwest Genomics Center, New York Genome Center, Illumina Genomic Services, PSOMAGEN [formerly Macrogen], Baylor College of Medicine Human Genome Sequencing Center and McDonnell Genome Institute [MGI] at Washington University). In most cases, all samples for a given study within a given Phase were sequenced at the same center (Supplementary Text). The reads were aligned to human genome build GRCh38 using a common pipeline across all centers (BWA-MEM).

The TOPMed Informatics Research Core at the University of Michigan performed joint genotype calling on all samples in Freeze 8. The variant calling "GotCloud" pipeline (https://github.com/statgen/topmed\_variant\_calling) is under continuous development and details on each step can be accessed through TOPMed website for Freeze8 (https://www.nhlbiwgs.org/topmed-whole-genome-sequencing-methods-freeze-8)<sup>17</sup>. The resulting BCF files were split by

study and consent group for distribution to approved dbGaP users. Quality control was performed at each stage of the process, poor variant quality was indicated by missing rate >20%, mappability score <0.8, mean depth of coverage >500X, and Ti/Tv ratio, by the Sequencing Centers, the IRC and the TOPMed Data Coordinating Center (DCC). The VCF/BCF files were converted to GDS (Genomic Data Structure) format by the DCC and were deposited into the dbGap TOPMed Exchange Area.

The genetic relationship matrix (GRM) is an N\*N matrix of relatedness information of the samples included in the study and was computed centrally using 'PC-relate' R package (version: 1.24.0)<sup>67</sup>. Using the 'Genesis' R package (version:2.20.1)<sup>68</sup> we generated subsetted GRM for the samples with plasma lipid profiles. The GDS files with the variants were annotated internally by curating data from multiple database sources using Functional Annotation of Variant–Online Resource (FAVOR (http://favor.genohub.org)<sup>13</sup>. This study used the resultant aGDS (annotation GDS) files.

The MGB Biobank replication cohort was genotyped using three different arrays (Multiethnic Exome Global (Meg), Human multi-ethnic array (Mega), and Expanded multi-ethnic genotyping array (Megex)), and we separately imputed the data using TOPMed imputation server with default parameters<sup>69,70</sup>. This study applied the Version-r2 of the imputation panel, it includes 97,256 reference samples and ~300M genetic variants. The Illumina Global Screening array was used to genotype the Penn Medicine Biobank. Penn Medicine Biobank TOPMed imputation was performed using EAGLE<sup>70</sup> and Minimac<sup>71</sup> software. For this study we downloaded variants that passed a min R<sup>2</sup> threshold of 0.3.

## **Single Variant Association**

We performed genome-wide single variant association analyses for autosomal variants with minor allele frequency (MAF) greater than 0.1% across the dataset with each of the four lipid phenotypes. We implemented the SAIGE-QT<sup>72</sup> method, which employs fast linear mixed models with kinship adjustment, in Encore (<a href="https://encore.sph.umich.edu/">https://encore.sph.umich.edu/</a>) for single variant association analyses. We additionally adjusted the model for covariates (PC1-PC11, age, sex, age<sup>2</sup>, and study-groups [cohort-race subgrouping]).

We conducted single variant association replications for putative novel variants. After comparing the results with published lipid GWAS summary statistics, we filtered putative novel GWAS variants based on a stringent whole genomewide significant threshold (alpha = 5x10<sup>-9</sup>)<sup>73</sup>. Replication was performed in the MGB and Penn Medicine Biobanks where models were fitted as indicated above. Additionally, we adjusted the MGB Biobank for study recruitment center and array and Penn Medicine Biobank for ancestry and BMI. In the MGB Biobank, we selected lipid concentrations closest to the sample acquisition time point and adjusted for statins if prescribed within one year prior to sample acquisition. In the Penn Biobank, we utilized each participant's median lipid concentration for replication; statins prescribed prior to lipid concentration used were adjusted in the models. Additionally, we carried out meta-analysis using fixed effects model based on inverse-variance-weighted effect size for the two replication cohorts using METASOFT<sup>74</sup>.

