Pancreatic carboxyl ester lipase (CEL) hydrolyzes cholesteryl esters (CE), triglycerides (TG), and lysophospholipids, with CE and TG hydrolysis stimulated by cholate. Originally thought to be confined to the gastrointestinal system, CEL has been reported in the plasma of humans and other mammals, implying its potential in vivo to modify lipids associated with LDL, HDL (CE, TG), and oxidized LDL (lysophosphatidylcholine, lysoPC). We measured the concentration of CEL in human plasma as 1.2+/--0.5 ng/ml (in the range reported for lipoprotein lipase). Human LDL and HDL3 reconstituted with radiolabeled lipids were incubated with purified porcine CEL without or with cholate (10 or 100 microM, concentrations achievable in systemic or portal plasma, respectively). Using a saturating concentration of lipoprotein-associated CE (4 microM), with increasing cholate concentration there was an increase in the hydrolysis of LDL- and HDL3-CE; at 100 microM cholate, the present hydrolysis per hour was 32+/--2 and 1.6+/--0.1, respectively, indicating that CEL interaction varied with lipoprotein class. HDL3-TG hydrolysis was also observed, but was only approximately 5-10\% of that for HDL3-CE at either 10 or 100 microM cholate. Oxidized LDL (OxLDL) is enriched with lysoPC, a proatherogenic compound. After a 4-h incubation with CEL, the lysoPC content of OxLDL was depleted 57\%. Colocalization of CEL in the vicinity of OxLDL formation was supported by demonstrating in human aortic homogenate a cholate-stimulated […]

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Pancreatic Carboxyl Ester Lipase: A Circulating Enzyme That Modifies Normal and Oxidized Lipoproteins In Vitro

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Abstract

Pancreatic carboxyl ester lipase (CEL) hydrolyzes cholesteryl esters (CE), triglycerides (TG), and lysophospholipids, with CE and TG hydrolysis stimulated by cholate. Originally thought to be confined to the gastrointestinal system, CEL has been reported in the plasma of humans and other mammals, implying its potential in vivo to modify lipids associated with LDL, HDL (CE, TG), and oxidized LDL (lyso-phosphatidylcholine, lysoPC). We measured the concentration of CEL in human plasma as 1.2±0.5 ng/ml (in the range reported for lipoprotein lipase). Human LDL and HDL reconstituted with radiolabeled lipids were incubated with purified porcine CEL without or with cholate (10 or 100 μM, concentrations achievable in systemic or portal plasma, respectively). Using a saturating concentration of lipoprotein-associated CE (4 μM), with increasing cholate concentration there was an increase in the hydrolysis of LDL- and HDL-CE; at 100 μM cholate, the percent hydrolysis per hour was 32±2 and 1.6±0.1, respectively, indicating that CEL interaction varied with lipoprotein class. HDL-3-TG hydrolysis was also observed, but was only ~5–10% of that for HDL-2-CE at either 10 or 100 μM cholate. Oxidized LDL (OxLDL) is enriched with lysoPC, a proatherogenic compound. After a 4-h incubation with CEL, the lysoPC content of OxLDL was depleted 57%. Colocalization of CEL in the vicinity of OxLDL formation was supported by demonstrating in human aortic homogenate a cholate-stimulated cholesteryl ester hydrolytic activity inhibited by anti–human CEL IgG. We conclude that CEL has the capability to modify normal human LDL and HDL composition and structure and to reduce the atherogenicity of OxLDL by decreasing its lysoPC content. (J. Clin. Invest. 1996. 97:1696–1704.) Key words: low density lipoprotein • high density lipoprotein • oxidized low density lipoprotein • aorta • lysophosphatidyl choline

Introduction

Modification by lipases, such as hepatic lipase (1), lipoprotein lipase (2), and phospholipase A2 (3), alters the composition, structure, and function of lipoproteins. Neglected in this regard is the pancreatic carboxyl ester lipase (CEL) sometimes referred to as bile salt–dependent cholesteryl ester hydrolase, whose primary function is thought to be hydrolysis of dietary lipid esters in the intestinal lumen (for a recent review, see reference 4).

One reason for this neglect may be the lack of appreciation that CEL activity is expressed outside of the pancreas in a number of mammalian species. For example, there are reports of CEL activity and/or mRNA in rat liver (e.g., references 5–7), human breast (8), human hepatocarcinoma HepG2 cells (9), human placenta (10), rat (11) and rabbit heart (12), and rat and rabbit aorta (13, 14). Particularly intriguing is the evidence that CEL is found in the blood plasma of a number of mammals, including humans (15), rats (11), dogs, and goats (16). Implanted, then, is that lipoproteins in the circulation or interstitial could be potential targets for CEL and also that circulating CEL, which has a heparin binding site (17), may become associated with a variety of tissues, independent of their capacity to synthesize the enzyme, by associating with cell-surface proteoglycans. The consequences of CEL-mediated lipoprotein modification are suggested by recent reports that (a) the treatment in vitro of LDL by nonmammalian cholesteryl esterases depletes LDL of core cholesteryl esters (18) and leads to increased net transfer of LDL-cholesterol to cells (19), and (b) CEL secreted by HepG2 cells increases the cellular uptake of cholesteryl esters from HDL (20).

