HDLs constitute one of four major classes of particles that transport lipids between organs, tissues, and cells (Table 1). “Good cholesterol,” the layman’s term for HDL cholesterol, makes reference to the established role of HDL in transport of cholesterol from peripheral tissues to plasma, and from plasma to the liver, where the cholesterol is processed for packaging into bile acids for excretion. This putatively anti-atherogenic (and hence “good”) biological role of HDL in supporting excretion of excess cholesterol was first postulated by Glomset and colleagues (1) and has since been termed reverse cholesterol transport (2). Subsequently, significant evidence identified HDL as a therapeutic target to combat atherosclerosis. However, recent clinical trials designed to test whether increasing plasma HDL levels is therapeutically beneficial did not demonstrate efficacy (3–5), and genetic studies have refuted the prediction that higher levels of plasma HDL in humans associates with protection from cardiovascular disease (6). Harnessing the role of HDL in cholesterol clearance remains of great interest, but the field now recognizes that simple measurements of plasma HDL cholesterol do not capture its ability to support cholesterol efflux from cells, including cholesterol-loaded macrophages within atherosclerotic plaques. Indeed, the most important site from which HDLs act to promote cholesterol efflux from cells is in the extracellular matrix of tissues, not in plasma. It is plausible that some persons who display high levels of plasma HDL cholesterol have poor HDL recirculation and/or function within the interstitium of tissues, leading to poor cholesterol efflux at relevant sites. This Review considers what is known about HDL cholesterol in the interstitium and its transport through lymphatic vessels as a requisite part of its role in reverse cholesterol transport. To more comprehensively consider this topic, we also discuss chylomicron transport via lymphatics, a process that may provide insight into transport of HDL from other peripheral tissues.

**Transit of HDL through the interstitium and lymph**

The major apolipoprotein of HDL is apoA1. When the total volume of interstitial fluid is taken into account and its concentration therein considered, approximately half of all apoA1 in the body is extravascular and found within interstitial fluid of peripheral organs (7). Calculations based on lipoprotein particle numbers in plasma (8) and concentrations of apoA1 and apoB (the major protein of LDL) in plasma and peripheral lymph (9) indicate that in normal human interstitial fluid there are usually more than 50 HDL particles to one LDL particle. Nonetheless, in spite of its manifest importance in relation to lipid transport in health and disease, there is a dearth of information on HDL in interstitial fluid of peripheral tissues, owing to the difficulty of obtaining sufficient fluid under physiologic conditions.

Although subcutaneous implantation of microdialysis probes provides valuable information on movement of small molecules between plasma and interstitial fluid in humans, the molecular size cut-off of the probes is too low for recovery of lipoproteins (10, 11). Subcutaneous wicks (12) provide insufficient material for analysis of lipoprotein composition. Greater volumes are obtained from suction blisters (13), but suction blister fluid is non-physiologic, formed by separation of the dermis from epidermis (14). The only viable approach is collection of interstitial fluid as it drains from tissues via afferent lymphatic vessels before reaching LNs (Figure 1). An underlying tenet of this approach is that interstitial fluid travels down a pressure gradient from blood to lymphatic capillaries such that the composition of lymph is a close reflection of interstitial fluid itself (15). As observed in mice, a key feature enabling movement of fluid into lymph is the structure of inter-endothelial junctions, organized so that adhesions are discontinuous, like buttons on a coat, to generate flaps between the buttons (Figure 1) that can be pulled open under tension by anchoring filaments attached to the capillary, making the vessel accessible to large macromolecules (16). Transcytosis following uptake of fluid by micropinocytosis and/or receptor-mediated endocytosis may also contribute to entry of lymph into lymphatic capillaries (17).

The first data on human peripheral lymph lipoproteins were obtained by Reichl and colleagues, using fluid collected from a superficial vessel in the dorsum of the foot. In studies conducted between 1973 and 1989, they showed that lymphatic lipoproteins differ from those in plasma in concentration and composition. Concentrations of apoA1 and apoB were much lower than in plasma (18–20), and all apoB was in LDL (21). Studies with radiolabeled lipoproteins confirmed that lymph apolipoproteins were derived from plasma (22). Measurements of cholesterol-specific radioactiv-
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Several studies, reviewed elsewhere in detail (25), were consistent with the concept proposed by Glomset (1) that HDLs are the transport vehicles for cholesterol from tissues. Reichl et al. (26) subsequently found that peripheral lymph contains particles with apoA1, but no apoA2, of similar size and charge as pre-B-HDLs, small lipid-poor particles that Castro and Fielding (27) had shown to be the primary acceptors of cell-derived cholesterol from cultured fibroblasts. An intervention study showed that treatment with gemfibrozil, an oral fenofibrate activator of PPARα, increased apoA1 and cholesterol in lymph but not in plasma and demonstrated the potential of the method for obtaining unique insight into extravascular cholesterol transport (28).

