Oxidized Low-Density Lipoprotein Drives Dysfunction of the Liver Lymphatic System

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SUMMARY
Chronic liver disease results in lymphatic dysfunction in the liver which is rescued by administration of recombinant vascular endothelial growth factor C. The lymphatic dysfunction causes a decrease in lymphatic transport of molecules and occurs via oxidized LDL dependent loss of lymphatic identity.

BACKGROUND AND AIDS: As the incidence of nonalcoholic steatohepatitis (NASH) continues to rise, understanding how normal liver functions are affected during disease is required before developing novel therapeutics which could reduce morbidity and mortality. However, very little is understood about how the transport of proteins and cells from the liver by the lymphatic vasculature is affected by inflammatory mediators or during disease.

METHODS: To answer these questions, we utilized a well-validated mouse model of NASH and exposure to highly oxidized low density lipoprotein (oxLDL). In addition to single cell sequencing, multiplexed immunofluorescence and metabolomic analysis of liver lymphatic endothelial cells (LEC)s we evaluated lymphatic permeability and transport both in vitro and in vivo.

RESULTS: Confirming similarities between human and mouse liver lymphatic vasculature in NASH, we found that the lymphatic vasculature expands as disease progresses and results in the downregulation of genes important to lymphatic identity and function. We also demonstrate, in mice with NASH, that fluorescein isothiocyanate (FITC) dextran does not accumulate in the liver draining lymph node upon intrahepatic
injection, a defect that was rescued with therapeutic administration of the lymphatic growth factor, recombinant vascular endothelial growth factor C (rVEGFC). Similarly, exposure to oxLDL reduced the amount of FITC-dextran in the portal draining lymph node and through an LEC monolayer. We provide evidence that the mechanism by which oxLDL impacts lymphatic permeability is via a reduction in Proxl expression which decreases lymphatic specific gene expression, impedes LEC metabolism and reorganizes the highly permeable lymphatic cell-cell junctions which are a defining feature of lymphatic capillaries.

CONCLUSIONS: We identify oxLDL as a major contributor to decreased lymphatic permeability in the liver, a change which is consistent with decreased protein homeostasis and increased inflammation during chronic liver disease. (Cell Mol Gastroenterol Hepatol 2021;11:573–595; https://doi.org/10.1016/j.jcmgh.2020.09.007)

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Currently, in the United States, ~17,000 people are on the liver transplant waiting list and only 38% of those people will receive a transplant due to organ shortage (Organ Procurement and Transplant Network data as of 2019). The number of people on the waiting list is projected to increase by 23% in the next 15 years, with nonalcoholic steatohepatitis (NASH) projected to become the leading indication for liver transplantation in the United States.1,2 The current health care expenditure associated with liver transplantation is approximately $1.43 million per patient, which will increase to $2.1 million in the next 15 years. These data highlight the need for alternative treatment options for patients with chronic liver disease. Thus, the development of novel targets and therapeutic options for patients that currently have none is needed.

The liver lymphatic system, while largely understudied, holds an essential role in organ health. The primary function of the lymphatic system is to transport interstitial fluid (lymph) from the tissue to the circulatory system for removal from the body.3 The lymphatics also participate in the transport of proteins and lipoproteins to maintain protein homeostasis4 and trafficking of tissue resident immune cells such as dendritic cells to the lymph node (LN).5 Previous studies spanning the years of 1960–1985 demonstrated that during cirrhosis, the lymph draining from the liver is depleted of high-molecular-weight proteins in both human and animal models.6–8 These studies postulated that low lymph protein content in cirrhosis was a result of decreased permeability of the microvascular system. However, this interpretation was limited by the tools available at the time, and thus how lymphatic or vascular permeability contributed to these differences was unknown. Since 1985, new markers and regulators of the lymphatics have been discovered. One of these regulators is PROX1, a transcription factor required for lineage commitment of lymphatic endothelial cells (LECs),9,10 which regulates the expression of other lymphatic markers including podoplanin (PDPN), lymphatic vessel endothelial receptor 1 (LYVE-1),11 and vascular endothelial growth factor receptor 3 (VEGFR3).12

In the last 35 years, these markers have allowed investigators to distinguish between blood and lymphatic vasculature and develop assays to model LEC signaling pathways and specific LEC functions that occur in vivo.13–16 Perhaps one of the most striking findings in the last 15 years was the discovery of the functionally specialized button-like junctions in LECs found in the lymphatic capillaries.17 These button junctions allow for cells and proteins to pass freely from the tissue into the lymph for immune surveillance in the LN.18 Whether these lymphatic functions are required in the liver to mitigate inflammation is unknown and has been largely overlooked, particularly in the setting of chronic liver diseases, such as NASH.

While oxidized low-density lipoprotein (oxLDL) is a key player in the pathogenesis of atherosclerosis,19 mounting evidence suggests that oxLDL can contribute to the progression of liver disease.20,21 OxLDL accumulates as a result of free radicals generated by inflammation and can directly induce liver injury.14 Further, levels of oxLDL are elevated in the serum of patients with NASH,22 hepatitis C infection,23 alcoholic liver disease,24 and cholestasis.25 Antibodies specific to oxLDL may be a biomarker for people underlying NASH23 and neutralization of oxLDL ameliorates experimental nonalcoholic fatty liver disease (NAFLD), a precursor to NASH, in mice.27 Finally, lipid lowering medication has been demonstrated to reduce liver inflammation in both mice and humans.28–30 These findings point to oxLDL as being an important mediator in the progression of chronic liver disease, however the precise mechanism(s) by which oxLDL influences liver inflammation is unknown.

In this study, we demonstrate in both people and mice with fatty liver disease that the lymphatic system draining the liver is significantly altered. Specifically, using single-cell sequencing, we demonstrate that LECs from a diseased liver have a significant reduction of transcripts that are associated with LEC identity and function. Further, we find defects in the ability of the lymphatic vasculature to properly
transit fluorescein isothiocyanate (FITC) dextran to the portal lymph node during disease. A finding which can be rescued by treatment with a lymphatic growth factor, vascular endothelial growth factor C (VEGFC). We identify significant alterations in LECs in the presence of oxLDL, an inflammatory mediator of fatty liver disease, including changes in transcription, metabolism and permeability. Finally, we demonstrate defects in FITC-dextran transport from the liver to the portal LN could be directly induced by systemic treatment with oxLDL, an effect that was independent of liver inflammation. Together, these data combined with published findings provide evidence that protein homeostasis, maintained from chronic liver inflammation and ameliorate disease.

