Abstract Adipose TG lipase (ATGL) catalyzes the rate-limiting step in TG hydrolysis in most tissues. We have shown that hepatic ATGL preferentially channels hydrolyzed FAs to β-oxidation and induces PPAR-α signaling. Previous studies have suggested that liver FA binding protein (L-FABP) transports FAs from lipid droplets to the nucleus for ligand delivery and to the mitochondria for β-oxidation. To determine if L-FABP is involved in ATGL-mediated FA channeling, we used adenovirus-mediated suppression or overexpression of hepatic ATGL in either WT or L-FABP KO mice. Hepatic ATGL knockdown increased liver weight and TG content of overnight fasted mice regardless of genotype. L-FABP deletion did not impair the effects of ATGL overexpression on the oxidation of hydrolyzed FAs in primary hepatocyte cultures or on serum β-hydroxybutyrate concentrations in vivo. Moreover, L-FABP deletion did not influence the effects of ATGL knockdown or overexpression on PPAR-α target gene expression. Taken together, we conclude that L-FABP is not required to channel ATGL-hydrolyzed FAs to mitochondria for β-oxidation or the nucleus for PPAR-α regulation.

Liver FA binding protein (L-FABP) is an abundant cytosolic FA carrier involved in long chain FA transport throughout the cytoplasm and channeling of FAs and acyl-CoAs to the mitochondria for oxidation (12). L-FABP has been shown to interact with mitochondria (13) and, more specifically, carnitine palmitoyltransferase 1 (CPT1), the rate-limiting step in mitochondrial FA oxidation (12, 14, 15). These observations have been taken to imply that L-FABP may mediate the transport of FAs to the mitochondria. In support of this, L-FABP gene ablation in cultured primary hepatocytes suppresses mitochondrial FA oxidation (16). Additionally, in vitro studies have determined that L-FABP binds PPAR-α with high affinity and further demonstrate not VLDL secretion in vitro and in vivo (2, 3). Similar effects of ATGL on channeling hydrolyzed FAs to β-oxidation have also been observed in adipocytes, small intestine, and myocardial tissue (4–6). The mechanism through which ATGL channels FAs hydrolyzed at the surface of lipid droplets (LDs) to the mitochondria, however, is unknown.

In addition to its effects on FA trafficking, ATGL is also an important regulator of PPAR-α. We have shown that the expression of PPAR-α and its downstream target genes is decreased in mice or primary hepatocytes receiving ATGL shRNA (1). These results are consistent with microarray analysis of tissues in the whole-body ATGL KO mice (7) and work from others reporting that ATGL regulates PPAR-α in heart, small intestine, and brown adipose tissue (4–6). Although numerous metabolites can bind PPAR-α, FAs are thought to be the principal physiological ligands that activate PPAR-α (8–10). Given that ATGL produces FAs, it is assumed that ATGL regulates PPAR-α through the supply of FA ligands (11). However, data defining the mechanism through which ATGL regulates PPAR-α or how FAs are trafficked to PPAR-α are lacking.

Liver FA binding protein (L-FABP) is an abundant cytosolic FA carrier involved in long chain FA transport throughout the cytoplasm and channeling of FAs and acyl-CoAs to the mitochondria for oxidation (12). L-FABP has been shown to interact with mitochondria (13) and, more specifically, carnitine palmitoyltransferase 1 (CPT1), the rate-limiting step in mitochondrial FA oxidation (12, 14, 15). These observations have been taken to imply that L-FABP may mediate the transport of FAs to the mitochondria. In support of this, L-FABP gene ablation in cultured primary hepatocytes suppresses mitochondrial FA oxidation (16). Additionally, in vitro studies have determined that L-FABP binds PPAR-α with high affinity and further demonstrate

Supplementary key words peroxisome proliferator-activated receptor-α • β-oxidation • liver fatty acid binding protein • adipose triglyceride lipase

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Hepatic ATGL mediates PPAR-α signaling and fatty acid channeling through an L-FABP independent mechanism

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Adipose TG lipase (ATGL) was originally characterized as the major TG hydrolase in adipose tissue and heart, and more recent studies show it also has a predominant role in other tissues such as liver (1, 2). In addition to its effects on TG catabolism, ATGL influences the downstream partitioning of its FA products. We have shown that hepatic ATGL selectively channels hydrolyzed FAs to β-oxidation without affecting VLDL secretion (1). Consistent with these results, several other studies have also observed that hepatic ATGL overexpression or deletion affects FA oxidation but
that manipulating L-FABP expression alters PPAR-α-dependent transcription of genes coding for FA oxidative enzymes (17, 18). Yet, studies performed in vivo are inconsistent in regard to the role of L-FABP in mediating PPAR-α activity, with some studies showing that PPAR-α target gene expression is regulated independently from L-FABP (16, 19) and others showing dependence on L-FABP (20, 21).