In addition to its activity against the dietary lipids cholesteryl ester (CE), triglyceride (TG), and retinyl ester, CEL has significant activity against lysophospholipids and, in fact, was originally cloned as the pancreatic lysophospholipase (21). Thus, it is plausible that CEL may modify not only the core CE or TG of a normal lipoprotein, but also the lysophosphatidylcholine (lysoPC) of oxidized LDL (OxLDL). Given the major roles attributed to lysoPC in the promotion of atherosclerotic disease (such as enhanced monocye chemotaxis [22], mitogenic stimulation of macrophages [23], and inhibition of endothelium-dependent arterial relaxation [24]) and the reports that CEL activity was found in rat and rabbit aortae (13, 14), CEL could serve as a protective factor in vascular tissue

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1. Abbreviations used in this paper: CE, cholesteryl ester; CEL, carboxyl ester lipase; HDL, HDL subclass 3; LpL, lipoprotein lipase; lysoPC, lysophosphatidylcholine; OxLDL, oxidized LDL; PC, phosphatidylcholine; TBARS, thiobarbituric acid–reactive substances; TG, triglyceride or triacylglycerol.
against the adverse effects of OxLDL. Supporting this hypothesis are the recent reports that the treatment of OxLDL with a bacterial lysophospholipase reduced the antagonism by lysoPC of the endothelial-dependent relaxation response to acetylcholine of aortic rings (24, 25). Since oxidation of LDL is thought to occur not in plasma but within arterial tissue (26), aortic CEL activity would be in the appropriate location to serve as a protective factor.

In the studies summarized in this report, we have demonstrated: (a) the ability of CEL to hydrolyze the core lipids of human LDL and HDL using bile salt concentrations that can occur outside of the intestinal lumen in healthy individuals; (b) the ability of CEL to reduce the lysoPC content of human OxLDL; (c) the presence of CEL in human plasma; and (d) the presence of an activity in human aortic homogenate with enzymatic and immunological properties indistinguishable from CEL. The results strongly suggest that CEL plays a wider role in lipid and lipoprotein metabolism than previously appreciated both in terms of enzymatic targets and tissue distribution.

Methods

All chemicals were the highest grade commercially available and purchased from Sigma Chemical Co. (St. Louis, MO), unless otherwise noted. Cholesterol [1-14C]oleate (supplied with specific activity of 53.4 mCi/mmol), [carboxyl-14C]triolein (112 mCi/mmol), and [palmitoyl-1-14C]lysocephatidylcholine (56.7 mCi/mmol) were purchased from New England Nuclear (Boston, MA). [1α,2α(n)-2H]cholesterol oleate (specific activity of 49.6 Ci/mmol) was purchased from Amersham (Arlington Heights, IL). Porcine pancreatic CEL was purified as reported previously (27). Reagents for lipoprotein electrophoresis were obtained from Beckman (Fullerton, CA).

The procedures for obtaining human blood samples have been approved by the Committee for Protection of Human Subjects at the Medical College of Pennsylvania and the donors have signed informed consent forms.

Reconstitution of human lipoproteins with radiolabeled lipids

Lipoprotein fractions were isolated from the plasma of normal human volunteers using sequential density gradient centrifugation (28, 29). HDL3 was delipidated with ethanol/diethyl ether (3:2 vol/vol) at 0°C (30) and the apoproteins were reserved for the reconstitution step (see below). The ethanol/diethyl ether lipid extracts were dried by rotary evaporation and stored in 2:1 chloroform/methanol. To incorporate cholesteryl [1-14C]oleate or [14C]triolein into HDL3, the labeled compounds were added directly to the lipid extract, which was then reconstituted with HDL3 apoproteins by sonication (31). The reconstituted HDL3 had a final specific activity of ~ 0.02 μCi of cholesteryl [1-14C]oleate per μmol CE or ~ 7 μCi of [carboxyl-14C]triolein per μmol TG.

[1α,2α(n)-2H]cholesterol oleate was incorporated into LDL by the potato-starch method of Krieger et al. (32). The final specific activity was ~ 0.10 μCi of [1α,2α(n)-2H]cholesterol oleate per μmol LDL-CE.

Analysis of reconstituted and native human lipoproteins

For reconstituted lipoproteins, lipids were extracted by the procedure of Bligh and Dyer (33). Phospholipid was determined by the method of Sokoloff and Rothblat (34). Protein was assayed by a modification of the Lowry technique (35). Cholesterol (free and total) was determined by gas-liquid chromatography using cholesteryl methyl ether as an internal standard (36). Esterified cholesterol was calculated by subtracting free cholesterol from total cholesterol. TG was determined by the Sigma triglyceride diagnostic kit, using glycerol as a standard. In some experiments, the cholesterol content (total, free, ester) of native LDL was determined as above. Agarose electrophoresis of native and reconstituted lipoproteins and gel staining by Sudan black were performed using Beckman Paragon Lipo Gel materials.

