Lipolytic Surface Remnants of Triglyceride-rich Lipoproteins Are Cytotoxic to Macrophages but Not in the Presence of High Density Lipoprotein

A Possible Mechanism of Atherogenesis?

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Abstract

Hypertriglyceridemic (HTG) serum, lipolyzed in vitro by purified bovine milk lipoprotein lipase, was found to be cytotoxic to cultured macrophages. Surviving macrophages contained numerous lipid inclusions similar to those found in foam cells. Individual lipoprotein fractions isolated from the lipolyzed HTG serum, including HDL, were also cytotoxic. Lipolysis of isolated lipoprotein fractions (either HTG or normal) allowed localization of cytotoxicity to postlipolysis remnant VLDL and chylomicron particles. The presence of a critical concentration of HDL in either the lipolysis mixture or the culture dishes inhibited the cytotoxicity. Below this critical concentration HDL itself became cytotoxic, producing lipid inclusions in surviving macrophages. The lipid fraction of the cytotoxic remnants contained the cytotoxic factor(s); neither FFA nor lyssolecithin alone could account for this cytotoxicity. Postprandial lipemic sera from subjects with a brisk chylomicron response, when lipolyzed in vitro, were cytotoxic to cultured macrophages; neither fasted sera from these subjects, nor postprandial sera from normolipidemic subjects with a normal chylomicron response, were cytotoxic. Postheparin (in vivo lipolyzed) serum and its isolated lipoprotein fractions obtained 30 min after heparin injection in subjects with HTG were shown to be cytotoxic to macrophages; by 60 min most of the cytotoxicity had disappeared. The postprandial and postheparin observations support an in vivo significance for remnant-associated cytotoxicity. We hypothesize that cytotoxic remnants of lipolyzed VLDL and chylomicrons may be one of the major atherogenic lipoproteins. Further, we suggest that inhibition of the cytotoxicity of these remnants may be one important way that HDL prevents atherosclerosis.

Introduction

Elevated levels of circulating lipoproteins have been implicated as risk factors for the development of atherosclerosis (1). Whereas most attention has been given to LDL as the atherogenic lipoprotein (2), considerable evidence exists that other lipoprotein species may be at least as important as LDL in the pathogenesis of atherosclerosis (3). The basic mechanisms for the putative atherogenicity of the different lipoproteins have yet to be clearly defined.

Dietary fat and cholesterol are absorbed from the intestine and secreted into the blood plasma in the form of triglyceride (TG)-rich chylomicrons. Most cholesterol synthesized in the liver enters the plasma in the form of TG-rich VLDL. The initial step of catalysis of both TG-rich lipoproteins in plasma is hydrolysis of the TG of the lipoproteins by lipoprotein lipase (LpL) bound to the endothelial surface of blood vessels (4). One consequence of this LpL-mediated hydrolysis of TG is a rise in the level of HDL through an episodic flux of excess surface components of TG-rich lipoproteins into the plasma HDL fraction (5).

The severity of atherosclerosis in patients with certain types of hyperlipoproteinemia and in cholesterol-fed rabbits has been correlated with the levels of circulating lipolytic remnants of TG-rich lipoproteins, including β-VLDL (6–10). β-VLDL and lipolytic remnants of TG-rich lipoproteins incubated in vitro with macrophages produce an accumulation of cholesteryl esters in these cells (11–14), mimicking the appearance of one of the hallmarks of the atherosclerotic plaque, the macrophage foam cell. Unmodified LDL incubated with macrophages under the same conditions fails to produce a similar cholesteryl ester accumulation (11, 12).

The pathogenesis of the atherosclerotic lesion involves an early stage in which the foam cell is the predominant lesion and a later stage in which necrosis is predominant. Published studies of the role of LDL or TG-rich lipoprotein remnants in atherogenesis have failed to explain adequately the inflammatory/necrotic nature of the atherosclerotic plaque. We show in this paper that lipolytic remnants of TG-rich lipoproteins are not only one cause of foam cell formation in macrophages but are also one cause of necrosis in these same cells, a necrosis that can be inhibited by HDL.

To examine the possible role of lipolytic remnants of TG-rich lipoproteins in atherogenesis, we produced lipolytic remnants of TG-rich lipoproteins in vitro by incubating HTG serum or isolated VLDL with purified bovine milk LpL or in vivo by injecting heparin into a hypertriglyceridemic (HTG) human subject. The interactions of these remnant products with cultured mouse peritoneal macrophages were then studied.

