Function and Distribution of Apolipoprotein A1 in the Artery Wall Are Markedly Distinct From Those in Plasma

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Background—Prior studies show that apolipoprotein A1 (apoA1) recovered from human atherosclerotic lesions is highly oxidized. Ex vivo oxidation of apoA1 or high-density lipoprotein (HDL) cross-links apoA1 and impairs lipid binding, cholesterol efflux, and lecithin-cholesterol acyltransferase activities of the lipoprotein. Remarkably, no studies to date directly quantify either the function or HDL particle distribution of apoA1 recovered from the human artery wall.

Methods and Results—A monoclonal antibody (10G1.5) was developed that equally recognizes lipid-free and HDL-associated apoA1 in both native and oxidized forms. Examination of homogenates of atherosclerotic plaque–laden aorta showed >100-fold enrichment of apoA1 compared with normal aorta (P<0.001). Surprisingly, buoyant density fractionation revealed that only a minority (<3% of total) of apoA1 recovered from either lesions or normal aorta resides within an HDL-like particle (1.063≤d<1.21). In contrast, the majority (>90%) of apoA1 within aortic tissue (normal and lesions) was recovered within the lipoprotein-depleted fraction (d>1.21). Moreover, both lesion and normal artery wall apoA1 are highly cross-linked (50% to 70% of total), and functional characterization of apoA1 quantitatively recovered from aorta with the use of monoclonal antibody 10G1.5 showed ≈80% lower cholesterol efflux activity and ≈90% lower lecithin-cholesterol acyltransferase activity relative to circulating apoA1.

Conclusions—The function and distribution of apoA1 in human aorta are quite distinct from those found in plasma. The lipoprotein is markedly enriched within atherosclerotic plaque, predominantly lipid-poor, not associated with HDL, extensively oxidatively cross-linked, and functionally impaired. (Circulation. 2013;128:1644-1655.)

Key Words: apolipoproteins • arteriosclerosis • cardiovascular diseases • plaque, atherosclerotic

The poor performance of several recent clinical trials targeting elevation of high-density lipoprotein (HDL) cholesterol1–3 and the recent Mendelian genetic studies questioning a causal link between genetic variants controlling HDL cholesterol levels and cardiovascular disease risk4 argue for a reappraisal of our understanding of HDL. Such a reappraisal demands that we question assumptions about the pathobiology of the lipoprotein, particularly in cases in which direct investigation is lacking. Much of what is known biologically about apolipoprotein A1 (apoA1) in human studies comes from investigations using isolated lipoprotein particles from the circulation (plasma or serum) with the use of buoyant density ultracentrifugation. As a known exchangeable lipoprotein, it is widely recognized that the vast majority of apoA1 within the circulation resides on spherical HDL particles, where it serves as the major structural protein of a complex macromolecular assembly of lipoprotein particles with defined buoyant density (1.063≤d≤1.21).

An unproven assumption is that the numerous biological functions observed with HDL or apoA1 recovered from the circulation will mirror what occurs elsewhere in vivo.

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The functional properties of apoA1 and HDL within the circulation, however, may not faithfully reflect what occurs within the artery wall. Early studies identified that lipoproteins isolated from the artery wall, particularly low-density lipoprotein (LDL), undergo assorted alterations including proteolysis, various oxidative modifications, and lipolysis to varying extents.6,8 Several years ago, we reported that apoA1...
recovered from human atherosclerotic arterial lesions was selectively targeted for oxidative modification by myeloperoxidase-generated and NO-derived oxidants and that oxidative modification of apoA1 and HDL in vivo to a comparable extent resulted in loss of cholesterol efflux activity of the lipoprotein. Parallel functional characterization and mass spectrometry studies of circulating HDL isolated by buoyant density ultracentrifugation revealed that higher apoA1 content of oxidative modifications specifically formed by myeloperoxidase- and NO-derived oxidants was associated with impairment in plasma membrane transporter ATP-binding cassette A1 (ABCA1)–dependent cholesterol efflux function of the lipoprotein, lecithin-cholesterol acyltransferase (LCAT) activity, and acquisition of proinflammatory activity. Similar findings have been replicated by other groups, and numerous additional proteomics studies have since mapped site-specific oxidative modifications to apoA1 recovered from the human artery wall. These studies collectively reveal that apoA1 is extensively oxidatively modified within an atherosclerotic-laden artery wall, and similar oxidative modifications to the lipoprotein in vivo are associated with proatherogenic changes in apoA1 function. However, no studies to date have directly examined the functional properties or the particle distribution of apoA1 recovered from human artery wall. The paucity of direct functional characterization studies is likely a result of the significant challenges that exist in obtaining sufficient quantities of fresh human arterial tissue for such biochemical and biological studies.

Herein we sought to examine both the distribution and the functional properties of apoA1 recovered from the human artery wall. The present studies demonstrate multiple remarkable findings, including direct evidence that the biological function and HDL particle distribution of apoA1 within both normal and atherosclerosis-laden human aortic tissues are markedly distinct from those of circulating apoA1 and HDL. These studies suggest that the historical focus thus far on circulating HDL cholesterol levels may not adequately reflect the actual situation with regard to apoA1 function and HDL particle distribution within the artery wall.

Materials and Methods

Materials

D2O was purchased from Cambridge Isotopes, Inc (Andover, MA). Chelex-100 resin, fatty acid–free bovine serum albumin, and crystalline catalase (from bovine liver; thymol-free) were purchased from Fisher Chemical Company (Pittsburgh, PA). Commercial apoA1 antibodies were from Abcam (donor: hydrogen peroxide, oxidoreductase, EC 1.11.1.7) was isolated from plasma and tissue homogenates, respectively, were isolated by sequential buoyant density ultracentrifugation at low salt concentrations with the use of D2O/sucrose. Protein concentrations were determined by Markwell modified protein assay with bovine serum albumin as standard. Human apoA1 used as control for cholesterol efflux and LCAT activity assays was purified as described. Reconstituted HDL (HDL) from isolated apoA1 was prepared by the cholate dialysis method with the use of a molar ratio of apoA1:1-palmitoyl-2-oleoyl-sn-glycerol-3-phosphocholine:cholesterol of 1:100:10. HDL particles were further purified by gel filtration chromatography with the use of a Sephacryl S300 column (GE Healthcare, Waukesha, WI) on a Bio-Rad Biologics DuoFio FPLC (Bio-Rad, Hercules, CA). Myeloperoxidase was purchased from Avanti Polar Lipids. All other buffers used were passed through a Chelex-100 column and supplemented with 100 μmol/L diethylenetriaminepentaacetic acid (DTPA) to remove any trace levels of reductase-active metals. All glassware used was rinsed with 100 μmol/L DTPA, pH 7.4, and then Chelex-100–treated distilled deionized H2O and baked at 500°C before use. SDS-PAGE was performed as described.