Results

The Lymphatic Vasculature Increases in the Liver During the Progression of NAFLD and NASH

Our previous publication demonstrated that lymphatic vessels were significantly expanded in the liver during cirrhosis, independent of disease etiology. However, it was unclear if the liver lymphatic vasculature expanded during the progression of disease or if this lymphatic expansion was merely a consequence of changes in the liver associated with cirrhosis. To answer this question, we examined the lymphatic vessel density (LVD) of the lymphatic vasculature in portal and fibrotic regions of livers. We assessed biopsies from 45 individuals with various stages of NAFLD and NASH who received liver biopsy during elective gastric bypass surgery or who were referred to our hepatology clinic with a preliminary diagnosis of NASH (Table 1). Prior to inclusion in our study, patient biopsies were staged for fibrosis utilizing Brunt staging by a pathologist blinded to other clinical parameters. Using this patient cohort, we found that the lymphatic vasculature in the human liver expands with the progression of disease as measured by PDNP staining (Figure 1A) and quantified based on lymphatic area divided by tissue area (density) (Figure 1B). These findings demonstrate that the liver lymphatic vasculature significantly increases in parallel with the progression of disease and the observed expansion of the liver lymphatic vasculature during end-stage liver disease is not merely a consequence of cirrhosis.

A Murine Model of NASH Induced Liver Lymphangiogenesis and Reduced Lymphatic Marker Expression

The previous findings demonstrate that the lymphatic vasculature expands in the human liver during disease. To validate that similar processes occur in murine models of chronic liver disease, we stained numerous murine models of liver disease with PDPN, LYVE-1, and cytokeratin-19 (CK19). Similar to our findings in human disease, we found that the lymphatic vasculature expands in a murine model of NASH, defined as PDPN+LYVE-1+CK19+ lymphatic vasculature and not PDPN+LYVE-1-CK19+ cholangiocytes (Figure 2A and B). As with our studies of human disease, the lymphatic expansion was caused by division of lymphatic endothelial cells over time, as we observed an increase in the frequency of Ki67+ lymphatic vasculature as disease progresses (Figure 2C). To better understand the effect of NASH on liver lymphatics, we used an established high-fat, high-cholesterol (HFHC) diet (45% fat and 2% cholesterol). We chose a diet high in cholesterol, as our previous findings demonstrated that cholesterol caused a significant decline in the expression of lymphatic lineage-specific genes, PROX1 and FLT4/VEGFR3, and led to impaired stability of lymphatic vessel like structures in vitro. Upon staining of the HFHC livers we noticed the mean fluorescence intensity of both PDNP and LYVE-1 was decreased in the lymphatic vasculature (Figure 2D) consistent with in vitro data demonstrating that Prox1, a regulator of these genes, was downregulated when exposed to oxLDL. Additionally, this increase of lymphatic density in murine livers during disease was independent of disease etiology (MDR2/-, Lieber-DeCarli diet and high-fat, low-carbohydrate diet).

Table 1. Patient Demographics

| Age, y | F0 (n = 7) | F1 (n = 9) | F2 (n = 9) | F3 (n = 10) | F4 (n = 9) | P Value |
|--------|-----------|-----------|-----------|------------|-----------|---------|
| Male   | 3 (43)    | 4 (44)    | 4 (44)    | 7 (70)     | 4 (44)    | .722    |
| Female | 4 (57)    | 5 (56)    | 5 (56)    | 3 (30)     | 5 (56)    | .999    |
| Race   | White/Caucasian | 7 | 9 | 9 | 8 | 9 | .999 |
|        | African American | 0 | 0 | 0 | 1 | 0 |    |
|        | Other      | 0 | 0 | 0 | 1 | 0 |    |
| Ethnicity | Non-Hispanic | 6 | 8 | 6 | 9 | 5 |    |
|        | Hispanic   | 1 | 1 | 3 | 0 | 4 | .155 |
|        | Other      | 0 | 0 | 0 | 1 | 0 |    |

Values are mean ± SD, n (%), or n.

*F0 vs F4 was significant (P = .0471), all other pairwise comparisons were nonsignificant (Kruskal-Wallis); others are Fisher’s exact test.
low-cholesterol diet) (Figure 2E and F).\(^{33}\) Using the HFHC diet model, we found that after 17–24 weeks on the diet, the mice developed steatosis and inflammation, increased aspartate aminotransferase (AST) and alanine aminotransferase (ALT), and increased total cholesterol, high-density lipoprotein, and LDL (Figure 3A–D). As our model of NASH demonstrates high liver LDL levels, these findings are also consistent with increased LDL exposure causing lymphatic-specific gene downregulation (Figure 2D). Finally, VEGFC transcript was increased in the liver, suggesting that the lymphatic expansion observed was caused by VEGFR3 engagement on the lymphatic vasculature (Figure 3E). Together, these data demonstrate that murine models of chronic liver disease can accurately recapitulate the lymphatic changes we find in human LECs exposed to cholesterol and in human chronic liver disease.

**Chronic HFHC Diet Feeding Induces Transcriptional Changes in LECs.** To determine if diet-induced liver disease, using our NASH model, alters the transcriptional profile of the lymphatic vasculature in the absence of cirrhosis, we utilized single-cell messenger RNA sequencing to determine differences in LEC transcriptional profile. Specifically, age-matched male mice were fed either a chow or HFHC diet for 22–32 weeks, and endothelial cells were enriched via flow cytometric sorting (Figure 4A). Sorted endothelial cells were then subjected to single cell sequencing as described.\(^{31}\) LECs from control-fed mice were first clustered using clusifyr and then validated by canonical LEC gene expression such as Lyve-1, Pdpn, Flt4/Vegfr3, and Prox1 (Figure 4B). Upon unsupervised clustering of the endothelial cells in the liver, we found that LECs and portal endothelial cells (PECs) cluster similarly to our human studies,\(^{31}\) while liver sinusoidal endothelial cells (LSECs) are slightly more transcriptionally different than LEC or PECs (Figure 4C). This unbiased clustering allows us to
compare the transcriptional profiles of the LECs isolated from control- and HFHC-fed mice independent of the other cell types captured (Figures 4D and 5). Upon comparison of control LECs to HFHC LECs, we found significant disease pathways upregulated in LECs from the HFHC-fed mice. Using Ingenuity software analysis
(IPA version 52921811; Qiagen, Germantown, MD), we discovered significant upregulation of catabolism, protein synthesis, interactions and binding of endothelial cells, lipid synthesis, and fatty acid metabolism among others (Figure 4D). When looking at differences in specific gene regulation, in control- vs HFHC-fed LECs, we evaluated LEC specific genes, genes involved in interactions and binding of endothelial cells, or lipid synthesis and fatty acid metabolism based on our pathway analysis. Upon evaluation of LEC specific genes, we saw a striking downregulation of Lyve-1, Pdpn, and Prox1 in HFHC-fed mice (Figure 5A), consistent with our immunofluorescence staining of PDPN and LYVE-1 protein (Figure 2D) and the downregulation of Prox1 in LECs exposed to oxLDL.31 These changes were specific to LECs in the liver as the transcriptional profiles of either LSECs (Figure 5B) or PECs (Figure 5C) did not indicate an upregulation of VEGFR2 (Kdr), downregulation of VEGFR3 (Flt4), or an upregulation of VE-cadherin (Cdh5). Furthermore, the ratio between Vegfr2 and Vegfr3, thought to be important for regulating permeable button-like structures between LECs,32 was different in HFHC-fed compared with control-fed mice (Figure 5D). When evaluating genes involved in interactions and binding of endothelial cells, we saw little difference in LEC expression of integrin alpha 9 (Itga9); however VE-cadherin (Cdh5) expression appeared to be increased in diseased LECs (Figure 5A). Also consistent with increased lipid synthesis in LECs we observed an increase in expression of Cd36, a gene also shown to be increased in response to oxLDL as well as in liver LECs from individuals with NASH.31 Finally, the proportion of LECs in active cell cycle (S/G2M), based on a defined set of genes known to regulate different stages of the cell cycle, was increased in HFHC-fed mice (Figure 5E), consistent with our findings in humans with NASH and our Ki67 staining (Figure 2C). As we saw Vegfc transcript upregulation using whole-liver reverse-transcription quantitative polymerase chain reaction (RT-qPCR) (Figure 3E) we asked about Vegfc expression by the cell types we evaluated. We found that of the cell types we collected, LSECs and PECs were the primary source of Vegfc in the liver (Figure 5F). However, as we sorted for endothelial cells, our data do not rule out Vegfc expression by other cell types not collected in this analysis. In summary, the transcriptional profile of LECs from HFHC livers was consistent with changes in the LEC transcriptome that reflect both proliferation and a potential de-differentiation of liver LECs in the context of disease, which may impact their functions, such as permeability and metabolism.