Methods

Materials. Fresh normolipidemic and HTG (type IV and V) plasma or serum was obtained from volunteers and from the Alabama Regional Blood Center, Birmingham, AL. Fresh serum was also obtained from

1. Abbreviations used in this paper: BHT, butylated hydroxytoluene; HTG, hypertriglyceridemic; IDL, intermediate density lipoprotein; LpL, lipoprotein lipase; SYS, single vertical spin; TG, triglyceride; VAP, vertical autoprobe.
normolipidemic volunteer subjects after an overnight fast (fasted serum) and 4 h after a fatty meal (postprandial lipemic serum). Two subjects, one with moderate hypertriglyceridemia (fasting TG and cholesterol concentrations of 660 and 308 mg/dl, respectively) and one with mild hypertriglyceridemia (fasting TG and cholesterol concentrations of 232 and 141 mg/dl), were recruited for the in vivo lipolysis studies. Lipoprotein cholesterol profiles in these plasmas were examined by the vertical autoprobe (VAP) method developed in this laboratory (15). VLDL, LDL, and HDL fractions were quantitatively fractionated from a portion of the above plasma by the single vertical spin (SVS) density gradient ultracentrifugation method (16). LpL was separated from raw bovine milk and purified by the heparin-Sepharose (Pharmacia Fine Chemicals, Piscataway, NJ) affinity chromatographic method described by Iverius et al. (17). Unstimulated macrophages were harvested from the peritoneal cavities of white mouse (CD1; Charles River Breeding Laboratories, Wilmington, MA) according to the method of Edelson and Cohn (18) and plated in 24-well culture dishes containing cover slips. Human peripheral monocytes and monocyte-derived macrophages were kindly provided by Dr. Marianne Egan at the University of Alabama. Monocytes were prepared from buffy coats of fresh blood by the method of Recalde (19). Cultured human umbilical vein endothelial cells, human skin fibroblasts and rat cardiac myocytes were kindly provided by Dr. Francois Booyse, Dr. Henning Birkenzal, and Dr. Larry Bugaisky, respectively, at the University of Alabama.

Lipolysis of serum and lipoproteins. Lipolysis of TG-rich lipoproteins in HTG or normolipidemic serum was performed in vitro by adding purified LpL into serum (50 μl/ml serum) and subsequently incubating the mixtures at 37°C for 60–90 min. For lipolysis of isolated lipoproteins in vitro, mixtures containing the isolated lipoproteins (40 mg/dl VLDL-cholesterol, 80 mg/dl LDL-cholesterol, or 30 mg/dl HDL-cholesterol) and 6% fatty acid–depleted BSA were incubated with purified LpL (50 μl/ml) for 60–90 min at 37°C. In certain experiments, butyalted hydroxytyolene (BHT) dissolved in ethanol was added to the lipolysis mixture to a concentration of 20 μM. Isolated HDL as an acceptor of lipolytic surface remnants of VLDL was also included in lipolysis mixtures containing HDL and albumin in certain experiments. The level of HDL in the lipolysis mixture was adjusted to give HDL-cholesterol to VLDL-cholesterol ratios of 0.25–2.0. Incubation mixtures containing heat-inactivated LpL (70°C for 20 min) served as the controls. The control and lipolyzed serum or pre- and postlipolysis mixtures containing lipoproteins and BSA were subjected to SVS density gradient ultracentrifugation to fractionate individual lipoprotein fractions or to separated lipoproteins from the BSA.

In vivo lipolysis of TG-rich lipoproteins in serum was performed by intravenous injection of heparin (50 IU heparin per kg body weight) into two fasting HTG volunteers. Blood samples were obtained from these subjects just before (control) and 30 min and/or 60 min after the injection of heparin, and serum was separated from the red blood cells. A portion of the control and lipolyzed serum samples were subjected to SVS density gradient ultracentrifugation to fractionate individual plasma lipoproteins.

All samples of serum and fractionated lipoproteins and lipoprotein-deficient plasma fractions were dialyzed against buffered saline (0.05 M sodium phosphate-0.15 M NaCl, or 0.01 M Tris-0.15 M NaCl, pH 7.4) to remove KBr and other salts and were then sterilized by filtering them through 0.45-μm membranes (Millipore Corp., Milford, MA). In certain experiments, the antioxidant, butyalted hydroxytyolene (BHT) was included in dialysis buffers.

Interaction of lipolyzed serum and lipoproteins with macrophages. Control or lipolyzed serum or lipoproteins (5–150 μg cholesterol) were added to culture dishes containing ~10⁶ resident macrophages and 1 ml serum-free DME (Gibco Laboratories, Grand Island, NY), and the dishes were incubated at 37°C in a humidified incubation chamber having 5% CO₂ in air for 18 h. At the end of incubation, cover slips containing cell monolayers were removed from the dishes and were washed briefly with buffered isotonc saline to remove the residual lipoproteins or serum components from the cover slips. Cells on the cover slips are then stained with trypan blue by covering the cover slips with 0.4% trypan blue solution for 3–4 min. After washing off the trypan blue with buffered saline, cover slips were mounted on glass slides, and the number of stained cells and nonstained cells in the several microscopic fields were counted (20). The viability of other cells in monolayer culture (fibroblasts, endothelial cells, and cardiac myocytes) was determined as described for macrophages. The viability of cells in suspension culture was determined as follows: the cells in culture medium were collected by centrifugation of the cell suspensions in conical tubes at 1,000 rpm for 5 min. The supernatant fluid was siphoned off and discarded. 0.5 ml of culture media devoid of serum components and 0.1 ml of 0.4% trypan blue were added to the cell pellet. The mixture was dispersed by use of a Pasteur pipette, and a drop of the suspension was placed on a hemocytometer. The number of stained cells and nonstained cells in a given area on the hemocytometer were counted after 3 min staining of the cells. A minimum of 100 cells were counted.

