Identification of a Gain-of-Function LIPC Variant as a Novel Cause of Familial Combined Hypocholesterolemia

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BACKGROUND: Atherosclerotic cardiovascular disease is the main cause of mortality worldwide and is strongly influenced by circulating low-density lipoprotein (LDL) cholesterol levels. Only a few genes causally related to plasma LDL cholesterol levels have been identified so far, and only 1 gene, ANGPTL3, has been causally related to combined hypocholesterolemia. Here, our aim was to elucidate the genetic origin of an unexplained combined hypocholesterolemia inherited in 4 generations of a French family.

METHODS: Using next-generation sequencing, we identified a novel dominant rare variant in the LIPC gene, encoding for hepatic lipase, which cosegregates with the phenotype. We characterized the impact of this LIPC-E97G variant on circulating lipid and lipoprotein levels in family members using nuclear magnetic resonance–based lipoprotein profiling and lipidomics. To uncover the mechanisms underlying the combined hypocholesterolemia, we used protein homology modeling, measured triglyceride lipase and phospholipase activities in cell culture, and studied the phenotype of APOE*3.Leiden.CETP mice after LIPC-E97G overexpression.

RESULTS: Family members carrying the LIPC-E97G variant had very low circulating levels of LDL cholesterol and high-density lipoprotein cholesterol, LDL particle numbers, and phospholipids. The lysophospholipids/phospholipids ratio was increased in plasma of LIPC-E97G carriers, suggestive of an increased lipolytic activity on phospholipids. In vitro and in vivo studies confirmed that the LIPC-E97G variant specifically increases the phospholipase activity of hepatic lipase through modification of an evolutionarily conserved motif that determines substrate access to the hepatic lipase catalytic site. Mice overexpressing human LIPC-E97G recapitulated the combined hypocholesterolemic phenotype of the family and demonstrated that the increased phospholipase activity promotes catabolism of triglyceride-rich lipoproteins by different extrahepatic tissues but not the liver.

CONCLUSIONS: We identified and characterized a novel rare variant in the LIPC gene in a family who presents with dominant familial combined hypocholesterolemia. This gain-of-function variant makes LIPC the second identified gene, after ANGPTL3, causally involved in familial combined hypocholesterolemia. Our mechanistic data highlight the critical role of hepatic lipase phospholipase activity in LDL cholesterol homeostasis and suggest a new LDL clearance mechanism.

Key Words: cholesterol, HDL ■ cholesterol, LDL ■ LIPC protein, human ■ LIPC protein, mouse ■ phospholipases

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Dijk et al Familial Hypocholesterolemia by a LIPC Variant

Clinical Perspective

What Is New?

- The first gain-of-function variant in the LIPC gene (LIPC-E97G) encoding hepatic lipase in a family with combined hypocholesterolemia (low low-density lipoprotein and low high-density lipoprotein cholesterol) was identified.
- This unique LIPC-E97G variant specifically increases the phospholipase activity of hepatic lipase without affecting triglyceride lipase activity.
- The hypocholesterolemic phenotype related to LIPC-E97G variant is attributable to an increased clearance of cholesterol within triglyceride-rich lipoprotein remnants predominantly by extrahepatic tissues.

What Are the Clinical Implications?

- This novel gain-of-function variant in LIPC potentially represents the second cause of familial combined hypocholesterolemia, after loss-of-function variants in ANGPTL3.
- This study highlights an unexpected and critical role of the phospholipase activity of hepatic lipase (encoded by LIPC) in low-density lipoprotein cholesterol metabolism and identifies it as a potential novel drug target.
- Additional data are warranted to clarify the impact of LIPC-E97G–related combined hypocholesterolemia on atherosclerosis and atherosclerotic cardiovascular disease, due to the occurrence of documented coronary stenosis and evolutive carotid atherosclerosis in the index case.

The development of atherosclerotic cardiovascular disease (ASCVD), the main cause of mortality worldwide, is strongly influenced by circulating lipoprotein levels. Elevated low-density lipoprotein (LDL) cholesterol (LDL-C) is a well-established causal risk factor for ASCVD, whereas the role of high-density lipoprotein cholesterol (HDL-C) in ASCVD development remains to be further characterized. Plasma LDL-C concentrations are strongly affected by genetics, with heritability explaining ≈ 40% to 60% of plasma levels. Genetically modulated LDL-C concentrations typically have a polygenic origin but can also be caused by a codominant disease attributable to biallelic or monoallelic pathogenic variants. So far, however, only a few genes with a strong causal relationship to plasma LDL-C levels have been identified.