Preparation and characterization of oxidized human LDL

Human LDL, isolated as above, was oxidized by a 7-h incubation with Cu2+ (5 μM above EDTA concentration [37]). Butylated hydroxytoluene (20 μM) was then added to stop any further oxidation. The cholesterol content of LDL and OxLDL was determined enzymatically by the Sigma cholesterol diagnostic kit No. 352. Protein content was assayed using a modification of the Lowry technique (35). The cholesterol contents of LDL and OxLDL were 1.4 mg and 1.2 mg/mg protein, respectively. The protein concentrations of LDL and OxLDL were 1.4 and 1.3 mg/ml, respectively. The extent of lipid peroxidation was estimated as thiobarbituric acid–reactive substances (TBARS) as described previously (38). Briefly, LDL and OxLDL (50 μg of protein of either) and various amounts of standard (1,1,3,3-tetramethoxyxylene) were mixed with 1 ml of 25% trichloroacetic acid and 1 ml of 1% 2-thiobarbituric acid containing 5 μM CuSO4. The mixture (final volume of 0.4 ml) was incubated at 95°C for 45 min and centrifuged at 1,500 rpm at 10°C for 30 min to remove any insoluble material. The OD436 of the supernatant was determined and the amount of TBARS was expressed as nanomoles of malondialdehyde equivalents per milliliter of reaction mixture.

Human tissue collection and preparation

Male and female aortic arch samples were obtained at autopsy and frozen immediately in liquid N2 and then stored at −70°C until analyzed. Under the Institutional Review Board guidelines autopsy samples were collected so that neither the identity nor cause of death of subjects was revealed to us. To prepare samples for enzymatic analysis frozen tissues were thawed on ice. Adherent vascular and membranous materials were dissected away and the remaining tissue dispersed at 4°C in 3 vol of 0.25 M sucrose at full speed for 1 min in a homogenizer (Tekmar Co., Cincinnati, OH). Aliquots were stored frozen at −70°C until enzymatic assays were performed (see below).

Assay of the hydrolysis of CE, TG, or lysoPC

All activity determinations were done in duplicate with a variability coefficient < 10%.

CE and TG. For reconstituted lipoproteins, the hydrolysis of CE or TG was determined based on radiometric procedures described previously (11, 39). For substrates dispersed in ethanol, the reaction mixture had a final volume of 0.2 ml; for substrates in LDL or HDL3, the final volume was 1 ml. The final concentration of Tris-maleate (pH 7.0) was 50 mM, and sodium cholate was either 0, 10, or 100 μM. Reactions were initiated by addition of purified CEL and labeled substrate (to the concentrations given in Results) and incubated at 37°C for the lengths of time also given in Results. The released [14C]oleate was extracted as described previously (11) and quantitated by scintillation counting. Results are typically expressed in units of 1 pmol fatty acid released/h/μg CEL. To examine the effect of CEL on the hydrolysis of CE of native LDL a similar protocol was performed, but the reaction volume was 5 ml and cholate (total, free, ester) was assayed as in the above section, Analysis of reconstituted and native human lipoproteins.

For analysis of aortic homogenates, 50 μl of an appropriately diluted homogenate was used as the enzyme source. The final reaction volume was 0.2 ml and the concentration of Tris-maleate (pH 7.0) was 50 mM. Cholate or deoxycholate was added at a concentration ranging from 0 to 100 mM as indicated in Results. The reactions were initiated by addition of 0.01 ml ethanol containing 2 nmol of cholesteryl [1-14C]oleate with a specific activity of 25 μCi/μmol. The tubes were incubated at 37°C for 30 min. The released [14C]oleate was extracted and quantitated as above. Results are expressed in units of 1 pmol [14C]oleate released/h/μg of tissue.

To investigate the inhibition of aortic homogenate CEL activity by a rabbit IgG directed against human CEL, the following experi-
ment was performed. Aortic homogenate was preincubated overnight at 4°C with 5 μg of either rabbit anti–human CEL (40, 41) or rabbit preimmune IgG. The samples were then assayed for CEL activity as described above.

**LysoPC.** Lysophospholipase activity was measured using the method described by Vanden Bosch et al. (42). Substrate and either buffered enzyme (porcine pancreatic CEL or bacterial 2-lysophosphatidylethanolamine acylhydrolyase [EC 3.1.1.5; Sigma]) or buffer alone (potassium phosphate 0.1 M) were diluted with D_{2}O to a final volume of 0.4 ml and a final potassium phosphate concentration of 200 mM. In some cases, to determine the presence of endogenous lysophospholipase activities of LDL and OxLDL, these lipoproteins were substituted for the enzyme source.

The reaction was started by adding 200 nmol (0.1 ml of a 2 mM aqueous solution) of the substrate [palmitoyl-1-14C]lysopalmitoyl phosphatidylcholine with specific activity of 26.1 × 10^{-3} μCi/μmol. After 10 min at 37°C, the reaction was stopped with 2.5 ml of modified Dole’s solution (43). Approximately 100 mg of silica gel (100–200 mesh) was added, followed by 1.5 ml of heptane and 1.5 ml of water. After each addition, the tube was vortexed for 15 s. 1 ml of the resulting upper phase, which contained the labeled fatty acid released from lysoPC by hydrolysis, was taken for scintillation counting. Results are expressed in nanomoles of fatty acid released per minute per milligram of protein. Hydrolysis of lysoPC at a final concentration of 80 μM (similar to the concentration of OxLDL-associated lysoPC in the experiments described below) was also assayed, but with 4 μg/ml of CEL and for incubation times up to 240 min. The percent hydrolysis at each time point was calculated from the scintillation counting data.