#### Rare variant association test

We performed rare variant association (RVA) using the Variant-Set Test for Association using Annotation infoRmation (STAAR) pipeline<sup>13</sup> from STARtopmed R package. STAARpipeline is a regression-based framework that permits adjustment of covariates, population structure, and relatedness by fitting linear and logistic mixed models for quantitative and dichotomous traits<sup>75</sup>, . We chose STAAR to leverage the annotation information and associated scores that were available for TOPMed Freeze 8 data to incorporate the analysis of rare non-coding variants from whole genome sequencing. The method implements genome-wide scanning of rare variants (MAF<0.01) in gene-centric and regionbased workflows. For each variant set, STAARpipeline calculates a set-based p-value using the STAAR method, which increases the analysis power by incorporating multiple in silico variant functional annotation scores capturing diverse genomic features and biochemical readouts<sup>13</sup>. We aggregated rare variants into multiple groups for coding and noncoding analyses. For the coding region, we defined five different aggregate masks of rare variants 1) plof (putative loss-offunction), plof-Ds (putative loss-of-function or disruptive missense), missense, disruptive-missense, and synonymous. For the non-coding regions, we used seven rare variant masks: 1) promoter-CAGE (promoter variants within Cap Analysis of Gene Expression [CAGE] sites<sup>77,78</sup>), 2) promoter-DHS (promoter variants within DNase hypersensitivity [DHS] sites <sup>79</sup>), 3) enhancer-CAGE (enhancer within CAGE sites<sup>78</sup>), 4) enhancer-DHS (enhancer variants within DHS sites<sup>80</sup>), 5) UTR (rare variants in 3' untranslated region [UTR] and 5' UTR untranslated region), 6) upstream, and 7) downstream. Detailed explanations of the regions defined based on these masks is discussed within STAARpipeline<sup>13</sup>.

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In the gene-centric workflows, for both coding (within exonic boundaries) and non-coding (promoter: +/- 3kb window of transcription starting site (TSS), enhancer: GeneHancer predicted regions) regions, we considered only genes with at least two rare variants (i.e., 18,445 genes in all 22 autosomes). In the region-based workflows, we implemented two protocols: 1) a 'sliding window' approach, where we aggregated rare variants within 2-kb sliding windows and with 1-kb overlap length, and 2) a 'dynamic window' approach, where we executed SCANG<sup>33</sup> method and aggregated dynamically variant-sets between 40-300 variants per set, where the method systematically scans the whole genome with overlapping windows of varying sizes. The STAARtopmed R-package implements multiple rare-variants aggregate tests including SKAT<sup>81</sup>, Burden<sup>82</sup> and ACAT<sup>83</sup> and integrates them as STAAR-O<sup>13</sup>. We performed gene-centric and region-based rare variant tests using annotated GDS files of TOPMed.

We completed aggregate tests as three-step process. In the first step, we fitted a null model using glmmkin() function in STAARtopmed. The null model was fitted for each of the four lipid phenotypes adjusted for all covariates and relatedness except the genotype of interest. In the second step, we ran genome wide gene-centric and region-based rare-variant aggregate tests. The third step directed conditional analyses, where the results were adjusted for previously known significantly lipid-associated (i.e., p < 5x10<sup>-8</sup> in external datasets) individual variants from GWAS Catalog<sup>84</sup> and Million Veterans Program (MVP)<sup>15</sup> GWAS summary statistics. To obtain effect estimates of significant aggregate sets, we associated the cumulative genotypes (binary scores) based on the variants forming the aggregates and used Glmm.Wald

test from GMMAT R package<sup>75</sup>(version 1.3.1). For significantly-associated window-based rare variant aggregations, we trimmed the exonic variants and estimated the effects with only non-coding variants.

# **CETP** gene expression and lipid trait colocalization

We studied the correlation of LDL-C and HDL-C effects with eQTL effects at chromosome 16q13, which includes *CETP*. We downloaded GTEx eQTL build 38 (version8) data for Liver, Adipose Subcutaneous and Adipose Visceral (Omentum) tissues from GTEx Portal on 16/APR/2020<sup>85</sup>. We selected eQTLs with nominal significance (p-value<0.05) and utilized the eQTL-gene pairs with the most significant p-values. Genes with at least 5 eQTLs were selected for the colocalization analysis. We selected variants with a suggestive significance (p-value < 5x10<sup>-7</sup>) for LDL-C or HDL-C effects within 500 kb of the lead locus variant. We performed Pearson correlation tests between the lipid effect estimates and gene expression effects (slope) from GTEX.

