Graded Effects of Proteinuria on HDL Structure in Nephrotic Rats

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Nephrotic syndrome is characterized by increased triglycerides resulting from decreased clearance of VLDL and chylomicrons. These triglyceride-rich lipoproteins are structurally altered by interaction with HDL derived from animals with proteinuria and not as a consequence of hypoalbuminemia. HDL isolated from rats with massive proteinuria is depleted in apolipoprotein E (apoE). It is unknown at what threshold of urinary albumin loss HDL structure is altered, and it is unknown what effects proteinuria has on apolipoproteins other than apoE. Two models of albuminuria were used in Sprague-Dawley rats: Adriamycin and passive Heymann nephritis (HN). The adriamycin group was divided into minimal albumin excretion (MAE) and intermediate albumin excretion (MAE, 1 to 40; intermediate albumin excretion, 60 to 210 mg/d per 100 g body wt). Urinary albumin excretion exceeded 300 mg/d per 100 g body wt in the HN rats. HDL apolipoprotein composition was analyzed with SDS-PAGE densitometry and liquid chromatography–time of flight mass spectrometer mass spectrometry. HDL apoA-IV content relative to apoA-I was reduced at all levels of albuminuria (P < 0.0001). ApoE was not reduced in MAE but was significantly reduced in IAE (72%; P < 0.001). By contrast, apoA-II and apoC-III were each significantly increased with increasing UAE. ApoA-IV and apoE were decreased to approximately 10% of control in HDL isolated from rats with HN, whereas apoA-II, apoC-II, and apoC-III were each significantly increased relative to apoA-I. HDL is structurally altered by levels of albuminuria that are insufficient to change serum albumin levels and is progressively altered as albuminuria increases.


Hyperlipidemia in the nephrotic syndrome is a result of increased synthesis (1,2) and decreased catabolism (1,3,4). VLDL levels are increased predominantly as a consequence of decreased catabolism both in rats with the nephrotic syndrome (4) and in humans (5). VLDL from nephrotic rat plasma does not bind normally to vascular endothelial cells, a consequence of reduced apolipoprotein E (apoE) content. Nascent VLDL functions normally when secreted by nephrotic livers but acquires a structural and functional defect through interaction with nephrotic HDL after VLDL secretion (6). The alteration in HDL structure and function results directly from proteinuria (3,6) and not from hypoalbuminemia.

We have shown that defective clearance of VLDL in nephrotic rats requires interaction with HDL from proteinuric animals and that hypoalbuminemia does not deleteriously affect HDL structure. Although we have analyzed the effect of nephrotic range albuminuria on HDL structure, the effects of lesser amounts of albuminuria have not been well examined. Even small amounts of albuminuria are associated with increased relative risk for vascular disease in both diabetes (7,8) and hypertension (9,10). This association is generally believed to reflect underlying endothelial dysfunction in patients with microalbuminuria, causing both the renal albumin leak and the association with vascular disease. Nevertheless, the potential effect of albuminuria on HDL structure could provide a causal link between albuminuria and cardiovascular risk.

Microalbuminuria in humans is often associated with both the metabolic syndrome (11,12) and reduced levels of HDL. As urinary albumin excretion increases among patients with diabetes, the HDL size decreases (13). Activity of systemic lupus erythematosus is associated with proteinuria and low levels of HDL (14,15), but clearly the relationship between disease severity and HDL concentration may be confounded by other variables, including systemic inflammation (14). Although it is known that a nephrotic range of proteinuria decreases HDL apoE content, thereby altering its interaction with VLDL, lower levels of proteinuria may also alter HDL apolipoprotein content. We designed this study to examine the interrelationship between HDL structure and graded albuminuria in the subnephrotic to nephrotic range.

