Low level quantification of cholesteryl ester transfer protein in plasma subfractions and cell culture media by monoclonal antibody-based immunoassay

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Abstract Sensitive immunoradiometric (IRMA) and ELISA assays for cholesteryl ester transfer protein (CETP) have been developed using two different monoclonal antibodies (MAbs). The MAbs were prepared against human plasma CETP and demonstrated specificity by their inhibition of cholesteryl ester transfer activity and by immunoblots of crude plasma fractions and whole media from transfected CHO cells. For these sandwich-type assays, one MAb, 2F8, is used for capture, and the second MAb, 2E7, is iodinated (IRMA) or conjugated with alkaline phosphatase (ELISA) and used for detection. Both assays are linear and provide sensitivities much greater than previously reported. The IRMA allows for the accurate quantification of CETP in the range of 0.5-20 ng/assay (5-200 ng/ml), the ELISA 0.05-5 ng/assay (0.5-50 ng/ml). Using the IRMA, the mean plasma CETP concentration in 44 normolipidemic individuals was determined to be 2.10 ± 0.36 μg/ml; 2.05 ± 0.33 for males (n = 25) and 2.16 ± 0.40 for females (n = 19). The distribution of CETP in human plasma was examined both by gel permeation fast protein liquid chromatography (FPLC) and by native gel electrophoresis. For FPLC using agarose resins, a sandwich immunoblotting, showed CETP migrating within a size range of 170-220 kilodaltons. This result is consistent with suggestions that plasma CETP is associated with small-sized HDL. Agarose gel electrophoresis showed plasma CETP, as well as purified recombinant CETP, to be prebeta migrating. For determining the concentration of CETP in the media of cultured HepG2 cells, advantage was taken of the high sensitivity of the ELISA. CETP levels were found to increase linearly over the 100-h culture period, reaching 8.0 ± 0.4 ng/ml (18.0 ± 1.3 ng/mg cell protein). These sensitive, direct immunoassays for CETP mass should be valuable aids for examining the behavior of CETP in plasma and other complex systems, as well as for studying the synthesis and secretion of CETP by different cells and tissues. - Clark, B. W., J. B. Moberly, and M. J. Bamberger. Low level quantification of cholesteryl ester transfer protein of plasma subfractions and cell culture media by monoclonal antibody-based immunoassay. J. Lipid Res. 1995. 36: 876-889.

Cholesteryl ester transfer protein (CETP) catalyzes the transfer of cholesteryl esters (CE), triglyceride, and phospholipid between lipoproteins in the plasma of many animal species, including humans (1). Most plasma CE is generated on high density lipoproteins (HDL) through the action of lecithin:cholesterol acyltransferase (LCAT). CETP acts to redistribute a portion of this CE to triglyceride-rich lipoproteins, which are transformed via lipolysis to remnants and low density lipoprotein (LDL), or to LDL directly. Because this exchange of CE for triglyceride causes a net loss of CE from HDL, CETP has been proposed to play an important role in determining the balance between LDL- and HDL-cholesterol levels. That CETP can play such a role in vivo is suggested by the very high HDL levels observed for subjects genetically defective for CETP expression (2), and by the reduced HDL levels displayed by CETP transgenic mice (3). These contrasting states represent potentially antiatherogenic and atherogenic lipoprotein profiles, respectively, and call for more detailed studies on the role of CETP in intravascular lipoprotein metabolism, as well as its function in different cells and tissues.

Supplementary key words sandwich blotting • immunoblotting • native gel electrophoresis • prebeta migrating • Superose • Superdex • HepG2 cells

Abbreviations: CETP, cholesteryl ester transfer protein; CE, cholesteryl ester; TC, total cholesterol; CO, cholesteryl oleate; CMC, critical micelle concentration; VLDL/HDL, very low density lipoprotein/intermediate density lipoprotein; LDL, low density lipoprotein; HDL, high density lipoprotein; IRMA, immunoradiometric assay; MAbs, monoclonal antibody; FPLC, fast protein liquid chromatography.
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For CETP to transfer neutral lipid it must associate with the lipoproteins involved and bind the lipid to be transferred. Much effort has been expended to determine the degree to which CETP is associated with lipoprotein subclasses in the plasma and how factors such as surface charge, apolipoprotein content, plasma proteins, and ionic strength may be involved in such interactions. CETP interactions in whole plasma or in defined assays with isolated lipoproteins or synthetic vesicles have frequently been examined using gel filtration techniques to separate components followed by assessment of CETP content by CE-transfer activity (4-6). Other studies have examined CETP interaction with lipoproteins immobilized on Sepharose (7) or used both gel filtration and immobilized lipoproteins in separate experiments (8). In one investigation, plasma lipoprotein subclasses were specifically removed by apolipoprotein A-I and A-II immunoaffinity and the amount of CETP activity remaining in the non-binding plasma fraction was determined (9). Single dimension native polyacrylamide gel electrophoresis (PAGE) (10, 11) and two-dimensional agarose/PAGE (12) have also been used to characterize the CETP-associated species in human plasma with regard to size classes and surface charge. In none of these studies, however, was an attempt made to determine CETP mass. In studying the effects of anti-CETP antibodies on CETP binding to lipoproteins or vesicles in artificial systems, Swenson et al. (13) determined relative mass levels in chromatography fractions by sodium dodecyl sulfate (SDS)/PAGE followed by scanning of immunoblots, but did not determine actual concentrations. Likewise, demonstrations of the secretion of CETP by cells in culture have either relied solely on CE-transfer activity and inhibition of such by specific antibodies as described, or combined this evidence with that derived by immunoprecipitation and immunoblotting (17).