Tissue Collection

Fresh surgical specimens of human aortic tissue were obtained as discarded material both at time of organ harvest from transplant donors and during valve/aortic arch (“elephant trunk”) replacement surgery. Tissue was immediately rinsed in ice-cold normal saline until free of visible blood, submerged in argon-sparged 65 mmol/L sodium phosphate buffer (pH 7.4) supplemented with 100 μmol/L DTPA and 100 μmol/L butylated hydroxytoluene, and stored at −80°C in screw-cap specimen containers in which head space was purged with argon. Butylated hydroxytoluene was omitted from buffer in specimens in which apoA1 was isolated for functional activity assays.

mAb 10G1.5 Generation, Specification, and Labeling

Hybridoma cell lines were generated by immunizing apoA1−/− mice with purified delipidated human apoA1 isolated from HDL recovered from healthy donors. Among the positive clones, subclones were screened until a mAb with desired binding specificity for equal recognition of all forms of apoA1 (see below) was identified. The subclone, mAb 10G1.5, was selected by screening for equal recognition of lipid-free and lipidated (in rHDL) apoA1 under native conditions, as well as after oxidation by exposure to multiple different systems including myeloperoxidase/H2O2/C1−, myeloperoxidase/H2O2/N02−, and CuSO4 (oxidized as outlined below). To produce sufficient levels of 10G1.5 for immunoaffinity purification of apoA1 from arterial tissues, hybridoma clones were injected into pristane-treated male BALB/c mice (8 weeks of age). Ascites fluid was collected, precipitated with ammonium sulfate, then bound and eluted from a protein A/G column (Thermo Scientific Pierce, Rockford, IL) to purify mouse mAbs. Isotypes of the mAbs were determined with the use of Pierce Rapid Antibody Isotyping Strips plus Kappa and Lambda–Mouse, Thermo Scientific Pierce, Rockford, IL).

Specificity of mAb 10G1.5 was tested with the use of apoA1 or rHDL either in native form or after incubation at 37°C in 60 mmol/L Na[PO4] buffer (pH 7.4) with multiple different oxidation systems. The different myeloperoxidase systems consisted of 19 nmol/L myeloperoxidase, 100 μmol/L DTPA, 40 μmol/L H2O2, and either 100 mmol/L H2O2, and either 100 mmol/L DTPA and 100 μmol/L butylated hydroxytoluene, or stored at −80°C in screw-cap specimen containers in which head space was purged with argon. Butylated hydroxytoluene was omitted from buffer in specimens in which apoA1 was isolated for functional activity assays.
for 90 minutes at 37°C, and the reactions were stopped by addition of 2 mmol/L methionine and 300 mmol/L catalase. All other oxidation reactions were performed for 24 hours at 37°C. Final concentrations of oxidants used were as follows: H$_2$O$_2$, 40 µmol/L; ONOO$^-$, 40 µmol/L; ONOOH/HCO$_3^-$, 40 µmol/L each; CuSO$_4$, 10 µmol/L; CuSO$_4$/H$_2$O$_2$, 10 and 40 µmol/L, respectively; FeCl$_3$, 10 µmol/L; and FeCl$_3$/H$_2$O$_2$, 10 and 40 µmol/L, respectively. ApoA1 or HDL (prepared from apoA1 or their various oxidized versions) was coated at 0.5 µg/mL into enzyme immunoassay plates and probed with 10 ng/mL anti-apolipoprotein A1 antibody 10G1.5 at room temperature for 1 hour. For Western blot analyses, mAb 10G1.5 was IRDye labeled (LI-COR Biosciences, Lincoln NE) with the use of the LI-COR IRDye 800CW high-molecular kit at a dye/protein ratio at 4:1 and visualized by infrared imaging. The IRDye 800CW dye bears an N-hydroxysuccinimide ester reactive group that couples to free amino groups on the antibody, forming a stable conjugate with antibody. Coupling was performed according to the manufacturer’s instructions.

**Human ApoA1 Quantification**

Human apoA1 was quantified by a Food and Drug Administration–approved apoA1 immunoassay on the Abbott ARCHITECT ci8200 Integrated Analyzer System (Abbott Labs, Abbott Park, IL). All other apoA1 was quantified by quantitative immunoblot analysis with the use of mAb 10G1.5 as the detecting antibody, as determined against a standard curve of known purified apoA1 standard. Immune-reactive bands were quantified with the use of Image Studio software (version 2, LI-COR) or Image J (version 1.46; http://rsbweb.nih.gov/). All nascent HDL particles, as well as isolated human HDL2 and HDL3, were further purified by gel filtration chromatography with the use of a Sephacryl S300 column (GE Healthcare, Waukesha, WI) on a Bio-Rad Biologics DuoFlo FPLC.

**Aortic Tissue Homogenization**

Atherosclerotic lesions from aortic tissues were from subjects (n=20) with an average age of 83±3 years. Normal human aortic tissues were obtained from transplant donors (n=5) from the Cleveland Clinic and had an average age of 23±7 years. All tissue homogenization and lipoprotein fractionation procedures were performed within a cold room to ensure maintaining tissue and sample temperatures <4°C. Frozen tissue blocks (submerged in 65 mmol/L sodium phosphate buffer, pH 7.4, under argon, within screw-cap containers) were thawed by placement of the containers in an ice/water bath. Immediately before complete thaw, ice-cold Ca$^{2+}$- and Mg$^{2+}$-free Chelx-100–treated PBS supplemented with 100 µmol/L DTPA, pH 7.4, was added to rinse the tissue 5 times to remove any residual blood from tissue. The aorta segment was cleaned of adventitial fat and again rinsed 3 times with ice-cold PBS supplemented with 100 µmol/L DTPA. Wet weight of the aorta was determined, and the tissue was cut into small pieces and then suspended in ice-cold Ca$^{2+}$- and Mg$^{2+}$-free PBS supplemented with both 100 µmol/L DTPA (pH 7.4) and a protease inhibitor cocktail (Sigma-Aldrich, catalog No. P8340), which was included in all subsequent solutions used for homogenization and lipoprotein isolation. Aortic tissues were homogenized in an ice/water bath with a motor-driven Brinkmann homogenizer for 30-second protein isolation. Aortic tissues were homogenized in an ice/water bath with a motor-driven Brinkmann homogenizer for 30-second protein isolation. Aortic tissues were homogenized in an ice/water bath with a motor-driven Brinkmann homogenizer for 30-second protein isolation.

