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Nuclear envelope–localized torsinA-LAP1 complex regulates hepatic VLDL secretion and steatosis

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Deciphering novel pathways that regulate liver lipid content has profound implications for understanding the pathophysiology of nonalcoholic fatty liver disease (NAFLD) and nonalcoholic steatohepatitis (NASH). Recent evidence suggests that the nuclear envelope is a site of regulation of lipid metabolism, but there is limited appreciation of the responsible mechanisms and molecular components within this organelle. We showed that conditional hepatocyte deletion of the inner nuclear membrane protein lamina-associated polypeptide 1 (LAPI) causes defective VLDL secretion and steatosis, including intranuclear lipid accumulation. LAPI binds to and activates torsinA, an AAA+ ATPase that resides in the perinuclear space and continuous main ER. Deletion of torsinA from mouse hepatocytes caused even greater reductions in VLDL secretion and profound steatosis. Mice from both of the mutant lines studied developed hepatic steatosis and subsequent steatohepatitis on a regular chow diet in the absence of whole-body insulin resistance or obesity. Our results establish an essential role for the nuclear envelope–localized torsinA-LAP1 complex in hepatic VLDL secretion and suggest that the torsinA pathway participates in the pathophysiology of NAFLD.

Introduction
Liver triglyceride (TG) metabolism consists of 3 broad processes. First is the accumulation of fatty acids by hepatocytes from the uptake of plasma fatty acids, the uptake of TG fatty acids in remnants of VLDL or chylomicrons, and de novo fatty acid synthesis from glucose. Second is the generation of energy via β-oxidation of fatty acids within hepatocytes. Third is the storage of TGs in adipose tissue. Potential links to lipid metabolism also exist for integral proteins of the inner nuclear membrane. The lamin B receptor is a polytopic inner nuclear membrane protein that is essential for cholesterol synthesis (12). Overexpression in hepatocellular carcinoma of a truncated variant of lamina-associated polypeptide 2α, a nucleocytoplasmic isoform of an inner nuclear membrane protein, alters fatty acid uptake (13). Deletion of the lamin-associated polypeptide 1 (LAPI), a monotopic protein of the inner nuclear membrane (actually 3 different isoforms with variable nucleoplasmic domains encoded by the same gene), causes an apparent expansion of the inner nuclear membrane, suggestive of altered lipid metabolism (14).

The nucleoplasmic domain of LAPI interacts with nuclear lamins and emerin, another integral protein of the inner nuclear membrane (15, 16). The luminal domain of LAPI, however, binds to torsinA, an AAA+ ATPase that resides in the perinuclear space and the continuous main ER (17). Consistent with these data, torsinA is enzymatically inactive unless it interacts with LAPI lining the inner aspect of the inner nuclear membrane, cause Dunnigan-type familial partial lipodystrophy (7–9). Almost all patients with Dunnigan-type familial partial lipodystrophy have hepatic steatosis; however, in this condition it occurs secondarily to adipose dysfunction and insulin resistance (10). Additionally, there are data suggesting that hepatocyte-intrinsic alterations in the nuclear envelope may also directly affect liver lipid metabolism. Conditional deletion of Lmna from hepatocytes causes steatosis and increased susceptibility to steatohepatitis, but for unknown reasons, the effect is only observed in male mice (11). These data implicate the nuclear envelope as a site of regulation of lipid metabolism, but the responsible molecular components and mechanisms within this organelle are poorly understood.

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within the perinuclear space or with the luminal domain of the transmembrane protein luminal domain–like LAP1 (ULL1) within the main ER (17, 18). Recessive mutations in the gene encoding LAP1 that disrupt the B isoform have been linked to familial cardiomyopathy and muscular dystrophy (19, 20). Total loss of LAP1B and LAP1C causes multisystem disease and death during childhood (21). A single amino acid deletion in torsinA causes the neurodevelopmental disease DYT1 dystonia, which is an autosomal dominant disease (22). Despite these multiple disease associations, the role of the torsinA-LAP1 complex in different tissues remains unestablished.

To unravel fundamental features of the torsinA-LAP complex, we have explored the molecular and cellular consequences of disrupting these proteins in diverse tissue environments (16, 23). Here, we report that conditional deletion of either LAP1 or torsinA from hepatocytes caused profound steatosis and establish a critical role for the torsinA-LAP1 complex in hepatic VLDL assembly and secretion. These data advance our understanding of nuclear envelope–localized processes in lipid homeostasis and are the first to our knowledge to implicate the torsinA-LAP complex in mammalian lipid metabolism.

Results

Conditional hepatocyte deletion of LAP1 causes hepatic steatosis. We reported previously that the body mass of L-CKO mice did not differ from that of controls up to approximately 2 years of age (16). Further body composition analysis of 4- to 6-month-old male and female L-CKO mice fed a chow diet did not show any differences in the percentage of body fat compared with controls (Supplemental Figure 2A). The percentage of lean body mass was also the same in L-CKO and control mice of both sexes (Supplemental Figure 2B). Collectively, our results indicate that depletion of LAP1 from hepatocytes of male and female mice causes hepatic steatosis and nuclear lipid droplets. Steatosis occurred in mice on a chow diet in the absence of obesity, glucose intolerance, whole-body insulin resistance, or changes in body composition. Because none of the identified phenotypes showed sexual dimorphism, in all subsequent experiments we analyzed L-CKO data from both sexes in aggregate.

Defective hepatic TG and apoB100 secretion in L-CKO mice. To gain a mechanistic understanding of hepatic steatosis in L-CKO mice, we investigated de novo hepatic lipogenesis, fatty acid oxidation, and lipoprotein secretion. Analysis of the expression of selected genes encoding proteins involved in liver lipid metabolism failed to identify a specific pathway to explain steatosis in the L-CKO mice. Examination of the expression of key genes in de novo lipogenesis provided a mixed picture: Fasn, encoding fatty acid synthase, and Dgat2, encoding diacylglycerol O-acyltransferase 2, were significantly reduced, whereas expression of Srebf1, Srebf2, Acaca, Scd1, and Dgat1 did not differ significantly between L-CKO and littermate controls (Supplemental Figure 3). We therefore measured de novo hepatic lipogenesis in whole livers and fatty acid oxidation in isolated primary hepatocytes from L-CKO and control mice. We observed no significant abnormalities in either hepatic de novo lipogenesis (Figure 2A) or fatty acid oxidation in L-CKO mice (Figure 2B). It is unlikely that the supply of fatty acids from adipose tissue would have been changed in the L-CKO mice, as they have hepatocyte-selective gene deletion. Nonetheless, we examined plasma fatty acid concentrations to exclude increased peripheral delivery to the liver and found that the concentrations were the same in L-CKO and control mice (Figure 2C). These results indicate that increased synthesis, decreased oxidation, or increased delivery of circulating fatty acids was not contributing to hepatic steatosis in L-CKO mice.