# **Acknowledgments**

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### **Competing interests**

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## **Figure Legends**

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Fig1: Overall study schematic. The analyses were conducted using the multi-ancestral TOPMed freeze8 data to associate whole genome sequence variation with lipid phenotypes (i.e., LDL-C, HDL-C, TC and TG). A total of 66,329 samples with lipids quantified data from five ancestry groups were analyzed. Single variant GWAS were carried out using SAIGE on the Encore platform using SNPs with MAC >20. Both trans-ancestry and ancestry-specific GWAS were conducted. Genome-wide rare variant (MAF < 1%) gene-centric and region-based aggregate tests were grouped and analyzed using STAARtopmed. Finally, single variant and rare variant associations at Mendelian dyslipidemia genes were investigated in further detail. TOPMed – Trans-Omics for Precision Medicine; HDL-C – High-Density Lipoprotein Cholesterol; LDL-C – Low-Density Lipoprotein Cholesterol; TC – Total Cholesterol; TG – Triglycerides; GWAS – Genome Wide Association Study; SAIGE – Scalable and Accurate Implementation of GEneralized mixed model; MAC – Minor Allele Count; MAF – Minor Allele Frequency: SNPs – Single nucleotide polymorphisms. Fig2: Summary of single variant genome wide association. Representation of the single variant GWAS results from TOPMed Freeze 8 whole genome sequenced data of 66,329 samples. Each quarter represents a different lipid phenotype, and dots extending in clock-wise fashion represent variants with increasing evidence of association as noted by -log10(p-value), which was truncated at 200. The outer three circles show the GWAS data from TOPMed freeze8 where variants binned to nominally significant (p-value 0.05 - 5x10<sup>-07</sup>), suggestive significant (p-value 5x10<sup>-07</sup> - 5x10<sup>-09</sup>)

and genome wide significant (p-value < 5x10<sup>-09</sup>). The inner three circles compare our TOPMed results with known significantly associated lipid loci and variants from the MVP summary statistics and GWAS catalog to the identified novel variants and loci that are genome-wide significant from the current study, respectively. TOPMed – Trans-Omics for Precision Medicine; GWAS – Genome Wide Association Study; MVP – Million Veteran Program. Fig3: Comparison of effects estimates for HDL-C and LDL-C among variants in the CETP locus. The color scale of the data points was based on -log10 p-values from HDL-C association and the size of each data point was based on log10 p-values of LDL-C association. Variants which are genome wide significant with LDL-C are represented as chromosome:position:reference allele:alternate allele. HDL-C – High-Density Lipoprotein Cholesterol; LDL-C – Low-Density Lipoprotein Cholesterol. Fig4: Conditional analysis of coding rare-variants from the same gene and a near-by gene. Non-coding rare variant sets significantly associated with TC and TG after the conditional analysis on known variants are shown with additional adjustment on rare-coding variants. The additional adjustment for rare-coding variants were carried out for the same gene of the aggregate set and for certain gene aggregates (SPC24) the conditional analysis was carried out with a nearby Mendelian gene. After adjusting for rare-coding variants and known variants, EHD3 signal drops minimally, whereas signal from PCSK9 (promoter-DHS, enhancer-DHS), LDLR-loci (enhancer-DHS, SPC24 enhancer-DHS) enhances significantly. APOB1, SPC24 (enhancer-CAGE), HBB and APOE signal drops after the conditional analysis on rare-coding

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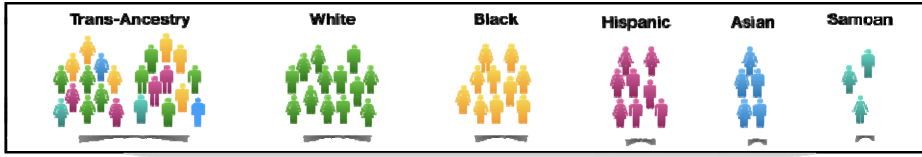
1003 variants. The different colored dots on the plot represents the conditional STAAR-O p-values when adjusting for known 1004 variants (Set1) and rare-coding variants of the same or near-by gene. 1005 STAAR – variant-Set Test for Association using Annotation information; TC – Total Cholesterol; TG – Triglycerides; CAGE 1006 - Cap Analysis of Gene Expression; DHS - DNase hypersensitivity. Fig5: Influence of common and rare variants with hypercholesterolemia. In addition to monogenic contributions from 1007 1008 rare variants in Mendelian hypercholesterolemia genes, multiple genome-wide significant LDL-C-associated common 1009 variants also yield a polygenic basis for hypercholesterolemia. In the present work, we now identify rare non-coding variants in proximity of Mendelian hypercholesterolemia genes, specifically LDLR and PCSK9, that also contribute to the 1010 1011 genetic basis of hypercholesterolemia. 1012 LDL-C - Low-Density Lipoprotein Cholesterol