Recombinant VEGFC (cys156ser) Promotes Lymphangiogenesis of LECs of the Liver

Access to the prolymphangiogenic cytokine, VEGFC, has been shown to promote lymphatic function,36 the expression of Prox1,37 and thus Vegfr3.37,38 Because we found a decrease in Vegfr3 expression by LECs in the HFHC diet we assessed if treating mice with rVEGFC (cys156ser) could promote lymphatic growth and differentiation. rVEGFC containing the cys156ser mutation selectively binds VEGFR3 homodimers found predominantly on lymphatic vessels.39,40 LSECs, the primary endothelial cell found in the liver, express both VEGFR2 and VEGFR3 but in the form of a VEGFR2/3 heterodimer and utilize VEGFR2 for regeneration.41 Thus, we asked if the cys156ser rVEGFC protein caused increased division of the LECs. We confirmed that rVEGFC (cys156ser) treatment resulted in increased lymphangiogenesis by quantification of the LVD in livers of treated mice (Figure 6A and B). This was in contrast to cholangiocytes, which also express VEGFR3 and do not change in frequency with rVEGFC (cys156ser) but do increase in frequency in the liver of HFHC mice (Figure 6C). Indeed, the injection of rVEGFC (cys156ser) to mice fed the HFHC diet for 3 weeks resulted in a significant increase in the frequency of Ki67+ LECs in the liver, but not Ki67+ LSECs (Figure 6D). While the rVEGFC (cys156ser) was specific to LEC proliferation and not LSEC proliferation we cannot rule out the possibility that the rVEGFC could affect other LSECs functions.

HFHC Diet Results in Defective Lymphatic Function in the Liver, Which Can Be Rescued by Administration of Recombinant VEGFC (cys156ser)

We observed lymphangiogenesis during disease progression in both humans (Figure 1) and mice (Figure 2), a transcriptional profile of liver LECs that was consistent with the loss of lymphatic identity (Figures 4 and 5), and lymphangiogenesis in the liver in response to rVEGFC (cys156ser). Therefore, we predicted that lymphatic

Figure 2. (See previous page). Liver lymphatic vessels increase in frequency during disease. (A) Mice were fed a control or HFHC diet for 24 weeks. Staining of liver sections was performed with DAPI (white), LYVE-1 (cyan), PDPN 8.8.1 (red), and CK19 (yellow). Scale bar is 100 μm. Shown is a representative image from 1 experiment with 5–7 mice per group repeated 2 additional times with similar results and statistical values. (B) Lymphatic vessels were designated as PDPN+CK19− structures with at least 2 nuclei per vessel. LVD was calculated as described in the Materials and Methods. (C) Number of Ki67+ lymphatic vessels per area of tissue over a 12-week to 24-week period. Few to no Ki67+ lymphatic vessels were detected prior to feeding HFHC diet. Three mice per time point were evaluated. Experiment was repeated with similar results. (D) Mean fluorescence intensity as a measure of protein levels of PDNP and LYVE-1 using inform software in indicated group acquired from images in panels A and B. (E) Staining and quantification of liver sections from control, 12-week-old mdr2−/− mice (a model of primary sclerosing cholangitis) and mice gavaged with ethanol (2% v/v) for 5 weeks (Lieber-DeCarli diet model). Sections were stained with PDNP, CK19, and DAPI. Scale bar is 20 μm. Statistical analysis was performed using an unpaired t test in which *P < .05, **P < .01, and ***P < .001. ALD, alcoholic liver disease; BD, bile duct; LV, lymphatic vessel.
Figure 3. HFHC diet induces liver disease and increases cholesterol levels in mice. (A) Representative hematoxylin and eosin staining from control mice or mice fed an HFHC diet for 17 weeks (HFHC: 60% fat and 2% cholesterol). Scale bar is 20 μm. (B) Inflammation score (left) and steatosis score (right) for control or HFHC diet–fed mice. (C) Analysis of AST or ALT levels in the serum of mice fed either a control or HFHC diet for 17 weeks. (D) Quantitation of high-density lipoprotein (HDL), LDL, and total cholesterol (free and cell associated) by enzyme-linked immunosorbent assay from the liver or serum of mice fed a control or HFHC diet. (E) Whole-liver RT-qPCR of VEGFC transcript from control of HFHC diet. Shown are data from 1 independent experiment repeated at least 3 additional times. Statistical analysis was performed using an unpaired t test in which *P < .05, **P < .01, and ****P < .001.
Figure 4. Single-cell sequencing of liver LECs identifies transcriptional changes associated with HFHC diet-induced liver disease. (A) Gating strategy used for flow sorting of endothelial cells from control- or HFHC-fed mice. Non-parenchymal cells from the liver were isolated and stained for indicated markers after running through a live/dead selection. All CD31 positive cells were sorted into a single tube and then processed for single-cell RNA sequencing. (B) Heatmap of genes typically associated with LECs are shown as a validation of our cell classification. Yellow is genes upregulated and purple is genes downregulated. Shown are cells from control mice. (C) Uniform manifold approximation and projection (UMAP) plot to visualize the similarities between cell types acquired using the 10x Genomics single cell sequencing platform. Cells were classified using clustifyr. Inset box shows endothelial cell populations acquired. Shown are cells from control and HFHC mice. (D) Gene expression data from LECs only were entered into Ingenuity software for pathway analysis. Shown are pathways with z-scores ≥2 in LECs from HFHC diet–fed compared with control diet–fed. Green color represents highly activated pathways while red scores represent the downregulation of pathways. Data in figure are combined data from 2–5 mice per capture with 4 independent captures acquired by the 10x Genomics 3’ kit following cell sorting. FSC-H, forward scatter-height; FSC-W, forward scatter-width; HSC, hepatic stellate cell; IM, infiltrating macrophage; KC, Kupffer cell; SSC-W, side scatter-width.
function, required to maintain tissue homeostasis by accommodating for increased fluid and immune infiltrate in the liver, was impacted by disease and could potentially be rescued by rVEGFC. To better understand if diet modulated liver lymphatic function, we utilized a FITC-dextran assay. Specifically, we measured the ability of intrahepatically injected fluorescently labeled dextran to reach the liver-draining portal LN compared with a non-liver-draining LN (inguinal) (Figure 7A). We found that when 500-kD FITC-dextran (the approximate size of LDL that should normally traffic through the lymphatics) was injected into the liver parenchyma of control C57Bl/6 mice, that the FITC-labeled dextran reached the portal LN within 5 minutes as demonstrated. However, when we examined mice fed the HFHC diet for 20–24 weeks, we found a significant reduction in the amount of FITC-dextran in the portal draining LN