The lack of quantitative information on CETP mass in the above studies appears due in large part to lack of a sensitive and reliable assay for measuring CETP at the low levels found in chromatographic fractions and cell culture media. We have developed a series of monoclonal antibodies (MAb) to human CETP. Two MAbs, identified as 2F8 and 2E7, have been found to function well in double-MAb sandwich immunoassays. Immunoassays (IRMA) and ELISA assays have been standardized using 2F8 as the capture antibody and either iodinated or alkaline phosphatase-conjugated 2E7 as the detection MAb. Conditions for optimizing the detection of CETP under different conditions, including the selection and concentration of appropriate detergents, are described, as well as the use of these assays to measure CETP in gel filtration fractions and cell conditioned media.

The IRMA and ELISA for CETP described in this report are direct noncompetitive assays that take advantage of the high specificity and low background of monoclonal antibodies, and are unaffected by widely varying lipid and lipoprotein levels. For quantification of CETP at low levels, the ELISA provides a sensitivity 40-50 times greater than that previously reported for both indirect competitive (10) and immunoradiometric (18, 19) assays.

**MATERIALS AND METHODS**

**Materials**

Non-labeled lipids, alkaline phosphatase and, routine chemicals were purchased from Sigma. [3H]cholesterol olate was from New England Nuclear, 125I-iodine and 125I-labeled streptavidin were from Amersham. All chromatography resins, columns, and standards were obtained from Pharmacia, except butyl Toyopearl 650M which was from TOSOHAAS. Reagents for SDS-PAGE were from Bio-Rad, those for agarose electrophoresis were from Beckman (Paragon system), and precast acrylamide minigels for native protein electrophoresis were from Daiichi. For the enzymatic determinations of cholesterol and triglyceride, enzymes were purchased from Boehringer Mannheim. The microBCA protein assay kit, lodo-gen and all detergents were from Pierce. Costar High Binding flat-bottom, stripwell plates were used for the IRMA and ELISA. Cell culture media and other reagents were from Gibco, except supplements for hybridoma culture which were from Sigma.

Human hepatoma HepG2 cells were obtained from the American Type Culture Collection, Rockville, MD. Chinese hamster ovary (CHO) cells stably transfected with the human CETP cDNA were acquired from Columbia University and the laboratory of Dr. Alan Tall.

**Purification of CETP**

For the initial immunization of mice, CETP was partially purified from a d > 1.21 g/ml fraction obtained from human plasma after sequential ultracentrifugation. This plasma fraction was processed by FPLC using hydrophobic interaction chromatography followed by cation and then anion exchange. The plasma subfraction was loaded onto a butyl Toyopearl 650M column in 10 mM Tris, 500 mM NaCl, pH 7.4, and CETP was eluted with a linear gradient beginning with 50 mM Tris and ending with H2O (20). CETP activity was determined by measuring [3H]CE transfer from LDL to HDL. CETP-containing fractions were pooled, dialyzed against 50 mM acetate, pH 4.5, and loaded onto a CM-Sepharose column (21). CETP was then eluted using a 0-1.0 M NaCl gradient in acetate buffer. The pooled CETP fraction was then dialyzed against 10 mM Tris, pH 7.4, applied to a Mono-Q column, and eluted with 0-1.0 M NaCl. The three purification steps increased CETP specific activity by 175 x, 10 x, and 12 x, respectively.
resulting in a specific activity approximately 21000 times that of the starting material and a purity of 60–70%.

For calibration of the IRMA and ELISA and for other analytical studies, CETP was purified from cell culture media conditioned by CHO cells expressing full-length human CETP. Purification using hydrophobic interaction and anion exchange chromatography, as described above, was sufficient for obtaining CETP that appeared pure by SDS-PAGE/silver staining and by amino acid analysis (see Analytical Methods). A standard was compared to fresh lots of purified recombinant human CETP. Purification using hydrophobic interaction and anion exchange chromatography, as described above, was sufficient for obtaining CETP that appeared pure by SDS-PAGE/silver staining and by amino acid analysis (see Analytical Methods). A standard was compared to fresh lots of purified recombinant CETP every 3–4 months and found to be stable over a 1-year period.

**Preparation of lipoprotein substrates**

Labeled and unlabeled lipoprotein substrates were prepared according to the method described by Morton and Zilversmit (22). Blood was obtained from fasted donors using EDTA as anticoagulant. After centrifugation the plasma was adjusted to 3.5 mM N-ethylmaleimide to inhibit lecithin:cholesterol acyltransferase. To produce labeled lipoproteins, 3.0 ml of unilamellar liposomes, labeled with \[^3H\]cholesteryl oleate (CO), was added to 150 ml plasma. The liposome solution consisted of egg phosphatidylcholine, CO, triolein, and BHT at a mole composition of 82:12:5.6:0.4, respectively, and contained 4.0 mCi[^3H]CO (250 nmol). The label containing plasma was incubated for 18 h at 37°C to allow the endogenous CETP activity to incorporate [^3H]CO into all the lipoprotein classes. Lipoprotein subfractions were obtained from both labeled and unlabeled plasma by sequential ultracentrifugation using potassium bromide to adjust density. VLDL/IDL, LDL, and HDL were isolated from the d < 1.019 g/ml, the 1.019-1.063 g/ml, and the 1.10-1.21 g/ml ranges, respectively. Isolated lipoproteins were dialyzed extensively against 100 mM sodium phosphate, pH 7.4, containing 1 mM EDTA and 0.02% NaN₃.