Materials and Methods

Animals

All protocols were approved by the Animal Resources Service committee at University of California Davis and VA Northern California Health Care System. Sprague-Dawley rats were purchased from...
Charles River (Wilmington, MD). For minimizing effects generated by the experimental model, two methods were used to induce proteinuria. Mild to moderate proteinuria was induced by the introduction of adriamycin by tail-vein injection at 2.5, 5.0, or 7.5 mg/kg as described previously (6). Proteinuria in the nephrotic range was induced by intraperitoneal injection of FX1A antibodies at 0.2 ml/kg. All rats were kept in 24-h light/dark rooms and fed standard rat diet ad libitum until the evening before the experiments. Five weeks after adriamycin injections, animals were housed in metabolic cages that allowed 24-h urine collections. Before all experiments, fasted rats were anesthetized with an intraperitoneal injection of 0.75 g/kg (control) or 0.40 g/kg (nephrotic) sodium pentobarbital and exsanguinated by aortic puncture. Urinary albumin excretion (UAE) was measured using the albumin blue 580 method (16,17). Four animal groups were generated (see Results section): Control (n = 6), minimal albumin excretion (MAE; n = 7), and intermediate albumin excretion (IAE; n = 7). The animals with Heymann nephritis were killed in a separate experiment (n = 8) along with independent controls (n = 4).

Chemical Analysis

Enzymatic kits were used for the determination of plasma triglyceride concentrations (kit 2780–400H; Thermo DMA, Chatsworth, CA), plasma cholesterol (kit TR12351; Thermo DMA), phospholipids (kit 990–54009; WAKO Chemicals, Richmond, VA), and lecithin-cholesterol acyltransferase (LCAT) (Roar Biomedicals, New York, NY).

Isolation and Content of HDL

HDL was isolated using the sequential flotation method described by Schumaker and Puppione (18). Plasma was handled using the method recommended by Edelstein and Scanu (19).

Apolipoprotein content of HDL was first determined by SDS-PAGE under denaturing conditions on 4 to 20% polyacrylamide gels and stained using SYPRO orange (Molecular Probes, Eugene, OR) and scanned using a STORM imager (Amersham Biosciences, Piscataway, NJ). The resulting images were analyzed using the accompanying ImageQuant software, version 5.0.

Alternatively, HDL composition was analyzed using HPLC (Waters 2795) coupled to an electrospray ionization (ESI) source and time of flight mass spectrometer (TOF; Micromass LCT, Manchester, UK). A 50-μl aliquot of each isolated fraction was separated using reverse-phase column chromatography (2.1 x 250-mm, 5-μm silica, 300-Å pore with polymeric bonded C18; Vydac 218TP52). Samples were separated using a gradient of Solvent A (98:2 water/acetonitrile [vol/vol] with 0.1% trifluoroacetic acid) and Solvent B (2:98 water/acetonitrile [vol/vol] with 0.1% trifluoroacetic acid). Initial conditions were 75% A, linearly ramped after injection to 42% B at 16 min and held for 4 min.

Figure 1. Experimental animals distributed into groups on the basis of proteinuria and albuminuria. Injection of adriamycin at 2.5, 5.0, and 7.5 mg/kg resulted in animals with levels of urinary protein that could be categorized into two groups. Some animals had low albuminuria levels that were insufficient to result in reduced albumin, whereas a second group had levels of albuminuria that were sufficient to result in a reduction in serum albumin as well (68% of control or minimal albumin excretion [MAE]).
before returning to the initial conditions and reconditioning the column for 2 min. Mass spectral data (800 to 2000 Da) were acquired from 7 to 20 min using the following conditions: Capillary voltage (3.25 kV), extraction cone (3.0 V), and sampling cone voltage (30.0 V). The mass spectrometer was calibrated using a solution of polyalanine and cyclo-dextrin before analysis, and leucine enkephalin was infused as a lock mass standard throughout the analysis to ensure accurate mass calibration.

