Endothelial lipase provides an alternative pathway for FFA uptake in lipoprotein lipase–deficient mouse adipose tissue

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Lipoprotein lipase (LPL) is thought to be the only enzyme responsible for the catabolism of triglycerides (TGs) associated with TG-rich lipoproteins in adipose tissue (AT). However, LPL deficiency in humans and induced mutant mice is not associated with decreased fat mass. We investigated whether endothelial lipase (EL), a recently discovered phospholipase, might represent an alternative mechanism for the uptake of phospholipid-derived fatty acids in murine lipoprotein-deficient AT. When LPL was expressed in AT and isolated murine adipocytes, EL mRNA was not detectable. In contrast, mouse AT and isolated adipocytes that lacked LPL expressed large amounts of EL mRNA. The cellular phospholipase activity in LPL-deficient fat pads was increased 4-fold compared with control fat pads and could be inhibited to control levels by a specific EL antibody. Fatty acids produced by EL activity were absorbed by adipocytes and incorporated into the TG moiety of AT. Our results suggest that EL activity in AT and other peripheral tissues might contribute to the tissue uptake of free fatty acids, which could have important implications for the metabolism of plasma lipoproteins.

Introduction

Lipoprotein lipase (LPL) is responsible for the generation of fatty acids from circulating triglyceride-rich (TG-rich) lipoproteins. The enzyme is predominantly found in adipose tissue (AT) and muscle, where it is bound to the capillary endothelium by its interaction with glucosaminoglycans. In AT, the released fatty acids are absorbed and reesterified, and TGs are stored within lipid droplets (1, 2). Complete absence of LPL, which occurs in human individuals, cats, and minks that are homozygous for mutations in the LPL gene (3–5), results in massive accumulation of plasma TGs, consistent with the view that the absence of LPL impedes the normal clearance of circulating TGs (6). In contrast to humans, cats, and minks, homozygous LPL-knockout mice (L0 mice) (7, 8) or mice that lack functional LPL due to a mutation in the id locus (9) do not survive the suckling period and die between 18 and 24 hours after birth. However, the transient expression of LPL by adenovirus infection during the suckling period (10) or the transgenic expression of LPL exclusively in skeletal muscle (SM) (11, 12), cardiac muscle (CM) (13), or liver (14) can rescue L0 mice from neonatal death.

The remarkable phenotype observed in LPL-deficient humans, mice, cats, and minks strongly supports the view that LPL is essential for the clearance of TG-associated fatty acids and their subsequent uptake by peripheral tissues. Unexpectedly, however, LPL-deficient humans or animals exhibit essentially normal AT mass, and their fat cells are filled with lipid (4, 5, 12, 13, 15–17). Importantly, the FFA composition of the TG moiety in LPL-deficient AT differed markedly from that in control AT, exhibiting a drastic decrease in polyunsaturated, essential fatty acids (12, 18). This finding suggested that preservation of AT mass in the absence of LPL was achieved by de novo biosynthesis of endogenous fatty acids. Recently we demonstrated that lipogenesis is indeed activated in LPL-deficient AT through an SREBP-1–mediated process (19). Lipogenesis, however, could not explain the presence of low levels of polyunsaturated fatty acids in LPL-deficient AT. Therefore, we speculated that alternative fatty acid uptake mechanisms must exist when LPL is absent. These mechanisms could involve the induction of alternative lipases that are capable of providing fatty acids from plasma lipoproteins and/or the uptake of FFAs from HDL/albmin complexes.

Endothelial lipase (EL) is a plausible candidate to provide FFAs for subsequent uptake in peripheral tissues in addition to LPL. The EL gene encodes for an enzyme that belongs to the family of TG lipases together with LPL, hepatic lipase, and pancreatic lipase (20, 21). EL is expressed in many tissues and organs, such as the liver, lung, kidney, steroid hormone–producing organs, thyroid, and placenta. Additionally, EL is also expressed in vascular endothelial cells. This is a unique feature among the lipase family, because despite the fact that LPL and hepatic lipase are active on the luminal surface of the endothelium, neither is synthesized by endothelial cells (1, 22). EL exhibits a 45% amino acid homology with LPL with putative heparin-binding and hydrophobic regions, which suggests that EL, like LPL and hepatic lipase, is located on the capillary endothelium (23–26). In contrast to LPL, however, EL was shown to be a phospholipase with little TG lipase activity (less than 0.1 μmol FFA/ml/h) in the presence of serum (20, 21, 27). The primary lipoprotein substrate of EL is HDL (27), and the enzyme has been assigned an important role in HDL metabolism (26, 28–30). Specifically, mice that lack EL have increased HDL cholesterol levels (28, 29), whereas EL overexpressing mice exhibited decreased HDL cholesterol levels (31). Additionally to its enzyme function as phospholipase, EL has been shown to facili-
tate the cellular uptake of lipoprotein particles by a nomenclature mechanism similar to that of LPL or hepatic lipase (23, 25).