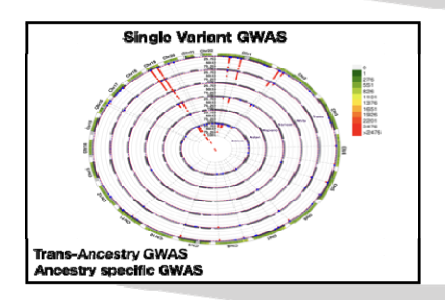
| Associated<br>Lipid<br>Phenotype | Novel<br>variant<br>class | Variants (Gene)                        | TOPMed Freeze8 (N=66329) |                        |        | MGB Biobank (N=25137) |                        |                | Penn Medicine Biobank (N=20079) |                             |                             | Meta Analysis<br>(METASOFT) |                         |
|----------------------------------|---------------------------|--|--------------------------|------------------------|--------|-----------------------|------------------------|----------------|---------------------------------|-----------------------------|-----------------------------|-----------------------------|-------------------------|
|                                  |                           |  | TOPMed                   | TOPMed                 | TOPMed | MGB<br>Biobank        | MGB<br>Biobank         | MGB<br>Biobank | Penn<br>Medicine<br>Biobank     | Penn<br>Medicine<br>Biobank | Penn<br>Medicine<br>Biobank | Beta P-va                   | P-value                 |
|                                  |                           |  | Effect<br>Estimate       | P-value                | MAF    | Effect<br>Estimate    | P-value                | MAF            | Effect<br>Estimate              | P-value                     | MAF                         |                             |                         |
| LDL-C                            | Novel<br>locus            | 12:97352354:T:C                        | -12.439                  | 4.88x10- <sup>09</sup> | 0.003  | 1.055                 | 8.08x10 <sup>-01</sup> | 0.002          | 11.441                          | 3.19x10 <sup>-01</sup>      | 0.001                       | 2.357                       | 5.62 x10 <sup>-01</sup> |
| LDL-C                            | Novel variant             | 16:56957451:C:T<br>( <i>CETP</i> )     | -1.568                   | 2.88x10 <sup>-09</sup> | 0.283  | -1.375                | 1.53x10 <sup>-04</sup> | 0.309          | -2.35                           | 1.54x10 <sup>-04</sup>      | 0.578                       | -1.624                      | 2.21 x10 <sup>-07</sup> |
| LDL-C                            | Novel<br>locus            | 4:176382171:C:T                        | -16.086                  | 2.82x10 <sup>-09</sup> | 0.002  | -13.340               | 1.71x10 <sup>-01</sup> | 0.001          | 4.716                           | 3.52x10 <sup>-01</sup>      | 0.005                       | 0.882                       | 8.44 x10 <sup>-01</sup> |
| TC                               | Novel variant             | 13:113841051:T:C<br>( <i>GAS6</i> )    | 1.731                    | 1.12x10 <sup>-09</sup> | 0.278  | 0.890                 | 3.94x10 <sup>-02</sup> | 0.304          | 0.416                           | 5.50x10 <sup>-01</sup>      | 0.563                       | 0.758                       | 3.89 x10 <sup>-02</sup> |
| тс                               | Novel<br>variant          | 7:137875053:T:C<br>( <i>CREB3L2</i> )  | -4.106                   | 7.54x10 <sup>-11</sup> | 0.045  | -4.755                | 1.06x10 <sup>-02</sup> | 0.013          | -3.365                          | 7.62x10 <sup>-03</sup>      | 0.118                       | -3.803                      | 2.69 x10 <sup>-04</sup> |
| TG                               | Novel<br>locus            | 11:69219641:C:T                        | 0.232                    | 1.98×10 <sup>-09</sup> | 0.002  | -0.047                | 7.33x10 <sup>-01</sup> | 0.000          | 0.202                           | 7.82x10 <sup>-02</sup>      | 0.001                       | 0.101                       | 2.53 x10 <sup>-01</sup> |
| TG                               | Novel<br>variant          | 13:107551611:C:T<br>( <i>FAM155A</i> ) | 0.052                    | 6.78x10 <sup>-10</sup> | 0.045  | 0.016                 | 4.68x10 <sup>-01</sup> | 0.014          | 0.039                           | 3.26x10 <sup>-01</sup>      | 0.016                       | 0.021                       | 2.62 x10 <sup>-01</sup> |