Apolipoproteins in HDL were analyzed by positive mode ESI/TOF-mass spectrometry (MS). The molecular mass of each identified protein was established by deconvolution of the associated multicharge envelope using MassLynx v 4.0 (Micromass), and individual apolipoproteins were identified by their calculated molecular mass from published sequence information. A published sequence for rat apoC-II is not available, and in this case, the mouse homolog is included for comparison. For allowing estimates of each observed protein, unique ions were selected from the multicharged envelopes and extracted from the total ion chromatograms. Ion selection was made to minimize interferences from closely eluting compounds. The resulting extracted ion chromatograms then were integrated, and the area counts were used to estimate the apolipoprotein content of each analyzed sample. Because quantitative calibration standards were not available, the relative content of each apolipoprotein was normalized to apoA-I in HDL and to total phospholipids in VLDL.

Statistical Analyses
Apolipoprotein abundance was normalized to apoA-I content to assess the apolipoprotein relative abundance per particle. A principal components analysis was performed on these data using Multivariate Analysis Add-in for Excel, Version 1.2 (Bristol Chemometrics, www.chm.bris.ac.uk/org/chemometrics) using mean centered data and allowing for four principal components. Significant differences were determined using linear regression analysis, ANOVA, or t test where appropriate.

Results
Animals
Regardless of the administered dose of adriamycin, animals that received adriamycin were divided into two levels of albuminuria: MAE and IAE. In all cases, including the group that received 7.5 mg/kg adriamycin, UAE was lower than albuminuria levels previously demonstrated to alter HDL structure and function (3,20). Administration of FX1A antibody did not produce graded levels of proteinuria. UAE was significantly increased in the MAE, IAE, and HN groups. Serum albumin was decreased similarly in the IAE and HN rats. UAE, however, was significantly greater in HN than in IAE as were serum triglycerides and serum cholesterol (Figure 1).

HDL Composition by SDS-PAGE
Apolipoproteins were identified on SDS-PAGE by their apparent molecular weight as described previously (6). By SDS-PAGE, only apoA-I, apoE, and apoA-IV were well resolved and identified with confidence. Low molecular weight apolipoproteins, apoC and apoA-II, were summed to determine whether there was a cumulative effect of proteinuria on this subset. The apoA-I/total protein ratio was positively correlated with ln(UAE) \( r^2 = 0.701, P < 0.0001 \), indicating that with increasing albuminuria, HDL becomes relatively enriched in apoA-I and has less accompanying apoproteins (Figure 2). However, after segregating the adriamycin-treated animals on the basis of albumin excretion, we found that the apoA-I/protein ratio was increased only in the IAE animals (control, 0.403 ± 0.034; MAE, 0.403 ± 0.020; IAE, 0.645 ± 0.025; \( P < 0.001 \); Figure 2). The apoA-I/cholesterol and the apoA-I/phospholipid ratio were unchanged by albuminuria in animals that had less than nephrotic-range proteinuria (adriamycin group).

ApoE and apoA-IV were normalized to apoA-I levels to account for the per-particle abundance content. Both apoE and apoA-IV decreased in abundance with increasing albuminuria. The relationship between apoA-IV and ln(UAE) was strongest \( (r^2 = 0.786, P < 0.0001) \), but apoE also significantly correlated with ln(UAE) \( (r^2 = 0.558, P = 0.0002) \; \text{Figure 3A}) \). This strong relationship between apoA-IV and urinary albumin was reflected in ANOVA. Compared with control, apoA-IV was consecutively reduced in MAE and IAE animals (0.248 ± 0.023,
Figure 3. HDL apoE and apoA-IV content decreases as albuminuria increases. Content of HDL apoA-IV and apoE were first measured by SDS-PAGE (A). ApoA-I, apoE, and apoA-IV were clearly resolved. ApoA-IV and apoE were normalized to apoA-I content and analyzed by linear regression and two-way ANOVA. There was a strong linear relationship for both apoA-IV ($r^2 = 0.7855, P < 0.0001$) and apoE ($r^2 = 0.5577, P = 0.0002$). After the animals were sorted into control, MAE, and IAE groups, two-way ANOVA were performed (B). These results demonstrated that whereas apoA-IV decreased continuously with increasing UAE, with apoE, the MAE group was not reduced compared with control when expressed relative to apoA-I, the unit structural apolipoprotein of HDL. In each case, MAE animals have reduced HDL apoA-IV content compared with control, but the HDL content of apoE is not reduced. In IAE animals, the apoE levels are reduced compared with control and MAE, and apoA-IV levels are reduced again compared with control and MAE animals. $^\dagger P < 0.05$ versus control; $^{\ddagger} P < 0.05$ versus control and MAE.