The present study was designed to investigate whether EL is expressed in AT of mice that lack LPL activity in adipocytes. We show that EL expression is induced on the mRNA level and increased enzyme activities are found in LPL-deficient AT. The FFAs generated by the increased phospholipase activity are taken up by AT and incorporated into the lipid droplets.

Results

tissue LPL activities and LPL mRNA abundance in AT. LPL enzyme activities in AT and muscle as well as LPL mRNA levels in AT for all mouse genotypes used in this study are summarized in Table 1 and Figure 1A, respectively. Normal mice with 2 LPL wild-type alleles (L2 mice) were used as control animals. L2-MCK mice are transgenic animals that overexpress a human LPL minigene under the control of the muscle specific mouse-creatine kinase promoter (32). Accordingly, these animals have increased LPL activities in SM and CM, whereas LPL activity in AT is identical to control mice (Table 1). L2 and L2-MCK mice show similar LPL mRNA expression levels in AT (Figure 1A).

L0-MCK mice resulted from the cross-breeding of L2-MCK mice with heterozygous L0 mice (11). L0-MCK animals lacked both mouse LPL alleles but expressed the human LPL transgene. Accordingly, these animals have detectable LPL activities exclusively in SM and CM but lack enzyme activity (Table 1) and detectable mRNA levels (Figure 1A) in AT.

Finally, L0 mice that had been rescued from postnatal death by the transient expression of LPL during the suckling period utilizing adenovirus-mediated gene transfer were used in this study (10). Rescued adult L0 mice exhibited no detectable LPL enzyme activity (Table 1) or mRNA (Figure 1A) in any tissue analyzed.

Plasma TG levels. As shown in Table 2, increased muscle LPL activities in L2-MCK mice resulted in a highly significant decrease of plasma TG levels compared with controls (36%). Additionally, HDL cholesterol concentrations were also decreased (16%). The physiological response to the altered tissue-specific LPL expression pattern in L0-MCK mice resulted in decreased plasma TG levels (41%) and decreased plasma HDL concentrations (29%). The complete loss of LPL activity in all tissues of L0 mice resulted in extremely high plasma TG levels (20-fold increase) and essentially undetectable plasma HDL cholesterol levels.

FFA composition of AT lipids. The variation of AT-LPL activities among the mouse genotypes resulted in marked differences in the fatty acid composition of the TG and phospholipid (PL) moieties (Table 3). Compared to AT that expressed normal LPL activities (L2 and L2-MCK mice), AT that lacked LPL (L0-MCK mice) exhibited a highly significant reduction of 18:2 and 18:3 fatty acids in both the TG (88% and 87%, respectively) and the PL (60% and 71%, respectively) moiety. Conversely, the concentrations of 16:1 and 18:1 were increased in the TGs (247% and 57%, respectively) and the PL (174% and 33%, respectively) content.

EL mRNA expression in AT and liver. Northern blot analysis (Figure 1B) revealed that no EL-specific mRNA was found in AT of control (L2) and L2-MCK mice. In contrast, high levels of mouse EL mRNA were detected in AT that lacked LPL activity. Compared with L2 and L2-MCK mice, both L0-MCK mice and rescued adult L0 mice showed increased EL mRNA levels, which indicates that the induction of AT-EL expression was not affected by the expression of LPL in other tissues but solely by the absence of LPL in adipocytes. Similarly to what is shown here, an induction of EL mRNA expression was also found in AT of mice that expressed LPL exclusively in CM (13) (data not shown). The upregulation of EL mRNA expression in AT of L0-MCK mice was additionally confirmed by quantitative real-time PCR and revealed a 7.9-fold induction compared with control adipose tissue (Figure 1C).

Hepatic EL mRNA concentrations were found to be similar in L2 and L0-MCK mice by real-time PCR (Figure 2) and in Northern blot experiments (Figure 2, inset). Interestingly, EL mRNA concentrations were markedly higher (2-fold) in LPL-deficient AT than in liver samples independent of the mRNA quantitation method.