Even fewer monogenic causes of low LDL-C, familial hypobetalipoproteinemia (FHBL), have been identified. These monogenic disorders were recently classified into 2 classes based on molecular mechanisms: FHBL attributable to lipoprotein secretion defects (FHBL-SD1, -SD2, and -SD3 for MTPP, APOB, and SAR1B-related FHBL) or to enhanced catabolism of apolipoprotein B (apoB)–containing lipoprotein (FHBL-EC1 and -EC2 for ANGPTL3- and PCSK9-related FHBL). Furthermore, pathogenic variants in the ANGPTL3 gene have been identified in cases of familial combined hypolipidemia, a condition characterized by very low concentrations of circulating very-low-density lipoprotein (VLDL), LDL, and HDL. Despite the suspected polygenic origins of unexplained cases of extremely elevated or reduced LDL-C concentrations, polygenic risk scores have so far yielded lower effect sizes compared with the effect of monogenic variants. This suggests that additional monogenic variants that cause very low plasma LDL-C concentrations or combined hypolipidemia remain to be characterized.

Knowledge of the molecular mechanisms and genes implicated in extreme lipoprotein phenotypes is of direct clinical relevance. Accordingly, many of the recent therapeutics developed to lower plasma LDL-C levels and that provided breakthroughs in ASCVD risk reduction involved the products of genes implicated in FHBL and familial combined hypolipidemia such as PCSK9 and ANGPTL3. Despite the effectiveness of these new therapies in lowering LDL-C, a need to identify additional mechanisms to effectively lower plasma LDL-C remains.

To identify new causative genes in LDL-C metabolism, we have built up large cohorts of patients with extreme

Nonstandard Abbreviations and Acronyms

- apoB: apolipoprotein B
- ASCVD: atherosclerotic cardiovascular disease
- CHOPIN: Cholesterol Personalized Innovation
- CO: cholesteryl oleate
- eGFP: enhanced green fluorescent protein
- EL: endothelial lipase
- FHBL: familial hypobetalipoproteinemia
- GENELIP: From Known to New Genes in Dyslipidemia
- HYPOCHOL: A Genetically-Based Strategy to Identify New Targets in Cholesterol Metabolism
- HDL: high-density lipoprotein
- HDL-C: high-density lipoprotein cholesterol
- HL: hepatic lipase
- LDL: low-density lipoprotein
- LDL-C: low-density lipoprotein cholesterol
- LIPC: lipase C, hepatic type (hepatic lipase)
- LIPC-WT: wild-type human LIPC
- LPL: lipoprotein lipase
- TBG: thyroxin-binding globulin
- TGRL: triglyceride-rich lipoprotein
- VLDL: very-low-density lipoprotein

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LDL-C phenotypes as part of the CHOPIN (Cholesterol Personalized Innovation) French national research program. In the present study, we identified and characterized a novel rare variant in the LIPC gene, encoding for hepatic lipase (HL), in a family who presents a dominant familial combined hypcholesterolemia. This first gain-of-function variant identifies LIPC as the second gene, after ANGPTL3, causally involved in familial combined hypcholesterolemia.

METHODS

Extended methods are available in the online Supplemental Material. The data, methods, and study materials used to conduct the research are available from the corresponding author on reasonable request.

Family Studies

The proband and his relatives were recruited in Lyon to investigate the genetic cause of a familial combined hypolipidemia (GENELIP study [From Known to New Genes in Dyslipidemia]; ClinicalTrials.com registration No. NCT03939039). GENELIP aims to decipher the mechanisms involved in the occurrence or modulation of dyslipidemia in patients referred for primary dyslipidemia. After the identification of 1 potential candidate gene and a confirmative cosegregation analysis, the members of the pedigree were recruited to the HYPOCHOL study (A Genetically-Based Strategy to Identify New Targets in Cholesterol Metabolism; ClinicalTrials.com registration No. NCT02354079) of the CHOPIN program for functional studies. Written informed consent was obtained from all participants in accordance with the principles of the Declaration of Helsinki and the French bioethics law. GENELIP and HYPOCHOL protocols were approved by the University Hospital Center of Lyon (France) and Nantes (CHU Nantes, France), respectively. The GENELIP study also obtained the agreement of the ethical committee of the Commission Nationale de l’Informatique et des Libertés (No. 920434).

Targeted Next-Generation Sequencing

After genomic DNA extraction, DNA from the proband and patient IV.2 was sequenced and analyzed as previously detailed with the DysliSEQ custom design.10 This panel includes coding exons and intron/exon junctions of 311 genes selected from published literature: (1) genes identified in monogenic dyslipidemia, (2) genes identified in genome-wide association studies in lipid metabolism through direct or indirect effects, (3) genes associated with dyslipidemia in mice, and (4) single-nucleotide polymorphisms already described in familial hypercholesterolemia genetic risk scores and in genome-wide association studies (with \( P < 5 \times 10^{-8} \)). Among these genes, a first intention panel was defined for FHBL (APOB, PCSK9, ANGPTL3, abetalipoproteinemia (ABL), OMIM 200100; MTTP), and chylomicron retention disease (CMRD, OMIM No. 246700; SAR1B). In the absence of a variant in the first intention panel, relevant variants were selected as previously described10 (Table S1.3). The only relevant variant common to the proband and his granddaughter and that cosegregated with the combined hypcholesterolemia observed in the family after Sanger sequencing was the E97G variant in the LIPC gene. In addition, the patients exhibited a combined hypolipidemia and a decrease of phospholipid concentration, as observed in mice after LIPC overexpression.11