Another potential mechanism for steatosis is a defect in hepatic lipoprotein secretion. Consistent with this possibility, we found that plasma TG concentrations were reduced in 4-month-old chow-fed L-CKO mice at all time points measured following i.v. administration of tyloxapol to block uptake of circulating TG-rich lipoproteins (Figure 3A). The hepatic TG secretion rate calculated from those data was significantly reduced by 20.1% ± 7.4% in L-CKO mice (Figure 3B). apob is essential for the assembly and secretion of VLDL. Although human livers synthesize and secrete only apob, rodents also synthesize and secrete an alternatively translated, truncated form, apob48, which is also made in human and rodent small intestines. The role of hepatic apob48 is unclear, but it can be the dominant form in rodent livers (24). Analysis of plasma proteins by SDS-PAGE followed by gel autoradiography 120 minutes after injection of 35S-methionine together with tyloxapol showed decreased levels of radiolabeled apob100 in L-CKO mice, but similar amounts of
Figure 1. Hepatic steatosis without evidence of insulin resistance in male L-CKO mice fed a chow diet. (A) Electron micrographs of liver sections from male control (Tor1aip1fl/fl) and L-CKO (AlbCre Tor1aip1fl/fl) mice. White arrowheads indicate intranuclear lipid droplets. Scale bars: 2 μm. (B) Confocal micrographs of isolated hepatocytes. Lipids were stained with BODIPY (green) and nuclei with DAPI (blue). The right panel is a zoomed image of the dashed-line square region. White arrowheads indicate intranuclear lipid droplets. Scale bars: 10 μm (zoom, 10 μm). (C) Representative light micrographs of H&E- and Oil Red O–stained liver sections from chow-fed mice. Scale bar: 50 μm. (D) Liver TG and cholesterol content; mice were fasted for 4 to 5 hours before livers were collected (n = 5 mice per group). *P < 0.05 and ***P < 0.001, by Student’s t test. (E) Plasma TG and cholesterol concentrations. Mice were fasted for 5 hours before collection of plasma (n = 5 mice per group). *P < 0.05, by Student’s t test. (F) Blood glucose concentration versus time after injection of a glucose bolus in overnight-fasted mice. Values represent the mean ± SEM (n = 6 mice per group). Results were not significantly different at any time point by ANOVA. (G) Serum insulin concentrations. Mice were fasted for 5 hours before collection of plasma (n = 4 mice per group). P = NS, by Student’s t test. In D, E, and G, the values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. The control and L-CKO mice used were 4–6 months of age.
the alternatively translated product apoB48 (Figure 3C). Secretion of newly synthesized apoB100 was reduced by 33.1% ± 5.4% in L-CKO mice compared with controls, and apoB48 secretion was normal in L-CKO mice (Figure 3D). To exclude extrahepatic influences on apoB100 secretion in L-CKO mice, we incubated primary cultures of hepatocytes isolated from control and L-CKO mice. Oxidative products of *14*C-oleic acid (14CO2 and 14C-labeled acid-soluble metabolites) were measured and normalized to milligrams of cellular protein (n = 3 mice per genotype). P = NS, by Student’s t test. (C) Free fatty acids (FFAs) were measured from snap-frozen plasma samples from control and L-CKO mice after a 4- to 5-hour fast. The values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months of age.

Figure 2. De novo lipogenesis, fatty acid oxidation, and plasma free fatty acids in livers of L-CKO mice. (A) De novo lipogenesis in the livers of control and L-CKO mice. Newly synthesized hepatic lipids were measured by incorporation of *14*H into fatty acids (FFAs) 1 hour after i.p. injection with *14*H2O (n = 6 mice per genotype). P = NS, by Student’s t test. (B) Oxidation of fatty acids in hepatocytes isolated from control and L-CKO mice. Oxidative products of *14*C-oleic acid (*14*C02 and *14*C-labeled acid-soluble metabolites) were measured and normalized to milligrams of cellular protein (n = 3 mice per genotype). P = NS, by Student’s t test. (C) Free fatty acids (FFAs) were measured from snap-frozen plasma samples from control and L-CKO mice after a 4- to 5-hour fast. The values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months of age.

Figure 4. De novo lipogenesis, fatty acid oxidation, and plasma free fatty acids in livers of L-CKO mice. (A) De novo lipogenesis in the livers of control and L-CKO mice. Newly synthesized hepatic lipids were measured by incorporation of *14*H into fatty acids (FFAs) 1 hour after i.p. injection with *14*H2O (n = 6 mice per genotype). P = NS, by Student’s t test. (B) Oxidation of fatty acids in hepatocytes isolated from control and L-CKO mice. Oxidative products of *14*C-oleic acid (*14*C02 and *14*C-labeled acid-soluble metabolites) were measured and normalized to milligrams of cellular protein (n = 3 mice per genotype). P = NS, by Student’s t test. (C) Free fatty acids (FFAs) were measured from snap-frozen plasma samples from control and L-CKO mice after a 4- to 5-hour fast. The values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months of age.

The alternatively translated product apoB48 (Figure 3C). Secretion of newly synthesized apoB100 was reduced by 33.1% ± 5.4% in L-CKO mice compared with controls, and apoB48 secretion was normal in L-CKO mice (Figure 3D). To exclude extrahepatic influences on apoB100 secretion in L-CKO mice, we incubated primary cultures of hepatocytes isolated from control and L-CKO mice with *35*S-methionine and performed autoradiography of SDS-polyacrylamide gels in which proteins in cell lysates and media were separated (Figure 3E). As with the in vivo findings, we observed a significant and selective 23.4% ± 6.4% decrease in apoB100 in media harvested from these cells. Analysis of the whole-cell lysates showed a small but significant decrease in apoB48 as well as a larger, significant decrease in apoB100 (Figure 3F). These data demonstrate that L-CKO mice have a selective defect in the secretion of apoB100 VLDL.