In certain occasions upon which large number of cells detached from the cover slips, the total number of cells in several microscopic fields of the cover slips from the dishes containing control and lipolyzed was counted and compared. The index of cytotoxicity of lipoproteins to the cells was expressed as a percentage of cells that have excluded trypan blue and a percentage of the cells detached from the cover slips at the termination of the incubation. The inclusion of lipid drops within the cytoplasm of cells was determined after staining the cells with oil red O.

Analysis of lipolyzed serum or lipoproteins. The extent of hydrolysis of TG in lipolyzed samples was determined by measuring TG by the enzymatic TG assay kit (Boehringer-Mannheim reagent set 348202; Biodynamics/BMC Co., Indianapolis, IN) in the pre- and postlipolysis samples after extensive dialysis of the samples. Cholesterol concentration was assayed by the enzymatic method with the reagent set (Set No. 12408; Boehringer Mannheim Biochemicals). FFA concentrations in lipolyzed serum or lipoproteins were assayed by the colorimetric method of Itaya (21). Concentrations of oxidized lipids in lipolysis samples of lipoproteins or serum were estimated by assayng the level of thiobarbituric acid–reacting material as described by Kosuki et al. (22). Lysolecithin was quantitated by measuring phosphorus-containing lipid (23) after separation of lysolecithin from the other phospholipid classes by TLC (24).

Lipid from control and lipolyzed serum or VLDL was extracted with a mixture of chloroform:methanol (2:1, vol/vol). Organic solvents in a portion of the lipid extract was evaporated by use of nitrogen gas, and liposomese were then prepared by sonication the lipid extract after addition of buffered saline. The other portion of the lipid extract was applied to an HPLC containing latex beads (Astec Co., Whippany, NJ), and the column was eluted sequentially with the mixtures of hexane/chloroform/methanol/water/phosphoric acid (28:2:60:300:30:4), methanol/water (88:12), and gradient mixtures of cyclohexane and ethyl acetate to fractionate the different classes of lipid. This fractionation was kindly performed by Dr. Walter Shaw at Avanti Polar Lipids, Inc., Birmingham, AL.

Liposomes containing either FFA, lysolecithin, or an individual lipid fraction obtained from the lipid extracts of control and lipolyzed VLDL were prepared by mixing the lipid component with egg phosphatidyicholine and sonicating the mixtures as described earlier. Apolipoproteins in pre- and postlipolysis samples of VLDL were examined by SDS gradient gel electrophoresis (25). The morphology of lipolytic remnants of VLDL was examined by negative staining with 2% potassium phosphotungstate and examining the grids on a Philips 400 transmission electron microscope.

Results
Cytotoxicity of in vitro lipolyzed serum and its lipoproteins to macrophages. Incubation of HTG serum in vitro with purified LpL for 90 min at 37°C usually resulted in the hydrolysis of
> 70% of the TG in serum. Incubation of this lipolyzed serum with peritoneal macrophages produced an unexpected result. Compared with control serum (incubated with heat-inactivated LpL) the lipolyzed serum was highly cytotoxic to the cells. More than 90% of the macrophages were killed by lipolyzed serum containing 5–10 μg cholesterol per culture dish; control serum at concentrations of up to 60 μg per dish had no detectable effect on cell viability (Fig. 1A).

Figure 1. Effects on mouse peritoneal macrophages of HTG plasma lipolyzed in vitro with purified bovine milk LpL. (A) Viability of macrophages incubated with lipolyzed HTG plasma (▲) versus control HTG plasma (●). Aliquots of the control or lipolyzed serum containing 1–60 μg cholesterol were added to culture dishes containing 1 ml culture medium. After an 18-h incubation of the dishes at 37°C in a humidified incubation chamber, the viability of the cells was examined by counting the percent of cells excluding trypan blue. (B) Changes in viability of macrophages during the incubation after an addition of 10 μM cholesterol of control (-) or lipolyzed (▲) serum to culture dishes. The culture dishes were withdrawn at the indicated time and the viability of cells was measured as described. (C) Appearance by phase-contrast light microscopy of macrophages after incubation with HTG plasma. (Left) Control HTG plasma (10 μg/dish); (center) lipolyzed HTG plasma (10 μg/dish); (right) lipolyzed HTG plasma (2 μg/dish). The cells were placed on glass cover slips and stained with trypan blue. (D) Viability of macrophages incubated with lipolyzed normolipidemic plasma supplemented with preisolated VLDL. Normolipidemic plasma (VLDL cholesterol < 10% of total plasma cholesterol) was supplemented with preisolated authentic VLDL equal to 0% (control) (●), 10% (▲), 20% (●), and 40% (▲), of the total plasma cholesterol. The supplemented plasma was incubated with active (bottom) or heat-inactivated (top) LpL. Aliquots of the mixtures (10, 30, or 60 μg cholesterol/dish) were then incubated with macrophages as described above.
approximately 50% of the cells were killed within 2 h and 100% of cells were killed within 9 h after addition of the lipolyzed serum to the culture dishes (Fig. 1 B).