Mouse Studies

All mouse studies were performed with APOE*3-Leiden. CETP mice that have a humanized plasma lipid profile12 and that were bred at Leiden University Medical Centre, Leiden, the Netherlands. All mice experiments were approved by the ethics committee of Pays de la Loire (France, 006) and the Ministère de l'enseignement supérieur et de l'innovation (France; APAFIS 26862) or the Central Committee on Animal Experimentation of the Netherlands (AVD11600202010187) and Animal Welfare Body of the Leiden University Medical Center and conducted in accordance with institutional guidelines. At \( \approx 10 \) weeks of age, male mice were placed on a diet containing 0.5% cholesterol and 15% cocoa butter (Ssniff, No. S8854-E035 EF 4021-04T) for 3 to 5 weeks. Mice were injected intravenously with \( 3 \times 10^{10} \) or \( 3 \times 10^{11} \) genome copies of adeno-associated viruses (AAV8) under the thyroxin-binding globulin (TBG) promoter and containing enhanced green fluorescent protein (eGFP), wild-type lipase C, hepatic type (LIPC), or LIPC-E97G (Vector Biolabs). Mice were monitored weekly for changes in plasma cholesterol and triglyceride levels and body weight. Four weeks after injection, experiments to phenotype characterize the mice were started as described in the Supplemental Material, each with at least a 1-week interval.

Statistical Analyses

For mice studies, a 1-way ANOVA with Tukey correction for multiple comparisons was used, with a 2-sided \( P \) value cutoff set at \( P < 0.05 \). For cell culture studies, a nonparametric Mann-Whitney test was used with a 2-sided \( P \) value cutoff set at \( P < 0.05 \).

RESULTS

A Novel Variant in LIPC Cosegregates With Familial Combined Hypcholesterolemia

A 61-year-old patient was admitted to the lipid clinic of the University Hospital in Lyon (France) to explore a combined hypercholesterolemia inherited as a dominant phenotype over 4 generations. He was referred for low LDL-C before statin treatment while presenting with coronary artery disease requiring a first coronary stenting. He needed subsequent iterative coronary stentings at 62 and 73 years of age. In addition, he had evolving silent carotid artery plaques, which progressed from 7% (NASCET [North American Symptomatic Carotid Endarterectomy Trial]) with an increased intima-media thickness (1.23 mm) when he was 61 years of age to 75% when he was 73 years of age. He was a former heavy smoker between 18 and 36 years of age and had well-controlled type 2 diabetes since 53 years of age. He had a body mass index of 25.3 kg/m\(^2\) and android adiposity. A phenotypic characterization of the
patient (patient II.8, Figure 1) revealed extremely low values of circulating cholesterol, LDL-C, HDL-C, phospholipids, and apolipoprotein A1 levels (below the fifth percentile for age and sex; Figure 1 and Table S1.1). In contrast, circulating triglycerides and apoB levels were within the low but normal range. Liver transaminases were mildly increased (aspartate aminotransferase, 54 UI/L; alanine aminotransferase, 54 UI/L when he was 71 years of age), but the fibrosis-4 indexes (a liver fibrosis score) were within the normal interval range (fibrosis-4 index, 1.5 and 1.0 when he was 62 and 74 years of age, respectively), suggesting the absence of liver fibrosis. To clarify the origin of the hypocholesterolemic phenotype, family members of the patient were recruited for plasma and DNA collection. Plasma lipid analyses revealed that a combined hypocholesterolemia below the fifth percentile for age and sex was present in multiple family members and followed an autosomal dominant mode of inheritance (Figure 1 and Table S1.1). More detailed nuclear magnetic resonance lipoprotein profiling of 3 affected family members (III.9, IV.4, IV.5) revealed profound changes in lipoprotein composition and sizes compared with age- and sex-matched individuals (Figure 2 and Table S1.2). Whereas VLDL particle numbers of affected family members were within the normal range, VLDL particle size was reduced compared with control subjects (Figure 2). In contrast, LDL particle size was normal, but LDL particle numbers were low in affected family members, concordant with circulating apoB levels, which were within the lower range in these individuals (Figure 2 and Table S1.1). Furthermore, we found an elevated number of large HDL particles and a near absence of small and medium HDL particles in affected family members, re-

Figure 1. A novel variant in \textit{LIPC} cosegregates with familial combined hypocholesterolemia. Pedigree of family with familial combined hypocholesterolemia. Squares indicate male family members; circles, female family members. Slashes indicate deceased individuals. Roman numerals to the left of the pedigree indicate the generation; numerals to the upper left of each symbol indicate the individual family member. Basic lipid parameters of the recruited family members are indicated in the table below the pedigree. Values of total cholesterol, triglyceride, and low-density lipoprotein cholesterol (LDL-C), and high-density lipoprotein cholesterol (HDL-C) levels below the fifth percentile for age and sex are in bold.
resulting in an elevated average HDL particle size (Figure 2). These large HDL particles are relatively triglyceride enriched, as indicated by the low HDL-C in these individuals and the elevated HDL-triglycerides/HDL-C ratio (Figure 2). In terms of ASCVD, only the proband (II.8) had ASCVD, with no knowledge of clinical ASCVD in the other family members.