Miller and coworkers (29) improved the lymph collection procedure, based on an original method of Engeset et al. (30), in which lymph is collected from a larger afferent lymphatic vessel in the lower leg, allowing collection at flow rates of 0.25–2.0 ml/h for several days. The lymph in this vessel is derived from skin, adipose tissue, and connective tissue. Consistent with Reichl et al. (18–21), lymph was essentially devoid of VLDL. With all apoB in LDLs (9, 29), the apoA1/apoB and sphingomyelin/phosphatidylcholine ratios were greater than in plasma. Several weeks after infusion of intravenous radiolabeled cholesterol, the specific radioactivity of lymph cholesterol exceeded that of plasma cholesterol (31). The HDLs of lymph were enriched in large apoA1-containing particles with a high content of unesterified cholesterol, phospholipids, and apo E (7, 9, 29). At the other end of the size spectrum, the smallest apoA1-containing particles were enriched in phospholipid. Lymph also contained discoidal HDL particles (29). Such HDLs are never seen in plasma except in subjects with familial lecithin-cholesterol acyltransferase (LCAT) deficiency. Of particular relevance to cholesterol clearance from the periphery, total cholesterol concentration in lymph HDL was about 30% greater than could be explained.
by the transendothelial transfer of HDL from plasma, indicating that HDLs acquire cholesterol from cells within the extravascular compartment. Pre-β-HDL concentration in lymph was positively and independently related to both plasma pre-β-HDL and lymph α-β-HDL (mature, spheroidal cholesteryl ester-rich [CE-rich] HDL) concentrations, suggesting that lymph pre-β-HDL particles result not only from transport out of plasma, but also from remodeling of plasma-derived α-β-HDL in interstitial fluid (Figure 2). Concordant results were obtained when the same particles were quantified by chromatography (32). Subsequent in vitro incubation studies showed that lipoprotein remodeling in interstitial fluid generates pre-β-HDL from α-β-HDL, in contrast to plasma in which there is net conversion in the reverse direction (7). That the duration of these incubations was no greater than the apparent average residence time of HDL in the extracellular matrix in humans (33) suggests that this process may be an important source of pre-β-HDLs in vivo.

By comparing cholesterol contents of size subclasses of HDLs in lymph with those in plasma, the rate of whole body cholesterol transport via lymph was estimated to average 0.89 mmol (344 mg) per day, which was compatible with published estimates of whole body cholesterol turnover by isotope dilution analysis (34). Taken together, these studies suggest that the interstitium is a major site for generation of pre-β-HDL, and implicate the lymphatic vasculature as the main transit route for movement of HDL from the interstitium to the bloodstream and liver. The concept that HDL relies on lymphatics to return to the blood during reverse cholesterol transport was demonstrated in mice, in which the lymphatic vasculature was surgically or genetically disrupted in skin, leading to marked reductions in the appearance of labeled cholesterol in plasma that originated from implanted tissue macrophages (35, 36).

Whether the number of HDL particles in interstitial fluid affects the rate of reverse cholesterol transport was assessed by intravenous infusion of reconstituted apoA1/lecithin discs, known to produce a rapid rise in plasma HDL concentration (37), into healthy humans previously given radiolabeled cholesterol (31). During seven days of continuous lymph collection, the infusion produced sequential increases in plasma HDLs, lymph pre-β-HDLs, lymph cholesterol-specific radioactivity (consistent with efflux of cholesterol from tissues), and fecal bile acid excretion. No changes occurred in lymph concentrations of enzymes that collectively remodel HDL (38), including LCAT, which esterifies free cholesterol for packaging into the central core of spherical HDLs, or phospholipid transfer protein (PLTP) and CE transfer protein (CETP), which catalyze the exchange of lipids between HDL particles or to other lipoprotein subclasses. Indeed, cholesterol esterification rate and CETP activity are both very low in lymph compared with plasma (7). On the other hand, PLTP has a higher specific activity in lymph than in plasma, owing to a greater ratio of active to inactive forms (7). High specific activity of PLTP, along with absence of cholesterol esterification, which promotes conversion of pre-β-HDL to α-β-HDL, likely contributes to the propensity of interstitial fluid to generate pre-β-HDL particles with a high activity for removing cholesterol from cells (7). In light of the need to better understand the impact of CETP inhibition on cholesterol transport in man (3), the effect of CETP activity or its inhibition on the efficiency with which HDL cycles through the interstitium is of paramount importance to investigate.