HDL Composition by ESI-TOF-MS

To confirm our SDS-PAGE data and overcome the difficulty in resolving low molecular weight apoproteins by SDS-PAGE, we used ESI-TOF-MS to validate and complement the SDS-PAGE analyses. Apolipoproteins were identified by their molecular mass relative to sequences contained in the references described in Table 1. The correspondence between the predicted and measured masses of proteins was within 1% in all but two cases, confirming the stability of the instrument mass calibration and the validity of the applied mass spectral deconvolution routines. An example of a raw and deconvoluted spectrum is shown in Figure 4. The predicted mass for rat apoC-II is not available and had to be compared with mouse apoC-II. The resultant error was 2.06%. The molecular mass of hamster apoC-II is 8204.1 (21), which is a difference of 3.29% with mouse apoC-II and a 1.25% difference with the mass observed in rats. We therefore concluded that the 8308.1 $\pm$ 13.9-Da protein observed at 12.45 min was rat apoC-II. The published mass for rat apoA-II is 8950.4 Da. We could not locate a protein with a similar mass. We did locate a protein

0.199 $\pm$ 0.017, and 0.069 $\pm$ 0.013, respectively; $P < 0.0001$; Figure 3B). Conversely, apoE was not reduced with low levels of urinary albumin loss. ApoE content was not reduced in MAE HDL but was reduced in IAE rats (control, 0.558 $\pm$ 0.060 versus 0.584 $\pm$ 0.077 and 0.176 $\pm$ 0.029, respectively; $P < 0.0001$).
with a mass of 8804.6 ± 7.0 Da, which corresponds to a mass error of 1.63% with apoA-II and 0.98% with apoC-III. Despite the apparent agreement with apoC-III, we designated this protein as apoA-II because apoC-III was identified with a precision of 0.06%. Furthermore, the signal sequence of apoA-II is assumed to be ALVRRQ rather than ALVRR, the resultant error of 0.06% reduces the error to within acceptable limits. This protein could also be apoC-III; however, a correlation exists between the proteins assigned as apoA-II and apoC-III ($r^2 = 0.850, P = 0.001$), and the mass correspondence for apoC-III was excellent. Furthermore, two alternative apoC-III signal lengths that do not correspond to any observed plasma proteins have been reported.

with their respective groups, the apoA-I/total protein ratio was increased only in the IAE animals (control, 0.685 ± 0.033; MAE, 0.859 ± 0.057; IAE, 1.104 ± 0.045; $P < 0.006$; Figure 5B).

ApoE and apoA-IV were normalized to apoA-I levels to account for the per-particle abundance content. Both apoE and apoA-IV decreased in abundance with increasing albuminuria; however, although the negative correlation between apoA-IV and ln(UAE) was greater in this analysis ($r^2 = 0.946, P < 0.0001$), there was no similar relationship with apoE (Figure 6A). With ANOVA, apoA-IV again was consecutively reduced in MAE and IAE animals (0.251 ± 0.028, 0.169 ± 0.026, and 0.059 ± 0.011, respectively; $P < 0.0001$). Conversely, apoE content was not reduced in MAE rats (0.569 ± 0.093 versus 0.540 ± 0.118) but was reduced in IAE rats (0.161 ± 0.030; $P < 0.0001$; Figure 6B).