**LCAT Activity**

Human recombinant LCAT was prepared and purified from culture medium of a CHO cell line (generously provided by John Parks, Wake Forest University, Winston-Salem, NC) with stable expression of human LCAT. LCAT activity was determined by calculating the conversion efficiency of [H]cholesterol to [H]cholesterol ester after lipid extraction of the reaction mixture followed by thin-layer chromatography. Fractional cholesterol esterification was calculated as disintegrations per minute in cholesterol esters divided by disintegrations per minute in cholesterol esters plus free cholesterol. The fractional cholesterol esterification rate was expressed as units of activity (nanomoles of cholesterol ester formed per hour per nanogram of LCAT) per microgram apoA1 protein.

**Cholesterol Efflux Activity**

Cholesterol efflux experiments were performed according to established procedures. The cholesterol efflux was calculated as the total radioactivity in the medium/(medium radioactivity plus cell radioactivity). Results are expressed as a percentage relative to cholesterol efflux measured with the use of human apoA1 isolated from healthy donors (n=5) as control.

**HDL Imaging**

HDL was dual-labeled and incubated with mouse peritoneal macrophages to individually monitor the fate of phospholipid versus protein components of the particle as follows. Briefly, the protein component of isolated human HDL was first labeled with Alexa Fluor 633 (ca 2 mg/mL) and Alexa Fluor 488 (ca 2 mg/mL) and antibody concentration in the flow-through was determined to calculate cross-linking efficiency, which was >90%. The gel was then rinsed extensively with 1 mol/L TRIS, pH 7.4, and 1 mol/L NaCl and then equilibrated in 1× PBS, pH 7.4, before use or storage (0.002% sodium azide was added if stored). Individual 1-time use affininty columns (1 mL, drained resin) were prepared with immobilized 10G1.5, and artery wall apoA1 was purified from individual samples of aortic homogenates under conditions that quantitatively recovered apoA1, as confirmed with the use of Western blot analyses of column fractions.

**Proteomic Analyses**

To confirm that the protein recovered after immunoaffinity isolation (with the use of mAb 10G1.5) from normal aortic tissue homogenate was apoA1, the major SDS-PAGE gel bands at molecular weights 25 and 50 kDa were excised. The samples were first treated with dithiothreitol/iodoacetamide (Sigma, St Louis, MO) to carbamidomethylate any cysteines in the protein(s), and then proteins were digested with the use of Mass Spec grade trypsin (Promega, Madison, WI) at 37°C overnight. Tryptic peptides were loaded onto an IntegrAFlit sample trap (ProteoPep C18, 300 Å, 150 µm×2.5 cm, New Objective, Woburn, MA) at 1 µL/min with 5% acetonitrile and 0.1% formic acid to desalt the samples. The peptides were subsequently eluted through a column (75 µm×15 cm) packed in-house with XePerTek 218TP, C18, 300 Å pore size, 150 µm particle size (Cobert Associates, St Louis, MO) at 200 nL/min with the use of a Proxeon Easy-nLC II system (Thermo Scientific, Waltham, MA).
with a gradient of 5% to 65% acetonitrile, 0.1% formic acid over 120 minutes into a LTQ-Orbitrap Velos mass spectrometer (Thermo Scientific, Waltham, MA). Peak lists were generated with the use of Proteome Discoverer 1.1 (Thermo Fischer Scientific, Waltham, MA). The resulting Unified Search Files (*.srf) were searched against the Uniprot FASTA of all apolipoproteins and also against a human protein database downloaded from the European Bioinformatic Institute (release: 2013_02). Modifications used for searches included carbamidomethylated cysteine (fixed), oxidized methionine and tryptophan (variable), 3-chlorotyrosine, and 3-nitrotyrosine (variable). Only strictly tryptic peptides with a maximum of 2 missed cleavage sites were allowed in the database searches. Monoisotopic precursor ions were searched with a tolerance of 100 ppm with 0.8 Da for the fragment ions on the data obtained from the hybrid LTQ-Orbitrap Velos mass spectrometer. Unidentified fragment ions in all fragmentation spectra were manually validated with the use of Protein Prospector (University of California, San Francisco).

**Statistical Analysis**

Nonparametric statistical methods were used to determine statistical differences attributable to sampling numbers. The Wilcoxon rank sum test was used for 2-group comparisons, and the Kruskal-Wallis test was used for multiple-group comparisons (>2 groups). In cases in which the Kruskal-Wallis test was performed for multiple-group comparisons and found to be significant (P<0.05), multiple-comparison procedures such as the Dunn test were used for pairwise comparison between groups and controls. The Wilcoxon rank sum test was also used for pairwise comparison. Where indicated, the 1-sample robust Hotelling T2 test was used to determine statistical significance when enzymatic activity between the control group and experimental group was compared.