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**Table 1**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>LPL activity in AT (μmol FFA/g/h)</th>
<th>CM-LPL activity (μmol FFA/g/h)</th>
<th>SM-LPL activity (μmol FFA/g/h)</th>
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</thead>
<tbody>
<tr>
<td>L2</td>
<td>8.6 ± 3.1</td>
<td>18.4 ± 4.3</td>
<td>2.2 ± 0.5</td>
</tr>
<tr>
<td>L0-MCK</td>
<td>0.09 ± 0.05a</td>
<td>3.7 ± 0.8a</td>
<td>7.9 ± 2.7a</td>
</tr>
<tr>
<td>L2-MCK</td>
<td>16.5 ± 6.5</td>
<td>25.8 ± 3.9d</td>
<td>7.9 ± 2.4d</td>
</tr>
<tr>
<td>L0</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

All values represent means ± SD. Age of the animals at the time of analysis was 10–14 weeks. *P ≤ 0.001, **P ≤ 0.01, ***P < 0.05 vs. controls (L2). n.d., not detected.

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**Figure 1**

**A** Murine LPL and EL mRNA expression in adipose tissue. Relative abundance of (A) LPL and (B) EL mRNA in AT of mice expressing (L2, L2-MCK) or lacking (L0-MCK, L0) LPL in AT. Total RNA was isolated from murine AT in the absence or presence of LPL and subjected to Northern blot analysis. Ten micrograms of total RNA were separated on a formaldehyde/agarose gel, blotted onto a nylon membrane, and hybridized with a murine-specific [32P]-labeled cDNA probe. rRNA, ribosomal RNA. (C) EL mRNA concentrations in AT were confirmed by fluorescent real-time PCR (see Methods). EL mRNA quantities were normalized to those of β-actin. Data are expressed as the mean values of 3 experiments performed in duplicates relative to the expression of EL mRNA in control mice (L2). Bars represent mean ± SD of the hybridization intensity signals. *P < 0.05.
EL mRNA in differentiated 3T3-L1 cells and isolated mouse adipocytes. Next, we investigated whether the transcriptional activation of EL gene expression might occur during the differentiation of 3T3-L1 mouse embryonic fibroblasts into adipocytes similarly as observed for LPL. Total RNA from 3T3-L1 cells at different stages of differentiation was isolated, and EL mRNA concentrations were determined by real-time PCR analysis. In contrast to LPL mRNA expression, which was markedly induced within the initial phase of differentiation (Figure 3A), EL was not expressed in undifferentiated cells or cells at 4 and 8 days of differentiation (Figure 3B). To elucidate whether isolated adipocytes express EL mRNA, epididymal fat pads from AT of control and L0-MCK mice were digested by collagenase treatment, and fat cells were isolated by flotation. This procedure enabled the separation of adipocytes from other contaminating cell types such as endothelial and blood cells. Quantitative real-time PCR analysis revealed a 26-fold induction of EL mRNA expression in isolated adipocytes of L0-MCK mice (Figure 3B). Thus, in the absence of LPL, isolated adipocytes are capable of synthesizing EL mRNA.

Phospholipase activity levels in LPL-deficient AT. Consistent with the increased mRNA levels, the phospholipase activity was also 4.0-fold higher in fat pads isolated from L0-MCK mice as compared with L2 mice (Figure 4). Preincubation of fat pads with a specific anti-EL IgG totally inhibited EL activity in L0-MCK AT, which resulted in decreased phospholipase activity levels and displayed levels similar to those in L2 mice. In contrast, the total phospholipase activities in preheparin or postheparin plasma of control and L0-MCK mice revealed no significant differences in the 2 genotypes (0.8 and 0.7 μmol FFA/ml/h, respectively).

Incorporation of PL-derived fatty acids into the TG moiety of LPL-deficient AT. The increased EL activity in LPL-deficient AT suggested a potential mechanism for the cellular uptake of FFAs and their subsequent intracellular reesterification and storage. To test this hypothesis, we incubated isolated epididymal fat pads of L2 and L0-MCK animals with [14C]phosphatidylcholine-labeled ([14C]PC-labeled) HDL, and the incorporation of the fatty acid–associated radioactivity into TGs was examined. As shown in Figure 5, the amount of PC-derived FFAs absorbed by AT and incorporated into TGs was 2.8-fold higher in AT-LPL–deficient adipocytes than in control cells.