To assess whether the defect in TG and apoB100 secretion was specific, or reflected a general abnormality of hepatic secretory function, we measured the secretion of newly synthesized serum albumin and apoAI by autoradiography in the same in vivo studies in which we measured apoB secretion (Supplemental Figure 4A). We observed no significant differences for secretion of either protein in L-CKO mice (Supplemental Figure 4B). The steady-state serum albumin concentration was also normal in L-CKO mice (Supplemental Figure 4C). These data exclude a general secretory defect in L-CKO hepatocytes and point to a specific defect of apoB100 secretion caused by the absence of LAP1.

The *Alb-Cre* transgene used to delete LAP1 in L-CKO mice closely mimics the expression of endogenous *Alb*, which begins on E10.5 (25). To exclude the possibility that the phenotypes of these mice result from developmental effects, we deleted LAP1 from hepatocytes of adult mice by injecting a Cre-expressing adenovirus (AAV) construct (AAV-Cre) into the tail veins of adult mice homozygous for floxed alleles of the gene encoding LAP1. LAP1 levels were reduced by approximately 50% three weeks after virus injection and were largely undetectable 8 weeks after injection (Figure 4A). As seen with hepatocytes from L-CKO mice, primary hepatocytes isolated from mice injected with AAV-Cre exhibited increased numbers of larger cytoplasmic lipid droplets, compared with control hepatocytes, as well as intranuclear lipid droplets (Figure 4B). Synthesis and secretion of apoB in primary cultures of hepatocytes from AAV-Cre–injected mice after incubation with *35*S-methionine also mimicked the findings in L-CKO mice (Figure 4C). We observed a significant decrease in apoB100 in cell lysates and media (72.8% ± 9.2% and 56.2% ± 12.9%, respectively) of AAV-Cre–injected primary hepatocytes, whereas apoB48 was slightly but significantly decreased (23.4% ± 6.4%) in cell lysates but unchanged in the media (Figure 4D). These data confirm an essential role for LAP1 in hepatocyte lipid metabolism and exclude the possibility that the phenotype of L-CKO mice results from an early developmental defect.

Conditional hepatocyte deletion of *torsinA* causes hepatic steatosis. The luminal domain of LAP1 binds to the AAA+ ATPase torsinA within the perinuclear space of the nuclear envelope (17). LAP1 binding is required for torsinA ATPase activity (18). We first excluded the possibility that loss of LAP1 decreases torsinA levels or dramatically alters its localization in hepatocytes. When we examined the livers of L-CKO mice for torsinA expression, there was actually a slight but significant increase in torsinA expression compared with that in the livers of control mice (Supplemental Figure 5, A and B). TorsinA continued to have an ER-like distribution in hepatocytes lacking LAP1, as in control hepatocytes (Supplemental Figure 5C). We therefore hypothesized that loss of LAP1 induces hepatic steatosis by decreasing torsinA activity at the nuclear envelope. We tested this hypothesis through conditional deletion of *torsinA* from hepatocytes by intercrossing *Tor1afl/fl* (23) and *Alb-Cre*–transgenic mice (referred to herein as A-CKO mice), predicting that these mice would also have hepatic steatosis and decreased VLDL section.

A-CKO mice were born at the expected Mendelian ratios and were indistinguishable from their littermate controls at birth. We confirmed the efficiency of torsinA depletion by immunoblotting of proteins in lysates of liver and primary hepatocytes from A-CKO mice (Supplemental Figure 6, A and B). Adult male A-CKO mice fed a chow diet exhibited grossly enlarged, whitish livers (Figure 5A). H&E-stained sections of livers from A-CKO mice had grossly enlarged hepatocytes with mixed macroversicular and microversicular steatosis, which was confirmed with Oil Red O staining (Figure 5B). Livers from A-CKO mice contained dramatically increased amounts of TGs and cholesterol compared with littermate controls; levels of both lipids were much higher than those in the livers of L-CKO mice (Figure 5C). Concomitant with the much greater steatosis, A-CKO mice had markedly reduced plasma TG and cholesterol concentrations (Figure 5D). Despite the more dramatic hepatic steatosis and reductions in plasma lipid levels
Defective hepatic TG and apoB100 secretion in A-CKO mice. To further assess the hypothesis that the phenotype of L-CKO mice reflects deficient torsinA function, we examined whether A-CKO mice exhibit similar defects in apoB metabolism. These analyses demonstrated that, concomitant with the greater steatosis and more marked reductions in plasma lipid levels in the A-CKO mice, they had greater reductions in plasma TG levels at all time points measured following i.v. administration of tyloxapol to block the uptake of circulating TG-rich lipoproteins (Figure 6A). The calculated hepatic TG secretion rate for A-CKO mice was markedly and significantly reduced (by 63.8% ± 12.3%; Figure 6B). We analyzed plasma protein levels after injection of 35S-methionine into A-CKO mice (Figure 6C). Compared with control mice, A-CKO mice exhibited significantly decreased apoB100 secretion compared with controls (reduced by 61.3% ± 11.3%). As with L-CKO mice, secretion of apoB48 in A-CKO mice did not differ from that detected in controls (Figure 6D). The reduction in apoB100 secretion in A-CKO mice was clearly greater than that observed in L-CKO mice. We also analyzed the synthesis and secretion of apoB in primary hepatocyte cultures from A-CKO mice after incubation with 35S-methionine (Figure 6E). This analysis showed sig-
nificantly decreased levels of newly synthesized apoB100 (86.7% ± 11.8%) and apoB48 (30.6% ± 6.5%) in media (Figure 6F). In cell lysates, we detected a significant decrease in apoB100 (by 62.0% ± 9.3%), but no significant changes in newly synthesized apoB48 (Figure 6F). These results indicate that the secretion of both apoB100 and apoB48 is decreased in hepatocytes of A-CKO mice.

Like L-CKO mice, A-CKO mice exhibited no abnormalities in the rate of hepatic secretion of albumin or apoA1 (Supplemental Figure 9, A and B). Serum albumin concentrations were also normal in A-CKO mice (Supplemental Figure 9C). These data indicate that, as in L-CKO mice, there was no general synthetic or secretory defect in the hepatocytes of A-CKO mice.