In pilot experiments, the potential for CEL to act as a phospholipase was investigated using both radiolabeled phosphatidylethanolamine (PE) and HDL reconstituted with radiolabeled PE. No activity above background was detected in either case.

**Effect of CEL on the lysophospholipid content of OxLDL.** Human lipoproteins (LDL, OxLDL) containing ~0.5 mg protein/ml were incubated with either 4 μg CEL protein/ml or 1 μg/ml bacterial 2-lysophosphatidylethanolamine acylhydrolyase or buffer alone (potassium phosphate, 200 mM final concentration) for 4 h at 37°C. At time points (0, 1, 2, and 4 h) the reaction was stopped with 1 ml methanol and lipids were extracted using the procedure of Bligh and Dyer (33). The phospholipids were separated by TLC on silica gel G (250 μm, Analtech Inc., Newark, DE) with two solvent systems used in the same dimension: chloroform/methanol/water, 65:35:6 (vol/vol/vol) and chloroform/acetic acid/water, 6:8:2:2:1. After visualization with iodine vapor, PE and lysoPC bands were scraped from the plates and assayed for inorganic phosphorus content according to the method of Sokoloff and Rothblat (34). Each TLC plate had lanes containing unlabeled and labeled standards to localize the desired bands in the sample lanes. Quantitative recovery of lysophospholipids was confirmed in some experiments by adding background was detected in either case.

Detection of circulating CEL in human serum

The amount of CEL in human serum samples was determined by ELISA as described previously (15). Briefly, wells of immunoplates (Nunc, Roskilde, Denmark) were coated with a 10 μg/ml solution (0.1 M carbonate buffer, pH 9.5) of anti–human CEL polyclonal antibodies raised in rabbit (40, 41). After overnight incubation at 4°C, wells were washed three times for 30 min at 37°C with 0.05% Tween 20 in PBS (PBS/Tween). Then, 100-μl portions of each sample (either human serum or pure human CEL standard), appropriately diluted in PBS/Tween, were added to the wells and incubated for 30 min at 37°C. After three washes with PBS/Tween, 100 μl of biotin-labeled rabbit polyclonal antibodies to human CEL (2 μg/ml was added to each well and incubated for 2 h at 37°C. After three washes with PBS/Tween, dilute alkaline phosphatase–avidin conjugates were added (100 μl/well) and incubated for 2 h at 37°C. After three washes with PBS/Tween, 100 μl of p-nitrophenyl phosphate (1 mg/ml in 0.2 M Tris/HCl, pH 8.5, 1 mM CaCl$_2$, and 1 mM MgCl$_2$) was added to each well and incubated for 1 h at 37°C. The degree of color development was determined by reading the plate in an MR 5000 spectrophotometer (Dynatech Laboratories Inc., Chantilly, VA) at 405 nm and comparing the serum values with the standard curve based on the assay results for serially diluted samples of the pure human CEL.

Data analysis

Unless otherwise indicated, data are expressed as mean values±SD and statistical comparisons made by the Student’s t test or ANOVA. The software program InStat (GraphPAD Software for Science, San Diego, CA) was used for testing statistical significance.

Results

**CEL is a circulating lipase in humans.** CEL activity or mass has been reported in the plasma of rats (11), dogs, goats (16), and humans (15). We were interested in confirming that CEL is a circulating enzyme in humans. Plasma CEL mass was measured in six normal volunteers by a quantitative ELISA assay (15). The average concentration was 1.2±0.5 ng/ml (range 0.6–2.0), consistent with literature values (15) and in the range reported for lipoprotein lipase (LpL) (44).

The effects of CEL on normal lipoproteins

**Properties of reconstituted human HDL$_3$ and LDL.** The reconstitution of normal human HDL$_3$ and LDL included radiolabeled CE. Since human HDL also contains small amounts of TG, some HDL$_3$ was also reconstituted with [3H]TG as the only labeled core lipid. As shown in Table I, the chemical compositions of the reconstituted lipoproteins resembled the reported values for native human lipoproteins (45). In addition, we tested whether the cholate concentrations used in the experiments described below would significantly affect the physical properties of the reconstituted lipoproteins. The agarose gel electrophoretic mobilities of reconstituted LDL and HDL$_3$ were not affected by treatment with 10 or 100 μM cholate (data not shown), indicating that the bile acid did not cause major disruptions of the structure of these lipoproteins.