Given the range of possible functional and physiological roles of L-FABP in FA oxidation, we questioned whether L-FABP is involved in the transport of a specific pool of FAs, ATGL-hydrolyzed FAs, to the mitochondria for oxidation and/or to the nucleus to regulate gene expression. To investigate the role of L-FABP in mediating the effects of ATGL, we used a strategy in which ATGL was either overexpressed (adenovirus expressing (Ad)-ATGL) or suppressed (ATGL shRNA) in both WT and L-FABP KO mice. Our findings reveal that despite its importance in hepatic FA transport, L-FABP is not required for the channeling of ATGL-hydrolyzed FAs to either the mitochondria for oxidation or to the nucleus for transcriptional regulation of PPAR-α.

MATERIALS AND METHODS

Animals, diets, and adenoviral administration

All animal protocols were approved by the University of Minnesota Institutional Animal Care and Use Committee. Male L-FABP KO mice on a C57BL/6J background were generated as previously described (22). All mice were housed under controlled temperature and lighting (20–22°C; 12:12 h light-dark cycle). Eight- to ten-week-old WT or L-FABP KO mice were injected via the tail vein with adenovirus expressing green fluorescent protein (Ad-GFP) or Ad-ATGL for overexpression studies or adenoviruses encoding ATGL shRNA or control shRNA targeting a nonspecific mRNA. Adenoviruses were generated as described previously (1, 23). All mice had free access to water and were fed a purified diet (TD.94045) from Harlan Teklad Premier Laboratory Diets (Madison, WI) for 1 week following adenoviral administration. The control diet contained protein (19% of total calories), carbohydrates (64%), and fat (17%), and the fat source was soybean oil (70 g/kg). Exactly 1 week after adenoviral administration, mice were euthanized for liver tissue and serum collection after a 4 h or overnight fast.

Primary hepatocyte isolation

Mouse primary hepatocytes were isolated by the collagenase perfusion method described previously (1). Hepatocytes were plated on collagen-coated multi-well tissue culture plates (Nunc) for 4 h with M199 plating media (Invitrogen) that contained 23 mM HEPES, 26 mM sodium bicarbonate, 10% FBS, 1% penicillin/streptomycin, 100 nM dexamethasone, 100 nM insulin, and 11 mM glucose. M199 maintenance media contained 23 mM HEPES, 26 mM sodium bicarbonate, 1% penicillin/streptomycin, 5.5 mM glucose, 100 µM carnitine, 10 nM dexamethasone, and 10 nM insulin. Cells were maintained in a humidified incubator at 37°C, 5% CO₂.

Cell adenoviral infection and radiolabeling

Ad-ATGL or Ad-GFP, which serves as a control virus, was generated as described previously (1). After 4 h of plating, primary hepatocytes were exposed to either Ad-ATGL or Ad-GFP at 100 multiplicities of infection for 24 h. Cells were pulsed with 500 µM oleate and 1 µCi [1-14C]oleate bound to FA-free BSA in a 3:1 molar ratio for 1.5 h (pulse period). Some cells were harvested for measurement of radiolabel incorporation into cellular lipid fractions. Parallel wells were washed with phosphate-buffered saline and new medium, void of FAs, was added for an additional 6 h (chase period) followed by collection of cells for lipid extraction. Lipid samples were separated into different fractions by TLC on 0.25 mm silica gel G plates in a hexane:ethyl ether:acetic acid (80:20:2, v/v/v) solvent system. TG fractions were identified and measured by AR-2000 radio-TLC imaging scanner. Total radiolabeled lipids were quantified by scintillation counter (LS6000IC, Beckman) after addition of Bio-Safe II cocktail.