The depletion of non–apoA-I apolipoproteins was not characterized by uniform apolipoprotein depletion (Figure 7). HDL apoC-I, apoC-II, and apoC-IV content was unrelated to urinary albumin loss. Alternatively, HDL apoC-III and apoA-II content correlated positively with urinary albumin loss ($r^2 = 0.731, P = 0.0068$; $r^2 = 0.604, P = 0.023$, respectively). The results indicate that albuminuria-dependent depletion of non–apoA-I apolipoproteins was due to a depletion of apoA-IV and apoE, whereas the content of some small molecular weight apolipoproteins in HDL actually increased.

### Apolipoprotein Composition in HN

We found that animals with nephrotic-range proteinuria exhibited a greater change in HDL apolipoprotein composition than observed in the adriamycin model. The changes observed in the MAE and IAE rats were amplified in HN. The apoA-I protein content was increased as was apoA-II and apoC-III, whereas apoA-IV and apoE content were greatly reduced, each

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$a$ apo, apolipoprotein.
$b$ The retention time of proteins using the described chromatographic method.
$c$ The reported mass of pro-apoC-I is the closest reported mass to unknown A and B and is used as a comparison.
$d$ The mouse apoC-II is used for a comparison because the rat apoC-II mass has not been previously reported.

$*a$apo, apolipoprotein.
by approximately an order of magnitude relative to control (Table 2).

**Discussion**

We report here that HDL apolipoprotein content is related directly to urinary albumin loss and that even mild albumin loss results in altered HDL structure. In the rat, nephrotic-range proteinuria results in reduced HDL apoE content (3,6). It was reported previously that in rats with puromycin-induced nephrotic syndrome, HDL apoA-IV content is also reduced to approximately 10% that observed in control animals (22). We find similar results in extremely nephrotic rats with HN reported here. Our main finding, however, is that even small levels of urinary albumin loss, insufficient to change serum albumin or lipid levels, result in a decrease in HDL apoA-IV content, whereas HDL apoA-II and apoC-III content are increased.

These changes are unlikely due simply to changes in apolipoprotein gene expression in response to hypoalbuminemia. ApoA-I gene transcription and synthesis rate are increased approximately threefold in nephrotic rats (23). Although this might be expected to yield a particle that is relatively enriched in this apolipoprotein in comparison with others, such an explanation does not explain the relative increases in apoA-II, apoC-II, and C-III that were observed. Marshall reported an increased transcription of apoA-II of approximately 1.9-fold, significantly less than the 6.1-fold increase in apoA-I transcription that they observed (24). Thus, if these same changes in gene expression translate into protein synthesis, then a relative depletion of apoA-II in comparison with apoA-I should have been observed in our model. In addition, apoA-I transcription is increased in rats with hereditary analbuminemia (25), yet they have normal HDL structure. It has been reported that apoA-II displaces apoA-I from the HDL particle (26); however, we found that HDL apoA-I content increased concurrently with increasing apoA-II content.

HDL and components of HDL have been identified in the urine of nephrotic subjects (27). Finding specific apolipoproteins in the urine is not necessarily translated into the changes seen in serum levels of these proteins. Low molecular weight proteins are freely filtered by the kidney; thus, although renal failure might result in their accumulation (28), no change in their clearance would be anticipated to occur after changes in permselectivity. ApoA-II, apoC-II, and apoC-III are examples of such proteins; however, apoA-II and apoC-III are increased in all animals with proteinuria, and among HN animals, even apoC-II is increased relative to apoA-I.