**Assay of cholesteryl ester transfer**

For determining CETP activity of fractions isolated during purification of CETP, and for the screening of hybridoma supernatants for anti-CETP activity, an assay measuring [^3H]CE transfer from LDL (4 nmol CE) to HDL (2.5–5.0 nmol CE) was performed (170 µl assay volume). The source of CETP activity in assays of hybridoma supernatants was the d > 1.21 g/ml human plasma fraction (150 µg protein). Samples were incubated at 37°C for 1.0–18 h (20–30% transfer of label) and LDL was precipitated with Na₂H₇PO₄/MnCl₂ (100 mM/10 mM final concentrations). Radiolabel present in the HDL-containing supernatant, obtained by centrifugation, was determined by liquid scintillation.

To determine the relative CETP activity of different human plasma samples, an HDL to LDL transfer assay was set up in which the variation due to the lipoprotein content of the sample (2.5 µl plasma, 8–14 nmol total cholesterol) was minimized by using much larger amounts of exogenous donor [^3H]HDL (30 nmol TC) and acceptor LDL (125 nmol TC). For assay of transfer activity in HepG2 conditioned media the donor was [^3H]HDL (5 nmol TC) and the acceptor was a mix of VLDL/IDL (50 nmol TC) and LDL (25 nmol TC). For both assays acceptor lipoprotein was precipitated at the end of the incubation period by adding PO₄³⁻/MnCl₂ as described above.

The inhibition of CE-transfer by anti-CETP monoclonal antibodies in whole human plasma was determined by measuring [^3H]CO transfer from exogenous HDL (5 nmol CE) to the non-HDL lipoproteins of the sample. Non-HDL lipoproteins were precipitated at the end of the assay by addition of an equal volume of 20% (wt/vol) PEG 8000 (23).

**Preparation of anti-CETP monoclonal antibodies**

For the initial immunization, 10 µg of partially purified CETP in complete Freund's adjuvant was injected intraperitoneally into Balb/c mice. After 1 month the animals were boosted with 10 µg in incomplete adjuvant. Spleen lymphocytes were fused with mouse myeloma cells (SP2/0) using PEG 1000 (24). Media conditioned by the resulting hybridomas were screened for CETP inhibition in an assay measuring CE transfer from labeled LDL to unlabeled HDL (see above). Many media were screened simultaneously in an ELISA using immobilized partially purified CETP. Positive hybridomas were cloned by limiting dilution and were injected intraperitoneally into pristane-primed Balb/c mice for ascites production. After delipidation the ascites fluid was processed by Protein A Superose (Pharmacia) chromatography for isolation of purified monoclonal antibodies. For MAbs of the IgG₁ subclass, such as 2E7 and 2F8, a high salt buffer was used for binding to Protein A.

**IRMA and ELISA for determination of CETP mass**

A series of monoclonal antibodies to CETP were examined for their ability to function as capture and detection antibodies in double-MAb sandwich assays. Two antibodies, 2F8 and 2E7, served these functions well and allowed for the development of sensitive and linear assays. For the data shown in this paper, 2F8 functioned as the capture MAb and iodinated or alkaline phosphatase-labeled monoclonal antibodies. For MAbs of the IgG₁ subclass, such as 2E7 and 2F8, a high salt buffer was used for binding to Protein A.

[^3]Requests for samples of the 2F8 and 2E7 MAbs should be directed to R. W. Clark.
conjugated 2E7 as the detection MAb, although similar performance was seen with the antibodies in reverse roles. Use of the same MAb as both the capture and detection antibody resulted in complete elimination of CETP detection. This result, for both the 2E7 and 2F8 MAB, suggests that the determinants recognized by these MAbs are monovalent.

IRMA and ELISA assays were performed as follows. Capture MAb was added to Costar High-Binding plates at 1 μg protein/100 μl PBS (pH 7.8) per well. The plates were left at 4°C overnight. Free antibody was removed, the wells were rinsed 4 x with 0.05% Tween 20–PBS, and 0.05% Tween–1% BSA–PBS block solution was added. The block was left for 3 h at room temperature, then removed, and the wells were rinsed once. The samples, diluted in block solution, were then added at 100 μl/well and allowed to stand at room temperature for 2.5 h. The antigen solutions were removed, the wells were rinsed 4 x, and 2E7 detection MAb was added. For the IRMA, 0.225–0.25 μg 125I-labeled 2E7 (1.25–5.0 μCi/μg) was added per well; for ELISA 2E7–alkaline phosphatase conjugate was added (0.4 μg 2E7/well). After 1.25 h, free 125I-labeled 2E7 was removed, the plate was washed 4 x, and the individual wells were separated and counted. For the ELISA, after the 1.25-h incubation period and removal of free 2E7 conjugate, the wells were washed 3 x with Tween–PBS and once with 0.05% Tween 20–276 BSA–PBS, pH 7.8, detergent. After transfer the membranes were blocked with 0.05% Tween 20–2% BSA–PBS, pH 7.8, for 3 h, then incubated with iodinated 2E7 detection MAb as described in the legends for individual figures. For detection of apoA-I, untreated NC was used.