Light microscopic examination of cultured macrophages incubated with a sublethal dose of lipolyzed serum (2 µg cholesterol/dish) showed the presence of numerous lipid inclusions, resembling those seen in foam cells, within the cytoplasm of the surviving macrophages (Fig. 1 C).

Normolipidemic serum was lipolyzed under identical conditions as HTG serum and incubated with macrophages. Neither cytotoxicity nor foam cell formation was observed (Fig. 1 D). However, when this plasma was supplemented with preisolated VLDL and the supplemented plasma incubated with LpL, the resultant product was both cytotoxic and foam cell inducing; cytotoxicity was found to be dependent on the level of VLDL supplementation (Fig. 1 D).

Control and lipolyzed HTG serum were fractionated by SVS into VLDL, LDL, HDL, and free protein, and each of these fractions incubated with macrophages. All of the lipoprotein fractions from in vitro lipolyzed plasma (10 µg cholesterol) were both cytotoxic and foam cell inducing (Fig. 2 A); the free protein fraction was not cytotoxic but induced a modest number of lipid inclusions. Corresponding control plasma fractions, at four times greater concentrations (40 µg cholesterol), were neither cytotoxic nor foam cell inducing (Fig. 2 A). Based on the relative cytotoxicity of these individual lipoprotein fractions, HDL was the most cytotoxic fraction in the lipolyzed HTG serum (Fig. 2 B).

Cytotoxicity of postprandial lipemic serum from normolipidemic subjects. Ingestion of a fatty meal by normotriglyceridemic subjects resulted in an ~380 mg/dl increase in plasma TG and a 10 mg/dl increase in plasma cholesterol 4 h after the meal in the subject with a brisk chylomicron response but resulted in only a 33 mg/dl increase in TG with little or no change in plasma cholesterol in the subject with a normal chylomicron response (Fig. 3 A). The increase in cholesterol was associated exclusively with an increase in cholesterol in the VLDL density region of the plasma (Fig. 3 A). In vitro lipolysis produced little or no cytotoxicity in fasted or postprandial serum from the normal responder and in fasted serum from the brisk responder. However, the postprandial serum from the brisk responder after in vitro lipolysis was highly cytotoxic to cultured macrophages (Fig. 3 B).

Cytotoxicity of in vivo lipolyzed serum and its lipoproteins to macrophages. Heparin injection into mild and moderately severe HTG subjects resulted in hydrolysis of 65 and 68% of serum TG within 30 min after injection, respectively. The extent of serum TG hydrolysis 60 min after injection of heparin in the moderately severe HTG subject was similar to that seen at 30 min (68 vs. 71%). Lipoprotein cholesterol profiles of pre- and postheparin serum showed that ~40–50% of VLDL-cholesterol in serum from the moderately severe HTG subject (Fig. 4 A) and ~75% of VLDL cholesterol in serum from the mildly HTG subject (Fig. 4 D) were moved into the intermediate density lipoprotein (IDL), LDL, and HDL density regions.

When pre- or postheparin serum was added to the cultured macrophages, the postheparin serum obtained at 30 min from the moderately severe HTG subject was highly cytotoxic and foam cell inducing; >90% of the macrophages were killed with 10 µg (cholesterol) of serum (Fig. 4 B). The preheparin serum, even at five times the concentration of postheparin serum, was...

Figure 2. Cytotoxicity of lipoproteins isolated from in vitro lipolyzed HTG serum to macrophages. (A) Viability of macrophages incubated with individual lipoprotein fractions isolated by density gradient ultracentrifugation from pre- and postlipolysis HTG plasma. The pre- and postlipolysis plasma samples were subjected to SVS ultracentrifugation, and aliquots containing 40 µg cholesterol (nonlipolyzed plasma) and 10 µg cholesterol (lipolyzed plasma) were removed from the even numbered gradient fractions and added to the culture dishes of macrophages. Aliquots of fractions 2 and 4, containing lipoprotein-free plasma proteins, were based on volume (50 µl). Cell viability was then measured after 18 h incubation as described earlier. (B) Viability of macrophages incubated with varying levels of VLDL (•), LDL (●), and HDL (▲), separated from in vitro lipolyzed HTG serum. Viability after incubation with control lipoproteins separated from unlipolyzed serum is indicated with corresponding open symbols.