It is equally difficult to explain most alterations in higher
molecular weight apolipoproteins by urinary albumin loss. Approximately 25% of apoA-I is cleared by or eliminated by the kidney in normal rats as well as after development of the nephrotic syndrome (25), suggesting that changed permselectivity has little effect on glomerular sieving of this 27-kD apolipoprotein. Although HN represents an extreme of urinary protein loss, we find that even minimal levels of albuminuria result in a significant decrease in the content of apoA-IV. ApoA-II is intermediate in size between apoA-I and LCAT and is approximately evenly distributed between HDL and free fractions (29), a distribution that potentially protects it from urinary loss regardless of dysfunction of the glomerular barrier. We observed here reductions in HDL apoA-IV content at very low levels of albuminuria, levels that are too low to result in a change in serum albumin concentration, thus suggesting that processes other than urinary protein loss may play a role in reducing HDL apoA-IV content. Furthermore, even massive levels of proteinuria do not result in decreased serum apoA-IV levels (30), suggesting that the affinity of HDL for apoA-IV may be reduced under conditions of microalbuminuria. These changes were independent of serum albumin concentration. Our observation that apoA-II and C-III also accumulate in HDL in a linear relationship with albuminuria suggests that processes other than urinary protein loss or dilution of all other apolipoproteins by an increasing apoA-I pool are responsible for changes in HDL structure.

At greater levels of urinary albumin loss, we observed a decrease in HDL apoE as previously observed. It is instructive to note that serum triglyceride or cholesterol levels did not increase at all at low levels of albuminuria, when HDL apoE content was no different from control, supporting our earlier hypothesis that HDL apoE depletion plays a role in hypertriglyceridemia (3).

The changes that we have demonstrated are important in understanding important risk factors associated with albuminuria. Although the role of apoA-IV as an activator of LCAT has been documented (31) and in humans as an activator of cholesterol ester transfer protein (32), its precise function is not well understood. Important for vascular function, apoA-IV reduces atherogenic risk (33,34) and oxidative stress (35). Mice that express human apoA-IV are less susceptible to atherosclerotic lesions even when expressed on a highly pro-atherogenic background such as the apoE knockout mouse (36). Vaziri et al. (37) demonstrated that LCAT activity was reduced in rats with nephrotic syndrome and albuminuria sufficient to reduce serum albumin levels to 55% of control. The disappearance of LCAT in plasma and the resultant appearance in the urine of LCAT activity \((r^2 = 0.62)\) lead Vaziri et al. (37) to speculate that glomerular protein loss explained LCAT depletion and connected urinary protein losses to increased lipid levels through that mechanism. The molecular mass of LCAT is roughly 66 kD (38,39), approximately that of albumin and of a size that would be most affected by changes in glomerular permselectivity. LCAT is loosely bound to HDL, dissociating on average once every catalytic cycle from HDL (40,41) and thus is accessible for filtration; however, apoA-IV levels are reduced in the presence of proteinuria too low to produce urinary loss of LCAT sufficient to anticipate any change in plasma level.

Elevated apoA-II is associated with insulin resistance and reduces the affinity of the HDL particle for the CD36 scavenger receptor (41). ApoA-II also inhibits LCAT activity (42,43) and cholesterol ester transfer protein (44) but increases hepatic lipase activity (45). Whereas increased hepatic lipase activity is associated with reduced atherogenic risk, reduced LCAT is pivotal in the maturation of HDL in this model. Thus, the changes in HDL structure associated with increasing urinary albumin loss should render HDL a less effective substrate for LCAT.

ApoC-III is strongly related to hypertriglyceridemia by inhibition of lipoprotein lipase (LpL) (46,47) and interferes with apoE-mediated binding to LDL receptors (48). Consequently, apoC-III is associated with increased risk for aortic valve scler-
Figure 6. HDL apoE and apoA-IV by ESI-TOF-MS. Two control, three MAE, and three IAE rats were additionally analyzed by liquid chromatography (LC) ESI-TOF-MS (A). ApoA-I, apoE, and apoA-IV were re-analyzed. ApoA-IV and apoE were normalized to apoA-I content and analyzed by linear regression and two-way ANOVA. The improved quantification by MS demonstrated a strong linear relationship for apoA-IV ($r^2 = 0.9460$, $P < 0.0001$); however, the relationship between UAE and apoE was not as strong ($r^2 = 0.4773$, $P = 0.0578$). After the animals were sorted into control, MAE, and IAE groups, two-way ANOVA replicated the findings using SDS-PAGE. Whereas apoA-IV decreased continuously with increasing UAE, among apoE, the MAE group was not reduced compared with control (B). † $P < 0.05$ versus control; ‡ $P < 0.05$ versus control and MAE.