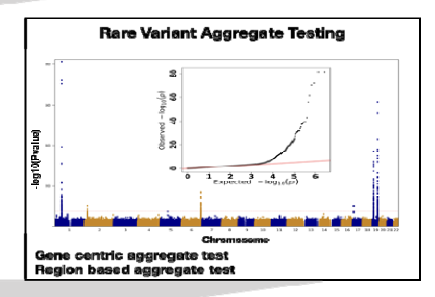
bioRxiv preprint doi: https://doi.org/10.1101/2021.10.11.463514; this version posted October 12, 2021. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license.

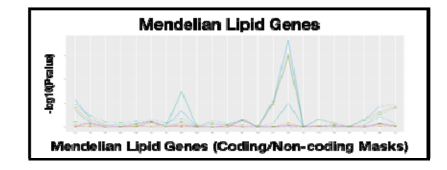
Table 1. Putative novel variants identified in TOPMed and evidence for replication. Variants identified as novel after comparing with the GWAS catalog and MVP summary statistics for associations with lipid phenotypes, including LDL-C, TC, and TG. All effect estimates are in mg/dL units, except for TG which was log-transformed in analysis thereby representing fractional change. Variants are categorized as novel loci or novel variant (i.e., known locus associated with another lipid phenotype) and the genes assigned to the variants per TOPMed whole genome sequence annotations (WGSA) are listed. Data is provided for the discovery (TOPMed freeze8) and replication cohorts (MGB Biobank and Penn Medicine Biobank). Meta-analysis with the replication cohorts was carried out and the corresponding beta and p-values are provided.

GWAS – Genome Wide Association Study; MVP – Million Veteran Program; LDL-C – Low-Density Lipoprotein Cholesterol; TC – Total Cholesterol; TG – Triglycerides; TOPMed – Trans-Omics for Precision Medicine; WGSA – Whole Genome Sequence Annotations.









Overall study schematic. The analyses were conducted using the multi-ancestral TOPMed freeze8 data to associate whole genome sequence variation with lipid phenotypes (i.e., LDL-C, HDL-C, TC and TG). A total of 66,329 samples with lipids quantified data from five ancestry groups were analyzed. Single variant GWAS were carried out using SAIGE on the Encore platform using SNPs with MAC >20. Both trans-ancestry and ancestry-specific GWAS were conducted. Genomewide rare variant (MAF < 1%) gene-centric and region-based aggregate tests were grouped and analyzed using STAARtopmed. Finally, single variant and rare variant associations at Mendelian dyslipidemia genes were investigated in further detail. TOPMed – Trans-Omics for Precision Medicine; HDL-C – High-Density Lipoprotein Cholesterol; LDL-C – Low-Density Lipoprotein Cholesterol; TC – Total Cholesterol; TG – Triglycerides; GWAS – Genome Wide Association Study; SAIGE – Scalable and Accurate Implementation of GEneralized mixed model; MAC – Minor Allele Count; MAF – Minor Allele Frequency; SNPs – Single nucleotide polymorphisms.