To elucidate the genetic origin of the hypocholesterolemia in the family, DNA from the proband was sequenced with the DysliSEQ custom next-generation sequencing panel design based on ontology of lipid disorders. No rare single nucleotide or copy number variation was found in coding exons of genes involved in FHBL (ANGPTL3, APOB, MTTP, PCSK9, SAR1B), and the polygenic risk score (0.91, 49th percentile) did not support a polygenic hypobetalipoproteinemia. Our rare variant filtering criteria (Table S1.3) enabled the detection of a rare missense variant, P(Glu97Gly) or E97G, in the LIPC gene. The LIPC gene encodes for HL, a protein known to be involved in lipid metabolism. The LIPC-E97G variant is not reported in any large genetic data set (gnomAD and >300,000 individuals of the UK Biobank database) and cosegregated with the combined hypocholesterolemia in the family (Figure 1). To further confirm the absence of another candidate gene, we performed a whole-genome sequencing analysis of all recruited family members, leading to the identification of 3 variants that cosegregate with the hypocholesterolemic phenotype (Table 1). Of these variants, the LIPC-E97G variant was predicted to be the most conserved (Genomic Evolutionary Rate Profiling score, 4.6) and the most deleterious by 9 prediction algorithms (Table S1.4). This result confirmed that
the combined hypocholesterolemia might be related to the LIPC-E97G variant.

### LIPC-E97G Modifies the Structural Conformation of HL

HL is part of a family of glycerol-sn-1-fatty acid hydrolases that includes lipoprotein lipase (LPL) and endothelial lipase (EL).\(^{13}\) HL has intermediate triglyceride lipase and phospholipase activities and hydrolyses triglycerides and phospholipids on circulating lipoproteins such as HDL, intermediate-density lipoprotein, and chy-10mic remnants.\(^{14}\) Besides its lipolytic actions, HL also functions as a ligand to facilitate the uptake of lipoproteins by heparan sulfate proteoglycans or other cell surface receptors.\(^{15}\) To determine how the E97G variant might affect HL functionality, we performed an in silico analysis using homology modeling. A homology model for HL was created from a recently established crystal structure of LPL (6O80,\(^{16}\) Figure S1). Using this model, we observed that E97 is not part of the catalytic triad (S168, D194, and H279) but lies in relatively close proximity to the triad owing to protein folding (Figure 3A).

Furthermore, we observed the existence of a salt bridge between E97 and K276, an amino acid present in the lid region of HL (residues 244–277; Figure 3B). This lid region has been shown to determine substrate access to the catalytic site.\(^{17}\) The conformation of the lateral chain of K276 was found to be similar to the corresponding lateral chain in LPL (Lys265, lid region from residues 233–266), indicating a conservation of this particular motif between the 2 lipases (Figure S1). The importance of this structural motif is underlined by the observation that K265 is hydrogen bonded to the inhibitor present in the catalytic site of an experimental structure of LPL (Figure S1).

We next virtually mutated E97G in the HL homology model and optimized the geometry/energy of the resulting structure. The E97G variant significantly changes the conformation of K276 (Figure 3C). Instead of with E97, K276 is now involved in a salt bridge with the carboxylate group of D280 [d(Hz3...OE1)=1.80 Å] in the mutant model. These results show that E97 and K276 might be important for structural features of the catalytic site. By significantly affecting the conformation of K276, E97G alters the HL lid region and might consequently modify the substrate access to the HL catalytic site.

### LIPC-E97G Alters HL Substrate Specificity

We next set out to determine the impact of the E97G variant on the HL triglyceride lipase and phospholipase activities. We overexpressed wild-type human LIPC (LIPC-WT), LIPC-E97G, or LIPC-S168G, a catalytically inactive mutant, in immortalized human hepatocytes and determined triglyceride lipase and phospholipase activities in the medium after treatment with heparin, which releases HL from cell surface heparan sulfate proteoglycan binding sites (Figure S2A). Triglyceride lipase and phospholipase activities of the released HL were identical between mock-treated and LIPC-WT–overexpressing cells when corrected for the amount of released HL (Figure 4A and 4B) but were nearly absent in the LIPC-S168G–overexpressing cells. A striking finding was that cells overexpressing LIPC-E97G had a moderately reduced triglyceride lipase activity level but a 4-fold increased phospholipase activity level (\(P<0.01\); Figure 4A and 4B). To investigate whether the heterozygous presence of E97G, as found in our family, similarly affects triglyceride lipase and phospholipase activities, we introduced the E97G variant into immortalized human hepatocytes using CRISPR-Cas9 and recorrected the variant to the wild-type allele (Figure S2B). After heparin treatment, the heterozygous presence of E97G did not affect triglyceride lipase activity in the medium compared with wild-type cells (Figure 4C and 4D). However, we observed a 7-fold increase in HL phospholipase activity (\(P<0.01\)) with heterozygous E97G presence (Figure 4C and 4D). This effect was completely reversed when E97G was corrected to the wild-type allele (Figure 4C and 4D). These data suggest that the E97G variant specifically increases HL phospholipase activity without increasing triglyceride lipase activity.