Table 2
TG and HDL cholesterol plasma levels

<table>
<thead>
<tr>
<th>Genotype</th>
<th>n</th>
<th>TGs (mg/dl)</th>
<th>HDL-C (mg/dl)</th>
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<tbody>
<tr>
<td>L2</td>
<td>9</td>
<td>98 ± 27</td>
<td>107 ± 16</td>
</tr>
<tr>
<td>L0-MCK</td>
<td>9</td>
<td>58 ± 8a</td>
<td>76 ± 15b</td>
</tr>
<tr>
<td>L2-MCK</td>
<td>9</td>
<td>43 ± 10c</td>
<td>90 ± 12c</td>
</tr>
<tr>
<td>L0</td>
<td>5</td>
<td>2.007 ± 375b</td>
<td>2 ± 0.6c</td>
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</tbody>
</table>

All values represent mean ± SD. The age of animals at the time of analysis was 9 weeks. *P ≤ 0.01, **P ≤ 0.001, #P < 0.05 vs. controls (L2).

Discussion
Lipases are essential for the generation of lipid-derived fatty acids and their subsequent tissue uptake. In peripheral tissues such as AT, LPL is considered the only enzyme capable of this function. Yet LPL deficiency, as observed in patients with type I hyperlipoproteinemia or in L0 mice, is not associated with a complete absence of polyunsaturated fatty acids in AT. In fact, a significant proportion of essential fatty acids (10–30% of normal) are still found in the lipid moiety of LPL-deficient AT, which suggests that in the absence of LPL, uptake mechanisms for plasma fatty acids must exist. Two processes are conceivable. First, FFAs could be absorbed from plasma FFA/albumin complexes, and indeed our results demonstrate that LPL-deficient AT absorbs unesterified fatty acids more efficiently than normal LPL-expressing AT. Second, alternative lipases might become activated and generate fatty acids from lipids, which are subsequently taken up by the target tissue. Considering the potential involvement of alternative lipases, we analyzed LPL-deficient AT for the expression of EL and compared it to AT from normal control mice. EL seemed a reasonable candidate because it is structurally closely related to LPL and, similar to LPL, it is found at the capillary endothelium bound to glycosaminoglycans (23, 25, 26). Unlike LPL, however, EL hydrolyzes predominantly PLs, and the major lipoprotein substrates are not TG-rich lipoproteins but HDL (21, 27). The enzyme is expressed in endothelial cells and a variety of other cell types and tissues. However, expression of EL in AT or muscle, the major sites of LPL expression, has not been reported.

When fat tissue was analyzed for EL mRNA in Northern blot experiments, large amounts of EL mRNA were found in AT that lacked LPL, whereas no EL mRNA was detected in LPL-expressing the ability of LPL-deficient adipocytes to take up PC-derived FFAs and incorporate them into the TG moiety to control levels.

Incorporation of FFAs into the TG moiety of LPL-deficient AT. It is conceivable that adipose tissue lacking LPL not only induces EL expression as a compensatory uptake mechanism of fatty acids but also increases the uptake of albumin-bound unesterified fatty acids. To investigate this premise, we incubated isolated epididymal fat pads of L2 and L0-MCK animals with [3H]oleate, and the incorporation of the fatty acid–associated radioactivity into TGs was examined. As shown in Figure 6, the amount of FFAs absorbed by AT and incorporated into TGs was 4.3-fold higher in AT-LPL–deficient adipocytes than in control cells.

Table 3
Fatty acid composition of PLs and TGs in AT of control (L2) and AT-LPL–deficient (L0-MCK) mice

<table>
<thead>
<tr>
<th>Fatty acid</th>
<th>PLs (%)</th>
<th>TGs (%)</th>
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<tbody>
<tr>
<td>16:0</td>
<td>17.00 ± 2.24</td>
<td>17.47 ± 1.71</td>
</tr>
<tr>
<td>16:1</td>
<td>3.00 ± 0.13</td>
<td>6.22 ± 2.33a</td>
</tr>
<tr>
<td>18:0</td>
<td>17.33 ± 2.17</td>
<td>14.48 ± 3.37</td>
</tr>
<tr>
<td>18:1</td>
<td>31.15 ± 4.10</td>
<td>41.32 ± 3.34a</td>
</tr>
<tr>
<td>18:2</td>
<td>18.76 ± 1.03</td>
<td>7.50 ± 0.97a</td>
</tr>
<tr>
<td>18:3</td>
<td>2.31 ± 1.10</td>
<td>0.68 ± 0.39a</td>
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</table>