Because of the profound steatosis and decreased VLDL secretion in A-CKO mice, we examined livers from mice heterozygous for \( T_{or1a} \) deletion, with 1 allelic expression of torsinA in hepatocytes (A-CKO het). Control and A-CKO het mice did not show significant hepatocyte lipid accumulation, in contrast to A-CKO mice (Supplemental Figure 10A). Liver TG and cholesterol content and plasma concentrations were also similar to those of WT controls (Supplemental Figure 10, B and C). Secretion of hepatic TG and apoB was also identical to that seen in control mice (Supplemental Figure 10, D and E). Hence, heterozygous loss of torsinA from hepatocytes did not lead to steatosis or significantly decreased liver VLDL secretion.

As with L-CKO mice, we examined the expression of selected genes encoding proteins involved in liver lipid metabolism as well as de novo lipogenesis and fatty acid oxidation in the livers of A-CKO mice. Whereas L-CKO mice had reduced hepatic expression of \( Fasn \) and \( Dgat2 \), both of which are involved in de novo lipogenesis, A-CKO mice showed only a small but statistically significant increase in the expression of another gene involved in the same metabolic process, \( Dgat1 \), which encodes diacylglycerol \( O \)-acyltransferase 1 (Supplemental Figure 11). However, we observed no difference in de novo hepatic fatty acid synthesis in vivo in A-CKO mice compared with their littermate controls (Figure 7A). In contrast to L-CKO mice, however, the fatty acid oxidation rate in primary hepatocytes isolated from A-CKO mice was significantly reduced by 31.8% ± 10.4% compared with the rate detected in control mice (Figure 7B). As in L-CKO mice, plasma fatty acid concentrations were also not altered in A-CKO mice (Figure 7C).

To exclude the possibility that the phenotypes of A-CKO mice result from developmental effects, we deleted torsinA in adult mice by injecting AAV-Cre into the tail veins of 4-month-old mice homozygous for \( T_{or1a} \)-floxed alleles. We found that torsinA
levels in liver were reduced by approximately 65% compared with levels in control mice 4 weeks after virus injection (Figure 8A). Primary hepatocytes isolated from mice injected with AAV-Cre showed increased lipid droplets in ER-appearing structures (Figure 8B). We also measured newly synthesized and secreted \(^{35}\)S-labeled apoB in primary hepatocytes from AAV-Cre–injected mice (Figure 8C). This analysis demonstrated a significant reduction in apoB100 (59.5% ± 14.7%) but not apoB48 in the cell lysates; media from these cultures had a significant decrease in the amount of both apoB100 and apoB48 (91.1% ± 4.7% and 63.0% ± 6.0%, respectively; Figure 8D). These results demonstrate that depletion of torsinA from adult hepatocytes causes hepatocyte lipid accumulation and defects in VLDL synthesis and secretion.

**ER stress in livers of L-CKO and A-CKO mice.** An association exists between ER stress and defective apoB-mediated TG secretion (26). We therefore examined the ER stress response in livers from L-CKO and A-CKO mice. We performed immunoblotting of protein extracts from livers of control and L-CKO mice at 6 months of age to detect glucose-regulated protein 78 (GRP78), C/EBP homolog protein (CHOP), phosphorylated eukaryotic initiation factor 2α (p-eIF2α), and total eIF2α (Supplemental Figure 12A). This analysis revealed significant increases in the expression of GRP78, but not CHOP or the ratio of p-eIF2α to total eIF2α (Supplemental Figure 12B). We also measured these proteins by immunoblotting liver lysates from A-CKO mice at 6 months of age (Supplemental Figure 12C). GRP78 and p-eIF2α exhibited increases; CHOP and total eIF2α were unchanged (Supplemental Figure 12D). IRE1α activates XBP1 by catalyzing the splicing of its mRNA in response to ER stress (27). We performed an XBP1 splicing assay using cDNAs synthesized from total RNAs isolated from livers of L-CKO mice (Supplemental Figure 12E). We observed no significant differences between L-CKO and control mice (Supplemental Figure 12F). Similar to L-CKO mice, A-CKO mice exhibited no abnormalities in XBP1 splicing of RNA isolated from livers of A-CKO mice (Supplemental Figure 12, G and H). Considered together, these experiments indicate that disruption of the torsinA-LAP1 complex in hepatocytes is accompanied by a modest degree of ER stress.

**Hepatic insulin resistance in the livers of L-CKO and A-CKO mice.** Although glucose tolerance and plasma insulin concentrations were normal in L-CKO and A-CKO mice, indicating the absence of whole-body insulin resistance, we further examined hepatic insulin responsiveness by assessing the activation (phosphorylation) of insulin receptors and AKT2 in livers under physiological conditions. Insulin receptor activation was decreased in the livers of L-CKO mice, but AKT2 activation was not significantly different from that in control mice (Supplemental Figure 13, A and B). In the livers of A-CKO mice, insulin receptor activation was not significantly different than that detected in controls, but AKT2 phosphorylation was significantly decreased (Supplemental Figure 13, C and D). These results suggest that L-CKO and A-CKO mice may have modest alterations in hepatic insulin responsiveness.

**Phosphatidylcholine/phosphatidylethanolamine ratios in livers of L-CKO and A-CKO mice.** The hepatic phosphatidylcholine/phosphatidylethanolamine (PC/PE) ratio influences the dynamics and size limit of lipid droplets, and alterations in it have been linked to impaired VLDL secretion and fatty liver disease (28). Because of the impaired hepatic VLDL secretion in L-CKO and A-CKO mice and the mixed microvesicular and macrovesicular steatosis in A-CKO mice, we examined the livers of both types of mice for altered PC/PE ratios. Although the PC/PE ratios were not different in L-CKO mice compared with ratios in littermate controls, the ratios were significantly decreased in A-CKO mice (Supplemental Table 1). Hence, a decreased PC/PE ratio in the livers of A-CKO mice could have contributed to the impaired VLDL secretion and the microvesicular steatosis we observed.