**Characterization of the action of CEL on the CE of HDL$_3$.** Since CE is the major core lipid in HDL, and this lipoprotein is relatively simple to reconstitute, we used [3H]CE-labeled HDL$_3$ to establish appropriate reaction conditions for examining lipoprotein modification by CEL. Previous studies of lipoproteins modified by cholesteryl esterases have typically used enzymes in the milligrams per milliliter range (e.g., references

| Table I. Chemical Compositions of Human HDL$_3$ and LDL Reconstituted with Labeled Core Lipids |
|-----------------------------------------------|-------------------------------|-------------------------------|-------------------------------|
| Protein                                      | 27.4±2.1                      | 48.0±0.07                     | 41.0±0.05                     |
| Phospholipid                                  | 29.8±1.6                      | 32.2±0.04                     | 32.4±0.90                     |
| CE                                           | 23.7±1.4                      | 15.4±1.20                     | 20.3±1.53                     |
| Cholesterol                                  | 15.3±1.5                      | 2.90±0.05                     | 3.78±0.11                     |
| TG                                           | 3.84±0.63                     | 1.57±0.03                     | 2.52±0.18                     |

The chemical composition was determined in triplicate and is given as percentage±SD of total weight.
Table II. Hydrolysis by Pancreatic CEL of CE and TG Dispersed in Ethanol or Associated with Lipoproteins (pmol/h/µg CEL Protein ±SD)

<table>
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<tr>
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<th>No cholate</th>
<th>10 µM cholate</th>
<th>100 µM cholate</th>
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<tr>
<td><strong>Ethanol</strong></td>
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<td></td>
</tr>
<tr>
<td>CE</td>
<td>ND</td>
<td>110±9</td>
<td>4049±490</td>
</tr>
<tr>
<td>TG</td>
<td>ND</td>
<td>3980±1044</td>
<td>8518±2168</td>
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<tr>
<td><strong>LDL</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CE</td>
<td>27±1</td>
<td>185±16</td>
<td>5818±345</td>
</tr>
<tr>
<td>TG</td>
<td>4±1</td>
<td>20±2</td>
<td>23±1</td>
</tr>
<tr>
<td><strong>HDL₃</strong></td>
<td></td>
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</tr>
<tr>
<td>CE</td>
<td>0.3±0.0</td>
<td>1.2±0.3</td>
<td>2.5±0.2</td>
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<tr>
<td>TG</td>
<td></td>
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</table>

CE activity was assayed at 37°C for 60 min using (a) CE (4 µM) dispersed in ethanol, (b) TG (4 µM) dispersed in ethanol, (c) reconstituted LDL (4 µM CE), (d) reconstituted HDL₃ (4 µM CE), (e) reconstituted HDL₃ (4 µM TG). Purified CEL was used at 4 µg/ml except for substrates a and c when the cholate concentration was 100 µM; in these cases, the CEL concentration was 0.25 µg/ml. ND. Activity above background not detected.

18 and 19). However, given the similar concentrations in human plasma of LpL and CEL noted above, we decided to use a concentration in the micrograms per milliliter range, similar to those used in studies of LpL-lipoprotein interactions in vitro (e.g., references 46 and 47). The next choice was the range of cholate concentration. The concentration of bile acid in human plasma is in the range of 10 µM (48) and is ~10-fold higher in postprandial portal plasma (49). Thus, cholate concentrations of 0, 10, and 100 µM were used.

With the chosen CEL and cholate concentrations, the dependences of HDL₃-CE hydrolysis on incubation time and substrate concentration were examined. Independent of cholate concentration, the hydrolysis of CE did not approach a plateau until after 60 min (Fig. 1 A), but clearly saturated at a CE concentration of 4 µM (Fig. 1 B). These parameters of incubation time (60 min) and lipid concentration (4 µM) were maintained in subsequent testing of the effects of CEL on the rates of hydrolysis of HDL₃-TG and LDL-CE in order to make comparisons among results obtained under identical conditions.

The effects of substrate presentation on CEL-mediated hydrolysis of CE and TG. Using the conditions defined above, CE was presented to CEL in one of the following forms: (a) dispersed in EtOH, (b) associated with HDL₃, or (c) associated with LDL. In addition, TG was presented in either form a or b.

The results are summarized in Table II. Note that for all forms of substrate presentation, 10 µM cholate significantly stimulated hydrolysis in comparison to the no cholate control. In all but one case (HDL₃-CE), increasing the cholate concentration from 10 to 100 µM further stimulated hydrolytic activity. It is also interesting to note that for EtOH-CE or LDL-CE, at either concentration of cholate, the hydrolysis was comparable and greatly exceeded that observed for HDL₃-CE. Similarly, the hydrolysis of EtOH-TG was significantly greater than that of HDL₃-TG.