**Figure 5.** Liver LECs from mice fed an HFHC diet downregulate lymphatic genes and upregulate metabolic pathways. Violin plot of gene expression by (A) LECs, (B) LSECs, and (C) PECs. Shown are changes in gene expression in LECs from the livers of mice fed HFHC diet compared with Control diet. (D) Ridge plot showing that the distribution of per-cell Vegfr2/Vegfr3 ratios differs between the LECs from control- and HFHC-fed animals. A cell with a score of 0 in this metric has the average Vegfr2/Vegfr3 ratio, while a cell with a positive score has a higher than average Vegfr2/Vegfr3 ratio. (E) Cell cycle analysis of LECs (control/HFHC) based on gene expression from single cell sequencing. (F) Uniform manifold approximation and projection (UMAP) representation of Vegfc expression in single cells where purple indicates increased expression. Cells from control and HFHC livers are shown as in Figure 4C. Exact $P$ values are shown.
(dLN) compared with the non-draining inguinal LN (Figure 7B), suggesting a disruption in lymphatic drainage from the liver of HFHC diet-fed mice that have liver disease (Figure 3A–D). Importantly, treatment with rVEGFC (cys156ser) restored the drainage to the portal LN while not altering dextran drainage to the inguinal LN (Figure 7C) or plasma (Figure 7D). Furthermore, we found that this effect was not restricted to 500-kD FITC-dextran as mice fed a
HFHC diet also had less of 70 kDa FITC-dextran (the approximate size of albumin) in the portal dLN after intrahepatic injection (Figure 7E). We again saw a significant decrease in the expression of PDPN protein by lymphatic vessels in the liver of HFHC-fed mice compared with control-fed mice, by fluorescence intensity; however, the difference was minimized between control and HFHC mice treated with rVEGFC (cys156ser) (Figure 7F) suggesting the rVEGFC (cys156ser) not only promoted lymphangiogenesis (Figure 6), but also stabilized lymphatic gene expression. While this rVEGFC (cys156ser) treatment did not result in significant changes in ALT and AST or cholesterol levels, we did observe a moderate decrease in liver inflammation score by histological analysis (Figure 7G). This is consistent with an improvement in the ability of the lymphatic vasculature to allow exit of immune cells from the tissue and suggests, consistent with other reports,36 that rVEGFC treatment improves lymphatic function. Together, these studies demonstrate a significant defect in FITC-dextran transport from the liver to the portal LN of both 70- and 500-kDa FITC-dextran during diet-induced liver disease that is rescued by treatment with rVEGFC.

oxLDL Directly Impacts Lymphatic Function in the Liver. Because we have previously published that oxLDL is significantly elevated in the serum32 of mice fed an HFHC diet and that oxLDL reduces human LEC (hLEC) expression of PROX1 and FLT4/VEGFR3 in vitro,41 we next asked if oxLDL was increased in the liver tissue of mice fed the HFHC diet compared with control mice. We found a significant increase in the amount of oxLDL in the liver of mice fed the HFHC diet compared with age matched control fed mice (Figure 8A). As we had noticed similar transcriptional changes in LECs between mice fed the HFHC diet and LECs treated with oxLDL in vitro,31 we next asked if oxLDL was directly contributing to the changes in lymphatic trafficking of FITC-dextran to the portal dLN. To test this, we administered highly oxidized LDL to mice intravenously over a 2- to 3-week period and then performed the FITC-dextran assay (Figure 8B). Strikingly, there was a significant reduction in 500-kDa FITC-dextran in the portal dLN, similar to mice fed the HFHC diet, when oxLDL was given intravenously (Figure 8C). Moreover, this reduction in FITC-dextran accumulation in the portal dLN following oxLDL administration occurred in the absence of any pathological changes (Figure 8D) or inflammatory gene expression in the liver (Figure 8E). Furthermore, this difference in FITC-dextran was dependent on the oxidation state of the LDL, as we saw no difference in the amount of FITC-dextran in the portal LN when mice were administered native LDL (Figure 8F). These findings demonstrate that oxLDL, even in the absence of inflammation, can directly affect lymphatic function in the liver.