Data from 4 mice are shown as mean ± SD. *P ≤ 0.01; **P ≤ 0.001, #P < 0.05 vs. controls (L2).
control AT. The induction of EL expression was observed in LPL-deficient AT of mouse lines that lacked LPL completely (L0) or expressed LPL exclusively in SM (L0-MCK) or CM (data not shown), which suggests that the location and the quantity of LPL expression in other tissues of the body do not affect EL expression in AT. The comparison of the signal intensities determined by quantitative real-time PCR indicated that higher relative amounts of EL mRNA were present in LPL-deficient AT than in liver, an organ of high EL abundance. This observation made it likely that EL mRNA was actually produced by the adipocytes in AT rather than by contaminating endothelial cells. To test this hypothesis experimentally, we treated fat pads from control and LPL-deficient AT with collagenase, isolated adipocytes by standard techniques, and analyzed total RNA for EL mRNA levels by quantitative real-time PCR. As expected, isolated adipocytes from LPL-deficient AT produced high amounts of EL mRNA, whereas no signals were detected in RNA derived from control AT. In differentiated 3T3-L1 cells, which express high amounts of LPL during the course of differentiation, EL mRNA was undetectable at all time points of the differentiation period. These findings are consistent with the observations in AT. It follows that the absence of LPL is a prerequisite for the uptake and reesterification of fatty acids released from HDL-associated PC.

EL enzyme activity is inversely correlated to plasma HDL cholesterol levels (21). Recent studies have also shown that the inhibition of EL causes increased plasma HDL cholesterol concentrations in vivo identifying EL as an important drug target for the treatment of patients with low HDL cholesterol levels (28, 31). Whether AT-specific EL expression in LPL-deficient mice affects plasma HDL levels is presently unknown and remains to be determined. Additionally, it will be important to clarify whether patients affected with familial LPL deficiency express EL in AT and whether EL expression contributes to the excessively low HDL cholesterol levels in these patients.

**Figure 2**

Murine EL mRNA expression in liver. Relative abundance of EL mRNA in livers in the presence (L2) or absence (L0-MCK) of AT-LPL were compared with EL expression in AT of L0-MCK mice. Total RNA was isolated from fat pads of fed L2 and L0-MCK mice. mRNA concentrations were determined by fluorescent real-time PCR. EL mRNA quantities were normalized to those of β-actin. Data are expressed as the mean values of 3 experiments performed in duplicate. Bars represent mean ± SD of the hybridization intensity signals. Inset: Representative Northern blot analysis of 10 μg total RNA per lane, hybridized with a murine-specific [32P]-labeled EL cDNA probe. Lane 1, L0-MCK AT; lane 2, L2 liver; lane 3, L0-MCK liver.

**Figure 3**

LPL and EL mRNA expression in differentiated 3T3-L1 cells and isolated mouse adipocytes. Fat pads of mice in the presence (L2) or absence (L0-MCK) of LPL were homogenized and subjected to collagenase digestion, and total RNA of adipocytes was isolated. Total RNA from 3T3-L1 fibroblasts was isolated at 50% cell confluence (day −2), at confluence (day 0) and during (days 2, 4, 6, and 7), and at the end of differentiation (day 8) to adipocytes. (A) Northern blot analysis of 10 μg total RNA per lane, hybridized with a murine-specific [32P]-labeled LPL probe. (B) EL mRNA concentrations were determined by fluorescent real-time PCR. EL mRNA quantities were normalized to those of β-actin, and control values from L2 mice were arbitrarily set to 1. *P < 0.05.
Taken together, our data are consistent with the concept that peripheral EL expression on the mRNA level and the level of enzyme activity can be initiated in murine AT when LPL is absent. In LPL-deficient AT, both the metabolic switch from the catabolism of chylomicron and VLDL TGs (which is LPL mediated) to HDL PLs (which is EL mediated) and the observed increased uptake of albumin-bound fatty acids are likely to supply those essential fatty acids that are needed to maintain essential adipocyte functions, such as accurate membrane fluidity and the biosynthesis of prostaglandins and leukotrienes. However, the markedly reduced content of polyunsaturated fatty acids in both the TG and the PL moieties of LPL-deficient AT indicated that the EL-mediated FFA import and the increased uptake of albumin-bound fatty acids are not sufficient to completely normalize fatty acid import in adipocytes when LPL is absent. This deficit is additionally counteracted by the massive induction of lipogenic processes in LPL-deficient AT (19). Whether EL expression in AT has important implications for the cellular lipid metabolism and the metabolism of plasma lipoproteins remains to be elucidated. In this context, it will be important to reveal conditions other than LPL deficiency that can modulate EL expression in AT and possibly other peripheral tissues.