Analytical methods

FPLC was performed using a Pharmacia LKB system as described for the individual experiments. Free and total cholesterol were quantified by enzymatic methods using cholesterol oxidase and esterase from Boehringer Mannheim. Esterified cholesterol was calculated by difference between free and total cholesterol. Protein concentration was assayed using the Pierce microBCA assay kit and included heating at 60°C for 60 min to minimize protein to protein variation (27). BSA was used for the standard curve. A comparison of protein concentration determined for two purified samples of CETP by microBCA assay versus amino acid analysis was performed. The amino acid composition for both samples was shown to be in excellent agreement with the published composition for CETP (28). The microBCA gave results for the two samples of 51.3 and 359 μg/ml compared to 51.0 and 359 μg/ml by amino acid analysis. Therefore, determination of purified CETP mass by either method appears equivalent.

RESULTS

Development of IRMA and ELISA for determination of CETP mass

Monoclonal antibodies purified by Protein A affinity chromatography were tested in pairs for suitability in double-Mab sandwich IRMA and ELISA. Based on IRMA/ELISA studies, using partially purified human plasma CETP and highly purified CETP derived from transfected CHO cells (29) as antigen, two antibodies were selected. Identified as 2F8 and 2E7, these MAbs inhibit CE transfer in both whole human plasma, as shown in Fig. 1, and in dilute [3H]LDL–HDL assays using highly purified recombinant CETP. In both assays, at Mab:CETP molar ratios of 3:1, the antibodies inhibited CE transfer by over 90%. For calibration of the IRMA...
and ELISA, CETP was purified from culture media conditioned by Chinese hamster ovary cells stably transfected with human CETP cDNA (29). Purified CETP, processed by SDS-PAGE and visualized by silver staining (Fig. 2A), showed a 71/66 kilodalton doublet characteristic for CETP expressed by this cell line, with no other bands apparent. The purity of the CETP preparation was verified by amino acid analysis (see Material and Methods). In Fig. 2B–C, SDS-PAGE immunoblots show the specific detection of 73/68 and 71/66 kilodalton doublets for plasma and recombinant CETP, respectively, consistent with earlier reports (20, 29, 30).

In the experiment shown in Fig. 3 the IRMA and ELISA are compared. A serial dilution of CETP was made in Tween/BSA block solution and used to create standard curves of 0.5–20 ng CETP/well for the IRMA and 0.05–5 ng CETP/well for the ELISA. As shown in Fig. 3A, the IRMA is linear over the entire range and is suitable for the accurate determination of CETP mass down to levels of 0.5 ng CETP per IRMA well (5 ng/ml). For accurate quantification, this represents a several-fold increase in sensitivity compared to previously described immunocompetitive (10) and immunoradiometric (18, 19) assays for CETP. The ELISA is also linear (Fig. 3B), especially in the lower range of the standard curve after extended development. Using the ELISA, CETP levels as low as 0.05 ng CETP/well (0.5 ng/ml) can be quantified. In the same experiment, the level of CETP in a pooled human plasma sample was determined by the two assays and compared. The sample was serially diluted in ten steps to form a sample set of 0.1x to 0.00025x whole plasma concentration. The IRMA was then used to measure CETP levels for the seven samples at the high end of the dilution series (ca. 0.5–20 ng CETP/well) and the ELISA for the seven samples at the low end of the series (0.05–5 ng CETP/well). The mean CETP concentration (μg/ml ± SD) for the sample was determined to be 2.05 ± 0.06 by IRMA and 1.98 ± 0.10 by ELISA, demonstrating that the two assays yield equivalent results.

Because these assays were developed for measuring CETP from a variety of lipid-containing sources, the effects of lipids and lipoproteins on the IRMA were evaluated. Preincubation of purified CETP with liposomes containing phospholipid, cholesteryl ester, and triglyceride was found to greatly inhibit CETP detection, while the same treatment with various plasma lipoprotein subfractions had little or no effect (Fig. 4). Detergents, including Tween 20 and 80, Triton X-100 and X-114, Brij 56, CHAPS, octyl glucoside, and sodium cholate were tested, at concentrations above and below the CMC, for their ability to restore IRMA CETP detection in the presence of liposomes. CHAPS, when used at 0.75% in preparing the standard curves, was found to completely restore the IRMA signal and Tween 20 at 0.05% was also effective to a large extent (Fig. 4). Triton X-100 and X-114 at 0.025% performed similarly to 0.05% Tween 20 (data not shown), while other detergents proved less effective. We found the concentrations of Triton X-100 (0.5–1.0%), reported by Marcel et al. (10) to improve the specificity of a competitive RIA using the TP2 anti-CETP Mab and utilized in three other mass assays for CETP (18, 19, 31), not only to be unnecessary but caused a large reduction in detection of CETP by the 2F8/2E7 IRMA. This suggests that the antigenic sites for the 2E7 and 2F8 MAb are such that apoA-I does not compete with their binding to CETP as apoA-I under standard assay conditions.
Fig. 2. Purification of CETP and detection in conditioned media by western blotting. CETP was purified from culture media conditioned by transfected CHO cells expressing full length CETP in a two-step procedure using hydrophobic interaction and anion exchange chromatography. To assess purity, 1.0- to 1.5-µg aliquots were separated by SDS-PAGE using 5–15% gels and protein bands were visualized by silver staining (A). Lane 1 shows molecular weight standards, lanes 2–4 pooled CETP-containing fractions eluted from Q-Sepharose at mean NaCl concentrations of 170, 155, and 130 mM, respectively. The purified CETP doublet has an apparent molecular mass of 71/66 kDa (albumin ran at a higher molecular weight than expected relative to the other standards). For western blot detection of CETP in whole conditioned media (B), nitrocellulose (NC) sections were pretreated with the 2F8 capture Mab and blotting was conducted as described in Materials and Methods. After blocking, the blot was incubated with 75,000 and 3,400 cpm/cm² 1*5I-labeled 2E7 detection Mab and 125I-labeled streptavidin, respectively, for 1.5 h. For the experiment shown, 350 ng CETP (650 µg total media protein) was added to lane 1, and 2 µg biotinylated SDS-PAGE standards (Bio-Rad) to lane 2. To obtain similar intensities lane 1 was photographed after an 11-day film exposure, and lane 2 after a 3-day exposure. C. For comparison of recombinant CETP to human plasma CETP, aliquots of media and plasma were electrophoresed in adjacent lanes (C). Plasma CETP was partially purified by butyl toyopearl chromatography, and an aliquot containing 225 ng CETP and 27 µg total protein was applied. Whole CHO media (350 ng CETP) was applied to lane 2. Electrophoresis and immunoblotting were conducted as for B.