Overall, the collective evidence from these studies is consistent with the scheme in Figure 2. Passage of α-β-HDLs into the interstitium across endothelium may occur by transcytosis through endothelial cells, as suggested in in vitro studies (39, 40), but ongoing work in vivo suggests a 2-pore model of ultrafiltration rather than transcytosis (our unpublished observations). When the known radii of different HDL subclasses are compared with those of the pores, the shift from small to large HDL subclasses that occurs when scavenger receptor B1 (SR-B1) and CETP are reduced would be expected to produce a major reduction in total α-β-HDL transport into interstitial fluid, consistent with the observation that large lipoproteins do not enter the arterial wall (41). Small, lipid-poor apoA1-containing HDLs with pre-β electrophoretic mobility are generated in interstitial fluid by remodeling of spheroidal α-β-HDLs derived from plasma. The pre-β-HDLs in interstitial fluid then interact with ABCA1 transporters on extravascular cells to acquire unesterified cholesterol and phospholipid, resulting in formation of discoidal HDL, which travel to blood via the lymphatic system along with other macromolecules that exceed the radius of TNF-α (3.24 nm) (42). HDL ranges in radius from 3.82 to 5.43 nm (43, 44).

Upon re-entering blood at the thoracic duct, discoidal HDLs act as efficient substrates for LCAT, generating spheroidal α-β-HDLs rich in CE. Transfer of CEs to the liver occurs by two processes: direct uptake from α-β-HDLs via SR-B1 and transfer to VLDLs and LDLs via CETP. The cyclical extravascular-intravascular remodeling of HDL is critical to maintaining a flow of cholesterol from peripheral cells to the liver for re-utilization and elimination as bile acids.

Table 1
Major classes of lipoproteins

<table>
<thead>
<tr>
<th>Class</th>
<th>Origin</th>
<th>Major function</th>
<th>Diameter (nm)</th>
<th>Density (g/ml)</th>
<th>Major protein</th>
<th>Concentration in interstitial fluid</th>
<th>Average composition by weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chylomicrons</td>
<td>Ileum</td>
<td>Transport ingested fat and fat-soluble vitamins</td>
<td>200–1,000</td>
<td>&lt;0.95</td>
<td>apoB48</td>
<td>Absent</td>
<td>TG 85 CE 3 PL 8 Chol 2 Protein 1</td>
</tr>
<tr>
<td>VLDL</td>
<td>Liver</td>
<td>Transport synthesized glyceride</td>
<td>30–90</td>
<td>0.95–1.006</td>
<td>apoB100</td>
<td>Absent</td>
<td>TG 55 CE 18 PL 20 Chol 9</td>
</tr>
<tr>
<td>LDL</td>
<td>Lipolysis of VLDL</td>
<td>Deliver cholesterol to cells</td>
<td>22–28</td>
<td>1.006–1.063</td>
<td>apoB100</td>
<td>~9% that in plasma</td>
<td>TG 10 CE 50 PL 29 Chol 11 Protein 20</td>
</tr>
<tr>
<td>HDL</td>
<td>Liver and ileum</td>
<td>Reverse cholesterol transport</td>
<td>7–11</td>
<td>1.063–1.21</td>
<td>apoA1</td>
<td>~20% that in plasma</td>
<td>TG 6 CE 40 PL 46 Chol 7 Protein 50</td>
</tr>
</tbody>
</table>

TG, triglyceride; PL, phospholipid; Chol, cholesterol.
Lipoprotein entry into lymphatic vasculature: lessons from chylomicrons