Methods

Mice. The generation of all mouse lines used as well as their detailed characterization was published previously (10–12). Genotypes were identified from tail tip DNA by PCR analysis, as reported (12). All animals were maintained on a regular light-dark cycle (14 h light, 10 h dark) and kept on a standard laboratory chow diet. All animal experiments were performed in accordance with the standards established by the Austrian Federal Ministry of Education, Science and Culture, Division of Genetic Engineering and Animal Experiments (Vienna, Austria).

Cell culture. 3T3-L1 mouse embryonic fibroblasts were cultured in DMEM containing 10% FCS, 1% t-glutamine, and 1% streptomycin/penicillin at 37°C with 5% carbon dioxide in air. At confluence, the medium was changed and supplemented with 8 μg/ml biontin and 4 μg/ml pantothenic acid for 2 days. Differentiation was initiated by the addition of 10 μg/ml insulin and 0.4 μg/ml dexamethasone (day 0, 2 days after confluence). After 48 hours (day 2), the medium was changed to DMEM supplemented with 8 μg/ml biontin, 4 μg/ml pantothenic acid, and 10 μg/ml insulin. Subsequently, the medium was replaced every 48 hours with DMEM containing 8 μg/ml biontin, 4 μg/ml pantothenic acid, and 0.2 μg/ml insulin. Adipocyte differentiation was examined microscopically. RNA was isolated (RNase K; QIAGEN) every 2 days starting at 50% confluence and ending 8 days after initiation of differentiation (day 8).

Isolation of mature adipocytes from mouse epididymal fat pads. Fat pads were removed surgically, weighed, minced with scissors, and washed in Krebs-Ringer buffer plus HEPES (KRBH) (120 mM NaCl, 4 mM KHPO₄, 700 μM MgSO₄, 10 mM NaHCO₃, 1 mM CaCl₂, 30 mM HEPES, 200 μM adenosine, 1% BSA, pH 7.4) at 37°C. Collagenase digestion was performed as described previously (33). Per 100 mg of AT, 400 μl KRBH and 25 μl collagenase A (20 mg/ml; Roche Diagnostics) were added, and the mixture was incubated in a shaking water bath at 37°C for 1 h. After collagenase digestion and the addition of 250 μl of KRBH at 37°C, the cells were mixed and gently passed through a 250-μm nylon mesh filter. The cells were washed twice with 750 μl KRBH at 37°C and briefly centrifuged at 200 g. Adipose cells floating on top of the aqueous buffer were collected. Total RNA was isolated using an RNase K (QIAGEN).

Plasma lipids and AT fatty acid analysis. Blood was collected from fed animals by retroorbital bleeding, and EDTA-plasma was prepared. Plasma TG concentrations (GPO-Trinder 20) and HDL cholesterol concentrations (HDL-Cholesterol Kit 352-3) were measured enzymatically (Sigma-Aldrich).