experiment was performed in which a plasma pool was supplemented with VLDL/IDL, LDL, or HDL plasma subfractions. The plasma was first diluted to 0.25x whole plasma concentration with 1% BSA/PBS block solution, containing either 0.05% Tween 20 or 0.75% CHAPS, supplemented with lipoprotein such that the added VLDL/IDL, LDL, and HDL concentrations were 100 and 50, 200 and 100, and 80 and 40 mg cholesterol/dl, as shown in Table 1. The samples were then diluted further, with block solution only, to 0.04, 0.02, and 0.01x whole plasma concentrations, and CETP levels were determined by IRMA. Values shown are the mean for the final three dilutions and indicate no significant effect of added lipoprotein on CETP determination by the IRMA. The results also demonstrate that use of 0.05% Tween 20 or 0.75% CHAPS gave equal results. Therefore, although there may be an advantage in using CHAPS as detergent when performing experiments with synthetic lipid vesicles or in other specialized studies, Tween 20 appears sufficient for routine assay of CETP in plasma. Omission of all detergent from the assay results in plasma CETP values 10–40% less than those obtained with Tween or CHAPS, indicating that detergent is required to fully expose antigenic determinants otherwise masked by interaction with lipid or other plasma proteins.

Use of the IRMA for determination of CETP concentration in plasma

Having calibrated the IRMA, CETP concentrations were determined in human plasma samples obtained from 44 normolipidemic individuals. Total and HDL-cholesterol levels for these fasted donors were 178 ± 32 and 46 ± 11 mg/dl, respectively. Mean plasma CETP levels for these subjects were determined by IRMA to be 2.10 ± 0.36 µg/ml. Males had an average value of 2.05 ± 0.33 µg/ml (n = 25), females 2.16 ± 0.40 (n = 19). Values ranged from 1.28 to 2.97 µg/ml and mass values correlated well with CETP activity (Fig. 5, r = 0.914). This mean value of 2.1 µg CETP/ml is similar to the 2.1 to 2.7 µg/ml reported by Fukasawa, Arai, and Inoue (31) for 20 normolipemic Japanese adults. It is also within 20–25% of the 1.7 µg/ml (n = 50) and 1.8 µg/ml (n = 79) determined by Marcel et al. (10) and McPherson et al. (32), respectively, using the same competitive RIA.

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Examination of plasma CETP distribution using sandwich immunoassays

We were interested in using these immunoassays not merely to determine CETP concentrations, but to study the protein’s interaction with lipoproteins in native plasma. Initially, plasma CETP distribution was examined by FPLC gel permeation chromatography. Superose 6 (Pharmacia) is an agarose-based medium that is often used for generating plasma lipoprotein profiles. It allows for the separation of VLDL, LDL, and HDL into distinct peaks (33), and has been used to study the location of cholesteryl ester transfer protein activity in human plasma (6). When human plasma was fractionated by Superose 6 chromatography using a typical 150 mM NaCl buffer system (Fig. 6A), the total recovery of CETP, determined by IRMA, was found to be incomplete (73%). A trailing of the CETP peak towards a size range much smaller than expected was also observed, suggesting that hydrophobic interactions of CETP with the agarose resin were occurring. Such interactions may explain the partial recovery of CETP activity (76%) reported previously (6) for Superose 6 chromatography. To minimize such interactions, a low ionic strength buffer consisting of 65 mM sucrose, 225 mM mannitol, and 2.5 mM Tris, pH 8.1 (SMT buffer) was tested. Using SMT buffer the recovery of CETP for the same plasma sample was 93% (Fig. 6B). Also, the peak for recovered CETP mass was centered at a molecular mass of approximately 150 kilodaltons, suggesting a complex larger than monomeric CETP alone. CETP has the potential to exist in the plasma as free monomeric protein or as HDL-bound or self-associated dimeric or oligomeric protein. To better resolve these potential forms, plasma was fractionated using Superdex 200. Superdex 200 is a dextran/agarose composite medium with a molecular weight size exclusion limit of $1.3 \times 10^6$ versus $40 \times 10^6$ for Superose 6. Although this resin is only part agarose, plasma fractionated using NaCl buffer yielded only 54% recovery of CETP mass (data not shown).
same high-HDL individual as for Superose study, but for plasma collected during a different month. The main CETP peak is centered at 183 kilodaltons. This is a somewhat greater size class than that seen for the Superose experiment and may be related to actual differences in CETP distribution for the samples collected at different times, and/or to differences in the degree to which CETP inacts with the Superose and Superdex resins at low ionic strength. It has been shown previously that the ability of purified CETP to associate with isolated lipoproteins (8) or with lipid vesicles (13) is greatly reduced in low ionic strength/hypotonic buffers. In contrast, the above results suggest that once CETP is already associated with lipoprotein, presumably HDL, the complex may in large part resist dissociation in a low ionic strength/isotonic buffer.