Peripheral lymph differs in function and composition from intestinal lymph. The latter transports newly synthesized chylomicrons, which are at least an order of magnitude greater in radius than HDL, from the ileum to the bloodstream during the absorption of ingested fat. Together with the liver, the ileum is a significant source of newly synthesized apoA1, which appears in intestinal lymph partly as a component of nascent chylomicrons and partly in combination with phospholipid and cholesterol as discoidal nascent HDL (45, 46). The absorptive lymphatic capillaries for lipoproteins in the intestine extend as a single lymphatic vessel in each villus, termed lacteals (Figure 1). These drain into mesenteric collecting lymphatic vessels that run through muscular and neural control (47–49). This lymph runs through mesenteric LNs and ultimately into the thoracic duct that drains the transported chylomicron-rich lymph into the bloodstream at the left subclavian vein. It is notable that this pattern of transport results in passage of lymph-derived lipoproteins, including the nutrients collected as “fatty meal,” through not only mesenteric LNs (prior to entering the thoracic duct), but also through the heart and vasculature of the lung and subsequently other organs, where they are degraded by lipoprotein lipase. The remnant particles derived from these lipoproteins have access to the liver, strongly contrasting with non-fatty nutrients that directly enter the portal venous circulation from intestinal villi. One consequence of this route of transit is that the lung can be exposed to lipopolysaccharides and other components from intestinal microbiota with an affinity for lipoproteins that enter lymph (50). Therefore, when gut leakage of microbiota is sufficiently great to threaten organ failure, the lung is particularly susceptible (50). It is thus important that LNs, through which all lymph runs, respond to and filter absorbed lymph to protect the host against inflammatory lipoproteins. Mice lacking the lipoprotein lipase inhibitor angiopoietin-like 4, which is expressed in mesenteric LN macrophages, are susceptible to lethal inflammation within mesenteric LNs exposed to chylomicrons bearing saturated fats (51). Secretion of apoA1 by the intestine itself may also protect the host from mesenteric inflammation, given the anti-inflammatory properties of HDL (52). It would be interesting to test whether adverse responses to gut leakage are heightened when apoA1 secretion by enterocytes is selectively lost.

Figure 2
The intravascular/extravascular cycle of HDL remodeling that maintains reverse cholesterol transport. (i) Transfer of HDL across vascular endothelium. (ii) Production of small, lipid-poor apoA1-containing preβ-HDLs in interstitial fluid through the remodeling of spheroidal CE-rich α-HDLs. (iii) Conversion of preβ-HDLs to discoidal HDLs through uptake of unesterified cholesterol (cholesterol) and phospholipid (PL) via the ABCA1 transporters of peripheral cells. (iv) Transport of the discs via the lymphatic system to the blood via the thoracic duct. (v) Conversion of the discs to spheroidal CE-rich α-HDLs in plasma through the action of LCAT. (vi) Transfer of CE from α-HDLs to liver cells, directly via SR-B1 receptors and indirectly via CETP and apoB-containing lipoproteins (VLDLs and LDLs) that are endocytosed by apoB100 receptors. The principal function of LCAT is to generate CEs for delivery to the liver. The net production of preβ-HDLs in interstitial fluid appears to be maintained by a high ratio of active to inactive PLTP in the presence of a near-zero cholesterol esterification rate, in contrast to a high esterification rate and lower active/inactive PLTP ratio in plasma. Black arrows represent the path of apoA1 as a component of different HDLs as they move between the intravascular and extravascular compartments. Red arrows represent the flow of cholesterol, initially as unesterified cholesterol in interstitial fluid and lymph, and then as CE in blood.
nisms like macropinocytosis contribute to entry of fluid into the lymphatic vessel (17). In opposition to the concept of receptor-independent entry, Lim et al. reported that SR-B1 is required for the uptake of HDL in the skin lymphatic vasculature (36), implying that HDL could become trapped in tissues by loss of SR-B1 on lymphatic capillaries. If this finding is widely applicable, the premise that assessment of lipoproteins in afferent lymph mirror those in interstitial fluid may not hold under at least some circumstances. By contrast, the presence or absence of SR-B1 does not modulate chylomicron absorption in the intestine (53). Perhaps alternative explanations exist for the proposed role of SR-B1 in HDL transit out of the skin. For instance, the SR-B1−deficient mouse accumulates very large HDL particles in the circulation (54, 55), such that the plasma may be unable to supply the interstitium with HDL acceptors, leading to an alternative explanation for the failure of reverse cholesterol transport of exogenously administered cholesterol. Furthermore, SR-B1 plays a pivotal role in platelet function (56). Platelets are critical mediators in development of the lymphatic vasculature through their expression of CLEC2 (57–59), a key C-type lectin receptor for podoplanin that is widely expressed on lymphatic endothelial cells. Thus, the skin lymphatic vasculature may be abnormal in SR-B1−deficient mice, thereby negatively affecting reverse cholesterol transport without direct receptor-mediated uptake of HDL into lymphatics. On the other hand, as evidence indicates an extrahepatic role for SR-B1 in promoting cardiovascular disease (60), the possibility that either the uptake of HDL into tissue via vascular endothelium, which may at least partially require SR-B1 (61), or its egress from tissue via lymphatics (36) might contribute to poor reverse cholesterol transport, and therefore to atherosclerosis, is plausible and intriguing.