The fatty acid composition of PLs and TGs in AT of L2 and L0-MCK mice was determined by GC-MS analysis from AT of 4 L2 and 4 L0-MCK mice. Fat pads of fed mice were removed surgically, weighed, and homogenized in PBS in sterile plastic vials. Immediately afterward, the homogenate was transferred to glass vials with Teflon stoppers. Lipids were extracted with 5 ml of chloroform/methanol (2:1, vol/vol) on ice, the organic phase was collected, and the aqueous phase was reextracted with 3 ml chloroform/methanol (2:1, vol/vol). The organic phases were pooled and dried under nitrogen. Per 100 mg of AT, 1 ml of chloroform was added, and 100 μl of the redissolved lipids were separated by thin-layer chromatography (silica gel 60; hexane/diethyl ether/acetic acid, 70:29:1). PL- and TG-specific spots were scraped from the plates and dissolved in 500 μl toluene containing 30 μg 15:0 fatty acid as internal standard and 40 μl of [14C]PC-labeled HDL (50 μg HDL-associated protein/sample) at 37°C for 6 hours. The incorporation of FFAs into the TG moiety of AT was performed in the absence (control) or presence of 50 μg anti-EL IgG, 50 μg/ml THL or/and 2 U/ml heparin in a total volume of 150 μl. Results are mean ± SD from 5 mice of each genotype performed in duplicate. **P ≤ 0.01.
RNA isolation and Northern blot analysis. Fat pads of adult L2 and L0-MCK mice were removed surgically, weighed, and subsequently frozen in liquid nitrogen. Total RNA was isolated using the TRI Reagent procedure according to the manufacturer’s protocol (Molecular Research Center Inc.). For Northern blot analysis, 10 μg of total RNA was separated by 1% formaldehyde/agarose gel electrophoresis and blotted overnight onto nylon membranes (Hybond N; Amersham Biosciences). RNA was immobilized to membranes by ultraviolet irradiation. The blots were prehybridized for 4 hours at 65°C in a buffer containing 0.15 M sodium phosphate (pH 7.2), 1 mM EDTA, 7% (wt/vol) SDS, and 1% (wt/vol) BSA. Membranes were hybridized in the same buffer at 65°C overnight with appropriate radiolabeled murine EL and LPL probes. After hybridization, the blots were washed in 2× SSC (0.15 M NaCl, 0.015 M sodium citrate) containing 0.1% SDS for 20 minutes at room temperature, followed by 2 additional washes in 0.1× SSC/0.1% SDS for 10 minutes at 65°C each. Specific hybridization signals were visualized after exposure to PhosphorImager Screens on a Storm PhosphorImager and analyzed using ImageQuant software (Amersham Biosciences).

A full-length murine EL cDNA clone was generated from total mouse liver RNA by RT-PCR. After reverse transcription of liver mRNA using oligo-dT primers, the resulting cDNA was PCR amplified with 2 EL-specific primers — 5′-AAGGATGCGAAACACGGTTT-3′ (forward) and 5′-GCACGAGGATCAGAGTTGATG-3′ (reverse) — using the Advantage RT-for-PCR Kit (BD Biosciences — Clontech). The reaction resulted in a 1.5-kb EL cDNA fragment, which was subcloned into pBluescript. After we confirmed the correct amplification product by DNA sequencing, an 800-bp BamHI-XhoI fragment was radioactively labeled with [32P]dCTP (NEN, PerkinElmer Life and Analytical Sciences; Prime-a-Gene Kit, Promega Corp.) and used to detect murine EL mRNA in Northern blot experiments. To detect murine LPL mRNA, we used a 700-bp PstI fragment from exon 10 of mouse LPL as a probe.

Real-time PCR. RT-PCR and quantitative real-time PCR were performed according to the manufacturer’s instructions (TagMan One-Step RT-PCR Kit; Applied Biosystems) on a 5700 ABI PRISM Sequence Detection System instrument (35, 36). Murine EL and β-actin TaqMan primers and probes were designed using Primer Express software version 2.0 (Applied Biosystems). Primers and probes were: EL: forward: 5′-TCTCGCATACCTTACACCCTGTC-3′, EL reverse: 5′-GCAAGGATGCGAAAGATTACTCTG-3′, EL probe: 5′-CCAGATGTCGCTTCCCTGT-3′, β-actin forward: 5′-GACAGGATGCAGAGTGT slider-3′, β-actin reverse: 5′-GCAACCGATACAGATG-3′, β-actin probe: 5′-CAAGATCATGCTCCTCTGTAGCGCA-3′. Sequence-specific EL amplification was detected by an increasing fluorescence signal of FAM (reporter dye) during the amplification cycles. Amplification of β-actin was performed on all samples as an internal control for variations in mRNA amounts. Levels of the different mRNA amounts were normalized to β-actin mRNA levels.