To confirm a change in HL substrate specificity in carriers of the E97G variant, we carried out detailed lipidomics on plasma of family members (III.9, IV.4, IV.5) and compared these data with data from normolipidemic individuals (Table S1.5). HL has phospholipase A1 activity and hydrolyzes phospholipids at the SN-1 position to form a fatty acid and a lysophospholipid. In agreement with increased HL phospholipase activity in carriers of the E97G variant, plasma lipidomics data of the E97G carriers showed a clear reduction in overall fatty acids compared with control subjects, with a moderate reduction in fatty acid/apoB ratios (Figure 4E and Table S1.5). Polyunsaturated fatty acids were significantly more

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**Table 1. Genetic Variants Identified With Whole-Genome Sequencing That Cosegregate With the Hypcholesterolemic Phenotype of our Family**

| Chr  | Start   | Ref | Alt | ID         | Gene | Consequences |
|------|---------|-----|-----|------------|------|--------------|
| chr15| 58834000| A   | G   | NA         | LIPC | NM_000236:exon3:c.A290G:p.E97G |
| chr19| 36223172| G   | C   | rs370189638| KMT2B| NM_014727:exon2:c.G5722C:p.A1908P |
| chr19| 42814166| G   | A   | rs143542522| PRR19| NM_199285:exon2:c.G430A:p.V144M |

Alt indicates alternative; Chr, chromosome; ID, identifier; NA, not assigned; and Ref, reference.
**Figure 3. E97G modifies the structural conformation of HL.**

A. Global structure of the homology model of hepatic lipase (HL). The N and C termini are on the **top left** and **right**, respectively. Dashed circle indicates the zone of particular interest in this study, with the location of the E97 and K276 residues and of the amino acids of the catalytic triad (S168, D194, and H279).

B. Closer view of the structural motif (salt bridge and hydrogen bond) involving E97, W99, and K276. Residues of the lid domain are shown in blue.

C. Detailed view of the surroundings of the K276 residue in the E97G mutant model (orange) and the HL homology model (purple). The name of some residues is indicated for clarity. Dashed lines represent the interactions (salt bridge, hydrogen bonds) between the various residues. Residues of the lid domain are shown in blue.
reduced than monounsaturated fatty acids, in agreement with the literature on in vitro HL substrate preference (Figure 4F). A more striking finding was that phospholipid concentrations were significantly reduced in plasma of E97G carriers, and a concomitant 3- to 5-fold increase in lysophospholipids/phospholipids ratios was observed (Figure 4E and 4G). Furthermore, an increased ratio of lysophospholipids and a decreased ratio of phospholipids versus circulating apoB levels were found in E97G carriers compared with control subjects, suggesting significant changes in the composition of circulating apoB-containing lipoprotein particles (Figure S2C). Together, these data suggest that the E97G variant alters substrate specificity by significantly increasing HL phospholipase activity levels.

**Overexpression of the Human LIPC-E97G Variant Markedly Lowers LDL-C and HDL-C in APOE*3.Leiden.CETP Mice**

To determine how the increased phospholipase activity of the E97G variant causes combined hypercholesterolemia, we overexpressed LIPC-WT, LIPC-E97G, and an eGFP control in hepatocytes of the humanized...
Functional Consequences of the Overexpression of the LIPC-E97G Variant

We first hypothesized that the reduced LDL-C and HDL-C levels associated with the LIPC-E97G variant were caused by an increased hepatic cholesterol clearance such as an increased hepatic cholesteryl ester uptake or increased hepatic receptor–mediated endocytosis of circulating lipoproteins as a result of hydrolysis of phospholipids on the lipoprotein surface.19,20 A moderate reduction in the hepatic expression of 3-hydroxy-3-methylglutaryl-CoA reductase Hmgcr, the rate-limiting enzyme in cholesterol synthesis (Figure S4), was found. However, no significant enrichment of either hepatic triacylglycerides or free or esterified cholesterol was observed in the livers of mice overexpressing the E97G variant (Figure S4 and Table S2.2). Consistently, after an intravenous injection of D7-labeled cholesterol 3 days before death, we found no increase in hepatic D7-cholesterol and fecal D7-cholesterol excretion in LIPC-E97G–high–overexpressing mice (Figure 5G and 5H). Given that D7-cholesterol labels were cleared more rapidly from the blood of LIPC-E97G–high–overexpressing mice, this suggests that hepatic catabolism and reverse cholesterol transport are unlikely to be the main cause of the reduced plasma cholesterol levels (Figure 5I). In addition, no impact of the overexpression of LIPC-E97G on VLDL secretion rates compared with overexpression of LIPC-WT or eGFP was found (Figure S4C).