The distribution of CETP-containing particles in plasma was next studied by native agarose and acrylamide electrophoresis followed by western blotting. Double MAb sandwich blotting was performed as described in Materials and Methods. For agarose gel electrophoresis, approximately 5–7.5 ng CETP was loaded per lane (Fig. 7, top gel). The autoradiogram shows band intensity after a 2-day exposure. CETP present in whole plasma, a d > 1.21 g/ml plasma subfraction or purified and applied with albumin or apoA-I as carrier, migrates with prebeta mobility. An apoA-I western for the same samples (Fig. 7, bottom gel) shows both alpha- and prebeta-migrating bands. This prebeta mobility of plasma CETP differs

**Fig. 4.** Effects of liposomes and lipoproteins on detection of CETP by IRMA. A mixed multi- and unilamellar liposome preparation was formed by partial clarification of a lipid vesicle suspension by sonication. The liposomes consisted of egg phosphatidylcholine–cholesteryl oleate–triolein–BHT 81:7:12:0.6:0.04. Human VLDL/IDL, LDL, and HDL fractions were isolated by sequential centrifugation as described in Materials and Methods. For the experiment shown, purified CETP (100 ng/100 μl) was preincubated in 0.25% BSA/PBS containing liposomes, VLDL/IDL, LDL, or HDL at concentrations of 10, 30, 150, and 50 mg total cholesterol/dl. CETP in unsupplemented BSA/PBS served as the control. After 60 min at room temperature the samples were serially diluted to 10, 5, 2.5, and 1.0 ng CETP/100 μl using liposome containing (2 mg cholesterol/dl) BSA/PBS for liposome-preincubated samples and BSA/PBS only for the lipoprotein-preincubated samples. Diluent solutions contained no detergent or Tween 20 or CHAPS, such that all samples forming the standard curves contained either no detergent or 0.05% Tween 20 or 0.75% CHAPS. The IRMA was performed as described in Fig. 3, and the standard curves for samples supplemented with liposomes or lipoproteins (with or without detergent) were compared to those for samples lacking both supplements and detergent. No supplement, no detergent (●); (+) supplement, no detergent (▲); (+) supplement (+) Tween 20 (▲); (+) supplement (+) 0.75% CHAPS (▼).

**Table 1.** Lack of effect of added lipoprotein on IRMA determination of CETP in plasma

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<tr>
<th>Addition to 25% plasma</th>
<th>Plasma CETP Levels</th>
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<tr>
<td></td>
<td>μg/ml ± SD</td>
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<tr>
<td>Control, no addition</td>
<td>2.36 ± 0.09</td>
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<tr>
<td>VLDL/IDL, 100 mg/dl</td>
<td>2.45 ± 0.10</td>
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<tr>
<td>VLDL/IDL, 50 mg/dl</td>
<td>2.43 ± 0.10</td>
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<tr>
<td>LDL, 200 mg/dl</td>
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<td>LDL, 100 mg/dl</td>
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<td>HDL, 80 mg/dl</td>
<td>2.48 ± 0.04</td>
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<td>HDL, 40 mg/dl</td>
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Typically, for the IRMA determination of plasma CETP, the samples are diluted with block solution to 0.025–0.020× whole plasma concentration. In this experiment the potential effects of high concentration of lipoprotein were evaluated by supplementing a human plasma pool (180 mg total cholesterol/dl) with additional lipoprotein subfractions. Lipoprotein ± isotonic salt were added such that the initial 0.25x diluted plasma contained additional lipoprotein at the levels indicated. The samples were incubated at 37°C for 30 min, then serial-diluted with 1% BSA/PBS ± detergent to 0.04, 0.02, and 0.01x. IRMA results are the mean for the 0.04, 0.02, and 0.01x dilutions (assayed in duplicate) ± standard deviation. CETP present in the added VLDL/IDL and LDL was negligible, but was significant for the HDL-supplemented samples (18% of total for 80 mg/ml samples) and IRMA results are corrected for this supplemented CETP.
somewhat from the alpha-2 mobility described by Marcel et al. (10). The presence of apoA-I did not alter the pre-beta mobility of CETP but did greatly enhance detection by the 2E7 MAbs. It appears that cotransfer of CETP and apoA-I to nitrocellulose preserves the availability of the epitope for 2E7 recognition, that is otherwise lost when CETP alone is transferred. As both plasma-associated and purified CETP appear as pre-beta species, such single dimension agarose gels are uninformative in distinguishing between lipoprotein bound and free CETP. Native acrylamide gel electrophoresis, however, does demonstrate that CETP in whole plasma exists within a relatively narrow size class (Fig. 8A) compared to that of apoA-I (Fig. 8B). Based on the molecular weight standards, the bulk of CETP in the plasma from the six individuals tested in this experiment migrated at apparent molecular masses of 170-220 kilodaltons.