containing a high concentration of the lipolyzed serum (>40 µg/dish) a large number of cells in the culture dish were detached from the cover slips and presented in the culture dish as cell debris (data not shown).
not cytotoxic to the cells (Fig. 4 B); only a few lipid inclusions were detectable in these cells. The postheparin serum obtained at 60 min was only marginally cytotoxic to the cells (Fig. 4 B), although the extent of hydrolysis of TG in this serum was slightly greater than that in the postheparin serum obtained at 30 min. These results suggest that the cytotoxic components of lipolyzed serum can be produced in vivo but are removed rapidly from the circulation. However, macrophages incubated with postheparin serum obtained at 60 min had approximately as many lipid inclusions as macrophages incubated with 30 min postheparin serum (data not shown). Postheparin serum from the mildly HTG subject was also cytotoxic to cultured macrophages, but the cytotoxicity of this serum was considerably less than the postheparin serum from the moderately severe HTG serum (Fig. 4 E).

Pre- and postheparin HTG sera from the moderately severe HTG subject were fractionated by SVS into VLDL, LDL, HDL, and free protein and each of these fractions incubated with macrophages. HDL, IDL, and to a lesser extent VLDL were considerably more cytotoxic than the LDL fraction (Fig. 4 C). However, cells incubated with LDL fraction were shown to contain numerous lipid inclusions (data not shown). The lipoprotein-free plasma fraction from the postheparin serum was not cytotoxic but caused a few lipid inclusions within the cells (Fig. 4 C). None of the preheparin HTG serum fractions were cytotoxic (Fig. 4 C).

**Figure 3.** Effects of postprandial lipemia on lipoprotein cholesterol profiles and cytotoxicity of lipolyzed serum. (A) VAP lipoprotein cholesterol profiles of fasting and postprandial lipemic serum: brisk chylomicron responder (left) and normal chylomicron responder (right). (B) Viability of macrophages incubated with (top) prelipolysis serum or (bottom) postlipolysis serum: fasting (○) or postprandial lipemic (△) serum from the brisk responder and fasting (●) or postprandial lipemic (◆) serum from the normal responder.
Figure 4. Effects of postheparin in vivo lipolysis on serum and its lipoproteins from two subjects with differing degrees of HTG. Subject with moderately severe HTG. (A) VAP lipoprotein cholesterol profiles of preheparin and postheparin serum. (B) Viability of macrophages incubated with pre- or postheparin serum. Culture dishes containing increasing amounts of preheparin serum (●), or postheparin serum obtained at 30 min (●), and 60 min (▲), were incubated for 18 h and the viability of cells was determined. (C) Viability of macrophages incubated with various lipoprotein fractions isolated from pre- and postheparin HTG serum. Culture dishes containing increasing amounts of VLDL (●), LDL (▲), HDL (○), or 200-μl lipoprotein-free serum (●) obtained from postheparin (30 min) serum were incubated for 18 h and the viability of the cells was measured. Viability after incubation with control lipoproteins separated from preheparin serum is indicated with corresponding symbols and dotted lines. Subject with mild HTG. (D) VAP lipoprotein cholesterol profiles of preheparin and postheparin serum. (E) Viability of macrophages after incubation with preheparin serum (●), or postheparin serum (▲).

HTG and normal plasma is a lower HDL to VLDL ratio in HTG. Therefore, we considered the possibility that HDL would inhibit the cytotoxicity of lipolyzed remnant VLDL at higher HDL to VLDL ratios.

Effects of HDL on macrophage viability. To test the possibility that HDL would inhibit the cytotoxicity of lipolyzed remnant VLDL at higher HDL to VLDL ratios, we produced the lipolytic remnants of VLDL in the presence of increasing concentrations of HDL in the lipolysis mixture, and the cytotoxicity and structure or composition of the VLDL remanants were further determined. As shown in Fig. 6 A, HDL did indeed reverse the cytotoxicity of lipolyzed remnant VLDL. At HDL to VLDL ratios of 1.0 or lower, both the remnant VLDL and the HDL isolated from the lipolysis mixtures were cytotoxic, killing 60% or more of the cells. However, at ratios of 2.0 or greater, the cytotoxicity of both the remnant VLDL and the
postlipolysis HDL fractions approached that of the controls. Normolipidemic plasmas have HDL to VLDL ratios of four or greater. When HDL was added directly to the culture dishes (Fig. 6B), inhibition of cytotoxicity was even more impressive; complete protection, comparable to controls, was provided by the addition of HDL at a HDL to VLDL ratio of 2.0. The addition of LDL directly to the culture dishes at up to six times the effective concentration of HDL had no measurable effect on remnant VLDL cytotoxicity (Fig. 6B), indicating that inhibition of cytotoxicity is unique to HDL among the lipoprotein classes.