These results suggested that modest amounts of CEL could deplete the CE content of LDL to a greater extent than that of HDL₃, given not only the increased fractional hydrolysis of LDL-CE, but also that the CE core of LDL is much larger than that of HDL₃. In Table III are summarized experiments in which the fractional hydrolysis per hour of LDL-CE and HDL₃-CE by CEL was measured. As shown, there was indeed an appreciable percentage of LDL-CE hydrolyzed over time at either cholate concentration. That hydrolysis of LDL-CE by CEL and its stimulation by cholate were not artifacts of the lipoprotein reconstitution procedure was supported by finding similar results with native LDL studied under the same conditions (4 µM LDL-CE and 0.25 µg/ml CEL): at 0 and 100 µM cholate, there were reductions in CE mass of 1.3% (±0.7, n =

Figure 1. Dependence of CEL-mediated hydrolysis on incubation time and substrate concentration using reconstituted HDL₃. (A) CEL activity was measured at 37°C as described in Methods using reconstituted HDL₃ (4 µM CE). With 0 or 10 µM cholate, 4 µg/ml CEL was used. With 100 µM cholate, the concentration of CEL was 0.25 µg/ml. The amount of [14C]olate released was determined as described in Methods and expressed as picomoles per microgram of CEL. (B) CEL activity was measured as above, except the reaction time was 1 h and the amount of reconstituted HDL₃ varied to supply the indicated CE concentrations. □ no cholate; ▲ 10 µM cholate; ○ 100 µM cholate; * differs from 0 cholate at P < 0.01; ** differs at P < 0.001.

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Characterization of lipoproteins. The effects of CEL on OxLDL 
substrate with cholate at a concentration of 100 μM; in this case, the CEL concentration was 0.25 μg/ml.


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<tr>
<td>0</td>
<td>2.1±0.1</td>
<td>0.3±0.03</td>
<td>0.03±0.007</td>
</tr>
<tr>
<td>10 μM</td>
<td>14.6±1.3</td>
<td>1.61±0.18</td>
<td>0.06±0.01</td>
</tr>
<tr>
<td>100 μM</td>
<td>32.0±1.9</td>
<td>1.64±0.14</td>
<td>0.10±0.03</td>
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In a set of reactions separate from those summarized in Table II, hydrolysis by purified CEL was assayed using (a) reconstituted LDL (4 μM CE), (b) reconstituted HDL (4 μM CE), (c) reconstituted HDL (4 μM TG). Purified CEL was used at 4 μg/ml except for substrate with cholate at a concentration of 100 μM; in this case, the CEL concentration was 0.25 μg/ml.

4) and 35.0% (±5.8, n = 4), respectively.

In contrast to the LDL results, the percentage of the HDL[^3]C CE hydrolyzed under comparable conditions was relatively minor. Nonetheless, given the long half-life of HDL (≈3 d) in the human circulation (for review see reference 50), the extent of hydrolysis of HDL[^3]C CE in vivo may be underestimated by the results in vitro.

The effects of CEL on OxLDL. Lysophospholipase activity was determined using the assay described earlier (see Methods) under conditions of a saturating concentration of substrate. The endogenous activity of native or OxLDL was not different from background (data not shown). The activity of the porcine pancreatic CEL was 6,740±205 nmol of malondialdehyde equivalents/mg protein, respectively. As expected, compared with LDL, OxLDL demonstrated increased anodic electrophoretic mobility (data not shown), reflecting a surface more negatively charged. In the absence of lipases, the PC and lysoPC contents of LDL and OxLDL did not change during 4 h of incubation.

Lysophospholipase activities of LDL and CEL. Lysophospholipase activity was determined using the assay described earlier (see Methods) under conditions of a saturating concentration of substrate. The endogenous activity of native or OxLDL was not different from background (data not shown). The activity of the porcine pancreatic CEL was 6,740±205 nmol of malondialdehyde equivalents/mg protein, respectively. As expected, compared with LDL, OxLDL demonstrated increased anodic electrophoretic mobility (data not shown), reflecting a surface more negatively charged. In the absence of lipases, the PC and lysoPC contents of LDL and OxLDL did not change during 4 h of incubation.

**Figure 2.** Percent hydrolysis by CEL of lysoPC. Lysophospholipase activity was measured as described in Methods. The concentrations of lysoPC and CEL were 80 μM and 4 μg/ml, respectively. The reactions were stopped at the indicated times, then the released fatty acids were extracted and quantitated as described in Methods. Percent hydrolysis was determined as 100 (extracted cpm/total cpm) and is presented as mean±SD, n = 3.

creatic CEL was present in aortic homogenate (13, 14). To pursue whether there is a similar activity in human aorta, tissue samples were obtained at autopsy. There was significant activity present in 10/11 males (for n = 11: 28.9±10.3 nmol CE hydrolyzed/h/gram of tissue, mean±SE; range of 0–122) and 8/10 females (for n = 10: 11.1±4.7; range of 0–46.4). Although there was a higher mean level in males, because of the individual variability the gender difference was not statistically significant, unless the analysis was restricted to 10 males and 8
females with detectable activity, in which case the male average (n = 10; 31.8±11.0) was significantly higher (P < 0.04) than the female average (n = 8; 13.8±5.5).

That the homogenate activity represented bona fide CEL was supported by two experiments. In the first, the maximal stimulation of the reaction by millimolar concentrations of trihydroxy (e.g., cholate) and not dihydroxy (e.g., deoxycholate) bile salts, a characteristic feature of CEL activity measured in vitro (5), was determined. As shown in Fig. 4, human aortic homogenate CEL activity increased with increasing concentration of cholate, but not deoxycholate. In the second type of experiment, the ability of an antibody raised against human CEL (40, 41) to inhibit the homogenate activity was tested. As shown in Fig. 5, preincubation of human aortic homogenate with a small amount (5 µg) of rabbit anti-human CEL antibody significantly (P < 0.01) reduced the hydrolysis of CE compared with treatment with an equivalent amount of rabbit nonimmune serum.