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Fig. 1

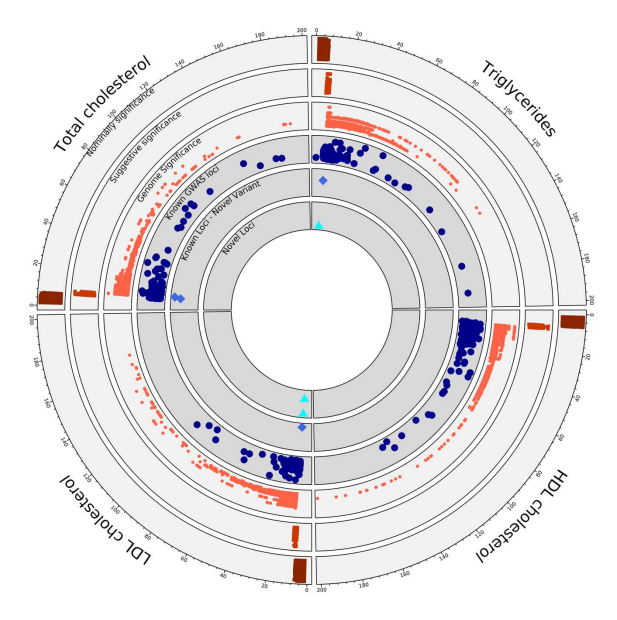


Fig. 2 Summary of single variant genome wide association. Representation of the single variant GWAS results from TOPMed Freeze 8 whole genome sequenced data of 66,329 samples. Each quarter represents a different lipid phenotype, and dots extending in clock-wise fashion represent variants with increasing evidence of association as noted by -log10(p-value), which was truncated at 200. The outer three circles show the GWAS data from TOPMed freeze8 where variants binned to nominally significant (p-value 0.05 - 5x10<sup>-07</sup>), suggestive significant (p-value 5x10<sup>-07</sup> - 5x10<sup>-09</sup>) and genome wide significant (p-value < 5x10<sup>-09</sup>). The inner three circles compare our TOPMed results with known significantly associated lipid loci and variants from the MVP summary statistics and GWAS catalog to the identified novel variants and loci that are genome-wide significant from the current study, respectively. TOPMed – Trans-Omics for Precision Medicine: GWAS – Genome Wide Association Study: MVP – Million Veteran Program.

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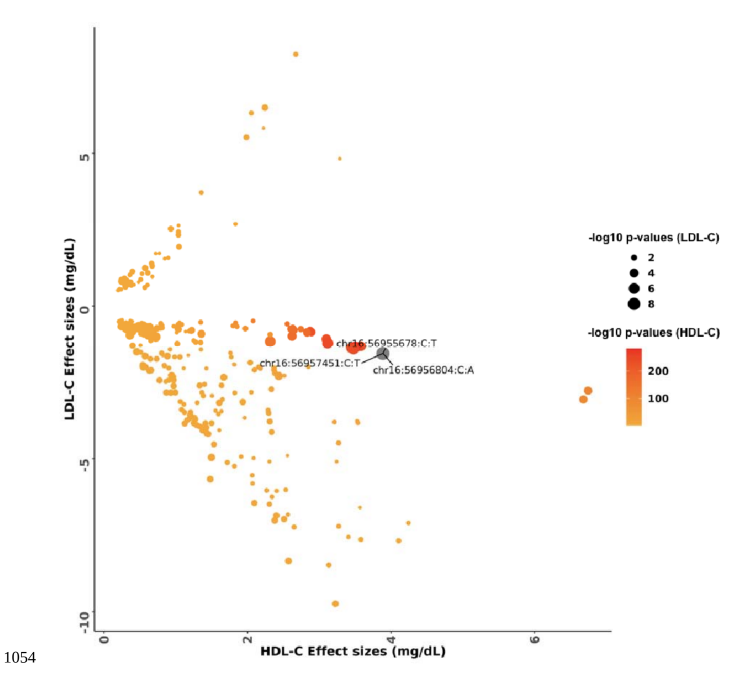
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| 1055<br>1056<br>1057 | Fig. 3   |  |  |  |  |  |  |  |  |
|----------------------|--|--|--|--|--|--|--|--|--|
|                      | Comparison of effects estimates for HDL-C and LDL-C among variants in the CETP locus. The color scale of the           |  |  |  |  |  |  |  |  |
| 1058                 | data points was based on -log10 p-values from HDL-C association and the size of each data point was based on -log10 p- |  |  |  |  |  |  |  |  |
| 1059                 | values of LDL-C association. Variants which are genome wide significant with LDL-C are represented as                  |  |  |  |  |  |  |  |  |
| 1060                 | chromosome:position:reference allele:alternate allele.   |  |  |  |  |  |  |  |  |
| 1061                 | HDL-C – High-Density Lipoprotein Cholesterol; LDL-C – Low-Density Lipoprotein Cholesterol.                             |  |  |  |  |  |  |  |  |