Additional experiments indicated that serum albumin at twice physiological concentrations provided minimal protection (20% viability) from cytotoxicity (Fig. 6A). Because the levels of albumin in plasma are less variable than those of TG-rich lipoproteins or HDL, because the physiological levels of albumin in whole HTG plasma failed to provide protection from post-lipolysis cytotoxicity, and because the postheparin HTG serum was cytotoxic at 30 min, the protection provided in vivo by albumin may be of considerably less physiological significance than that provided by HDL.

Characterization of cytotoxic components. As a first step in characterization of the cytotoxic factor(s) in lipolyzed VLDL, we extracted the lipid of control and lipolyzed VLDL with the mixtures of chloroform-methanol (2:1, vol/vol). After removal of the solvent and sonication of the lipid in aqueous suspension, the two lipid extracts were incubated with macrophages. The lipid from the lipolyzed VLDL on a per microgram cholesterol basis was somewhat more cytotoxic as compared with that of the original lipolyzed VLDL, whereas the lipid from the control VLDL was essentially noncytotoxic (Fig. 7A). The lipid-free apoprotein components of both VLDL fractions were noncytotoxic (data not shown).

We have begun a preliminary characterization of the factor(s) in the lipid components of the VLDL remnants (presumably localized to the vesicular surface material) responsible for macrophage cytotoxicity. A number of studies have suggested that LDL-containing peroxidized lipid is cytotoxic to cells in culture (27, 28). We have examined the level of oxidized lipids in control and lipolyzed VLDL or HTG serum and determined the effect of including antioxidant (BHT) during lipolysis and dialysis of the sample. We found that lipolysis has no effect on the level of oxidized lipid, as judged by the changes in level of thiobarbituric acid–reacting material in the dialyzed samples of control and lipolyzed VLDL or HTG serum (Table I). Further, inclusion of BHT during lipolysis and subsequent dialysis of the lipolyzed HTG serum had no effect on the cytotoxicity of lipolyzed HTG serum (Table I).

FFA and lysolecithin are known products of lipolysis by LpL (29) and are potentially cytotoxic. Unesterified cholesterol has been implicated as a potential cytotoxic and atherogenic substance (30). We have examined the levels of FFA, lysolecithin, and unesterified cholesterol in lipolyzed VLDL and the cytotoxic effect of these lipids to cultured macrophages.

The levels of FFA and lysolecithin in the remnants of VLDL produced in the presence of a physiological level of albumin in the lipolysis mixture are three to four times greater than those in control VLDL. The concentration of FFA associated per milligram cholesterol of control VLDL and remnant VLDL was 40 and 176 µg, respectively, and the concentration of lysolecithin was 23 and 73 µg, respectively. The level of unesterified cholesterol in VLDL remnants was not detectably different from that in control VLDL.

When the level of lipoproteins in culture dishes were standardized based on the contents of FFA or lysolecithin, only lipolyzed VLDL was cytotoxic to the macrophages (Fig. 7B). Sonicated egg lecithin liposomes containing FFA, lysolecithin, or unesterified cholesterol were not cytotoxic to the cultured macrophages even at two to three times the concentration of FFA, lysolecithin, or unesterified cholesterol in lipolyzed VLDL (Fig. 7B). The liposomes containing FFA were, however, foam cell producing (data not shown).
Analysis of FFA in pre- and postheparin HTG serum showed that the 30-min sample contained less FFA (556 \( \mu \text{mol/dl} \)) than the 60-min sample (612 \( \mu \text{mol/dl} \)) but was significantly more cytotoxic than the 60-min sample (Fig. 4 B). Further, the 60-min postheparin serum contained six times as much FFA as the preheparin control serum (124 \( \mu \text{mol/dl} \)) but showed very little cytotoxicity to macrophages (Fig. 4 B). Analysis of FFA concentration in the plasma fractions of pre- and 30-min postheparin serum showed that the plasma free protein fraction contained most of the FFA increase in the serum (~ 50%) but was not cytotoxic to the macrophages (Fig. 4 C).

At physiological concentrations, albumin and HDL were found to be equally effective in removal of both FFA and lysolecithin from VLDL remnants. Because HDL is considerably more effective than albumin in inhibition of cytotoxicity (Fig. 6 A), neither FFA nor lysolecithin seem likely to be the cytotoxic agent(s). Preliminary characterization of the cytotoxic component(s) in the lipid extract of lipolyzed VLDL by HPLC indicates that component(s) having a solvent mobility between phospholipid and FFA are most cytotoxic to the macrophages. The identity of these component(s) is presently unknown but are currently under investigation in this laboratory.

**Discussion**

We demonstrate in this study that lipolysis of TG-rich lipoproteins in vitro or in vivo results in remnants that are cytotoxic and foam cell inducing when incubated with cultured macrophages. Although incubation of isolated LDL or HDL with LpL fails to cause these lipoproteins to become cytotoxic, lipolysis of whole serum can cause the HDL fraction, and to a lesser degree, the LDL fraction, to become cytotoxic and foam...
cell inducing. This effect on HDL seems likely to be due to enrichment of HDL with surface remnant products of lipolyzed TG-rich lipoproteins.