Measurement of FA oxidation

During either the pulse or chase periods, as described above, radiolabeled media were transferred to Eppendorf tubes containing 20% BSA. Subsequently, perchloric acid was added to the tubes followed by vortexing. Following an overnight incubation at room temperature, the tubes were centrifuged and the supernatant fraction that contained [14C]-labeled acid-soluble metabolites (ASMs) was quantified via scintillation counting. In some experiments, HTS01037, a pan-FABP inhibitor, was dissolved in DMSO and applied to hepatocytes during the chase period at a final concentration of 100 µM.

RNA isolation, RT-PCR, and real-time quantitative PCR analysis

RNA was extracted with TRIzol from liver tissues followed by reverse transcription with SuperScript III First-Strand Synthesis SuperMix (Invitrogen). Gene expression was quantified using a SYBR GreenER Two-Step quantitative RT-PCR kit (Invitrogen) and an Applied Biosystems StepOne Plus real-time PCR system. Data were analyzed using the delta-delta CT method. For all analyses, gene expression was normalized to ribosomal protein L32. Melting curve analysis was performed on all samples to verify primer specificity.

Liver TG and FA analysis

TG was extracted from liver tissues or hepatocytes according to a method from Folch (24). Snap-frozen liver tissues were homogenized in sterile water and extracted twice with chloroform:methanol. The chloroform layer was isolated, dried under nitrogen gas, and redissolved in diluted Triton X-100. Colorimetric assays were used to quantify liver TG (Stanbio) and NEFAs (both from Wako Chemicals), and serum TG (Stanbio).

Statistical analysis

Results are expressed as mean ± standard error of the mean. Statistical analysis was performed using an unpaired Student’s t-test or, where appropriate, a two-way ANOVA test followed by Fischer’s protected least significant difference. Values of P < 0.05 were considered statistically significant.

RESULTS

ATGL knockdown increases liver weights and TG content

Given that the deletion of L-FABP has been shown to decrease hepatic TG accumulation (22) and ATGL is a major hepatic TG hydrolase (1), we investigated the effects of ATGL knockdown in L-FABP KO mice. ATGL shRNA administration suppressed protein expression of hepatic ATGL in both WT and L-FABP KO mice efficiently 7 days.
posttransduction (Fig. 1A). ATGL knockdown had no effect on body weight in WT or L-FABP KO mice when compared with control shRNA treatment (Fig. 1B). ATGL knockdown led to increased liver weight in WT mice and this effect was replicated in L-FABP KO mice (Fig. 1C). L-FABP ablation also resulted in reduced liver weights. ATGL knockdown led to enhanced hepatic TG accumulation in both WT and L-FABP KO mice, although the fold induction was less in the L-FABP KO mice (Fig. 1D). Consistent with previous findings, hepatic TG content was significantly lower in L-FABP KO mice compared with WT controls (21). Neither ATGL knockdown or L-FABP ablation affected the content of liver free FAs in either genotype (Fig. 1E).

**ATGL overexpression has minimal effects on liver weight and TG content in 4 h-fasted mice**

To further examine the importance of L-FABP in mediating the effects of ATGL on hepatic lipid metabolism, we also overexpressed ATGL in both WT and L-FABP KO mice. Administration of Ad-ATGL adenovirus increased the protein expression of hepatic ATGL in both WT and L-FABP KO mice 7 days postinfection (Fig. 2A). Consistent with the findings above, overexpressing ATGL did not alter body weight of WT or L-FABP KO mice (Fig. 2B). Increasing the expression of hepatic ATGL decreased liver weights only in WT mice (Fig. 2C). In addition, L-FABP ablation also resulted in lower liver weights. Hepatic TG content was reduced in L-FABP KO mice, whereas ATGL overexpression had no significant effect (Fig. 2D). The lack of effect of ATGL on liver TG may stem from the low basal TG levels in 4 h-fasted mice.

**ATGL overexpression increases serum BHBA regardless of genotype**

We next determined the effect of ATGL knockdown on serum TG, BHBA, and NEFA. ATGL knockdown had no effect on serum TG, BHBA, or NEFA in either genotype (Fig. 3A–C). However, serum TG levels in overnight-fasted L-FABP KO mice were lower than those of WT mice treated with control shRNA adenovirus after an overnight fast (Fig. 3A). Two previous studies showed no change in serum TG levels with L-FABP deletion, but the discrepancies in fasting duration and mouse gender may account for the differences (22, 25). The main effect of L-FABP KO on lowering serum BHBA approached significance ($P = 0.06$) in mice fasted for 16 h (Fig. 3B), but not 4 h (Fig. 3E). Overexpressing ATGL did not alter serum TG or NEFA in WT or L-FABP KO mice, but increased BHBA in both mouse lines (Fig. 3D–F).