**Results**

**mAb 10G1.5 Recognizes ApoA1 Equally Well in its Lipid-Free or HDL-Associated, Native, and Oxidized Forms**

We initially sought to accurately quantify and immunoreactivity isolate apoA1 from artery wall tissue homogenate (from both normal and atherosclerotic lesions). We reasoned that this would require a sufficiently tight binding antibody that demonstrated minimal recognition bias between lipidated versus nonlipidated forms of the lipoprotein, as well as oxidized versus nonoxidatively modified forms. Examination of every commercially available antibody we could find (both monoclonal and polyclonal) showed significant bias in recognizing one form or another (typically recognition of oxidized forms preferentially, and with inadequate affinity). Figure 1A illustrates the biases observed with 3 characteristic commercial antibodies (2 polyclonal and 1 mouse mAb). Despite equal mass of protein loaded into adjacent lanes from native versus oxidized apoA1, and HDL versus oxidized HDL, the commercial antibodies show varied intensity of staining (eg, commercial antibody 1 showed oxidized HDL>>oxidized apoA1>>apoA1 or HDL; commercial antibody 2 shows oxidized apoA1>oxidized HDL>>apoA1 or HDL; and commercial antibody 3 shows oxidized apoA1 or oxidized HDL>>apoA1 or HDL). We therefore initially sought to develop a suitable antibody that met our strict apoA1 recognition criteria. Purified delipidated human apoA1 (isolated from plasma HDL) was injected into several apoA1−/− mice. After screening >5000 hybridoma clones for their ability to recognize apoA1 forms equally well, a small number (4) met our screening program requirements. One mAb, 10G1.5, was selected on the basis of specific activity of recognition by ELISA, immunoblot analysis, its ability to immunoprecipitate apoA1, and the growth characteristics of the hybridoma clone. Figure 1B illustrates that mAb 10G1.5 recognizes native apoA1 and apoA1 reconstituted into HDL particles equally well. Furthermore, mAb 10G1.5 recognizes apoA1 in native versus oxidized forms equivalently, with the use of a wide variety of oxidation schemes (Figure 1B). We further examined the ability of mAb 10G1.5 to quantify different concentrations of purified apoA1 (lipid-poor) versus equivalent amounts of...
total apoA1 in either isolated human HDL (total) or the individual HDL subfractions HDL2 or HDL3 (Figure 1C). As can be seen, mAb 10G1.5 displayed nearly identical ability to quantify apoA1 in its varied lipid-free and lipidated forms over a range of masses. On the basis of the observed unbiased recognition of all apoA1 forms, we refer to this antibody as “anti-total” apoA1. This mAb was used throughout the studies described below to detect, immunoaffinity purify, and quantify apoA1 recovered from plasma, atherosclerotic lesion homogenates, and normal artery wall homogenates.

The Majority of ApoA1 Isolated From Lesions Is Highly Cross-Linked and Not HDL Associated

The particle distribution of apoA1 within human atherosclerotic lesions has not been reported. We therefore homogenized human aortic atherosclerotic lesions (n=10 different subjects) and used sequential buoyant density ultracentrifugation to initially remove the VLDL/LDL-like fraction (d≤1.063) and then recover both the HDL-like fraction (1.063<d≤1.21) and the LPD fraction (density >1.21), as described in Methods. Samples were first examined on gradient (5% to 15%) SDS-PAGE separations with the use of Sypro Ruby Red protein staining, which shows minimal protein-to-protein differences in staining; equally stains lipoproteins, glycoproteins, and other difficult-to-stain proteins; and does not interfere with subsequent mass spectrometry analyses. Visual inspection showed a complex protein mixture, with an unknown band migrating at ≈27 kDa, the molecular weight of apoA1 (Figure 2A). Western analysis with anti-total apoA1 antibody (mAb 10G1.5) of a membrane containing transferred proteins from a parallel-run duplicate gel readily detected within lesion homogenates a band at the molecular weight of the apoA1 monomer (Figure 2B). Remarkably, the vast majority of apoA1 within the aortic lesion was observed to be present not within the HDL-like fraction but rather within the LPD fraction (Figure 2B). After substantial increase in exposure of the immunoblot, apoA1 was detected within the HDL-like particle fraction (Figure 2C). Also notable within the immunoblots were prominent slower migrating bands of immunoreactive apoA1-containing protein bands at molecular weights of ≈50, ≈75, and ≈100 kDa present in particular within the starting material (homogenate) and the LPD fraction (d>1.21) but noticeably diminished in the HDL-like particle fraction (1.063<d≤1.21). The sizes of these slower migrating apoA1-immunoreactive bands are consistent with the sizes of oxidatively cross-linked dimeric and multimeric apoA1 forms. Quantification of the distribution of protein and apoA1 forms recovered within homogenates from multiple distinct human atherosclerotic plaque–laden aorta (n=10) is shown in Figure 3. The majority (81.0±5.6%) of the total protein in the lesion homogenate was found in the LPD fraction, whereas the HDL-like fraction contained only 1.7±0.2% (Figure 3A). Quantitative analysis of the anti-total apoA1–specific immunoblots indicated that nearly all of the apoA1 isolated from lesions was lipid-poor and found within the LPD fraction (d>1.21), where 0.7±0.4 mg apoA1 per gram wet weight of lesion material was recovered (Figure 3B), corresponding to 92.4±4.1% of total apoA1 in the artery wall (Figure 3C). Surprisingly, only a nominal amount (<3%) of apoA1 within the artery wall (lesions) was recovered in the HDL-like particle fraction (Figure 3B and 3C). Yet another remarkable finding was the abundance of slower migrating immunoreactive apoA1-containing bands migrating with molecular masses of dimeric, trimeric, and tetrameric forms of apoA1 within the lesion homogenates. Quantification of these apoA1-immunoreactive bands revealed that approximately two thirds (66±4%) of apoA1 within lesions was oxidatively cross-linked, and the cross-linked forms were preferentially present in the LPD fraction (Figure 3D). Proteomics analyses of anti-total apoA1 (mAb 10G1.5) immunoprecipitated higher-molecular-weight apoA1 forms confirmed that these bands also were predominately composed of apoA1 (see below). Total apoA1 present in the aortic tissues recovered is
Our initial studies focused on apoA1 within atherosclerotic lesions. However, given the surprising finding that virtually all apoA1 within aortic lesions was not on an HDL particle, and fully two thirds of all apoA1 within lesions was cross-linked, we decided to examine apoA1 within normal aortic tissue for comparison. Normal aortic tissue was obtained at time of organ harvest from transplant donors. For illustrative purposes, images of a typical normal aortic specimen and a typical atherosclerotic plaque–laden aortic specimen are shown in Figure 4A. Homogenates of normal aortic tissue (n=5) were prepared and fractionated by buoyant density ultracentrifugation as described in Methods. Fractionation of protein from normal artery and lesion homogenates on (5% to 15%) gradient reducing SDS-PAGE gels stained with Sypro Ruby Red for protein revealed that although there are similarities in the protein banding pattern, the pattern is noticeably different in normal versus lesion homogenates (Figure 4B). Notably, immunoblot probing with apoA1-specific anti-total apoA1 (mAb 10G1.5) of parallel SDS-PAGE gels transferred to membranes showed that compared with lesion-derived homogenates, there is very little immunoreactive apoA1 in homogenates prepared from normal aortic tissue (Figure 4C). Because extremely low levels of apoA1 were observed within normal artery wall tissue, and especially the HDL-like fraction, an increased protein amount was loaded onto SDS-PAGE gels to permit visualization by Western blot and comparison of HDL-like and LPD fractions from the normal artery wall homogenate (Figure 4D). The majority of protein in the normal artery wall homogenate was found to be in the LPD fraction (Figure 1 in the online-only Data Supplement). As observed for aortic lesion apoA1, a significant portion of apoA1 was lipid-poor and recovered within the LPD fraction. Furthermore, there was a high degree of immunoreactive apoA1 forms in the normal artery wall that migrate at higher molecular weights, consistent with that of oxidatively cross-linked apoA1 dimer and higher multimeric forms, particularly within the LPD fraction (Figure 4D).