In preliminary studies, a number of cell types in addition to mouse peritoneal macrophages have been examined (Table II). Lipolyzed TG-rich lipoproteins and HTG serum are cytotoxic to human monocyte–derived macrophages but, interestingly, not to the monocytes themselves. Further, lipolyzed VLDL is cytotoxic to the macrophage-like J774 cell line but only at a dose seven times higher than needed to kill mouse or human macrophages (Table II). Lipolytic remnants are not cytotoxic to human skin fibroblasts and rat cardiac myocytes.

Finally lipolyzed VLDL is cytotoxic to cultured human umbilical cord endothelial cells (31). Because injury of the arterial endothelial cell is considered an early event in the development of atherosclerosis (32), these preliminary studies suggest the possibility that the cytotoxic and foam cell–inducing lipolytic remnants of TG-rich lipoproteins may play a role in the development of atherosclerosis. It is apparent from Table II that lipolyzed lipoproteins and serum appear to be less cytotoxic to endothelial cells than to macrophages. It is also apparent that the in vitro or in vivo (postheparin) lipolysis experiments described here are exaggerations of both the rate and concentration of products of in vivo lipolysis. However, it must be remembered that atherosclerosis under most conditions is a subtle and slowly progressing disease. The chronic nature of this disease is, therefore, entirely compatible with an atherogenic role for lipolytic remnants.

What is the molecular basis for this cytotoxicity? Among the possibilities are the following: (a) FFA and lysolecithin, perhaps together with other lipid components, may have synergistic cytotoxic effects. We consider this possibility unlikely. (b) Unusual species of FFA and/or lysolecithin not present in the commercial preparations used in our liposome cytotoxicity studies may account for the cytotoxicity of lipolyzed TG-rich lipoproteins. This possibility will be explored further. (c) The physical chemical organization of the surface remnants of lipolyzed TG-rich lipoproteins may increase the sensitivity of macrophages to the cytotoxic effects of FFA, lysolecithin, or other compounds localized to the surface remnants such as...
peroxidized lipids. This is an appealing possibility considering the ability of HDL to provide protection against cytotoxicity and change the physical organization of surface remnants of lipolyzed TG-rich lipoproteins. (d) Lipid components other than FFA and lysolecithin, may be the primary mediators of cytotoxicity in the lipolyzed TG-rich lipoproteins.

Our data do not support peroxidized lipids as the cause of lipolysis-induced cytotoxicity because lipolysis has no measurable effect on the level of oxidized lipids in VLDL and/or HTG serum and inclusion of antioxidant (BHT) during lipolysis and dialysis of samples failed to inhibit cytotoxicity (Table I). However, we can not rule out at present the possibility that lipolyzed VLDL is more susceptible to cell-mediated oxidation than control VLDL.

It has long been suspected that lipolytic remnants of TG-rich lipoproteins in animals on an atherogenic diet are responsible for the development of atherosclerosis in these animals (10). Several studies by Kruth (33, 34) have shown that early atherosclerotic lesions in both humans and experimental animals contain numerous particles rich in unesterified cholesterol but poor in cholesteryl ester. These particles are seen even before the appearance of foam cells in the lesions. Recent studies by Simionescu et al. (35) showed that prefoam cell atherosclerotic lesions in cholesterol-fed rabbits contain numerous liposome-like intimal vesicles rich in phospholipid and unesterified cholesterol. Although these authors did not determine the origin of these vesicles, the morphology and lipid composition of the vesicles are suggestive of the TG-rich lipoprotein surface remnants that we propose are cytotoxic and foam cell inducing both in vitro and in vivo. Similar liposome-like intimal lipid deposits have been reported before in both humans (36) and other animal models of atherosclerosis, such as baboons and dogs (37, 38). Furthermore, the lipid incursions in the foam cells of both humans and animal models of atherosclerosis have been reported to contain liposome or surface remnant-like myelin bodies (36, 37).

Humans spend a great deal of time, albeit episodic, in the postprandial HTG state. We showed in Fig. 3 that postprandial serum, but not fasting serum, when lipolyzed in vitro, is cytotoxic to cultured macrophages (Fig. 3). Lipolysis of dietary-induced TG-rich lipoproteins occurs to a large extent at the LpL binding sites on the endothelial surface of blood vessels. Although not much is known about the density of LpL binding sites on the endothelial surface of large-to-medium-size arteries where the atherosclerotic plaque occurs, there is a definite possibility that an episodic concentration gradient of potentially atherogenic TG-rich remnants exists at the endothelial surface of these arteries. Under these conditions, the remnant to HDL ratio at the endothelial surface would be expected to be much higher than in the circulation. It may be significant that endothelial cells have recently been reported to have a higher concentration of the putative HDL receptor than any other cell type examined (39). Perhaps a function of the HDL receptor is to raise the HDL concentration at the endothelial cell surface to match that of cytotoxic TG-rich remnants.