Use of the ELISA to quantify low levels of CETP secreted by HepG2 cells

Advantage was taken of the sensitivity of the ELISA to quantify low levels of CETP found in media conditioned by HepG2 cells (Fig. 9). The accumulation of CETP in the media of the CETP expressing CHO cells was also measured for comparison. The cells were allowed to condition the media for the periods indicated, after which the media was removed for CETP determination by IRMA (CHO cells) or ELISA (HepG2 cells). There was a near linear accumulation of CETP into the media of both cultures during the course of the experiment. Values are expressed as μg CETP/mg cell protein for CHO cell conditioned media, ng/ml for HepG2 media. As was seen earlier for human plasma samples, the mass of CETP in HepG2 conditioned media, determined by ELISA, closely reflected the ability of these samples to promote [3H]CE transfer (Fig. 5).

DISCUSSION

Sensitive IRMA and ELISA for CETP using monoclonal antibodies

Immunoradiometric (IRMA) and ELISA assays for measuring CETP mass have been developed using two different MAbs for antigen capture and detection. These specific anti-CETP MAbs inhibit cholesteryl ester transfer in assays using both human plasma and HepG2 cell conditioned media as the source of CETP, as well as purified CETP derived from transfected CHO cells. The specificity of the monoclonal antibodies is also demonstrated by western blot analysis using a crude human plasma fraction and whole media conditioned by transfected cells. In both cases detection is limited to a doublet typical for CETP from these two sources (20, 29, 30).

At 5 ng CETP/ml the IRMA achieves a CETP-specific signal nearly 20-fold greater than background, ensuring accurate measurements at this level or below. The ELISA shows excellent linearity between 0.5 and 5 ng CETP/ml and due to the low and stable background allows for accurate determination of CETP down to 0.5 ng/ml. Previously published mass assays for CETP have reported limits for accurate quantification of 20 ng/ml or greater (10, 18, 19). Therefore, the IRMA and ELISA described in this report offer an increased sensitivity of 4- to 40-fold, and should be of advantage for measuring CETP at the low levels typical for analytical fractions and cell culture media.
Measurement of human plasma CETP concentrations

We have used the IRMA to determine CETP concentrations in 44 normolipidemic adult subjects. The mean CETP level was 2.10 ± 0.36 µg/ml and ranged from 1.28 to 2.97 µg/ml. This is similar to that determined by Fukasawa et al. (31) (2.1-2.7 µg/ml), and close to the 1.7-1.8 µg/ml reported by Marcel et al. (10) and McPherson et al. (32). It is, however, nearly twice the 1.1 µg/ml value found by Ritsch et al. (18). The mass of purified CETP preparations was determined by amino acid analysis in the studies of Marcel et al. (10), Ritsch et al. (18), and ourselves. Therefore, it is unlikely that these differences in mean plasma concentrations are related to the method of protein estimation. Whereas our assay and those of Fukasawa et al. (31) and Marcel et al. (10) use MABs, this latter assay uses a polyclonal antibody preparation for both capture and detection of CETP. To obtain the highly purified CETP necessary for polyclonal generation, an approximate 30,000-fold enrichment from human plasma was required. If this lengthy purification process generated a partially denatured or altered preparation of CETP, then a portion of the polyclonal preparation could be directed against antigenic determinants not present in native plasma. This would be one explanation for an underestimation of plasma CETP using purified CETP as standard. Use of monoclonal antibodies avoids these possibilities, first by requiring only partially purified CETP for immunization and, second by the fact that once a monoclonal preparation is shown to interact with native...
plasma CETP, clonal specificity makes it unlikely that any other antibody in the preparation reacts otherwise. Recently an IRMA for CETP, using a monoclonal and oligoclonal antibodies was described by Moutour et al. (19). The TP4 MAb, homologous to TP2 (10, 19), was used for antigen capture, and peptide antibodies against residues 290–306 of CETP for detection. Mean plasma CETP concentration in 40 subjects was determined to be 1.1 ± 0.4 μg/ml by this assay. As the calibration curve was constructed using a CETP-rich fraction that was itself calibrated by the previously described competitive RIA (10), this value is a relative one (35% less than the RIA). Until this IRMA is calibrated independently with a pure preparation of CETP it remains unclear how its results compare to those derived by other assays.

When we began development of these MAb sandwich assays we had considered the possibility that the high specificity of monoclonals could itself be a problem because all forms of CETP might not contain the specific determinants for two different MAbs. If this were the case one would expect an underestimation of plasma CETP concentration. However, the mean value we have determined in human plasma is as high or higher than that reported using different assays. Of course, if the reactivity of the purified recombinant CETP for the MAb is less than that of plasma CETP, then plasma CETP concentration would be overestimated. For calibration of the IRMA and ELISA we chose to use highly purified CETP derived from the transfected CHO cells (29). These cells have been stably transfected with full-length wild-type cDNA, therefore, the expressed CETP is identical in amino acid sequence and total protein molecular weight to human CETP. The advantage in using the conditioned media as the source standard CETP is that CETP comprises about 0.5–1.0% of the total protein in the media versus approximately 0.003% in human plasma (2.1 μg CETP/70 mg protein per ml plasma). Thus highly purified CETP can be obtained using an abbreviated two-step purification protocol that increases the chances of obtaining unaltered, non-denatured protein. The CETP secreted by the transfected CHO cell line does differ slightly from plasma CETP in the extent of glycosylation. Western blotting indicated an apparent molecular mass of 71/66 kilodaltons for the cell derived CETP doublet versus 73/68 kilodaltons for plasma CETP (Fig. 2C), a difference similar to that observed by Inazu et al. (29). We feel that such minor differences are unlikely to cause unequal responses in the 2E7/2F8 sandwich assays. For example, when highly purified cell derived CETP was used to construct standard curves for the IRMA, identical curves resulted when different Q-Sepharose fractions were used, even though these fractions had different ratios of the 71 and 66 kilodalton forms of the protein. This suggests that the 5 kilodalton difference in glycosylation state did not affect the reactivity of CETP to the two antibodies.