**L-CKO and A-CKO mice on a chow diet develop features of NASH.** Chronic hepatic steatosis can cause hepatomegaly. In some instances, chronic steatosis can progress to NASH, which is characterized by steatosis, ballooning degeneration of hepatocytes, and inflammation. Because of the marked steatosis observed in L-CKO and A-CKO mice, we explored whether they developed hepatomegaly and features of NASH. We assessed hepatomegaly by calculating liver-to-body mass ratios. L-CKO mice had a significantly increased liver-to-body mass ratio at 18 months of age (Figure 9A) but no significant difference in the heart-to-body mass ratio (Figure 9B). Serum alanine aminotransferase (ALT) activity, an indicator of hepatocyte damage or death, was nearly 4-fold higher in L-CKO mice than in controls (Figure 9C). Histopathological examination of livers from L-CKO mice revealed steatosis, hepatocyte ballooning, and lobular inflammation, all of which are characteristic histological features of NASH (Supplemental Figure 14A). We observed that hepatic fibrosis was also increased in L-CKO mice compared with controls (Figure 9D). For an objective assessment of NASH features, a liver pathologist blinded to the genotype of the mice assigned NAFLD activity and fibrosis stage scores used in clinical practice (29). Of the 7 L-CKO mice examined, 2 had NAFLD activity scores of 5 or higher (consistent with NASH), 4 had a score of 3–4 (borderline NASH), and 1 had a score of 2 (not NASH); all 4 control mice examined had NAFLD activity scores of 2 or lower (not NASH). The mean NAFLD activity score of L-CKO mice was significantly increased compared with that for littermate controls (Figure 9E). L-CKO mice also developed fibrosis by 18 months of age. All 4 control mice had a blindly assigned fibrosis stage of 1A (mild, zone 3, perisinusoidal), whereas 1 L-CKO mouse had stage 3 (bridging), 3 mice had stage 1B (moderate, zone 3, perisinusoidal), and the remaining 5 mice had stage 1A (Supplemental Table 2). Analyses of A-CKO mice demonstrated more severe, accelerated disease. The liver-to-body mass ratio was already significantly increased in 6-month-old A-CKO mice (Figure 9F), with no significant difference in heart-to-body mass ratio (Figure 9G). In A-CKO mice, serum ALT activity was nearly 12-fold higher than that in controls (Figure 9H). Histopathological examination demonstrated hepatic steatosis with a predominance of microvesicular fat, hepatocyte ballooning, and lobular inflammation (Supplemental Figure 14B). By 6 months of age, hepatic fibrosis was already apparent in the livers of A-CKO mice (Figure 9I). Moreover, 6 of 7 A-CKO livers received a NAFLD activity score of 6 or 7 (consistent with NASH) from a pathologist blinded to the genotype, whereas all control livers received NAFLD activity scores of 2 or lower (not NASH) (Figure 9J). Three A-CKO mice were judged to have stage 2 fibrosis, 2 had stage 1B, and 1 exhibited stage 1A, as did all controls (Supplemental Table 2). These data demonstrate...
Figure 5. Hepatic steatosis without evidence of insulin resistance in male A-CKO mice fed a chow diet. (A) Photographs of livers from male control (Tort1afl/+ and A-CKO (AlbCre Tort1afl–)) mice fed a chow diet. Scale bar: 1 cm. (B) Representative light micrographs of H&E- and Oil Red O–stained liver sections from chow-fed mice. Scale bar: 50 μm. (C) Liver TG and cholesterol content. Mice were fasted for 4 to 5 hours before livers were collected (n = 4 mice per group). **P < 0.01 and ***P < 0.001, by Student’s t test. (D) Plasma TG and cholesterol concentrations (n = 4 mice per group). **P < 0.01, by Student’s t test. (E) Blood glucose concentrations versus time after injection of a glucose bolus into overnight-fasted mice (n = 5 mice per group). Results were not significantly different at any time point by ANOVA. (F) Serum insulin concentrations. Mice were fasted for 5 hours before plasma was collected (n = 3–5 mice per group). P = NS, by Student’s t test. (G) Electron micrographs of liver sections from 4-month-old control and A-CKO mice. Top panel shows low-magnification images of a single hepatocyte from control and A-CKO samples. The lower panel is a magnified image of the dashed-line square region in the upper panel. Scale bars: 10 μm (upper panel) and 500 nm (lower panel). N, nucleus. (H) Confocal micrographs of isolated hepatocytes. Lipids were stained with BODIPY (green) and nuclei with DAPI (blue). The right panel is a zoomed image of the micrographs of isolated hepatocytes. Lipids were stained with BODIPY (green) and nuclei with DAPI (blue). The right panel is a zoomed image of the dashed-line square region in the upper panel. Scale bars: 10 μm (zoom, 10 μm). In C, D, and F, the values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months old.

that both L-CKO and A-CKO mice develop features characteristic of human NASH. Considered together with our other results, these data strongly suggest that LAP1 and torsinA function in a common pathway, the disruption of which disrupts VLDL metabolism, which ultimately can progress to the core histopathological features of human NASH.

Discussion

NAFLD is highly prevalent and frequently progresses to NASH, representing a major challenge to public health. NAFLD occurs in 25% to 40% of the US and European populations, with rates as high as 70% in individuals with diabetes mellitus (3). Our studies establish a direct role for the torsinA/LAP1 pathway in the regulation of VLDL secretion and liver fat accumulation that progresses to the morphological equivalent of human NASH, including fibrosis. Remarkably, the lipid metabolic derangements resulting from disruption of the torsinA/LAP1 pathway occurred in mice on normal chow diets and in the absence of abnormalities of glucose homeostasis or obesity. These findings highlight nuclear envelope–localized events as key for hepatocyte-autonomous lipid metabolism defects, which are involved in NAFLD development. Consistent with a role in influencing mammalian liver and plasma lipid concentrations, the torsinA/LAP1 pathway was recently identified in an integrative systems genetic analysis of lipid metabolism in inbred mouse strains (30).

The prevalence of NAFLD/NASH is significantly higher in areas where insulin resistance, central obesity, and diabetes are common (31). These comorbidities are often merged under the umbrella of metabolic syndrome (32). There is broad consensus that these features of metabolic syndrome are strongly associated with NAFLD in the majority of affected individuals (33). In contrast, recent work has identified several genetic insults that predispose individuals to NAFLD, NASH, and even cirrhosis in a liver-autonomous manner, unaccompanied by the features of metabolic syndrome (34, 35). Our findings identify a nuclear envelope–localized event as a participant in such processes. The identified genes include those encoding proteins involved in lipid droplet biology, such as PNPLA3, encoding patatin-like phospholipase domain–containing protein 3 (36); in de novo lipogenesis, such as GCKR, encoding glucokinase regulatory protein (37); and in the assembly and secretion of VLDL, such as APOB, TM6SF2, encoding transmembrane 6 superfamily member 2, and MTPP, encoding microsomal triglyceride transfer protein (MTP) (38–40). All of these genes have been identified as liver-autonomous causes of NAFLD. Although the clinical course of NAFLD in these instances may be affected by the presence of insulin resistance, steatosis may occur even when there are no concomitant systemic abnormalities in carbohydrate or lipid metabolism (36, 41).