OxLDL Directly Affects Lymphatic Metabolism, Transcription, and Permeability In Vitro. The previous studies in concert with our previously published data demonstrate that oxLDL can have a significant effect on the in vivo function and transcriptional profile of LECs in the liver. As one of the main functions of the lymphatic vasculature is to allow for proteins and cells to pass through the highly permeable button-like junctions,17 we asked if oxLDL was impacting the permeability of the LECs. First, we confirmed that oxLDL injected in vivo was acquired by LECs using fluorescently labeled oxLDL (Figure 9A and B).32 Based on the acquisition of oxLDL by LECs in the liver, we next asked if oxLDL alone was enough to modify LEC function in vitro. It was recently published that LECs, in contrast to vascular endothelial cells, upregulate fatty acid oxidation (FAO) in a positive feedback loop that requires PROX1, p300, and CPT1a to promote lymphatic growth and lymphatic-specific chromatin modifications that differentiate LECs from vascular endothelial cells.42 As we observed an upregulation of metabolism, catabolism, and protein synthesis in our single-cell pathway analysis, we asked if and how oxLDL changes the metabolic state of LECs. Using mass spectrometry to evaluate metabolites, we discovered an apparent decrease in mitochondrial metabolism in the tricarboxylic acid cycle in LECs following oxLDL treatment (Figure 9C and D). In the oxLDL-treated LECs, we found a significant reduction in citrate, which is produced from acetyl-CoA via the glycolysis end-product pyruvate, and as a consequence of FAO. Late glycolytic intermediates and pyruvate were not significantly altered by oxLDL treatment, suggesting that the decline observed in citrate and downstream tricarboxylic acid cycle metabolites may be a result of decreased fatty acid metabolism in supplying carbons to the mitochondria (Figure 9C–E). Combined with our transcriptional data demonstrating a reduction in Prox1, Lyve-1, Pdpn, and Flt4/Vegfr3 in LECs from mice fed a HFHC diet, after in vitro treatment of LECs with oxLDL,37 as well as the requirement of Prox1 to upregulate FAO,45 these metabolic changes further demonstrate that LECs treated with oxLDL are unable to maintain LEC identity, perhaps making them more like the less permeable vascular endothelial cells, a
Figure 7. Liver lymphatic transport is impaired in HFHC-fed mice and rescued by rVEGFC. (A) Mice were fed with or without administration of rVEGFC (cys156ser). Prior to sacrifice, mice were anesthetized and livers were injected with 5 μL of 500-kD FITC-dextran into each of 3 liver lobes. Five minutes after administration of last injection, the portal-draining LN (PLN) and inguinal lymph nodes were removed. (B) Amount of FITC-labeled dextran was measured in portal at 5 minutes. Shown is the fold increase in FITC reading from portal LN to the inguinal lymph node from the same mouse. Fold increase over uninjected in labeled FITC-dextran from (C) the inguinal LN or (D) plasma. (E) Quantification of drainage of 70-kD FITC-dextran to the portal LN at 5 minutes in control or HFHC diet fed mice. (F) Mean fluorescence intensity of PDPN protein quantified from analyzed sections using InForm software as in Figure 2D. (G) Quantification of liver inflammation score by a pathologist blinded to the samples as in Lanaspa et al.53 Shown are combined data from 2 independent experiments with 3–6 mice per group. Statistical analysis in figure was performed using an unpaired t test in which *P < .05 and ***P < .001. CLN, celiac lymph node; dLN, draining lymph node; MLN, mesenteric lymph node; PV, portal vein.
consequence that could directly affect protein and cellular passage into the lymphatic capillaries.

We next asked if these transcriptional and metabolic changes in LECs could lead to differences in the ability of LECs to allow the passage of dextran (in lieu of proteins or cells) through their cell-cell junctions (i.e., permeability). To answer this question, we utilized a barrier transwell assay in which a confluent monolayer of hLECs were plated on the upper chamber of a transwell. Using this assay, we found that oxLDL treatment, for either 1 hour or 24 hours, reduced the ability of FITC-dextran to pass through the hLEC monolayer as measured by a decrease in the amount of FITC-dextran found in the lower well (Figure 10A). Upon staining with VE-cadherin of hLECs treated with oxLDL, we found a reorganization of the cellular junctions compared with vehicle-treated hLECs. In vehicle-treated hLECs we visualized highly permeable junctions, as demonstrated by distinct VE-cadherin positive junctions between cells. In contrast, oxLDL-treated hLECs had significantly increased staining between cell borders and less distinct junctions (Figure 10B). VE-cadherin protein expression by Western blot was significantly increased in hLECs treated with oxLDL (Figure 10C), consistent with the increase in fluorescence (Figure 10B) and transcription (Figure 5A) we found in liver...
LECs in mice fed the HFHC diets. We also observed an increase in the ratio of \textit{Vegfr2/Vegfr3} transcript in hLECs treated for 24 hours with oxLDL (Figure 10D), consistent with the increased ratio we found in LECs from mice fed the HFHC diet compared with control-fed mice (Figure 5D). In contrast, TNF\(\alpha\) treatment resulted in increased permeability of the hLECs as previously described (Figure 10E).46 These differences were specific to hLECs, as FITC passage through a layer of human primary umbilical vein endothelial cells (HUVECs) was unchanged following 24-hour treatment of
oxLDL (Figure 10F), and no changes were observed with HUVeCs in either VE-cadherin protein or transcript expression or junctions (Figure 10G and H). The increased ratio of Vegfr2/Vegfr3 was also specific to LECs, as we did not observe these changes in HUVeCs treated with oxLDL for 24 hours (Figure 10I) or LSECs or PECs from mice fed an HFHC diet compared with control (Figure 5A–C). These findings demonstrate a significant disruption in the ability of hLECs to maintain a lymphatic-specific transcriptional profile,11 metabolism, and highly permeable button-like junctions in the presence of oxLDL. When combined with our in vivo findings, these studies identify oxLDL as a critical factor that impedes lymphatic function in the liver during NASH and likely contributes to decreased protein homeostasis and inflammatory immune cell removal.

Discussion

Our data, in combination with historical clinical reports and published studies, support a model by which the liver lymphatic system plays a critical role in maintaining protein homeostasis and the exit of inflammatory immune cells from the liver. For decades, it has been appreciated that both the lymphatic vasculature in the liver and the outflow of lymph is dramatically increased in human and animal models of cirrhosis.64 However, it was unknown if these findings were merely a consequence of cirrhosis. The data in this study demonstrate that the liver lymphatic vasculature increases as disease progresses in humans with NAFLD and NASH. These findings are consistent with published observations that chronic liver disease in humans leads to increased expression of prolymphangiogenic growth factors in the liver.47 Importantly, we find that the LVD in the liver also increases in a well-validated mouse model of NASH; however, our studies demonstrate that even though the lymphatic vasculature expands, the role of the lymphatic vasculature in maintaining protein and immune cell clearance can be compromised by inflammatory mediators like oxLDL. These data increase our confidence that the role of the liver lymphatic vasculature is maintained across species and undergoes dramatic remodeling during the progression of chronic disease.

As noted previously, despite the increased lymphatic vasculature in the liver, and subsequent increased lymph drainage from the liver during cirrhosis, several reports demonstrated that the lymph draining the liver was depleted of high-molecular-weight proteins.7,48 A major function of the lymphatic vasculature is to allow the transport of proteins and immune cells through permeable button-like junctions, and thus we predicted that the expanded lymphatic vasculature was becoming less permeable during progressive stages of fibrosis and cirrhosis. Using single-cell messenger RNA sequencing we found that several transcripts critical for lymphatic function were decreased in the diseased liver LECs. Specifically, diseased liver LECs had decreased expression of Prox1, Lyve-1, Pdpn, and Flt4 (Vegfr3). These transcriptional changes are consistent with changes we saw after treatment of LECs with oxLDL.11 Furthermore, as Prox1 is required to upregulate FAO,49 we found that LECs treated with oxLDL had metabolic changes consistent with the decreased FAO and more similar to the metabolic profile of vascular endothelial cells. As Prox1 also regulates Vegfr3 expression11 it is perhaps not surprising that the ratio of Vegfr2/Vegfr3 is increased. Intriguingly, the balance of expression of the proteins VEGFR2 and VEGFR3, among others, have been shown to regulate the permeability of the lymphatic vasculature.54 Indeed, hLECs treated with oxLDL have both a significantly different VEGFR2/VEGFR3 ratio and are significantly less permeable than vehicle-treated LECs. This difference is also consistent in HFHC-fed compared with control-fed mice in which our single-cell analysis demonstrates an increase in the Vegfr2/Vegfr3 ratio and our in vivo data demonstrate a loss of FITC-dextran accumulation in the portal lymph node.