To more directly investigate the impact of LIPC-E97G on lipoprotein kinetics, we overexpressed high doses of eGFP, LIPC-WT, or LIPC-E97G in APOE*3-Leiden.CETP mice and injected these mice with VLDL-like particles or murine VLDL particles labeled with hydrolysable glycerol tri[3H]oleate ([3H]triolein) and [14C]cholesteryl oleate (CO), which is not hydrolysable by HL and is a tracer of triglyceride-rich lipoprotein (TGRL) core remnant particle uptake. Plasma decay of radiolabeled VLDL-like particles was followed, and mice were killed 10 minutes (VLDL-like particles) or 15 minutes (murine VLDL) after injection to determine tissue distribution of the radiolabels (Figure 6A and 6B). No major impact of LIPC-WT overexpression was found on [3H]triolein and [14C]CO decay, whereas [3H]triolein and [14C]CO plasma levels were cleared moderately faster in LIPC-E97G– compared with LIPC-WT–overexpressing mice when injected with VLDL-like particles but not when injected with murine VLDL (Figure 6A–6D). More remarkable differences were observed with a detailed analysis of tissue distributions. These data indicated that [3H]triolein uptake was significantly and [14C]CO uptake was moderately reduced in the liver after clearance of both types of particles (Figure 6E and 6F). Instead, [3H]triolein and [14C]CO labels were increasingly taken up by different adipose tissue depots and oxidative tissues (Figure 6G–6I). These data suggest that the LIPC-E97G variant is responsible for the combined hypocholesterolemic phenotype observed in our family by promoting the uptake of cholesterol-containing remnants by extrahepatic tissues. In support, a nonsignificant increase in cholesterol levels was observed in the muscle and gonadal white adipose tissue of mice overexpressing LIPC-E97G (Figure S5). It should be noted that these effects are likely not mediated through an apoB-mediated pathway because no significant differences in the tissue clearance profile were observed between the clearance of VLDL-like particles (without apoB) and endogenous murine VLDL (with apoB).
Figure 5. Overexpression of the LIPC-E97G variant markedly lowers LDL-C and HDL-C in APOE*3.Leiden.CETP mice. 
A. Experimental setup of mice study overexpressing AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G in APOE*3.Leiden.CETP mice. 
B. Human hepatic lipase (HL) levels as determined by ELISA in preheparin and postheparin plasma of mice overexpressing low or high doses of AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G. 
C and D, Triglyceride (TG) lipase activity and phospholipase A1 (PLA1) activity in plasma of APOE*3.Leiden.CETP mice overexpressing low or high doses of AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G and after heparin injection. 
E, Triglycerides (TG), phospholipids, and cholesterol concentrations in plasma of APOE*3.Leiden.CETP mice overexpressing low or high doses of AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G. 
F, Cholesterol and triglyceride (TG) concentrations in fast protein liquid chromatography (FPLC)-separated pooled plasma of APOE*3.Leiden.CETP mice overexpressing high doses of AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G. 
G through I, Cholesterol (CHOL) and D7-cholesterol (D7-CHOL) levels extracted from liver (G), feces (H), and plasma (I) of APOE*3.Leiden.CETP mice overexpressing high doses of AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G and injected with D7-cholesterol 3 days before death. n=5 or 6 per group. A 1-way ANOVA with Tukey correction for multiple comparisons was used for statistical analysis, with a P value cutoff at P<0.05. eGFP indicates enhanced green fluorescent protein; and LIPC, lipase C, hepatic type. *P<0.05; **P<0.01; ***P<0.001.
DISCUSSION

In the present study, we identified a rare gain-of-function variant in the LIPC gene, encoding HL, as a novel cause of monogenic dominant familial combined hypercholesterolemia. We show that this variant, E97G, modifies an evolutionarily conserved motif that determines substrate access to the HL catalytic site. This modification specifically increases HL phospholipase activity and enhances lipoprotein catabolism. Mice overexpressing LIPC-E97G recapitulate the combined hypercholesterolemic phenotype and demonstrate that the increased HL phospholipase activity caused by the variant promotes TGRL catabolism by extrahepatic tissues. Together, our data uncover a novel type of hypercholesterolemia that is caused by an increased peripheral lipoprotein clearance as a result of increased HL phospholipase activity. Whether this drastic reduction in circulating lipids translates into decreased atherosclerosis remains to be established. Indeed, whereas no linkage to ASCVD was found in the family, the proband had evolving documented coronary artery stenosis and carotid atherosclerosis.