Quantification of apoA1-specific immunoblots (with volumes of starting homogenate and density cut fractions recovered taken into account) revealed that the total apoA1 recovered per gram of wet weight aortic tissue from normal artery wall was 120-fold less than that recovered from lesion-laden aorta (Figure 4E; note log scale for y axis). Remarkably, the distribution of apoA1 within normal artery resembled those found in the same fractions from atherosclerotic lesions, with only 3% of apoA1 being HDL associated and 92% of the total apoA1 present in the normal artery wall residing in the LPD fraction (Figure 4F). Similar to our observation within lesions, nearly half (43.4±13.9%) of the total apoA1 within the normal artery wall was cross-linked and was observed to reside not on the HDL-like particle (3.6±2.6%) but in the LPD fraction (61.4±23.0%; Figure 4G).

Because the levels of apoA1 in the normal artery wall–derived homogenates were so low, we wanted to verify that the bands detected on Western blots with the use of mAb 10G1.5 were in fact apoA1. We therefore immunopurified apoA1 from normal aortic tissue homogenates (n=5) using 10G1.5 (as described in Methods), and immunoaffinity isolated proteins were separated on a gradient nonreducing (5% to 15%) SDS-PAGE gel and stained for protein. The major protein bands observed corresponded to doublet bands.
migrating at \( \approx 25 \) and \( \approx 50 \) kDa (Figure II in the online-only Data Supplement). These were individually excised and digested with trypsin for mass spectrometry analyses, as described in Methods. Tandem mass spectrometry analyses of tryptic peptides for each excised band revealed apoA1 as the dominant protein within each, with 40% to 62% peptide coverage in each of the gel-excised bands (Tables I through IV in the online-only Data Supplement).

**Plasma ApoA1 Is Predominantly HDL Associated and Has Decreased ApoA1 Cross-Links in the HDL-Like Fraction Compared With the LPD Fraction**

Given the surprising HDL particle distribution and cross-link prevalence of apoA1 found in the artery wall (Figures 2–4) and the known preponderance of apoA1 within the HDL fraction in plasma, we examined the distribution of apoA1 in plasma using the same anti-total apoA1 mAb 10G1.5 to show apoA1 and cross-linked apoA1. **Figure 5A** shows the dominant immunoreactive band observed migrates at \( \approx 27 \) kDa, corresponding to the apoA1 monomer, and is predominantly recovered within the HDL fraction (\( \approx 83\% \); Figure 5B and 5D). Unlike apoA1 recovered from the artery wall, there is little noticeable higher-molecular-weight cross-linked apoA1 in the starting plasma, particularly within the circulating HDL fraction. Of note, however, the content of apoA1 cross-linked within the LPD fraction was found to be 3-fold higher than that observed within the HDL fraction (\( \approx 83\% \); Figure 5B and 5D). Unlike apoA1 recovered from the artery wall, there is little noticeable higher-molecular-weight cross-linked apoA1 in the starting plasma, particularly within the circulating HDL fraction. Of note, however, the content of apoA1 cross-linked within the LPD fraction was found to be 3-fold higher than that observed within the HDL fraction (\( \approx 83\% \); Figure 5B and 5D).

**ApoA1 Isolated From Atherosclerotic Lesions Is Dysfunctional**

In a final series of experiments, apoA1 was isolated from additional atherosclerotic plaque–laden aortic tissues (n=10
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Volunteers. rHDL formed with lesion apoA1 demonstrated erated from apoA1 purified from plasma HDL from healthy examined for LCAT activity and compared with rHDL gen-

µ comparable amounts (5 g apoA1 mass) of each rHDL were by gel filtration fast protein liquid chromatography, and then purified HDL by cholate dialysis method and further purified from the lesion apoA1 samples was incorporated into reconstit-

tuted HDL by cholate dialysis method and further purified HDL from healthy donors (Figure 5). In parallel studies, each apoA1 sample recovered from lesions or apoA1 purified from plasma HDL (as lent amounts of apoA1 protein were recovered from lesions subjects) by individual immunoaffinity columns composed of immobilized anti-total apoA1 (mAb 10G1.5), as described in Methods. Immunoisolated apoA1 was quantified, and equivalent amounts of apoA1 protein were recovered from lesions from each sample, or apoA1 purified from plasma HDL (as a control) was incubated with cholesterol-loaded murine macrophage RAW264.7 cells to quantify cholesterol efflux activity. Remarkably, every apoA1 sample recovered from plaque-laden aorta demonstrated lower cholesterol acceptor activity, showing 78.9 ± 6.0% less total cholesterol efflux capacity compared with control apoA1 purified from plasma HDL from healthy donors (Figure 6). In parallel studies, each of the lesion apoA1 samples was incorporated into reconstituted HDL by cholate dialysis method and further purified by gel filtration fast protein liquid chromatography, and then comparable amounts (5 µg apoA1 mass) of each rHDL were examined for LCAT activity and compared with rHDL generated from apoA1 purified from plasma HDL from healthy volunteers. rHDL formed with lesion apoA1 demonstrated significantly less (89.6 ± 5.0%) LCAT activity compared with rHDL formed from control apoA1 (Figure 6). Thus, apoA1 in atherosclerotic plaque–laden aorta is markedly functionally impaired (ie, “dysfunctional”) with respect to both cholesterol acceptor and LCAT activities.