Another mouse model of significant hepatic steatosis without concomitant insulin resistance was created by antisense oligonucleotide–mediated knockdown of comparative gene identification 58 from hepatocytes. The absence of insulin resistance was attributed to increased compartmentalization of diacylglycerol and PKCs in lipid droplets versus the plasma membrane (42). Using the phosphorylation of insulin receptors and AKT2 as indicators of hepatic insulin signaling, we found evidence for modest insulin resistance in both L-CKO and A-CKO mice. A more detailed investigation of hepatic insulin signaling, including assessment of the subcellular distribution of diacylglycerol and PKCs, is warranted.

In addition to its activation by LAP1 in the perinuclear space, torsinA is also activated in the main ER by LULL1 (17, 18). The qualitative similarities but clear quantitative differences between mice lacking LAP1 or torsinA in hepatocytes fit with the hypothesis that LAP1 depletion causes only partial torsinA ATPase loss of function because of preserved activation by LULL1 in the main ER. This hypothesis predicts the more severe phenotype we observed with torsinA deletion, which ablated its function from both ER compartments. Our proposed model posits a role for nuclear membrane–localized torsinA ATPase in hepatic lipid homeostasis, particularly for VLDL assembly and secretion. Future studies targeting LULL1 and expressing a constitutively active form of torsinA in hepatocytes of mice with hepatic deletion of LAP1 or LULL1 will help to further test this hypothesis. A-CKO mouse livers also had a decreased PC/PE ratio, whereas the livers of L-CKO mice did not; this could also contribute to the more severely impaired VLDL secretion and greater degree of steatosis, as well as the altered size distribution and abnormal localization of lipid droplets in the A-CKO mice and requires further investigation.

Mice with hepatocyte conditional deletion of LAP1 or torsinA share similarities to mice with loss-of-function of proteins with established functions in VLDL secretion. Homozygous disruption of apoB is embryonically lethal, but heterozygous null mice exhibit decreased plasma cholesterol concentrations (43, 44). Mice with hepatocyte–specific depletion of MTP, which transfers TGs into the lumen of the ER for lipidation of apoB, have severely reduced plasma TG, VLDL, and apoB100 concentrations and hepatic steatosis when fed a chow diet, but lack the inflammation characteristic of NASH (45). Livers of these mice have near-complete loss of microsomal apoB100 and no evidence of ER stress (46). Likewise, mice treated with antisense oligonucleotides to inhibit MTP synthesis have reduced hepatic VLDL secretion and steatosis without evidence of ER stress (47, 48). In contrast, treatment of adult mice with
Although conditional hepatocyte deletion of LAP1 and conditional hepatocyte deletion of torsinA share essential phenotypes, the 2 situations are not identical. LAP1 hepatocyte deletion causes moderate steatosis, although this is particularly notable, as it occurs on a normal chow diet. These mice exhibit normal rates of hepatic de novo lipogenesis and fatty acid oxidation, as well as normal uptake of fatty acids from the circulation, as indicated by normal plasma fatty acids concentrations. The degree of steatosis therefore appears to be a direct result of the reduction in TG secretion observed in these mice. Unique to this model, however, is the accumulation of some nuclear lipid droplets. Nuclear lipid droplets in hepatocyte cell lines are associated with extension of the inner nuclear membrane into the nucleus (also referred to as the nucleoplasmic reticulum) (52). A potential explanation for nuclear lipid is that LAP1 deletion promotes invaginations of the nuclear envelope, an effect known to occur from alterations in lamins (which bind to LAP1) and other integral inner nuclear membrane proteins (53, 54).

There were no nuclear lipid droplets present in torsinA-null hepatocytes, but these mutants exhibited marked abnormalities in the number, size, and location of the cytoplasmic ones, with the vast majority being small and spherical. This phenotype is reminiscent of that of mice with deletion of fat storage-inducing transmembrane protein 2, which functions in the budding of lipid drop-
Valt P = NS, by Student’s t test. Free fatty acids from snap-frozen plasma samples from control and A-CKO mice were measured after a 4- to 5-hour fast.

** Results

Free fatty acids from snap-frozen plasma samples from control and A-CKO mice were measured after a 4- to 5-hour fast. P = NS, by Student’s t test. Values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months old.

Figure 7. De novo lipogenesis and fatty acid oxidation in livers of A-CKO mice. (A) De novo lipogenesis in livers of control and A-CKO mice. Newly synthesized hepatic lipids were measured by incorporation of H into fatty acids 1 hour after i.p. injection with H,O (n = 3 mice per genotype). P = NS, by Student’s t test. (B) Oxidation of fatty acids in hepatocytes isolated from control and A-CKO mice. Oxidative products of C-oleic acid (C, C-labeled acid-soluble metabolites) were measured and normalized to milligrams of cellular protein (n = 3 mice per genotype). **P < 0.01, by Student’s t test. (C) Free fatty acids from snap-frozen plasma samples from control and A-CKO mice were measured after a 4- to 5-hour fast. P = NS, by Student’s t test. Values for individual mice are shown, with longer horizontal bars indicating the mean and vertical bars indicating the SEM. Mice used were 4–6 months old.

Methods

Further information can be found in the Supplemental Methods.

Mice. The generation, maintenance, and genotyping of floxed alleles of Lap1 (Lap1<sup>fl/fl</sup>) and Torla (Torla<sup>fl/fl</sup>) mice have been previously described (16, 23). Alb-Cre<sup>+</sup>/– transgenic mice were purchased from The Jackson Laboratory (stock no. 003574). To generate L-CKO mice, Alb-Cre<sup>+</sup> mice were crossed with Lap1<sup>fl/fl</sup> mice to obtain L-CKO (Alb-Cre<sup>+</sup>/Lap1<sup>fl/fl</sup>) mice. The genetic background of L-CKO mice was C57BL/6. The breeding strategy for A-CKO (Alb-Cre<sup>+</sup>/Torla<sup>fl/fl</sup>) and control (Torla<sup>fl/fl</sup>) mice has been described previously (23). The genetic background of A-CKO mice was a combination of C57BL/6 and 129SvEv. All mice were fertile and produced at the expected Mendelian frequencies. Mice were housed in a climate-controlled room with a 12-hour light/12-hour dark cycle and fed a regular chow diet (Purina Mills, 5053).