Our data highlight, for the first time, the physiological importance of the phospholipase activity of HL in controlling lipoprotein metabolism. In our family, the increased phospholipase activity profoundly changes lipoprotein characteristics, with an increased presence of large triglyceride-rich HDL and low HDL-C, low quantities of LDL particles, and a small VLDL size. These changes are largely what would be expected from an LIPC gain-of-function variant based on literature, with the exception of the triglyceride enrichment of HDL. Indeed, individuals with an LIPC gain of function are characterized by elevated HDL-triglycerides, HDL-phospholipids, and intermediate-density lipoprotein levels. Our data suggest that the HL phospholipase activity is a pivotal factor controlling lipoprotein catabolism. With its increased preference for phospholipids, the E97G variant makes HL more closely resemble its family member EL in terms of substrate preference. EL has mainly phospholipase activity and has historically been linked to the modulation of plasma HDL levels. Recent studies have, however, suggested that an increased EL activity and subsequent accelerated LDL catabolism are involved in the LDL-C reduction induced by angiopoietin-like 3 inhibition. Furthermore, a study on individuals with rare EL variants causing hyperalphalipoproteinemia (ie, high HDL-C levels) indicated that many of these patients also had elevated LDL-C levels, and the inhibition of EL with a monoclonal antibody increased both HDL-C and LDL-C levels in nonhuman primates and humans. Together, these studies highlight the important role of intravascular phospholipases in VLDL and LDL metabolism, on top of their major role in HDL metabolism.

Our study suggests that the increased phospholipase activity of HL promotes peripheral catabolism of VLDL and LDL. It is well established that the phospholipase activity of HL promotes hepatic phospholipids uptake from circulating lipoproteins, and we initially hypothesized that an increased hepatic clearance of lipoproteins was responsible for the observed hypercholesterolemia. However, the reduced uptake of radiolabeled VLDL-like particles and murine VLDL by the liver and the absence of changes in D7-cholesterol in feces argue against this hypothesis. Instead, the LIPC-E97G variant seems to promote the increase in core remnant uptake in peripheral tissues such as adipose tissue and muscle. Peripheral clearance of TGRL has previously been demonstrated in the context of cold exposure: Local LPL-mediated remodeling of TGRL promoted whole-particle TGRL uptake through CD36 in brown adipose tissue and white adipose tissue for further lipolytic processing of lipoprotein by lysosomal acid lipase. Besides CD36, endothelial scavenger receptor B1 might also be involved in peripheral whole particle uptake, as was elegantly shown in the context of LDL transcytosis and atherosclerosis. In this respect, it is of interest to note that both CD36 and scavenger receptor B1 have a great affinity for anionic phospholipids such as phosphatidylserine, which have been previously shown to accumulate on the surface of HDL-remodeled lipoproteins. Further studies are warranted to further decipher the molecular

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Table 2. Plasma Parameters APOE*3.Leiden.CETP Mice

|                         | eGFP low | LIPC low | LIPC-E97G low | eGFP high | LIPC high | LIPC-E97G high |
|-------------------------|----------|----------|---------------|-----------|-----------|---------------|
| Cholesterol, mmol/L     | 3.10±1.05| 3.39±0.83| 1.92±0.49*    | 3.17±1.05 | 2.42±0.70 | 0.91±0.42‡    |
| HDL-C, mmol/L           | 0.77±0.19| 0.85±0.18| 0.48±0.17†‡   | 0.75±0.18 | 0.61±0.17§ | 0.22±0.10||    |
| Non-HDL-C, mmol/L       | 2.33±0.89| 2.54±0.77| 1.46±0.39*    | 2.42±1.11 | 1.82±0.56 | 0.69±0.33||    |
| Triglycerides, mmol/L   | 1.87±0.83| 1.82±0.93| 1.22±0.23     | 2.35±1.00 | 1.43±0.79 | 0.76±0.40‡    |
| Phospholipids, mmol/L   | 2.37±0.55| 2.70±0.63| 1.79±0.46*‡   | 2.72±0.66 | 2.17±0.56§ | 0.87±0.16||    |

Data are depicted as mean±SD. eGFP indicates enhanced green fluorescent protein; HDL-C, high-density lipoprotein cholesterol; and LIPC, lipase C, hepatic type.

*P<0.05 LIPC-E97G low vs LIPC low.
†P<0.05 LIPC-E97G low vs eGFP low.
‡P<0.05 LIPC-E97G high vs eGFP high.
§P<0.05 LIPC-E97G high vs eGFP high.
∥P<0.05 LIPC-E97G low eGFP high
LIPC-E97G high vs LIPC low.
LIPC-E97G low vs LIPC high.