Though more is known about uptake of chylomicrons into lacteals than about uptake of lipoproteins into peripheral lymphatics, there is a paucity of literature overall in this important area. Newborn mice deficient in pleomorphic adenoma gene-like 2 (Plag2) succumb to a wasting syndrome stemming from failed chylomicron absorption (62). In this study, PLAG2 was highly expressed by enterocytes, with at least some expression by lacteals as well. Oil red O staining indicated accumulation of lipid within enterocytes, while electron micrographs revealed that chylomicrons were released from the epithelium but could not enter the lacteal. Plag2 deficiency also impeded the uptake of cholesterol into other tissues from the plasma. There has been little follow-up to this study, and it remains unclear whether chylomicron uptake is coordinated by lacteals in a Plag2-dependent manner or if other changes related to chylomicron secretion, composition, or size account for the outcomes observed.

With regard to particle size, it is possible to develop a working model to explain why fats packaged in chylomicrons exclusively enter the lymphatic vasculature, whereas most nutrients traverse the portal venous system for primary delivery to the liver. The blood capillary network that surrounds each lacteal is fenestrated, particularly along the venous side. These fenestrations facilitate resorption of nutrients, even as they also allow ultrafiltration of molecules from the systemic vasculature into the lamina propria. However, the fenestrae are too narrow to permit passage of even the smaller size range of chylomicrons (63). Therefore, size exclusion, much like the mechanism postulated for HDL entry into skin lymphatics (42), likely accounts for why chylomicrons are directed to the periphery through the lymphatic vasculature, rather than to liver through the portal venous vasculature. The tip of the lacteal is thought to contain large pores that, in contrast to the nearby blood vessels, are sufficient in size to allow chylomicron entry (64). That the fenestrated blood vessels lie atop the lacteal around much of its exposed surface likely contributes to ensuring that smaller molecules, including hydrophilic nutrients and antigens not packaged in chylomicrons, primarily access the blood vasculature for transport, although there is a certain probability that a portion of these small molecules bypasses the fenestrated vasculature and enters the lymph (Figure 1), consistent with experimental observations. Furthermore, the rich macrophage and DC network in intestinal villi acquires many macromolecules that enter intestinal villi through robust endocytosis (65). Their collective endocytic activity protected the lacteal from absorption of tracer antigens, whereas depletion of these cells allowed increased absorption into the lacteal, with a resulting shift in the ensuing immune response (65). The study did not investigate whether the presence of macrophages and DCs affected chylomicron absorption or absorption of other nutrients. This issue deserves attention in future research because of its important implications. First, drugs engineered to target chylomicrons for transport into the lymphatic vasculature might avoid the liver toxicity sometimes associated with higher doses of drugs that are transported through the portal venous vasculature (66). On the other hand, the fact that environmental toxins like dichlorodiphenyltrichloroethane first gain access to the systemic circulation, rather than the portal circulation, where they could be detoxified by the liver, enhances the danger they pose to human health (67). Thus, for reasons ranging from maintenance of cardiovascular and immunologic health to the avoidance of drug toxicity, a better understanding is needed about how lymphatic vessels in the intestine absorb chylomicrons and other macromolecules. We believe these studies have merit in their own right and also provide a basis for future studies on HDL entry into lymphatics in the periphery.

Atherosclerosis and cholesterol transport through interstitium and lymph

Successful clinical interventions to improve cardiovascular health based on targeting HDL may require that we more thoroughly explore how the HDL cycle (Figure 2) is regulated and how it may differ in various tissues and organs. The propensity of macrophages to donate cholesterol to HDL during reverse cholesterol transport differs between different anatomical compartments (35). Yet the reason for this remains unclear. Is apoA1 more enriched in interstitial fluid at different anatomical sites? Or does the relative interstitial fluid space around macrophages influence reverse cholesterol transport, such that sites of inflammation, for example atherosclerotic plaque, where macrophages are aggregated would support a more sluggish rate of reverse cholesterol transport? Does the rate of interstitial fluid flow measurably influence the rate and extent of reverse cholesterol transport? Does the efficiency of lymph transport overall affect the development or reversibility of cholesterol-driven diseases like atherosclerosis?