**L-FABP ablation does not attenuate ATGL-mediated β-oxidation**

Our previous studies have shown that hepatic ATGL regulates β-oxidation in primary hepatocyte cultures (1). Thus, we performed pulse-chase experiments with [1-$^{14}$C]oleate in primary mouse hepatocytes isolated from WT or L-FABP KO mice to gain insights into the role of L-FABP in influencing ATGL-induced changes in β-oxidation. During the pulse period, overexpressing ATGL decreased incorporation of [1-$^{14}$C] oleate into the TG fraction of primary hepatocytes derived from both WT and L-FABP KO mice, but increased BHBA in both mouse lines (Fig. 4A). The lack of effect of ATGL on liver TG may stem from the low basal TG levels in 4 h-fasted mice.

ATGL overexpression has minimal effects on body weight in WT or L-FABP KO mice when compared with control shRNA treatment (Fig. 1B). ATGL knockdown led to increased liver weight in WT mice and this effect was replicated in L-FABP KO mice (Fig. 1C). L-FABP ablation also resulted in reduced liver weights. ATGL knockdown led to enhanced hepatic TG accumulation in both WT and L-FABP KO mice, although the fold induction was less in the L-FABP KO mice (Fig. 1D). Consistent with previous findings, hepatic TG content was significantly lower in L-FABP KO mice compared with WT controls (21). Neither ATGL knockdown or L-FABP ablation affected the content of liver free FAs in either genotype (Fig. 1E).

**ATGL overexpression has minimal effects on body weight in WT or L-FABP KO mice when compared with control shRNA treatment (Fig. 1B). ATGL knockdown led to increased liver weight in WT mice and this effect was replicated in L-FABP KO mice (Fig. 1C). L-FABP ablation also resulted in reduced liver weights. ATGL knockdown led to enhanced hepatic TG accumulation in both WT and L-FABP KO mice, although the fold induction was less in the L-FABP KO mice (Fig. 1D). Consistent with previous findings, hepatic TG content was significantly lower in L-FABP KO mice compared with WT controls (21).**
isoforms do not play a major role in mediating the oxidation of ATGL-derived FAs (Fig. 4E). Taken together, these data indicate that L-FABP is not a major carrier of ATGL-hydrolyzed FAs and may be more important for channeling exogenous FAs for the purpose of mitochondrial β-oxidation.

L-FABP ablation or ATGL knockdown alters the expression of other FABP isoforms

We quantified mRNA abundance of FABP2–7 to further test whether either L-FABP ablation or ATGL knockdown influenced expression of other FABP isoforms. Only FABP7 was significantly reduced in response to L-FABP ablation alone (Fig. 5). ATGL knockdown decreased the expression of several FABP isoforms in the WT or L-FABP backgrounds. In contrast, FABP3 was significantly increased following ATGL knockdown in the L-FABP KO mice, but not in the WT mice. Thus, based on mRNA abundance, if any compensation occurred in response to L-FABP ablation and ATGL knockdown it would likely be due to FABP3.

L-FABP is not required for the induction of PPAR-α target genes following ATGL overexpression

We have previously shown that hepatic ATGL promotes FA oxidation via changes in the expression of PPAR-α and its target genes (1). Other work has demonstrated that L-FABP binds directly to PPAR-α and may play a role in PPAR-α activation (17). To further examine whether L-FABP serves as a carrier of ATGL-hydrolyzed FAs to the nucleus to influence oxidative genes, we also quantified PPAR-α target gene expression in WT or L-FABP KO mice treated with Ad-ATGL adenovirus. If L-FABP mediates the effects of ATGL on FA trafficking, we would expect that the induction of FA oxidation and PPAR-α target gene expression following ATGL overexpression would be attenuated in L-FABP KO mice and that no reduction in the same outcomes would be observed following ATGL knockdown in L-FABP mice. In WT mice, hepatic ATGL overexpression

were not due to an effect of L-FABP on lipolysis, as ATGL increased the amount of radiolabeled TG lost during the chase period equally in WT and L-FABP KO hepatocytes (Fig. 4C).