Discussion

Besides its role in the digestive process (for review see reference 4), systemic actions of CEL must be considered, given that it circulates in a number of mammalian species (11, 16), including humans (15). Since other lipases that circulate have been shown to modify normal human HDL and LDL (e.g., LpL [1] and hepatic lipase [2]), it was natural to investigate whether the lipids on these lipoproteins could be hydrolyzed by CEL. However, one point we needed to take into account was that the millimolar concentrations of bile acid required for maximal hydrolysis of CE and TG in the intestine are not found in either plasma or interstitial fluid. Instead, in systemic plasma the concentration of bile acid is ~10 µM (48) and in postprandial portal plasma is ~10-fold greater (49). Since both of these concentrations are well in excess of the nanomolar levels shown to affect the conformation of CEL (51), it was certainly possible that, in the 10–100 µM range physiologically obtainable outside of the intestine, there would be sufficient enzymatic activity to hydrolyze significant amounts of lipoprotein-associated CE and TG.

As summarized in Tables II and III, this was indeed the case. Interestingly, the rates of hydrolysis (Table II) varied dramatically by mode of presentation (EtOH, HDL₃, LDL). The hydrolysis of lipoprotein-associated CE and TG in the absence of cholate is consistent with the studies of Brockman and colleagues (e.g., reference 52), who have shown that some CEL-mediated hydrolysis can occur in the absence of bile salts, depending on the characteristics of the substrate/enzyme interface. At either the 10 or 100 µM cholate concentration, the hydrolysis of both CE and TG associated with HDL₃ was much lower than when the corresponding substrate was presented in either EtOH or associated with LDL.

There are reports that with other lipases there was also differential hydrolysis of HDL and LDL lipids (e.g., references 18 and 53). For example, using a fungal cholesteryl esterase, under conditions in which LDL-CE (as percentage of total cholesterol) went from 75 to 6%, the corresponding values for HDL-CE were 79 and 65% (18). Factors such as packing density, surface charge, and apoprotein composition, all different between HDL₃ and LDL, probably affect the interaction of CEL (and the other enzymes) with lipoproteins as well as the availability to the enzymes of the substrates. In addition to the data in Tables II and III just alluded to, that such factors exert important influences on the interaction of CEL with HDL and HDL₃ is also indicated by some simple calculations (based on data in reference 54): The CE content of human LDL is 1,310 molecules/particle. Assuming a surface pool of CE of 3 mol%, the fraction of total LDL-CE that would be the functional substrate for the enzyme would be 20 surface molecules/1,310 total molecules, or 1.5%. The remaining 98.5% would be a “substrate reservoir.” A similar calculation for HDL₃ is a total CE content of 32 molecules/particle and 4 surface molecules, for a fractional functional substrate of 4/32, or 12.5%. Thus, in contrast to what was observed (Table II), the initial rate of hydrolysis of HDL₃-CE should have been 8 (i.e., 12.5%/1.5%) times

Figure 4. Cholesteryl ester hydrolase activity stimulated by trihydroxy bile salt in human aortic homogenate. CEL hydrolysis of CE in the presence of trihydroxy (cholate) or dihydroxy (deoxycholate) bile salt was determined in 50 µl of human aortic homogenate as described in Methods. Cholate (■) or deoxycholate (▲) was added at concentrations ranging from 0 to 100 mM. 1 unit is 1 nmol [14C]oleate released/h gram of tissue. Reactions were done in duplicate and the mean values are displayed.

Figure 5. Effect of anti-CEL antibody on bile salt–dependent CEL activity in human aortic homogenate. Diluted aortic homogenate (50 µl) was incubated overnight at 4°C with 5 µg of either rabbit anti–human CEL IgG or rabbit preimmune IgG. The samples were then assayed for CEL activity at 50 mM cholate as described in Methods. The reactions were done in triplicate and mean values±SD are displayed.
greater than that of LDL-CE. The fact that LDL-CE was the better substrate for CEL (Tables II and III) implies that the enzyme interacts with this particle better than with HDL.

The potential consequences of the action of CEL on the structural and functional properties of LDL and HDL are suggested by a number of studies. First, the core content of either CE or TG has been shown recently to affect the α-helical content of HDL—apo AI and the stabilities of both the apo AI and the HDL particle itself (Sparks, D.L., W.S. Davidson, S. Lund-Katz, and M.C. Phillips, manuscript submitted for publication). Such changes could, in turn, affect the ability of HDL to accept cellular cholesterol or to serve as a substrate for LCAT (55). This may be the basis for our previous finding that HDL added to medium containing CEL secreted from transfected rat hepatoma cells promoted greater clearance of cellular cholesterol compared with the clearance from nonsecreting control hepatoma cells (56).