Discussion

The present studies reveal multiple remarkable findings about apoA1 within the artery wall. First, the vast majority of apoA1 within both normal and atherosclerotic human arterial tissue, in contrast to within the circulation, is lipid-poor (ie, in the LPD fraction, d>1.21) and does not reside on an HDL-like (1.063 ≤ d ≤ 1.21) particle. Second, the content of apoA1 in atherosclerotic lesions is >100-fold higher than that observed within normal artery wall. Third, the majority of apoA1 within arterial tissues (both normal and atherosclerotic) is oxidatively cross-linked. Fourth, apoA1 within arterial tissues is dysfunctional, with ≥80% reduction in cholesterol acceptor activity and ≥90% reduction in capacity to activate LCAT. Fifth, the majority of oxidatively cross-linked apoA1 within the circulation is not HDL particle associated but rather resides within the LPD fraction (d>1.21). Collectively, the present studies thus argue that examination of total apoA1 and HDL function within the circulation may not adequately represent what is occurring within the artery wall, especially with respect to cholesterol acceptor and LCAT activities. Moreover, the overall strategy applied in the present studies may prove useful in the examination of other posttranslational modifications to apoA1, such as glycation, and site-specific oxidative modifications as antibodies become available. Finally, the present studies suggest that the common practice of isolating circulating HDL for study of its biological properties and discarding the lipid-poor (non–HDL associated) apoA1 may in fact be throwing out the very fraction (lipid-poor) that more closely
reflects the environment within the artery wall. Of note, the extent of oxidatively cross-linked apoA1 within the lipid-poor (LPD) fraction of plasma was 3-fold enriched relative to the HDL-like fraction (Figure 5E). On the basis of the cumulative results herein, one might speculate that dysfunctional forms of HDL monitored within the circulation will most likely reside not on the HDL particle itself but as lipid-poor forms in the LPD fraction (d>1.21). It is notable that the apoA1 found within the artery wall, which was predominantly within the LPD fraction in both atherosclerotic lesions and normal artery wall tissue, is remarkably highly cross-linked (50% to 70%). This value is even higher than previously observed in Western blots (though cross-linking was not quantified per se) in studies examining apoA1 oxidation levels in the artery wall based on buoyant density recovery of HDL-like particles. Through use of stable isotope dilution mass spectrometry–based approaches, these studies suggested that an upper boundary of up to 1 of every 2 HDL-like particles recovered from the artery wall carried an oxidative modification from either myeloperoxidase- or NO-derived oxidants. One question the present studies raise is how apoA1 becomes >100-fold enriched within atherosclerotic plaques compared with normal artery. A second question is how artery wall apoA1 is rendered lipid-poor when the vast majority of apoA1 that diffuses into the artery wall tissue from the circulation resides within the HDL particle. To address these questions, we performed preliminary confocal microscopy studies in which HDL was isolated from peripheral blood of healthy volunteers and subsequently doubly labeled (protein with Alexa Fluor 633 [red] and phospholipid with NBD-PE [green], as described in Methods). Brief incubation of the double-labeled HDL with macrophages led to virtually all of the phospholipid fluorophore being rapidly taken up within the macrophage, whereas the protein (predominantly apoA1) remained on the cell surface (data not shown). Such preliminary results provide a rationale to speculate that selective uptake of lipids may contribute as a mechanism for rapidly depleting the HDL particle of not just cholesterol but also phospholipid, leaving the lipid-poor apoA1 behind in the extracellular space. Lipid-poor or lipid-free forms of apoA1 are recognized as the preferred substrate of ABCA1. Therefore, a question that logically follows is why the lipid-poor apoA1 form within the artery wall is such a poor cholesterol acceptor (Figure 6). In past studies, we have shown that oxidative modification of apoA1 by myeloperoxidase-generated oxidants markedly impairs its ABCA1-dependent cholesterol efflux activity. Other studies have also reported that carboxy-terminal proteolized (ie, “clipped”) apoA1 can be recovered from aortic tissue that fails to efficiently bind lipid. In the present studies, however, we did not see substantial levels of lower-molecular-weight forms of apoA1 in tissue homogenates once we exercised extreme care in keeping tissues ice-cold and, more importantly, included an extensive cocktail of protease inhibitors during tissue dissection, homogenization, and fractionation, as outlined in Methods. Given the extensive oxidative cross-linking observed in total apoA1 within lesions, oxidative posttranslational modifications of apoA1 likely explain in large part why the lipoprotein, at least within lesions, remains lipid-poor. It should also be noted, however, that Parks and colleagues reported that interaction of apoA1 with ABCA1 results in lipid-poor pre-β HDL migrating forms of apoA1 that themselves are poor substrates for subsequent repeat or further lipidation by ABCA1, suggesting the requirement of an additional (non–ABCA1-mediated) process for further maturation into an HDL particle. Whether this contributes to the present results is unclear. Similarly, if an additional process is needed for further lipidation of lipid-poor apoA1 after initial interaction with ABCA1, whether this process is somehow lacking or inhibited within the artery wall requires further study. It also would have been interesting to test whether rHDL made with apoA1 isolated from the vessel wall was also defective in either ABCG1 or passive efflux compared with HDL made with plasma-derived apoA1. However, given the limitations in the amount of apoA1 able to be purified from human arterial tissues, these studies will need to await further investigation.

As for the mechanism(s) for apoA1 accumulation within lesions compared with normal artery, we have no clear answers from the present studies. We can speculate that the extensive oxidative cross-linking noted here may help to provide an answer, as well as mass spectrometry studies in the past. Oxidatively modified proteins tend to be less soluble and relatively protease resistant and might thus be “retained” within the subendothelial space, particularly within the hydrophobic environment of the atherosclerotic plaque. The phenomenon of lipoprotein retention within the subendothelial compartment of the artery wall has been suggested previously. Originally proposed for apoB lipoproteins, retention or trapping of LDL particles in the initial stages of atherosclerosis is suggested to cultivate formation of modified LDL, which may incite biological and inflammatory responses that initiate or advance the atherosclerotic process. Progressive apoB lipoprotein retention through the actions of secretory acid sphingomyelinase and lipoprotein lipase are thought to lead to accelerated lesion progression. Although we have no data to implicate lipase activation at present in depleting lipids from HDL within the artery wall leading to lipid-depleted forms, the presence of similar phospholipids on HDL suggests that a similar retention scheme for apoA1 lipoprotein retention is feasible and could thus contribute to the observed accumulation of apoA1 in artery wall lesions over time, in addition to enhanced lipid and sterol uptake into macrophages producing foam cells. Other studies have also shown that increased endothelial cell permeability in atherosclerotic lesions could lead to increased LDL and HDL migration into the diseased vessel wall and accelerate the retention process.