To ensure that this observation was not due to a compensatory effect by other FABP isoforms, we also utilized a pan-specific FABP inhibitor, HTS01037, during the chase period in hepatocytes isolated from WT and L-FABP KO mice. Although HTS01037 selectively inhibits FABP4, it inhibits all FABP isoforms with Kᵢ values of less than 10 μM (26), well below the 100 μM concentration used in the current study. HTS01037 resulted in a small, but significant, attenuation of the ATGL-mediated increase in ASM during the chase period regardless of the mouse line, suggesting that other FABP isoforms do not play a major role in mediating the oxidation of ATGL-derived FAs (Fig. 4E). Taken together, these data indicate that L-FABP is not a major carrier of ATGL-hydrolyzed FAs and may be more important for channeling exogenous FAs for the purpose of mitochondrial β-oxidation.
led to approximately a 0.4- to 2-fold increase in PPAR-α and its target genes (Fig. 6). Despite the absence of L-FABP, adenoviral-mediated overexpression of ATGL still induced gene expression of PPAR-α, acyl-CoA thioesterase (ACOT1), long chain acyl-CoA dehydrogenase (LCAD), and acyl-CoA synthetase 1 (ACSL1) (0.3- to 2.5-fold) in L-FABP KO mice. Thus, these data support the overexpression studies and show that L-FABP does not mediate the effects of ATGL on the expression of PPAR-α target genes, and imply that an alternate mechanism links changes in ATGL activity to transcriptional regulation.

**DISCUSSION**

It is well-established that L-FABP plays a central role in hepatic FA trafficking and energy metabolism. Its high expression level and ability to interact with mitochondria and nuclear transcription factors suggest that it may act as a FA sensor that alters metabolism directly via changes in intracellular FA handling as well as transcriptional regulation of metabolic pathways. To date, most studies have focused on the role of L-FABP in modulating exogenous FA metabolism, especially during fasting, high fat feeding, or in various disease models (20, 21, 25, 27, 28). The effects of L-FABP on the metabolism and signaling properties of endogenous FAs, such as those derived from lipolysis, are unknown. The present study sought to determine whether L-FABP is important in modulating the effects of ATGL on the signaling and channeling of hydrolyzed FAs.

Previous findings that show ATGL selectively channels hydrolyzed FAs exclusively to β-oxidation provide novel insights into the regulation of intracellular lipid trafficking (1, 2, 4–6). To better understand the mechanisms responsible for this selective partitioning of FAs, we tested the importance of L-FABP in mediating these effects. In contrast to its clear effects on channeling of exogenous FAs to oxidative pathways, L-FABP did not mediate the trafficking of hydrolyzed FAs derived from ATGL in β-oxidation. Consistent with the lack of effect on intracellular FA trafficking, L-FABP did not modulate serum TG, BHBA, or NEFA in mice with manipulated hepatic ATGL expression. The lack of effect of L-FABP ablation on oxidation of FAs derived from ATGL was surprising, and suggests that either another lipid carrier is responsible for channeling hydrolyzed FAs or that this pool of FAs does not require a protein carrier for their shuttling to the mitochondria. To address the former possibility, we measured oxidation of ATGL-derived FAs in the presence of a pan L-FABP inhibitor and found that the inhibitor only resulted in a small (~15–20%) attenuation of ATGL-induced FA oxidation. The results from this study suggest that there may be some compensation by other FABP isoforms, but it is unlikely to be a major contributor to the oxidation of hydrolyzed FAs. We also found that FABP3 was modestly increased in L-FABP KO mice treated with ATGL shRNA, although it is questionable whether this increase in a relatively low-abundance hepatic FABP isoform (29) could compensate for the lack of L-FABP. Previous findings show other L-FABP isoforms or putative FA transporters, such as sterol carrier proteins (SCP-2 and SCP-x), are not upregulated in L-FABP KO mice (20, 22). We found that only FABP7 was significantly decreased in the L-FABP KO mice. Although the current study does not
preclude the importance of other FA carriers, the data do not support a major role of other FABP isoforms in compensating for L-FABP ablation.