Together, these findings suggest that liver disease was directly causing a de-differentiation of lymphatic endothelial cells in the liver. Importantly, LECs from diseased liver did not completely adopt a vascular endothelial transcriptional profile. Upon comparison of control LECs to HFHC LECs we found 374 genes were significantly different with a P value of <.01 (Supplemental Table 1). Interestingly, while 774 genes were differentially regulated between control LECs and control LSECs (a blood endothelial cell) (Supplemental Table 2), 647 genes were differentially regulated between HFHC LECs and HFHC LSECs (Supplemental Table 3). While fewer genes had differential expression between LECs and LSECs from HFHC-fed mice compared with LECs and LSECs from control-fed mice, these cell types are still quite distinct. Thus, our results support the hypothesis that when LECs are exposed to an HFHC diet or oxLDL, they downregulate a
lymphatic gene program that defines at least some of their functions. Ultimately these cells do not become blood endothelial cells, but rather perhaps de-differentiate into an intermediate LEC-Blood endothelial cell phenotype, causing the LECs to perform in a similar manner as their blood progenitors in some respects such as cell-cell junctions and metabolic profile.

Indeed, when we tested if chronic liver disease altered the transport of FITC-dextran from the liver to the portal LN, we found a significant defect, which could be recovered by promoting lymphatic function with rVEGFC. It is important to note that using this assay, the dextran cannot be targeted only to the lymphatic vasculature, and previous studies have demonstrated a loss of LSEC fenestration.
during diet-induced liver disease.\textsuperscript{22} As fenestration of LSECs are believed to be key in regulating the protein content of the hepatic lymph, the results reported here are likely detecting changes in the permeability of both sinusoidal and lymphatic endothelium. However, the treatment of mice with the lymphatic growth factor VEGF-C (cys156ser), which did not affect LSEC division and thus may not influence LSEC fenestration, was able to restore lymphatic drainage from the liver of mice fed an HFHC diet. Thus, our studies indicate for the first time a defect in lymphatic function in the liver of mice with chronic liver disease, and that this defect can be rescued with a lymphatic growth factor. Furthermore, we demonstrate that treatment with rVEGFC improves liver inflammation, suggesting a positive link between lymphatic function and liver function.

Our studies focus in on the idea that decreased lymphatic function in the liver, observed during diet-induced disease, is likely a consequence of oxLDL signaling. Our data suggest that oxLDL directly results in a transcriptional and metabolic profile that points to a loss of LEC identity and thus LEC-specific functions. These differences are reflected in the inability of FITC-dextran to properly traffic through the lymphatic endothelial cell junctions, which contributes to the loss of intrahepatically injected FITC-dextran in the portal LN following a 2- to 3-week treatment with intravenous oxLDL. However, understanding how oxLDL affects lymphatic function in the liver, either directly or indirectly, via liver resident cells are the aims of future studies. Indeed, oxLDL was acquired by LECs, LSECs (Figure 9A and B) and macrophages.\textsuperscript{32} As both LSECs and macrophages are likely in close proximity to LECs and could produce or other additional factors which may alter LEC function. While this is a major consideration, the direct impact of oxLDL on LECs that we demonstrate here may provide an opportunity for understanding oxLDL sensing by LECs and the signaling mechanisms involved. Understanding these mechanisms is crucial for the development of effective therapies to improve liver lymphatic function as well as lymphatic function in other organs where the LECs may be exposed to oxLDL in the setting of ongoing disease.

As the rate of liver transplantation continues to rise with a marked increase in end-stage liver disease secondary to NASH, there is an increasing need for the development of therapeutics to slow disease progression. The data in this study highlight the liver lymphatic system as an untapped potential therapeutic target. Similar to published data in the setting of skin,\textsuperscript{49} adipose,\textsuperscript{50} heart,\textsuperscript{51} and kidney\textsuperscript{52} inflammation or ischemia-reperfusion injury in the liver,\textsuperscript{42} our data demonstrate that administration of recombinant VEGFC can reduce liver inflammation during diet-induced disease. These findings identify the lymphatic system as an attractive target to modulate liver inflammation, maintain protein homeostasis, and slow or reverse the progression of disease in combination with diet modification. Collectively, our data highlight the critical role of the lymphatic system in maintaining liver homeostasis and how disruption of lymphatic function during liver insult acts to potentiate mechanisms of disease.

**Materials and Methods**

**Immunohistochemistry, Microscopy, and Liver Histology**

For Human studies, liver biopsies were obtained from the University of Colorado Anschutz Research Histology Shared Resource. For patient demographics see table 1. Biopsies were cut in 5-μm-thick sections and adhered to a glass slide. Slides were deparaffinized and antigen retrieval
was done in TRS (Dako, Carpinteria, CA). Slides were blocked for 5 minutes in 0.3% hydrogen peroxide, followed by an hour block in 2% bovine serum albumin (BSA). Mouse anti-human PDPN (1:50, clone D2-40; Dako) was added to the biopsies for 2 hours. Anti-mouse secondary conjugated to HRP (Dako) was added for 30 minutes. DAB+ was used as the visualizing agent, and the slides were counterstained with hematoxylin. Whole slide images were taken using an Olympus IX83 microscope with a ZDC laser-based focusing system (Olympus, Tokyo, Japan). For analysis of lymphatic vessels in murine livers, at the time of harvest the median lobe of the liver was excised, the gallbladder was removed and tissue was fixed in 10% neutral buffered formalin at room temperature for 24 hours. Livers were then embedded in paraffin, cut into 5-μm-thick sections, placed on a glass slide, and stained as described.31 Briefly, slides were dewaxed with xylene, heat-treated in either pH 6 or pH 9 antigen retrieval buffer for 15 minutes in a pressure cooker, and blocked in antibody diluent (PerkinElmer, Waltham, MA). Sections were then sequentially stained for PDPN (8.1.1), Lyve-1 (ab33682), and CK19 (ab52625) primary antibodies followed by HRP-conjugated secondary polymer (anti-rabbit [PerkinElmer], anti-goat ab97110, or anti-hamster ab6892) and HRP-reactive OPAL fluorescent reagents (PerkinElmer). To image nuclei, slides were stained with spectral DAPI and coverslips were applied with Prolong Dimond mounting media (Thermo Fisher Scientific, Waltham, MA). Data were collected using the Vectra 3.0 Automated Quantitative Pathology Imaging system (PerkinElmer) as described.31 For quantification of PDPN and Lyve-1 fluorescence intensity, lymphatic vessels were identified, the lumen of the vessel was subtracted from analysis, and the median intensity of PDPN or Lyve-1 in the vessel area was calculated with inForm software (PerkinElmer). For Ki67 staining, anti-mouse Ki67 (clone SP6; Thermo Fisher Scientific), staining was performed with the previously mentioned antibodies to determine Ki67+ lymphatic vessels. Number of Ki67+ lymphatic vessels were manually counted from entire section and value per millimeter squared was determined. This study was approved by the institutional review board at the University of Colorado Anschutz Medical Campus.