Although multiple studies have noted extensive oxidative modification of apoA1 recovered from the artery wall, there has been disagreement about which residues are the main sites of oxidation. Our prior proteomic mapping studies used polyclonal antibodies (chicken anti-apoA1 or anti-HDL) to immunoprecipitate apoA1 from arterial tissue homogenates. In contrast, proteomic mapping studies from alternative groups typically have used buoyant density isolation to recover the HDL-like particle fraction within lesion homogenates. On the basis of the present studies using the mAb 10G1.5, which was developed specifically to allow both recovery and equal quantification of apoA1 in lipid-free versus lipiddated and native versus oxidized forms, it is now clear...
that analysis of recovered HDL-like particles from arterial tissues only examines a very small fraction (<3%) of the total apoA1 within the artery wall. Whether the small amount of apoA1 (and its associated proteome) recovered in an HDL-like particle from the artery wall provides a “snapshot” of the apoA1 on its way to particle “disintegration” and formation of the lipid-poor apoA1 that is the predominant form remains to be determined. The present studies suggest that results that focus on HDL-like particles recovered from the artery wall need to be interpreted within the context of recognizing their minor quantitative contribution to total apoA1 in the artery wall. An interesting question, although not examined in the present studies, is whether changes in the HDL-associated proteome within the circulation observed in subjects with cardiovascular disease or within subjects at heightened risk for cardiovascular disease\textsuperscript{13–41} have any relevance to the marked changes in the environment of apoA1 observed within the artery wall.

The complexity surrounding the role of the HDL particle in the pathogenesis of cardiovascular disease has been highlighted recently because of several high-profile clinical trials targeting raising of HDL cholesterol and recent Mendelian randomization studies on HDL cholesterol levels.\textsuperscript{12,24–47} The present studies suggest that traditional HDL-measured parameters within the circulation, such as HDL cholesterol and apoA1 mass, may not adequately reflect the biology of apoA1 occurring within the artery wall. Recent studies suggest that functional measures of cholesterol efflux activity within apolipoprotein B–depleted serum may serve as a superior surrogate for HDL function.\textsuperscript{48} However, this too has recently been questioned because the HDL particle was found to account for only a minority of the cholesterol acceptor activity in the cholesterol efflux activity assays performed.\textsuperscript{49} Furthermore, despite the reported inverse association between cholesterol efflux activity and prevalent cardiovascular disease, in a separate large clinical study of similar patients (ie, sequential subjects undergoing elective diagnostic coronary angiography), enhanced cholesterol efflux activity was observed to be associated paradoxically with increased prospective cardiovascular event risk.\textsuperscript{49} It is remarkable that what we now recognize as HDL was first described nearly a century ago.\textsuperscript{50} Yet, studies focusing on HDL still continue to surprise us and reveal how little we know about its complex biology. The present and recent studies suggest that measurement in the circulation of HDL cholesterol, apoA1, or even cholesterol efflux activity may not adequately reflect what is happening within the artery wall. Rather, development of dysfunctional HDL assays that detect structurally specific modified forms of apoA1 formed in the artery wall but that diffuse back out into the circulation may be what is needed to provide insights into the processes occurring within the artery wall.

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**Disclosures**

Dr Tang has previously received research grant support from Abbott Laboratories, Drs Hazen and Smith report being listed as coinventor on pending and issued patents held by the Cleveland Clinic relating to cardiovascular diagnostics. Dr Hazen reports having been paid as a consultant for the following companies: AstraZeneca Pharmaceuticals LP, Cleveland Heart Laboratory, Esperion, Lilly, Liposcience Inc, Merck & Co, Inc, Pfizer Inc, and Takeda. Dr Hazen reports receiving research funds from Abbott, Cleveland Heart Laboratory, and Liposcience Inc. Dr Smith reports having the right to receive royalty payments for inventions or discoveries related to cardiovascular diagnostics from Cleveland Heart Laboratory and being paid as a consultant for Esperion. Dr Hazen reports having the right to receive royalty payments for inventions or discoveries related to cardiovascular diagnostics and the companies shown below: Cleveland Heart Laboratory, Frantz Biomarkers, LLC, Liposcience Inc, and Siemens. The other authors report no conflicts.

**References**


Much of what is known about apolipoprotein A1 (apoA1) and high-density lipoprotein (HDL) comes from investigations using isolated HDL particles from the circulation with the use of buoyant density ultracentrifugation and not from HDL or apoA1 directly recovered from the human artery wall. A longstanding assumption is that measurement of circulating HDL cholesterol, apoA1, or even cholesterol efflux activity provides insights into biological processes relevant within the artery wall. ApoA1 recovered from human atherosclerotic lesions, however, is known to be highly oxidized, and ex vivo oxidation of apoA1 or HDL to an extent comparable to that observed in vivo both cross-links apoA1 and inhibits cholesterol efflux and lecithin-cholesterol acyltransferase activities of the lipoprotein. Herein we directly examined the function and distribution of apoA1 recovered from normal and atherosclerotic plaque-laden human aorta. Remarkably, the distribution and function of apoA1 in the artery wall were markedly distinct from those observed within the circulation. First, apoA1 was 100-fold enriched within atherosclerotic plaque compared with normal arterial tissue. Furthermore, arterial wall apoA1 was predominantly lipid-poor, not associated with HDL, extensively oxidatively cross-linked, and functionally impaired with respect to both cholesterol efflux and lecithin-cholesterol acyltransferase activities. Collectively, these findings argue that circulating levels of HDL cholesterol, HDL particles, or apoA1 inadequately reflect apoA1 biology within the artery wall. Instead, our findings suggest that examination of circulating lipid-poor apoA1 forms or detection of structurally specific oxidatively modified forms of apoA1 may better reflect processes important within the vessel wall and for rendering HDL dysfunctional.
Function and Distribution of Apolipoprotein A1 in the Artery Wall Are Markedly Distinct From Those in Plasma