Electron microscopy. Electron microscopy was performed as previously described (16). Briefly, mice were drop-fixed with 1% paraformaldehyde and 2.5% glutaraldehyde in 0.1 M cacodylate buffer (pH 7.4), postfixed with 1% osmium tetroxide in 0.1 M cacodylate buffer (pH 7.4), incubated with uranyl acetate, and dehydrated with ethanol. Tissues were subsequently rinsed with propylene oxide and embedded. Sections (60-nm thick) were counterstained with uranyl acetate and lead citrate and examined on a JEM-1200EX electron microscope (JEOL).

Confocal microscopy. Cultured primary hepatocytes on collagen-coated coverslips were stained with BODIPY (Thermo Fisher Scientific, D-3922) for 30 minutes. Subsequently, cells were fixed with 4% paraformaldehyde for 15 minutes at room temperature and then washed and stained with DAPI. Images were obtained using a multiphoton Ti Eclipse confocal microscope (Nikon).

Histology. The mice were euthanized, and the livers were immediately excised and blotted dry. The excised livers were fixed in 10% formalin for 48 hours. They were then transferred to the Columbia University Histology Core Laboratory for paraffin block preparation and sectioning into 5-μm slices. Sections were then stained with H&E or Picrosiris red (Polysciences, 24901). For Oil Red O staining, livers were transferred onto a 30% sucrose solution, and frozen sections were prepared at the Columbia University Histology Core. Sections (5-μm thick) were stained with Oil Red O (Polysciences, 25962). Stained sections were photographed using a BX53 upright light microscope attached to a DP72 digital camera (Olympus). To assess NAFLD activity and evaluate fibrosis, a liver pathologist blinded to the mouse genotypes examined sections of the livers stained with H&E, Oil Red O, and Picrosiris red. The NAFLD activity score has been described previously (29).

Quantification of liver TG and cholesterol. Liver lipids were extracted with a modified Folch method as previously described (58). Briefly, a snap-frozen piece of liver (~100 mg) was homogenized in

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PBS, and lipids were first extracted with chloroform/methanol (2:1 v/v) and a second time with chloroform/methanol/water (86:14:1 v/v/v). The organic layer was dried under nitrogen gas and resolubilized in 15% Triton X-100 (Sigma-Aldrich) in double-distilled H₂O. TGs and total cholesterol were measured using colorimetric assays (Wako Diagnostics, 461-08992 and 461-09092, respectively).

**Blood biochemistries.** Plasma TG and cholesterol concentrations were assayed using the colorimetric assays described above. Serum ALT activity and albumin concentration were measured using commercially available kits (Teco Diagnostics A526-120 and Sigma-Aldrich MAK124, respectively). Serum insulin concentrations were measured using the Mouse Insulin ELISA Kit (Crystal Chem, 90080). For glucose tolerance tests, mice were fasted for 12 hours and given 2 g glucose/kg body weight by i.p. injection. Blood glucose concentrations were measured 0, 15, 30, 60, 90, and 120 minutes after injection using a OneTouch Ultra Glucometer (LifeScan).

Plasma free fatty acid concentrations were measured in snap-frozen plasma samples obtained after a 4- to 5-hour fast using a commercially available kit (Wako Diagnostics, HR Series NEFA-HR).

**Hepatic de novo lipogenesis.** Methods to measure de novo lipogenesis have been described previously (59). In brief, mice were injected i.p. with 0.5 mCi of 3H₂O. After 1 hour, mice were anesthetized with ketamine/xylazine, and approximately 250 μL blood was collected. Mice were then euthanized and their livers harvested and flash-frozen. Liver tissue (500 mg) was incubated with 2.5 M KOH at 75°C for 2 hours. Saponified lipids were isolated by the addition of 80% ethanol and hexane (1:2 v/v). The bottom phase, containing fatty acid salts, was acidified with 3 M H₂SO₄. The liberated fatty acids were isolated with additional hexane, dried under nitrogen gas, and resolubilized in 500 μL chloroform. Fatty acids were separated by thin-layer chromatography, and the resulting spots were scraped and counted for 3H activity. 3H incorporated into fatty acids was calculated relative to the specific activity of 3H in total body water calculated from the blood sample collected prior to harvesting of the liver.

**Fatty acid oxidation.** The methods for measuring fatty acid oxidation in mouse primary hepatocytes have been described previously (60). Sixteen hours after plating primary hepatocytes in 6-well plates containing DMEM, the cells were incubated for 2 hours in labeling media (DMEM containing 1.5% BSA, 0.1 mM oleic acid, and 1 μCi/mL 14C-oleic acid). At the end of the labeling and chase periods, media were transferred into 25-mL sealed Erlenmeyer flasks. Lipid oxidation was stopped by adding 200 μL 70% perchloric acid to the bottom of the flask, thus driving the bicarbonate into CO₂. 14CO₂ was captured on a piece of KOH-soaked filter paper. After incubating the filter paper in the flask for 1 hour...
at room temperature, the filter paper was analyzed by liquid scintillation counting. Media remaining in the flasks were also collected and a fraction counted to measure ¹⁴C-labeled acid soluble metabolites. After removal of the chase media, cells were scraped off the plates and protein measured using the BCA Protein Assay Kit (Thermo Fisher Scientific). The data were plotted as cpm/mg cellular protein.