**Animal Studies and Feeding Regimens**

All experiments utilized 6- to 8-week-old male C57BL/6 mice purchased from Charles River Laboratories (Wilmington, MA). For diet studies, mice were randomly allocated to either a chow control or HFHC diet (#D17010101-03) formulated by Research Diets Inc. (New Brunswick, NJ) and as described in McGettigan et al.12 Mice were provided diet ad libitum for a period between 20 and 32 weeks. For treatment studies, mice were injected (intraperitoneally) with rVEGFC (cys156ser) (10 mg/kg per dose) or every 2 days for 3 weeks. For studies of oxLDL regulation of lymphatic permeability, mice were injected intravenously with 150 μg of highly oxidized LDL (Kalen Biochemicals, German-town, MD) or phosphate-buffered saline (PBS) as a vehicle control in a total volume of 200 μL. Injections of oxLDL or vehicle occurred twice weekly for a period of 1–2 weeks. All animal studies were approved by the University of Colorado Anschutz Institutional Animal Care and Use Committee.

**Quantification of Liver Pathology**

For quantification of liver histology, liver sections were stained with hematoxylin and eosin and scored semi-quantitatively for steatosis and inflammation as previously described.52 Briefly, liver sections were given a numeric score for each feature within a given pathology category. Within each category individual pathological features were given a numerical score (0-4) based on how prevalent it was. Thus, the total scores for each pathology category (steatosis (max score 5) and inflammation (max score 14)) were simply a sum of the individual scores for each feature. Histology scoring of blinded liver sections was performed by D.I.O., who was blinded to the identity of the liver section being scored.

**Quantification of Serum and Liver Cholesterol Levels and Liver Enzymes**

Age- and sex-matched mice were fed either a control chow or HFHC diet, and plasma was obtained during the course of feeding via submandibular bleed or at the time of harvest with cardiac puncture. Whole blood was allowed to clot at room temperature for 10 minutes and serum was isolated by centrifugation. For analysis of liver cholesterol, portions of livers were weighed and subjected to analysis. Determination of cholesterol levels was performed using the cholesterol assay kit (ab65390; Abcam, Cambridge, United Kingdom) exactly as described by the manufacturer. Measurement of serum AST and ALT (lower limit 10 U/l; Fuji Dri-Chem Slide GPT/AST) obtained using a DriChem 7000 Veterinary Chemistry Analyzer (Heska, Loveland, CO).

**Quantification of Liver oxLDL**

Liver tissue was obtained at the time of harvest, weighed, rinsed with PBS to remove excess blood and homogenized with a glass homogenizer. OxLDL levels were determined using a mouse oxLDL ELISA kit (MBS2512757; Mybiosource, San Diego, CA) per manufacturer instructions.

**Semi-Quantitative PCR**

Briefly, snap-frozen liver tissue or cultured cells were homogenized in Buffer RLT and total RNA was isolated from cell lysate using the RNeasy Mini Kit (Qiagen, Hilden, Germany), and complementary DNA was synthesized using the QuantiTect RT Kit (Qiagen) following standard protocols. PCR amplification was performed using either the QuantiTect Syber green (Qiagen) or TaqMan Fast Advanced Master Mix (Applied Biosystems, Foster City, CA) PCR kits. Quantitative PCR was performed on a QuantStudio 3 Real-time PCR machine (Applied Biosystems) and fold changes in messenger RNA levels were calculated. For each gene, all samples were normalized to the average fold change of the control treatment group (chow or PBS). The following
Qiagen QuantiTect primer assays were used: 18S ribosomal RNA (Rn18s; QT02448075), transforming growth factor beta 1 (Tgfβ1; QT00145250), CD36 (QT01058253), and IL13 (QT00099554) for mouse and VEGFC FWD: 5’ ccaagt-gagggtgtgtagatg 3’, VEGFC REV: 5’ ctgcttgacactgtgtaat 3’; for human: VEGFR2(KDR): QT00069818, VEGFR3(FLT4):QT000063637, VE-Cad (CDH5): QT000013244 and GusB (housekeeping gene): QT000046046.

**Quantification of LVD**

LVD is defined as the area of the lymphatics divided by the area of the defined tissue area. For lymphatic density in human biopsies, images were opened in photoshop. Approximately nine 600 × 500 pixel areas were used and portal triads were cut out of the image. Each area (75 mm²) contained 6–9 portal triads that were evaluated per biopsy in which the lymphatic vessel was masked, area calculated, and then divided by the total area. Lymphatics were colored in using the color tool and the whole 1800 × 1500 area was opened with ImageJ version 1.52T (National Institutes of Health, Bethesda, MD). The image was converted into an 8-bit image and the threshold was adjusted so only the drawn lymphatic area was seen. The analyze particles function was used to determine the area of the lymphatics. Area of lymphatics was then divided by the total area of cut out image. For analysis of murine livers, LVD was quantified with Nikon AR software version 4.60 (Nikon, Melville, NY) as described.31 Regions of approximately 1.3 mm² were randomly selected and the area of the lymphatic vessel was determined in the entire area, in which at least 2 portal triads were in the field of measurement. LVD was calculated as the area of the vessels divided by the area of the tissue.

**Liver Cell Isolation and Fluorescence-Activated Cell Sorting**

Briefly, murine livers were harvested and non-parenchymal cells were isolated as previously described.24 Following isolation, cells were washed 2× with PBS and dead cells were removed using the EasySep Dead Cell Removal Kit (Stem Cell Technologies, Vancouver, Canada). Viable cells were washed 2× with PBS containing 0.5% fetal bovine serum (Atlas Biologics, Fort Collins, CO) and stained with antibodies against CD45 (30-F11), CD31 (390), CD146 (ME-9F1), and PDPN (8.1.1) from BioLegend (San Diego, CA). Cells were washed twice and were sorted using an aria Fusion sorter (BD Biosciences, Franklin Lakes, NJ). Enriched stromal cells were then washed 2× with PBS and subjected to single cell sequencing as described subsequently.

**Single-Cell RNA Sequencing**

Approximately 10,000 viable enriched stromal cells were loaded on a 10x Genomics controller (10x Genomics, San Francisco, CA) to generate barcoded single-cell Gel Bead in-Emulsions (GEMs) using the 10x Genomics 3’ kit as previously described.31 Complementary DNA libraries were sequenced using a NovaSeq 6000 (Illumina, San Diego, CA) at the University of Colorado Genomics Shared Resource Core. To control for batch effects 3 independent sequencing experiments were performed which included both control- and HFHC diet–fed mice. Quantification and analysis of single cell sequencing was performed using the R packages Seurat and scanr as previously described.31 Ridge plots were generated using Seurat’s RidgePlot function. Cells were classified and correlation plots were generated usingclusfifyr.55 For cell cycle analysis, Seurat assigns each cell a score based on its expression of G2/M and S phase markers. These marker sets are anti-correlated in their expression levels, and cells expressing neither are assigned to G1 phase. Cells with fewer than 250 detectable genes or >20% of unique molecular identifiers (UMIs) derived from mitochondrial genes were excluded from the analysis to eliminate cells with insufficient expression data for clustering and dead cells, respectively.