Answering these questions requires more research on the role of lymphatics in clearing cholesterol from artery walls where atherosclerotic plaques occur. The adventitia of large arteries is supplied with lymphatic vasculature as part of the vasa vasorum, and advanced atherosclerotic plaques promote the growth of lymphatic vessels within the intima of plaques (68). Martel et al. employed a surgical technique that suggests lymphatic vessels mediate removal of cholesterol from the artery wall (35). This work, reviewed in greater detail elsewhere (69), needs to be verified in models that do not require lymphatic remodeling as part
of the experiment. This goal will likely require experimental models with larger lymphatic vessels than are observed in mice. Experimental surgical interventions in the pig do not require full aortic transplant. Remarkably, delivery of labeled cholesterol esters, either in the form of LDL or HDL, to a ligated and temporarily bypassed segment of the pig thoracic aorta revealed that HDL passes through the media and enters adventitia efficiently, leading investigators in the late 1980s to conclude that HDL was likely cleared through adventitial lymphatics (70, 71). Labeled HDL, by contrast, penetrated only into the intimal layer, indicating specificity in trafficking through the medial wall for labeled HDL.

Nonetheless, the transport of HDL in arteries is less well studied and may differ from skin, so caution should be exercised in extrapolating from studies of peripheral lymph to the interstitial fluid of diseased arteries. Furthermore, much of the apoA1 in plaque has been rendered dysfunctional (72). On the other hand, as skin is the largest organ in the body, a substantial fraction of apoA1 is continually found there, making skin a key player in the HDL cycle regardless of whether lipoproteins are transported from arteries. Indeed, hypercholesterolemic mice lacking apoA1 suffer from massive sequestration of cholesterol predominantly in skin (73). Hypercholesterolemia also impairs lymphatic transport from the skin (74), but quantifying transport from other body sites, including the artery wall, will require the development of novel assays.

Conclusion

Because it has proven more challenging than expected to understand the HDL cycle well enough to manipulate it therapeutically, significant attention should be focused on the half of apoA1-bearing HDL particles found within interstitium. This part of the HDL life cycle remains relatively inaccessible for study, yet transit through the interstitium is certainly as critical as the period that HDL spends in plasma. Although recent studies have recognized that simple measurements of plasma HDL cholesterol are insufficient to predict efficacy in promoting cholesterol efflux, new assays to measure HDL function still focus on HDL in the plasma (64), making it impossible to determine whether some individuals have defective trafficking or activity of HDL within the interstitium. However, a critical unanswered question is whether evaluation of HDL remodeling and passage through skin, by far the largest and most accessible tissue, would be valuable or detract from our understanding of HDL in the interstitium of the artery wall. On the other hand, focusing on mechanisms that regulate passage of HDL through any interstitium, including those that enhance passage of nascent HDL into tissues and support its ability to later enter lymph loaded with large amounts of cholesterol, would likely benefit our understanding of cholesterol uptake by HDL in all tissues, including the artery wall.

Easton et al. infused reconstituted HDL and observed clearance of apoA1 from the plasma in a biphasic manner (75), with a secondary rise in apoA1 between 24 to 48 hours, consistent with two pools of HDL (75, 76). The second pool is likely the interstitial pool of HDL, in which HDL has a mean residence time of approximately 29 hours (33). Thus, the second rise of apoA1 may mark the return of HDL to plasma after its transit through the interstitium, much of it likely in transit through the large organ of the skin. Nanjee et al. (31) observed a similar biphasic effect on plasma preβ-HDL concentration after intravenous infusion of reconstituted HDL, compatible with delayed appearance in plasma of preβ-HDLs generated from increased remodeling of plasma-derived α-HDLs in the interstitium. Detailed assessment of this biphasic clearance is warranted. If it serves as a readout of interstitial passage of HDL, an assay more accessible than lymph cannulation may emerge to allow estimates of HDL flux through the interstitium in large cohorts of people. These assessments in turn would make it possible to determine whether such information provides valuable predictors for coronary health.

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