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Figure 6. Overexpression of the LIPC-E97G variant promotes peripheral cholesterol uptake. A and C, Decay of plasma $^3$H activity (glycerol tri$[^3]$H-oleate, hydrolysable; A) and plasma $^{14}$C activity ([$^{14}$C]cholesterol oleate, nonhydrolysable) levels (C) in APOE*3.Leiden.CETP mice overexpressing AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G injected with very-low-density lipoprotein (VLDL)-like particles. B and D, Decay of plasma $^3$H activity (glycerol tri$[^3]$H-oleate, hydrolysable; B) and plasma $^{14}$C activity ([$^{14}$C]cholesterol oleate, nonhydrolysable) levels (D) in APOE*3.Leiden.CETP mice overexpressing AAV-TBG-eGFP, AAV-TBG-LIPC, and AAV-TBG-LIPC-E97G injected with radiolabeled murine VLDL. E, Liver $^{14}$C activity (cholesterol ester, nonhydrolysable; left) and liver $^3$H activity (triolein, hydrolysable) levels (right) in APOE*3.Leiden.CETP mice overexpressing AAV-TBG-eGFP, AAV-TBG-LIPC, (Continued)
The present study is not without limitations. Although we used the humanized APOE*3.Leiden.CETP mouse model, the moderate overexpression of LIPC-E97G in these mice is not fully comparable to the heterozygous presence of the E97G variant in our family. The hypocholesterolemic phenotype is overall recapitulated, but there are clear differences with respect to triglyceride levels and HDL size and composition. Indeed, the presence of large, triglyceride-enriched HDL particles in our family is peculiar, and we do not have a clear explanation for this phenotype. Previous studies have shown that HL-deficient individuals also have triglyceride-enriched HDL and that the inhibition of EL with a monoclonal antibody significantly increased HDL particle size in humans. Unfortunately, the phenotypic differences in HDL size and composition between mice and humans precluded more extensive studies in mice, and further studies could not be performed in family members for personal reasons. It should also be noted that we did not observe a major impact of a moderate LIPC-WT overexpression on plasma lipid levels or on plasma lipoprotein clearance, in contrast to previously published studies. The presence of endogenous mouse HL and the moderate level of overexpression might play a role here. We do feel that the strong and consistent phenotype on LIPC-E97G overexpression further highlights the major impact that this variant has on lipoprotein metabolism.

From a clinical perspective, the impact of our LIPC-E97G variant on ASCVD development remains unclear, which is a clear limitation of our study. Whereas the low LDL-C levels in our family would suggest a beneficial impact on ASCVD risk, the strong decrease in HDL-C, the near absence of small HDL particles, and the fact that our proband was initially referred after an acute coronary syndrome might suggest otherwise. It should be noted, however, that even the role of wild-type HL, but also EL, in ASCVD development remains a matter of debate because HL and EL have both proatherogenic and antiatherogenic properties. Furthermore, human genetic studies are not consistent, although overall data suggest that a reduction in HL activity is associated with a modestly increased ASCVD risk. Because low HDL-C does not seem to be causally linked to ASCVD, it could be that our complex clinical phenotype is an exception to the statement that any decrease in LDL-C translates into an ASCVD reduction. To confirm or refute such a hypothesis and to accurately study the impact of the LIPC-E97G variant on ASCVD risk would require a heterozygous transgenic model with a physiological expression profile, a human-like lipoprotein profile, and no endogenous Lpc, which was beyond the scope of the present study. In addition, the fact that the LIPC-E97G variant was associated with an increased catabolism of TGRLs in extrahepatic tissues suggests that this new pathway can promote the lowering of circulating lipids without increasing the risk of hepatic steatosis, an undesirable side effect that has halted the development of some new lipid-lowering drugs.

Conclusions

We identify a rare variant in the LIPC gene as a second cause of familial combined hypercholesterolemia by increased clearance. This study highlights the importance of the phospholipase activity of HL in the regulation of lipoprotein metabolism, as well as the need for an increased understanding of the link between lipase specificity (triglycerides/phospholipids) and lipoprotein catabolism (TGRL remnants/HDL). Future studies that combine the study of specific LIPC variants and corresponding molecular modeling investigations might shed further light on these aspects. Our observations are of direct clinical relevance because they highlight that an increased hepatic phospholipase activity can drastically lower plasma LDL-C levels. Because the proband developed ASCVD despite low LDL-C plasma levels, additional models and studies will be required to specify the impact of the E97G variant on atherosclerosis progression and to decipher whether this approach could be a promising area of research to lower ASCVD risk.
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Disclosures

Dr Carri reports personal fees from Abbott and Acoea, as well as grants and personal fees from Amgen, AstraZeneca, Bristol Myers Squibb, Gilead, Eli Lilly, Novartis, Novo Nordisk, Sanofi, and Regeneron, outside the submitted work. Dr Moulin reports grants, personal fees, and nonfinancial support from Acoea, Amgen, Boehringer, Ionis, and Novo, outside the submitted work. Dr Rensen reports grants from Eli Lilly, Lipigon, and Retrotope, outside the submitted work. Dr Amigo Gen, Boehringer, Ionis, and Novo, outside the submitted work. Dr Rensen reports grants, personal fees, and nonfinancial support from Akcea, Amgen, Boehringer, Ionis, and Novo, outside the submitted work. Dr Moulin reports grants, personal fees, and nonfinancial support from Akcea, Amgen, Boehringer, Ionis, and Novo, outside the submitted work. Dr Rensen is a stock owner of Biosfer Teslab and has a patent on the lipoprotein profiling

Supplemental Material

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