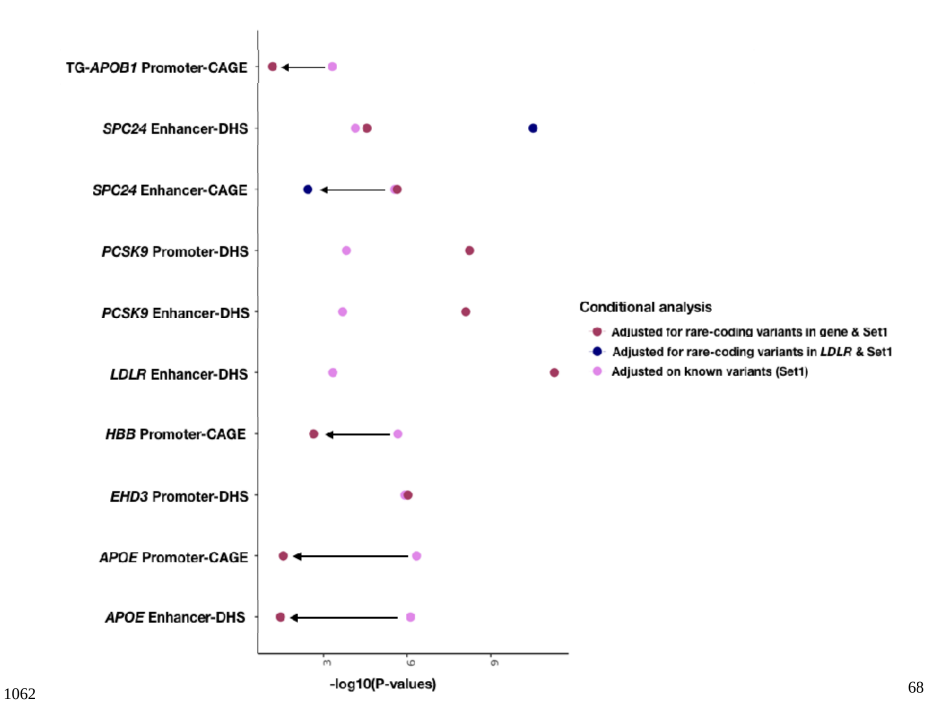


Fig. 4

Conditional analysis of coding rare-variants from the same gene and a near-by gene. Non-coding rare variant sets significantly associated with TC and TG after the conditional analysis on known variants are shown with additional adjustment on rare-coding variants. The additional adjustment for rare-coding variants were carried out for the same gene of the aggregate set and for certain gene aggregates (*SPC24*) the conditional analysis was carried out with a nearby Mendelian gene. After adjusting for rare-coding variants and known variants, *EHD3* signal drops minimally, whereas signal from *PCSK9* (promoter-DHS, enhancer-DHS), *LDLR*-loci (enhancer-DHS, *SPC24* enhancer-DHS) enhances significantly. *APOB1*, *SPC24* (enhancer-CAGE), *HBB* and *APOE* signal drops after the conditional analysis on rare-coding variants. The different colored dots on the plot represents the conditional STAAR-O p-values when adjusting for known variants (Set1) and rare-coding variants of the same or near-by gene.

STAAR – variant-Set Test for Association using Annotation information; TC – Total Cholesterol; TG – Triglycerides; CAGE – Cap Analysis of Gene Expression; DHS – DNase hypersensitivity.

# Hypercholesterolemia

Influence from monogenic rare coding variants

*LDLR* 

**APÓB** 

PCSK9

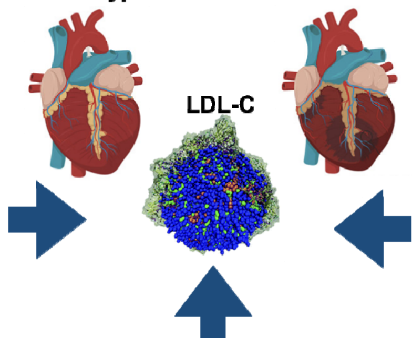
ABCG5

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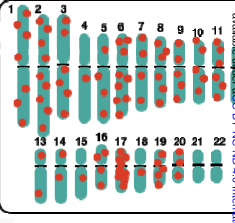
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**APOE** 

NPC1L1

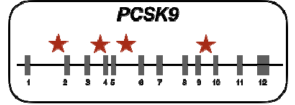


Influence from genome-wide polygenic common variants



# Influence from monogenic rare non-coding variants





Influence of common and rare variants with hypercholesterolemia. In addition to monogenic contributions from rare variants in Mendelian hypercholesterolemia genes, multiple genome-wide significant LDL-C-associated common variants also yield a polygenic basis for hypercholesterolemia. In the present work, we now identify rare non-coding variants in proximity of Mendelian hypercholesterolemia genes, specifically *LDLR* and *PCSK9*, that also contribute to the genetic basis of hypercholesterolemia.

LDL-C – Low-Density Lipoprotein Cholesterol