In vivo TG, apoB, albumin, and apoA1 secretion. In vivo TG and apoB secretion rates were determined as previously described (48). Briefly, mice were fasted for 4 to 5 hours prior to an i.v. injection of a mixture of 200 μCi ³⁵S-methionine and 500 mg/kg tyloxapol (Sigma-Aldrich, T8761-50G) in 0.9% NaCl. Tyloxapol inhibits both the lipolysis and tissue uptake of lipoproteins in mice, and the accumulation of TGs and ³⁵S-apoB in plasma can be used to estimate the rates of secretion. Blood samples were collected before injection and 30, 60, 90, and 120 minutes after injection of tyloxapol and ³⁵S-methionine. TG concentrations were measured using the colorimetric assay described above. For apoB secretion rates, whole plasma samples from the 120-minute time point were subjected to 4% SDS-PAGE. The volumes of the samples were adjusted as determined by trichloroacetic acid precipitable ³⁵S-labeled plasma proteins. The gel was dried and exposed to x-ray film to quantitate labeled apoB proteins by densitometry. Hepatic secretion of apoA1 and albumin was measured by subjecting the radiolabeled plasma proteins to 8% SDS-PAGE and performing similar analyses of the dried gels.

Primary hepatocyte isolation. Primary hepatocytes were isolated according to previously described methods (48). Briefly, mice were fasted for 4 to 5 hours prior to an i.v. injection of a mixture of 200 μCi ³⁵S-methionine and 500 mg/kg tyloxapol (Sigma-Aldrich, T8761-50G) in 0.9% NaCl. Tyloxapol inhibits both the lipolysis and tissue uptake of lipoproteins in mice, and the accumulation of TGs and ³⁵S-apoB in plasma can be used to estimate the rates of secretion. Blood samples were collected before injection and 30, 60, 90, and 120 minutes after injection of tyloxapol and ³⁵S-methionine. TG concentrations were measured using the colorimetric assay described above. For apoB secretion rates, whole plasma samples from the 120-minute time point were subjected to 4% SDS-PAGE. The volumes of the samples were adjusted as determined by trichloroacetic acid precipitable ³⁵S-labeled plasma proteins. The gel was dried and exposed to x-ray film to quantitate labeled apoB proteins by densitometry. Hepatic secretion of apoA1 and albumin was measured by subjecting the radiolabeled plasma proteins to 8% SDS-PAGE and performing similar analyses of the dried gels.

Primary hepatocyte isolation. Primary hepatocytes were isolated according to previously described methods (48). Briefly, mice were
perfused with HBSS without calcium (Thermo Fisher Scientific, 14175-079) and 8 mM HEPES (Thermo Fisher Scientific, 15630-080) via the abdominal vena cava after cutting the portal vein to allow outflow of the perfusate. The hepatocytes were perfused for 8 minutes at a rate of 5 mL/min at 37°C. This was followed by perfusion at the same rate as that for DMEM with 80 mg/100 mL collagenase type I (Worthington Biochemical Corporation CLS-1) for 6 minutes. The liver was removed and minced in a Petri dish containing 1 mL of the same warm DMEM collagenase mixture for an additional 2 to 4 minutes. Ice-cold DMEM was added and the digested tissue filtered through a nylon mesh and collected in a 50-mL conical tube. The suspension was centrifuged for 5 minutes at 50 × g, and the supernatant was aspirated and the cell pellet washed 3 times with 30 mL ice-cold DMEM. Hepatocytes isolated from A-CKO mice, because of their high-fat content, were handled without using serological pipettes to minimize mechanical stress on the cells. Viable cells were counted after staining with trypan blue, which revealed that more than 90% of cells were viable. The isolated cells were plated onto collagen-coated, 6-well plates at a density of 500,000 cells/well in 4 mL DMEM plus 10% FBS and cultured for at least 2 hours prior to use for subsequent experiments.

Synthesis and secretion of apoB in cultured hepatocytes. Isolated primary hepatocytes were plated on collagen-coated, 6-well plates in 10% FBS and DMEM and 5% CO₂, for 2 hours to allow attachment. Media were changed, and cells were incubated for an additional 12 hours, washed twice with PBS, and cultured for an additional hour in methionine-free DMEM (Thermo Fisher Scientific, 21013-024). Cells were then labeled for 2 hours in methionine-free DMEM containing 150 μCi/mL [35S]-methionine/cysteine (PerkinElmer,NEG-072014MC). Media were collected and cells washed twice with ice-cold PBS and then lysed in 1 mL lysis buffer (62.5 mM sucrose, 0.5% sodium deoxycholate, 0.5% Triton X-100, 50 mM Tris-HCl, 150 mM NaCl, 1 mM benzamidine, 5 mM EDTA, 2 units/mL aprotinin, 50 μg/mL leupeptin, 50 μg/mL pepstatin A, and 10 mM HEPES, pH 8.0). Immunoprecipitation of apoB100 and apoB48 in cell lysate and media was performed using an anti-apoB antibody (Calbiochem, 178467) followed by separation with 4% SDS-PAGE. The aliquot of lysate or media used for immunoprecipitation was adjusted to equal quantities of radioactivity as estimated by trichloroacetic acid precipitated [35S]-labeled proteins in the lysate. Gels were dried and exposed to x-ray films to quantify apoB100 and apoB48 by densitometry, or bands corresponding to apoB100 and apoB48 were cut from the gel, and radioactivity was quantified by scintillation counting.

AAV-mediated hepatic knockdown of LAP1 and torsinA. AAV8 vectors containing a control expression cassette (AAV-TBG-LacZ) or encoding Cre recombinase (AAV-TBG-Cre) were purchased from the University of Pennsylvania Vector Core. For “acute” knockdown experiments, Lap1-flxed (Lap1fl/fl) and Tor1a-flxed (Tor1afl/fl) mice were randomly assigned to receive either AAV-TBG-LacZ or AAV-TBG-Cre (1 × 1011 genome copies/mouse) via tail vein injection. Protein depletion was confirmed by immunoblotting with specific antibodies.

Statistics. All data are presented as the mean ± SEM. Statistical methods are described in the figure legends for each data set. An unpaired 2-tailed Student’s t test was used to compare differences between any 2 groups, and a 2-way ANOVA was used for multiple comparisons. Statistical significance was set at a P value of less than 0.05.

Study approval. The IACUCs of Columbia University and the University of Michigan approved all procedures, which were conducted in accordance with the NIH’s Guide for the Care and Use of Laboratory Animals (National Academies Press, 2011).

Author contributions JYS, AHO, TF, CÖ, SBG, and CCL conducted experiments. MJL performed histopathological analyses. JYS, AHO, TF, WTD, HNG, and HJW contributed to the study’s design and analyzed and interpreted data. JYS, WTD, HNG, and HJW wrote the manuscript. WTD, HNG, and HJW supervised the research. All authors reviewed and revised the manuscript.

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