Figure 7. Additional apolipoproteins resolved by MS. In addition to apoA-IV and apoE, the improved resolution of liquid chromatography (LC) ESI-MS-TOF allowed for analysis of apoA-II, apoC-I, apoC-II, apoC-III, and apoC-IV. The proportions of HDL apolipoproteins relative to apoA-I, the unit structural apolipoprotein of HDL, were measured by ESI-TOF-MS, and their relationship to UAE was determined by linear regression analysis. Two additional relationships were significant: apoA-II and apoC-III. The solid line represents the linear relationship, and the dotted line represents the 95% confidence interval. ApoA-II, $P = 0.0232$; apoC-III, $P = 0.0068$. 
Table 2. HDL structural changes in passive Heymann nephritis by ESI-TOF-MS

<table>
<thead>
<tr>
<th></th>
<th>apoA-I/Protein Ratio</th>
<th>apoA-II</th>
<th>apoA-IV</th>
<th>apoC-II</th>
<th>apoC-III</th>
<th>apoE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heymann</td>
<td>0.773 ± 0.023</td>
<td>72.6 ± 3.5</td>
<td>7.9 ± 1.3</td>
<td>23.1 ± 0.9</td>
<td>102.6 ± 7.9</td>
<td>52.0 ± 6.5</td>
</tr>
<tr>
<td>Control</td>
<td>0.642 ± 0.036</td>
<td>17.3 ± 5.5</td>
<td>72.1 ± 11.5</td>
<td>15.8 ± 1.7</td>
<td>34.6 ± 2.4</td>
<td>415.6 ± 47.1</td>
</tr>
<tr>
<td>Corrected P</td>
<td>&lt;0.0001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>0.576</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*aESI-TOF-MS, electrospray ionization time of flight mass spectrometry.

*bSee note e in Table 1.

...rosis (49) and likely plays a causative role in cardiovascular disease (50). ApoC-II is an LpL activator, and the apoC-II/C-III ratio has been observed to be decreased in VLDL obtained from nephrotic rats (51). We demonstrate here that this ratio in HDL shows a relationship to urinary albumin loss. It is unclear whether the alteration in VLDL apoC content is conferred upon it by HDL; however, the depletion of apoE from VLDL indeed does result from exposure to HDL obtained from proteinuric animals (6), and similar alterations in apoC composition of VLDL may well result from the changes that we find here in HDL. Because apoC-II is an LpL activator competitively inhibited by C-III, HDL may further contribute to delayed VLDL catabolism in this way. It is unclear what is responsible for the accumulation of these apolipoproteins as a result of proteinuria while the same process depletes HDL of other proteins.

A well-documented relationship exists between microalbuminuria and cardiovascular risk in both diabetic and hypertensive patients (7,8,10,52–54). In this study, the observed that changes in HDL structure associated with minimal albuminuria would mechanistically favor increased cardiovascular risk, suggesting that these changes themselves may indicate novel risk factors to the processes associated with increased cardiovascular disease in nephrotic subjects. It is believed that the association between microalbuminuria and cardiovascular risk is a reflection of either endothelial injury or changes in vascular compliance associated with vascular injury and reported by urinary albumin loss. It is possible, however, that even these small amounts of urinary protein loss may alter lipoprotein structure and perturb normal lipoprotein metabolism contributing to rather than simply reporting vascular injury.

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