**Liver Lymphatic Drainage Assay**

Control or HFHC diet–fed mice or mice injected with oxLDL (150 μg) or PBS were anesthetized with 2%–2.5% isoflurane or a solution of ketamine (30–60 mg/kg) and xylazine (3–6 mg/kg) and placed on a heating pad. An incision was made into the peritoneum to expose the liver and 5 μL of a solution of 10 mg/mL of FITC-labeled dextran (70 or 500 kD) in PBS was injected into left, median, and right lobes of the liver using a 28-gauge needle (BD Biosciences). Five minutes after FITC-dextran administration, the liver-draining LN (portal) were excised and placed into separate wells containing 400-μL PBS. As a control for vascular drainage of the dextran, the inguinal (skin draining) LN was also excised. LNs were minced with 22-gauge needles, and 200 μL of the minced LN in PBS was placed in a 96-well Costar Assay Plate (Corning, Corning, NY), and FITC was read using a Synergy H1 microplate fluorescence plate reader (BioTek, Winooski, VT). Data were normalized to the inguinal LN from the same mouse taken immediately after the portal LN.

**In Vitro Permeability Assay**

Briefly, 6000–10,000 human lymphatic endothelial cells (hLECs) (PromoCell, Heidelberg, Germany) or HUVECs were plated on a 0.2% gelatin-coated 24-well Costar Transwell permeable support (6.55-mm insert) with a 0.4 mm Polyster Membrane (Corning). hLECs on the transwell were then stained for VE-Cadherin by either immunofluorescence or Western blot. A total of 72 hours following plating, cells were incubated with oxLDL (100 μg/mL), TNFα (10 ng/mL), or vehicle control (PBS) for 1 hour or 24 hours. Following incubation, 500-kD FITC Dextran (100 μg/mL) in Hanks’ balanced salt solution was added to the top of the well. Media were removed from the bottom of the transwell and replaced with HBSS. Plates were incubated at 37°C with gentle rocking and migration of the FITC-labeled dextrans was measured using a Synergy H1 microplate reader by removing 60 μL from the bottom of the transwell at indicated time points at an excitation of 485 nm and emission of 528 nm. For quantification of
FITC, a standard curve of FITC-dextran was read along with experimental samples. The actual amount of FITC-dextran was then calculated using the equation of the line based on the standard curve generated on the same day.

**VE-Cadherin Staining**

Human dermal lymphatic endothelial cells (hdLEC) staining of cells from in vitro permeability assay in the 24-well transwell plate were fixed after assay was complete with 4% paraformaldehyde for 15 minutes at room temperatures. Cells were then rinsed and blocked with 10% donkey serum, 2% BSA, for 1 hour at room temperature. Staining was performed with rabbit anti-VE-cadherin 1:100 (Abcam) and DAPI 1:1000 (BioLegend). The filter was removed from the transwell insert and mounted on a microscope slide and read on an Olympus microscope using cellSens 1.16 software. Quantification of pixel intensity was performed using Photoshop (Adobe, San Jose, CA). Fluorescence from VE-cadherin was measured in the red color channel, and then 3 equally sized areas from each experiment were measured by mean gray value using the measurement log function for both the PBS- and oxLDL-treated samples. The 3 measurements from each sample were averaged and divided by the PBS value average to obtain fold increase over PBS (*P < .05).

**Western Blots**

hLECs were grown to >80% confluency in T25 flasks then treated with oxLDL at 100 mg/mL for either 4 or 24 hours. Treated as well as untreated cells were washed twice with cold PBS and cells were coated with 500 μL of lysis buffer (from RayBiotech kit AAH-MAPK-1-2 prepared with supplied protease and phosphatase inhibitors to manufacturer instructions). Flasks were then immediately moved to 80°C and froze overnight. Cells were thawed and scraped into the lysis buffer. Lysate was collected and clarified by spinning at >12,000 g for 20 minutes. Supernatant was collected and quantified using BCA kit (23225; Thermo Fisher Scientific). Gel samples were made by combining equal amounts of protein from each sample diluted to equal volume with lysis buffer. 4× Laemml buffer with 10% 2-mercaptoethanol was added for a final concentration of 1×, then samples were heated on a heat block of 90°C for 10 minutes. Samples were run on 10% acrylamide gels and transferred to 0.45-μm polyvinylidene difluoride membrane. Membranes were blocked with 5% BSA/Tris-buffered saline with 0.1% Tween 20 detergent (TBST) for 30–60 minutes at room temperature while rocking. Then membranes were incubated in anti-VE-cadherin (AB33168 [Abcam] diluted 1:1000 or 1 μg/mL in 3% BSA/TBST) or anti-GAPDH-HRP (MA5-15738-HRP; Invitrogen, Carlsbad, CA) overnight at 4°C while rocking. Membranes were then washed with TBST 3 × 5 minutes then for VE-Cadherin incubated with anti-Rabbit-HRP (AB7090 [Abcam] 1:10,000 in 3% BSA/TBST) at room temperature for 1 hour while rocking. Membranes were then washed 3 × 5 minutes in TBST and imaged using Pierce ECL western blotting substrate and the ChemiDoc MP imaging system (Bio-Rad, Hercules, CA). Quantification of Western blot was performed using ImageLab (Bio-Rad) analysis software. Pixel intensity was quantified for each band and then normalized to the loading control for each individual sample before dividing the values by the time zero values in order to obtain the fold increase over time zero.

**Metabolite Analysis**

Mass spectrometry-based metabolomics was performed on frozen cell pellets. Cells were extracted at 2 × 10⁶ cells per mL in ice-cold lysis/extraction buffer (methanol:acetonitrile:water 5:3:2) and analyzed using a 5-minute C18 gradient on a Vanquish UHPLC system coupled online to a Q Exactive mass spectrometer (Thermo Fisher Scientific). Sample preparation, data acquisition, and data analysis were performed exactly as described.

**Statistical Analysis**

All statistical analysis was performed using a Student’s t test or 1-way analysis of variance with multiple comparisons to obtain a P value. P values are designated by asterisks in which *P < .05, **P < .01, ***P < .001, ****P < .0001. The number of animals per group was determined based on a power calculation. Statistically significant differences between control and experimental groups (P ≤ .05 with determined number of animals per group based on power calculation) were obtained in 2 experiments (1+1 repeat). If in 1 of the 2 experiments the significance level did not attain at least P ≤ .05, a third and deciding repeat was performed. Based on our Institutional Animal Care and Use Committee (IACUC) protocol, unnecessary animal experiments were not performed. For in vitro experiments, all experiments were performed a minimum of 3 independent times with 3–5 replicates per group. All authors had access to the study data and had reviewed and approved the final manuscript.

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