In Aviram et al. (19), hydrolysis of ~1/3 of LDL-CE by a bacterial enzyme resulted in a smaller particle and a 30% reduction in lipoprotein binding and degradation by the J774 macrophage-like cell line. The authors concluded that these results were a consequence of conformational changes in apo B due not only to a reduction of core CE but also to a redistribution of the free cholesterol between the core and the surface of the lipoprotein. In Chao et al. (18), HDL and LDL were pretreated with trypsin, and CE was hydrolyzed by a fungal enzyme. Extensive hydrolysis of LDL-CE resulted in the formation of large multimellar free cholesterol-rich structures resembling particles that accumulate in atherosclerotic lesions.

Hydrolysis of core TG of HDL and LDL has also been investigated previously. Aviram et al. (2) treated LDL with LpL. In contrast to the results above, the modified LDL had a higher affinity for the LDL receptor. While the authors reconciled the results of their two papers by noting that predominately the core of the LDL is modified by LpL, whereas both the core and surface were modified by CE hydrolysis, a more likely explanation comes from later work of Williams et al. (47) and others (e.g., references 57 and 58) showing that LpL bound to LDL can act as a structural bridge to cell-surface proteoglycans. This would be consistent with the findings of Aviram et al. (2) that in the mathematical analysis of the binding of LpL-modified LDL to macrophages there was evidence for an increase in binding sites. It may be that a similar phenomenon will be observed for CEL-modified lipoproteins, since CEL has a heparin binding site that has been reported to promote the interaction of the enzyme with proteoglycans (17). This would explain the report that CEL enhanced the delivery of a nonhydrolyzable core lipid of HDL (cholesteryl ether) to hepatoma cells (20).

While many studies have focused on CEL as a triglyceride lipase and a cholesteryl ester hydrolase, it should be noted that it was originally cloned as a pancreatic lysophospholipase (21) and subsequently shown to be the only enzyme secreted by the pancreas with this activity (59). Unlike its other catalytic functions, there is no bile salt dependence under any condition for the hydrolysis of lysophospholipids. One major lysophospholipid with relevance to atherogenesis is lysoPC. OxLDL is enriched in lysoPC, which is thought to mediate deleterious effects in arterial tissue mainly by serving, for example, as a chemoattractant that recruits monocytes to the subintimal space (22) or as an antagonist of endothelial-dependent relaxation (24).

In a set of studies, Henry and colleagues (24, 25) have shown that the treatment of OxLDL with a bacterial lysophospholipase reduces not only the lysoPC content but also restores the endothelial-dependent relaxation response to acetylcholine of rabbit aortic rings. As shown in Fig. 3, we confirmed the ability of the bacterial enzyme to reduce OxLDL lysoPC content and also showed comparable effectiveness of CEL.

Since oxidation of LDL is not thought to occur in the plasma compartment but after LDL enters the subintimal space, to serve as a potential protective factor against lysoPC the optimal location of CEL for this function would be in arterial tissue. The partial purification of an enzymatic activity in rat and rabbit aortae (13, 14) with characteristics of pancreatic CEL encouraged us to examine human samples. That bona fide CEL was present was supported by (a) the stimulation of the cholesteryl ester hydrolase activity by only trihydroxy bile salt and not other detergent molecules (Fig. 4), a particularly characteristic feature of this enzyme (e.g., reference 5); and (b) the inhibition of homologene activity by preincubation with a specific antibody (40, 41) directed against human CEL (Fig. 5). The source of arterial CEL could be multiple; it could be derived from the circulating pool or it could be endogenously synthesized and secreted by one or more of the cell types (e.g., leukocytes [60], smooth muscle, endothelium) present in the tissue. Further studies using immunohistochemistry and in situ hybridization will be necessary to determine the source of aortic CEL activity. It should be noted that others have also reported a non-CEL cholesteryl ester hydrolase activity in the aorta or its component cells (e.g., references 2 and 61). This activity resembles that of hormone-sensitive lipase, in that it is maximally active at neutral pH and is stimulated by cAMP. We do not think that this contributed significantly to the results in human aortic homogenate, however, since a millimolar concentration of cholate inhibits hormone-sensitive lipase–like activity (reference 62 and Zolfaghari, R., and E.A. Fisher, unpublished results).

The relatively long half-lives of LDL and HDL (~3 d) in the human circulation (for review see reference 50) and the even greater half-lives of retained lipoproteins in the subintimal space of the aorta (for review see reference 63) imply that there may be significant modification of normal and oxidized lipoproteins by CEL, even for the more refractory substrates on HDL. For OxLDL, CEL is likely to be a protective factor, given its ability to reduce lysoPC content. For normal lipoproteins, it is not clear whether CEL modification would be beneficial or deleterious. For example, it could lead to the formation of LDL particles with increased cellular binding affinity (resulting in increased cellular cholesterol content) or denser LDL particles having increased access to the aortic subintimal space. On the other hand, LDL binding could be reduced (see reference 19) and/or efflux to HDL could be increased (56), both phenomena serving to decrease the net cellular cholesterol content. Nevertheless, our results clearly establish the potential for CEL to exert important effects on lipid metabolism outside of the intestinal lumen and support the undertaking of further studies to define the overall impact of the systemic activity on the development of atherosclerosis.

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References


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