Tissue lipase assays. AT, SM, and CM from mice were excised, weighed, and minced with scissors. For assay of the LPL activity, the tissue samples were transferred to ice-cold tubes containing 1 ml of DMEM, 2% BSA (wt/vol), and 2 μl of heparin and incubated at 37°C for 1 hour. LPL activity was assayed as described previously (37). PL activity was assayed in a mixture of 1 μg [14C]dipalmitoyl-PC (NEN; PerkinElmer Life and Analytical Sciences) and 13.3 μl lecithin (1 mg/ml) in 50 μl substrate buffer Tris-TCNB (100 mM Tris-HCl pH 7.4, 1% Triton X-100, 5 mM CaCl2, 200 mM NaCl, 0.1% BSA). This mixture, termed the PC substrate, was vortexed for 2 minutes and then evaporated under a stream of nitrogen. The dried PL was redissolved in 300 μl Tris-TCNB and vortexed for 10 seconds. Cell extracts from AT (in 250 μl DMEM, 2% BSA, 2 U/ml of heparin) were incubated at 37°C for 1 hour. To inhibit EL enzyme activity, we incubated AT cell extracts with 50 μg anti-EL IgG for 1 hour at 4°C (28). Subsequently, the conditioned media were incubated with 10 μl substrate at 37°C for 1 hour. The reactions were terminated by addition of 25 μl 1 N HCl, and lipids were extracted with 1 ml hexane/isopropanol (3:2, vol/vol; 0.1% HCl). Five hundred microliters of the upper phase were dried and reconstituted in 100 μl hexane/isopropanol (3:2, vol/vol). After separation by thin-layer chromatography (hexane/diethyl ether/acetonic acid, 70:29:1), the liberated [14C]FFAs were quantitated by scintillation counting.

HDL labeling with [14C]PC. HDL was isolated by ultracentrifugation in an SW41 rotor (38). The protein concentration was determined by a Protein Assay Kit (Bio-Rad Laboratories). Labeling of HDL was performed with 4 μg [14C]dipalmitoyl-PC labeled in the sn-1 and sn-2 positions. [14C]PC was dried under a gentle stream of nitrogen. After the addition of 4 mg HDL-associated protein/ml, the mixture was overlaid with argon and incubated overnight at 37°C under continuous shaking. Labeled HDL was separated from unincorporated [14C]PC and desalted by exclusion chromatography on PD-10 columns (Amersham Biosciences) using PBS as eluent.

Incubation of isolated fat pads with [14C]PC-labeled HDL. Fat pads were removed surgically, washed with DMEM, and incubated with [14C]dipalmitoyl-PC labeled HDL (50 μg HDL-associated protein/sample) in DMEM containing 10% lipoprotein-deficient serum (LPDS). To inhibit EL enzyme activity, we preincubated fat pads with 50 μg EL antibody for 1 hour at 4°C. To inhibit enzyme activity of both EL and LPL, we preincubated fat pads with 50 μg/ml of the lipase inhibitor THL (25) for 1 hour either in the absence or in the presence of 2 U/ml heparin. All incubations were performed in a total volume of 150 μl at 37°C for 6 hours. Subsequently, the fat pads were washed twice with PBS, dried, and weighed, and the lipids were extracted with 3 ml chloroform/methanol (2:1, vol/vol) on a rotary shaker for 30 minutes. The fat pads were removed, and the liquid was evaporated under a stream of nitrogen. The lipids were redissolved in 100 μl of chloroform and separated by thin-layer chromatography (hexane/diethyl ether/acetonic acid, 70:29:1). The TG-specific spots were excised, and the incorporated [14C]FFAs were quantitated by scintillation counting.

Incubation of isolated fat pads with [3H]FFAs. Fat pads were removed surgically, washed with DMEM, and incubated with 5 μCi [3H]oleate (NEN, PerkinElmer Life and Analytical Sciences) and 200 μM nonradioactive oleate

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**Figure 6**
Uptake and incorporation of albumin-bound FFAs by AT. Fat pads of mice in the presence (L2) or absence (L0-MCK) of LPL were incubated with DMEM/1% FFA-free BSA/200 μM oleate/[3H]oleate (5 μCi/ml) at 37°C for 6 hours. Results are mean ± SD from 3 mice of each genotype performed in triplicate. ***P ≤ 0.001.
in DMEM containing 1% FFA-free BSA for 6 hours at 37°C. After incubation, the fat pads were washed twice in PBS and dried, and the lipids were extracted with 1 ml chloroform/methanol (2:1, vol/vol) on a rotary shaker for 30 minutes. After evaporation of the solvent under a stream of nitrogen, lipids were redissolved in 100 μl chloroform and separated by thin-layer chromatography (hexane/diethyl ether/acetic acid, 70:29:1). The TG-specific spots were excised, and the incorporated [3H]FFAs were quantitated by scintillation counting.

Statistics. Results are expressed as mean ± SD. Two-tailed Student’s t test was used to calculate statistical significance among groups. Differences at P < 0.05 were considered to be statistically significant.

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