One line of evidence consistent with our determination of a mean plasma CETP concentration of 2.1 μg/ml is the fold-purification reported by those preparing highly purified CETP from human plasma. Based on the enrichment in CE transfer activity of the purified CETP, Fukasawa et al. (31), Kato et al. (34), and Ritsch et al. (18) obtained 24,000-, 28,000-, and 30,000-fold purifications, respectively. Assuming a mean plasma protein concentration of 70 mg/ml this would indicate CETP plasma concentration of 2.3–2.7 μg/ml which is close to the value we have obtained. Ohnishi, Yokoyama, and Yamamoto (20) and Hesler, Swenson, and Tall (35) obtained CETP preparations showing 43,000- and 55,000-fold purifications, corresponding to CETP concentrations of 1.6 and 1.3 μg/ml. Although these extrapolated values are lower than those discussed above, they are still higher than the 1.1 μg/ml value determined by Ritsch et al. (18). Of course such values will be overestimated if CETP activity is lost...
Fig. 8. Detection of CETP in human plasma by native gel electrophoresis and western blotting. A: Samples and standards were prepared in Tris/glycine buffer and remained on ice until electrophoresis. Electrophoresis was performed for 4 h at 100 volts (room temp) at which point the tracking dye reached the bottom of the gel. Electrophoresis was then continued for an additional 1 h and 45 min. Transfer of proteins to 2F8 Mab pretreated nitrocellulose, blocking, and detection with 125I-labeled 2E7 was as described in Fig. 2. Lanes 1-2 show the Coomassie blue staining profile for 1.0 and 0.2 μl, respectively, of a whole plasma pool from six individuals; lane 3 shows 12.5 μg of native molecular weight standards (Pharmacia). The autoradiogram for lanes 4-9 from the same gel shows CETP detection for 1.0 μl of plasma from each of the six individuals. Individuals for the first three lanes had total cholesterol/HDL-C ratios < 4.0; those for the last three lanes had ratios > 4.5. CETP values for the six subjects, determined by IRMA, ranged from 1.24 to 2.42 pg CETP/ml with the lowest and highest CETP plasma run on lanes 6 and 7, respectively. Based on the molecular mass standards, the bulk of CETP in the plasma of these individuals migrated at apparent molecular masses of 220-170 kilodaltons. B. ApoA-I was detected in 1.0-μl plasma samples from the same six individuals using a gel run in parallel to that for CETP. Proteins were transferred to untreated nitrocellulose and apoA-I was detected as described above.

during the purification, and underestimated if an inhibitor of CETP activity (36, 37) were removed during the process. The extent to which differences in reported CETP concentration can be explained by the use of different assays versus actual variations in CETP levels for different populations remains to be determined.

Use of the IRMA and ELISA for measuring CETP levels in analytical fractions and cell culture media

We have examined the distribution of CETP in human plasma using both gel permeation FPLC and native gel electrophoresis. Chromatography of human plasma on Superose 6, followed by determination of CETP concentration by IRMA, demonstrated that use of a typical isotonic salt buffer led to incomplete recovery and trailing of CETP from the column. This result appears due to hydrophobic interactions between CETP and the agarose matrix of the column, and may explain the partial recovery of CETP activity previously reported for Superose 6 (6). Both partial recoveries and trailing were also noted using Superdex 200. Use of an isotonic, low ionic strength buffer system, however, achieved near total recovery of CETP using both Superose and Superdex chromatography. Plasma from a high-HDL individual, fractionated by chromatography on Superdex 200, showed a main CETP peak centered at 183 kilodaltons (Fig. 6C). The distribution of CETP in plasma was also studied by native agarose and acrylamide electrophoresis followed by western blotting. Plasma fractionated by agarose electrophoresis demonstrated that CETP was present as a prebeta migrating species (Fig. 7). However, both purified CETP and CETP present in a d > 1.21 g/ml plasma fraction exhibited prebeta mobility. Therefore, single dimension agarose electrophoresis cannot discriminate between lipoprotein bound and free CETP. Native acrylamide gel electrophoresis, however, did show that plasma CETP from six individuals migrated within a relatively narrow apparent size range of 170-220 kilodaltons (Fig. 8A). Considering that the molecular mass of plasma CETP itself is about 70 kilodaltons, this result is consistent with the association of CETP with small-sized HDL as has been suggested previously (10). This size range also encompasses that observed for peak CETP fractions separated on Superdex 200 (Fig. 6C). The FPLC plasma CETP profiles differ from those seen by native gel electrophoresis, however, in that a significant portion of CETP is eluted at a size range extending into that for human albumin. One explanation may be in spite of the near total recovery of CETP, sufficient interaction of CETP with the column resin is occurring to retard its elution. Also, displacement of a portion of lipoprotein-
body, assays were incubated in the presence of 3 mg/ml amount of 3H-labeled HDL and the amount of radiolabel transferred was calculated from the decrease in supernatant (HDL) counts. For testing the effects of an anti-CETP antibody, assays were incubated in the presence (▲) or absence (▼) of 3 ng/ml 2F8 MAb.

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REFERENCES