Joseph A. DiDonato, Ying Huang, Kulwant S. Aulak, Orli Even-Or, Gary Gerstenecker, Valentin Gogonea, Yuping Wu, Paul L. Fox, W.H. Wilson Tang, Edward F. Plow, Jonathan D. Smith, Edward A. Fisher and Stanley L. Hazen

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Supplemental Material

The function and distribution of apolipoprotein A1 in the artery wall are markedly distinct from those in plasma

Joseph A. DiDonato, PhD¹*, Ying Huang, PhD¹*, Kulwant Aulak, PhD¹, Orli Even-Or, PhD², Gary Gerstenecker, PhD¹,³, Valentin Gogonea, PhD¹,³, Yuping Wu, PhD⁴, Paul L. Fox, PhD¹, W.H. Wilson Tang, MD¹,⁵, Edward F. Plow, PhD⁶, Jonathan D. Smith, PhD¹,⁵, Edward A. Fisher, MD², and Stanley L. Hazen, MD PhD¹,³,⁵

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⁶Department of Molecular Cardiology, Lerner Research Institute, Cleveland Clinic, Cleveland, Ohio

*Co-first authors
Supplemental Table 1: Peak Lists of Apolipoprotein A1 Peptides in Lower ~25 kDa band verified by MS/MS

~25 kDa MW Lower Band Protein Identification

<table>
<thead>
<tr>
<th>Observed Peptide Sequence</th>
<th>XCorr</th>
<th>Charge</th>
<th>m/z [Da]</th>
<th>ΔM [ppm]</th>
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ApoA1 was immuno-precipitated from normal artery wall homogenates and run on SDS-PAGE as described. The lower band migrating at ~25 kDa was excised (Supplemental Figure 2), digested with trypsin, and analyzed via nanospray-ESI on an Orbitrap-LTQ Velos. The resultant spectra were searched using Proteome Discoverer 1.1 against the Uniprot human database 2013_2. Shown is a listing of tryptic peptides confirmed by MS/MS for apoA1 with their respective SEQUEST correlation scores (XCorr), charge state, precursor mass and mass shift of the observed peptide from the theoretical mass (ΔM [ppm]).
Supplemental Table 2: Peak Lists of Apolipoprotein A1 Peptides in Upper ~25 kDa band verified by MS/MS

~25 kDa MW Upper Band Protein Identification

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<tr>
<th>Observed Peptide Sequence</th>
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ApoA1 was immuno-precipitated from normal artery wall homogenates and run on SDS-PAGE as described. The upper band migrating at ~25 kDa was excised (Supplemental Figure 2), digested with trypsin, and analyzed via nanospray-ESI on an Orbitrap-LTQ Velos. The resultant spectra were searched using Proteome Discoverer 1.1 against the Uniprot human database 2013_2. Shown is a listing of tryptic peptides confirmed by MS/MS for apoA1 with their respective SEQUEST correlation scores (XCorr), charge state, precursor mass and mass shift of the observed peptide from the theoretical mass (ΔM [ppm]).
Supplemental Table 3: Peak Lists of Apolipoprotein A1 Peptides in Lower ~50 kDa band verified by MS/MS

~50 kDa MW Lower Band Protein Identification

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ApoA1 was immuno-precipitated from normal artery wall homogenates and run on SDS-PAGE as described. The lower band migrating at ~50 kDa was excised (Supplemental Figure 2), digested with trypsin, and analyzed via nanospray-ESI on an Orbitrap-LTQ Velos. The resultant spectra were searched using Proteome Discoverer 1.1 against the Uniprot human database 2013_2. Shown is a listing of tryptic peptides confirmed by MS/MS for apoA1 with their respective SEQUEST correlation scores (XCorr), charge state, precursor mass and mass shift of the observed peptide from the theoretical mass (ΔM [ppm]).
Supplemental Table 4: Peak Lists of Apolipoprotein A1 Peptides in Upper ~50 kDa band verified by MS/MS

~50 kDa MW Upper Band Protein Identification

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<tr>
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ApoA1 was immuno-precipitated from normal artery wall homogenates and run on SDS-PAGE as described. The upper band migrating at ~50 kDa was excised (Supplemental Figure 2), digested with trypsin, and analyzed via nanospray-ESI on an Orbitrap-LTQ Velos. The resultant spectra were searched using Proteome Discoverer 1.1 against the Uniprot human database 2013_2. Shown is a listing of tryptic peptides confirmed by MS/MS for apoA1 with their respective SEQUEST correlation scores (XCorr), charge state, precursor mass and mass shift of the observed peptide from the theoretical mass (ΔM [ppm]).
Supplemental Figure 1. The distribution of protein recovered in normal artery wall homogenate before (100%) and following fractionation by ultracentrifugation. Normal artery wall homogenates (n=5) were fractionated by sequential buoyant density ultracentrifugation using D$_2$O/Sucrose mixtures to produce the indicated density fractions, as described under Methods, and then total protein was determined from starting material and LDL-like, HDL-like and lipoprotein-depleted (LPD) fractions. The percentage of total protein relative to homogenate (100%) in each sub-fraction is indicated. Error bars indicate standard deviation. Note that the majority of protein recovered from normal artery wall homogenate is within the lipoprotein-depleted fraction.
**Supplemental Figure 2.** ApoA1 is present in the normal artery wall and is cross-linked.

ApoA1 was immuno-precipitated from normal artery wall homogenates (n=5) and separated on a non-reducing (5-15%) SDS-PAGE gradient gel and stained with Coomassie Blue. Indicated bracketed bands (upper and lower ~25 kDa and ~50 kDa protein bands) were excised from the gel and subjected to trypsin digestion and mass spectrometry analysis. The major peptides recovered and sequenced from each indicated excised regions of the gel were observed by LC-MS/MS to be from apoA1 (Supplemental Tables 1-4).
Supplemental Figure 3. Plasma contains both monomeric (major) and slower migrating (minor) oxidized and cross-linked apoA1 forms. Indicated amounts of plasma and LPD fraction proteins (n=4) from those plasma samples were run on a reducing 12% SDS-PAGE gel. The proteins were then transferred to membrane for Western blot probing with mAb 10G1.5. The major bands noted within both plasma and the LPD fractions appear at the anticipated molecular weight of the apoA1 monomer (~25 kDa). Longer exposures (shown) of the Western blot reveal the presence of slower migrating apoA1 immuno-reactive bands (indicated by arrowheads), consistent with oxidatively cross-linked forms of apoA1.