A surface remnant model for atherogenesis has three interesting implications: (a) recent studies (40) support the notion that early atherosclerosis in humans begins in regions of the coronary artery subject to low hemodynamic shear stress. It is well known that arterial branch points, for example, the aortic intercostal ostia, are highly susceptible to early atherosclerotic plaques. A simple explanation for this correlation is that there is a concentration gradient of atherogenic (i.e., cytotoxic) substance(s) extending outward from the endothelial surface of the arterial wall; high shear stress would be expected to disrupt this gradient.

(b) Inherited or acquired regional differences in density and/or structure of the LpL binding sites or the HDL receptor on the endothelial surfaces of large-to-medium-size arteries could explain the otherwise unexplained individual variations

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**Table I. Relationship of Oxidized Lipids to Cytotoxicity**

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<th>Lipoproteins</th>
<th>Level of oxidized lipids</th>
<th>nM MDA/mg cholesterol</th>
<th>Cell viability (% of control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control VLDL</td>
<td>10</td>
<td>4.54</td>
<td>83</td>
</tr>
<tr>
<td>Lipolyzed VLDL</td>
<td>25</td>
<td>3.65</td>
<td>18</td>
</tr>
<tr>
<td>Control HTG serum</td>
<td>50</td>
<td>2.10</td>
<td>90</td>
</tr>
<tr>
<td>Lipolyzed HTG</td>
<td>100</td>
<td>1.90</td>
<td>0</td>
</tr>
<tr>
<td>+ BHT</td>
<td>500</td>
<td>1.79</td>
<td>6</td>
</tr>
</tbody>
</table>

Values of MDA are the means of duplicate determinations.

---

**Table II. Cytotoxicity of Control and Lipolyzed VLDL to Various Cultured Cells**

<table>
<thead>
<tr>
<th>Cells</th>
<th>Control VLDL</th>
<th>Lipolyzed VLDL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>25 50 100</td>
<td>10 25 50 75 100</td>
</tr>
<tr>
<td>Mouse peritoneal</td>
<td>92 93 89</td>
<td>9 0 0 --- 0</td>
</tr>
<tr>
<td>macrophages</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human macrophages</td>
<td>90 92 90</td>
<td>30 11 0 --- 0</td>
</tr>
<tr>
<td>Human blood monocytes</td>
<td>100 100 98</td>
<td>95 94 90 --- 90</td>
</tr>
<tr>
<td>J 774 macrophages</td>
<td>93 95 92</td>
<td>94 92 87 23 0</td>
</tr>
<tr>
<td>Human endothelial cells*</td>
<td>100 100 100</td>
<td>100 100 100 0</td>
</tr>
<tr>
<td>Human fibroblasts</td>
<td>100 100 100</td>
<td>100 100 100 100</td>
</tr>
<tr>
<td>Rat cardiac myocytes</td>
<td>100 100 100</td>
<td>100 100 100 100</td>
</tr>
</tbody>
</table>

observed in the location and severity of atherosclerotic plaques in the human arterial tree. Also explained could be the differing susceptibility of saphenous vein versus internal mammary artery coronary artery bypass grafts to accelerated atherosclerosis.

(c) Abnormalities in postprandial lipoprotein structure and metabolism may be at least as atherogenic as abnormalities in fasting lipoproteins. Furthermore, the surface remnant hypothesis suggests that an individual whose fasting lipoprotein levels are unresponsive to a high-fat diet may have an increased rate of development of atherosclerosis compared with another individual with the same lipoprotein profile but on a low-fat diet. Postprandial predictions of the surface remnant hypothesis are currently being tested in our laboratory.

A number of studies have shown that VLDL from HTG plasma is cytotoxic to cultured endothelial cells (41, 42). We have observed that circulating TG-rich lipoproteins isolated from certain HTG individuals are cytotoxic to macrophages even in the absence of prior in vitro lipolysis, and that the cytotoxicity of these lipoprotein fractions is further increased by in vitro lipolysis (data not shown). This is additional evidence that the cytotoxicity of remnant lipoproteins may be a factor in vivo.

In conclusion, as a working hypothesis we suggest that vesicular cytotoxic surface remnants of lipolyzed VLDL and chylomicrons may represent a major class of atherogenic lipoproteins that are exacerbated during postprandial hyperlipidemia. Further, we suggest that inhibition of the cytotoxicity of these remnants may be one important way that HDL prevents atherosclerosis. An appealing aspect of this hypothesis is that it can account for several unexplained features of atherosclerosis, such as anatomic differences in susceptibility to atherosclerosis in the vascular tree and the vesicular nature of lipid deposits in early atherosclerosis.

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References