An alternate potential mechanism explaining the selective channeling of hydrolyzed FAs to oxidative pathways may involve a direct interaction between LDs and mitochondria where FAs are “handed off” to downstream enzymes independent of FA carriers. In fact, mitochondria are often observed to be closely located to LDs in adipocytes, heart, and liver (30–35). Given that oxidative and metabolically active tissues often have higher rates of lipolysis and FA oxidation, conditions that increase energy demand may enhance these interactions. In exercise-trained dogs, goats, and humans, every LD in muscle tissue is observed to be in contact with at least one mitochondrion (36, 37). In addition, mitochondria and LDs also form chains of alternating organelles in type I fibers of human skeletal muscle (38). When lipolysis is activated with resveratrol in hepatocytes, more mitochondria are observed to directly attach to LDs than those treated with vehicle (39). Notably, perilipin 5, a LD protein commonly found in oxidative tissues, plays an important role in linking LDs to mitochondria in cardiomyocytes and muscle tissues (40, 41). However, despite increased LD-mitochondria interactions in response to perilipin 5 overexpression, oxidation of hydrolyzed FAs is not altered (40). Thus, the importance of LD-mitochondria interactions, or the mechanism through which hydrolyzed FAs are oxidized, remains largely unknown.

Several studies have shown that L-FABP mediates the effects of exogenous FAs, especially polyunsaturated FAs, on PPAR-α gene expression and activity (17, 42, 43). L-FABP physically interacts with PPAR-α and it is thought that this interaction facilitates transfer of FA ligands (18). Whether

**Fig. 5.** Changes in FABP isoform expression in response to ATGL knockdown or L-FABP ablation. WT or L-FABP KO mice, at 8–10 weeks of age, were infected with control or ATGL shRNA adenovirus (n = 8–10 per group) followed by harvesting of liver tissues 7 days after an overnight fast. Data are presented as mean ± SEM. *P < 0.05 versus control shRNA group; #P < 0.05 versus WT mice.

**Fig. 6.** L-FABP is not required for the induction of PPAR-α target genes following ATGL overexpression. WT or L-FABP KO mice, at 8–10 weeks of age, were transduced with Ad-GFP or Ad-ATGL adenovirus (n = 8–10 per group). Liver tissues were collected 7 days posttransduction after a 4 h fast. Data are presented as mean ± SEM. *P < 0.05 versus GFP; #P < 0.05 versus WT mice.
L-FABP is critical in regulating activation of PPAR-α by ATGL-hydrolyzed FAs is unknown. Herein, we reveal that L-FABP ablation does not mediate the effects of ATGL on PPAR-α target gene expression in comparison with WT mice, suggesting that L-FABP is not a major transporter of ATGL-hydrolyzed FAs to the nucleus. Moreover, ATGL overexpression or knockdown and L-FABP ablation did not influence FA levels in the liver. These data imply that additional factors beyond FABP and intracellular TG hydrolysis control intracellular FA levels, and that perhaps ATGL and L-FABP regulate distinct pools of FAs that are differentially channeled.

The finding that L-FABP ablation does not mediate the effects of ATGL on PPAR-α target gene expression is suggestive of a more complex mechanism linking lipolysis to transcriptional regulation. These data are in agreement with a previous study from our laboratory showing that administration of fenofibrate, a PPAR-α agonist, is unable to rescue PPAR-α target gene expression or normalize hepatic TG levels (1). In contrast, administration of Wy-14643 to ATGL KO mice rescues their cardiomyopathy and normalizes cardiac oxidative gene expression (6). It is unclear if these discrepancies between the two studies are due to differences between PPAR-α agonists or the target tissues evaluated. A recent study showed that ATGL was required for lipolytic stimulation of adipocytes to activate PPAR-α reporter constructs that were generated by fusing the ligand binding domain of PPAR-α to the C-terminal LD binding domain of perilipin 1 so that PPAR-α was bound to LDs (44). However, the physiological significance of this model compared with the predominant nuclear localization of PPAR-α remains to be determined.

In summary, this study demonstrates that L-FABP, despite being a major FA carrier and an abundant cytosolic protein, is not necessary for the shuttling of FAs hydrolyzed by ATGL to the mitochondria. Furthermore, L-FABP plays no essential role in delivering ATGL-hydrolyzed FAs to the nucleus to modulate oxidative genes. These findings allude to a more complex mechanism linking ATGL to downstream changes in hepatic energy metabolism. 

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