Diet-induced obesity and its serious consequences such as diabetes, cardiovascular disease, and cancer are rapidly becoming a major global health threat. Therefore, understanding the cellular and molecular mechanisms by which dietary fat causes obesity and diabetes is of paramount importance in order to identify preventive and therapeutic strategies. Increased dietary fat intake results in high plasma levels of triglyceride-rich lipoproteins (TGRL). Tissue uptake of TGRL has been shown to promote glucose intolerance. We generated mice with an adipocyte-specific inactivation of the multifunctional receptor LDL receptor–related protein–1 (LRP1) to determine its role in mediating the effects of TGRL on diet-induced obesity and diabetes. Knockout mice displayed delayed postprandial lipid clearance, reduced body weight, smaller fat stores, lipid-depleted brown adipocytes, improved glucose tolerance, and elevated energy expenditure due to enhanced muscle thermogenesis. We further demonstrated that inactivation of adipocyte LRP1 resulted in resistance to dietary fat–induced obesity and glucose intolerance. These findings identify LRP1 as a critical regulator of adipocyte energy homeostasis, where functional disruption leads to reduced lipid transport, increased insulin sensitivity, and muscular energy expenditure.

Introduction
The chronic consumption of meals rich in fat and carbohydrates is a major causative factor of obesity and diabetes. The prevailing view on the mechanism by which these dietary factors contribute to obesity and diabetes is that when energy intake surpasses expenditure, the excess calories are deposited as fat in adipose tissues and its subsequent mobilization to nonadipose tissues causes insulin resistance that ultimately leads to type 2 diabetes (1–3). High fat and carbohydrate intake also leads to plasma lipid abnormalities, including high plasma levels of nonesterified fatty acids (NEFAs) and triglyceride-rich lipoproteins (TGRL) as well as reduced plasma levels of HDL (2, 3). Numerous past studies have shown that elevated plasma NEFA levels directly induce insulin resistance and thus play a causative role in the pathogenesis of obesity-related diabetes (1, 4, 5). However, only sporadic attention has been paid to the role of TGRL in obesity and diabetes. The TGRL are generally thought of primarily as triglyceride carriers in the circulation, delivering substrates to tissues where lipoprotein lipase–catalyzed hydrolysis liberates NEFAs prior to their uptake by cells through CD36 and other pathways (6). It is important to note that TGRL as well as lipase-hydrolyzed TGRL remnants can also be internalized by cells directly via whole-particle uptake and provide triglyceride-derived fatty acids via mechanisms mediated by LDL receptor family member proteins (7). Results showing that Apoe−/− mice with defects in receptor-mediated tissue uptake of TGRL are less susceptible to diet-induced obesity and diabetes suggested the importance of TGRL whole-particle uptake in diet-induced obesity and diabetes (8). Indeed, a recent study reporting that TGRL uptake by skeletal muscle cells, independent of NEFA uptake, also directly modulates glucose metabolism and insulin sensitivity is supportive of this hypothesis (9).

One mechanism by which apoE modulates diet-induced obesity and diabetes is through its role in lipid transport and lipid redistribution among different cells and tissues. This process is dictated by the expression level and distribution of various apoE-binding receptors on tissues such as the liver, muscle, heart, and adipose tissues. The 3 apoE binding receptors prominently expressed in these tissues are LDL receptor, VLDL receptor, and LDL receptor–related protein–1 (LRP1). Although these receptors share the overlapping functions of binding and internalizing apoE-containing lipoproteins, there are distinct differences in preference for various lipoproteins. The LDL receptor binds remnants of chylomicrons and VLDL as well as LDL but interacts poorly with nascent chylomicrons and VLDL (10–12). The VLDL receptor appears to prefer intermediate-density lipoprotein (IDL) (13) and immobilizes LpL on the cell surface for hydrolysis of the triglycerides prior to fatty acid uptake into cells (14). In contrast, LRP1 prefers to interact with apoE-enriched chylomicron and VLDL remnants (15, 16). The latter receptor is also responsible for LpL-mediated endocytosis of whole lipoprotein particles (17–19). Thus, whereas VLDL receptor expression in high-energy metabolism tissues may modulate sensitivity to diet-induced obesity and diabetes via LpL-catalyzed fatty acid uptake (14, 20), the high level of LRP1 expression in adipocytes (21) suggests that LRP1 may partner with apoE and/or LpL in mediating the effects of TGRL in diet-induced obesity. In view of studies establishing the importance of adipose tis-
In controlling glucose homeostasis and energy balance (22–27), we hypothesized that adipocyte expression of LRPI is also a key modulator of glucose and energy homeostasis. Adipose-specific LRPI-knockout mice were produced to test these hypotheses.

Figure 1
Characteristics of 2 independently generated ad-Lrp1+/+ and ad-Lrp1−/− mouse colonies. Adipose tissue–specific LRPI gene inactivation was performed both on a mixed background (Dallas) and on an inbred C57BL/6 background (Cincinnati) by crossing ap2 Cre transgenic mice with Lrp1floxflox mice. (A) Detection of the Cre transgene in mice with the mixed background by PCR. A fragment of 400 bp was amplified from mouse tail genomic DNA when the Cre transgene driven by the ap2 promoter was present. A 200-bp fragment amplified from the ap2 gene was observed in both PCR products as control. (B) Detection of wild-type (+/+ ) and mutant (−/− ) mRNA of mice (mixed background) by quantitative real-time PCR. (C) Western blot identification of the β subunit of LRPI in epididymal WAT (eWAT), BAT, and brain of ad-Lrp1+/+ and ad-Lrp1−/− mice (mixed background). β-Actin was used as loading control. (D) Ponceau S staining and Western blot of the β subunit of LRPI in the stromal-vascular cells (sv) and mature adipocytes of subcutaneous (sWAT) and eWAT adipose tissue in ad-Lrp1+/+ and ad-Lrp1−/− mice. (E) Western blot of the β subunit of LRPI in BAT (lanes 1 and 2), epididymal WAT (eWAT) (lanes 3 and 4), liver (lanes 5 and 6), skeletal muscle (lanes 7 and 8), and peritoneal macrophages (lanes 9, 10) of ad-Lrp1+/+ (lanes 1, 3, 5, 7, and 9) and ad-Lrp1−/− (lanes 2, 4, 6, 8, and 10) mice (inbred background). Molecular size markers (Mr) were applied to the lane flanking the sample lanes as indicated.

Results
Adipose-specific LRPI inactivation was achieved independently at both the University of Cincinnati and University of Texas Southwestern Medical Center by crossing Lrp1floxflox mice (28) with ap2-Cre

Figure 2
Lipid transport to adipose tissue is impaired in ad-Lrp1−/− mice. (A) Fasting plasma cholesterol and triglyceride levels were similar in ad-Lrp1+/+ (open bars) and ad-Lrp1−/− (black bars) mice fed basal chow diet in both inbred and mixed-strain backgrounds. (B) Plasma triglyceride levels in ad-Lrp1−/− (open squares) and ad-Lrp1+/+ (filled circles) mice after an oral lipid load (inbred background). The inset shows the area under the curve (AUC) analysis of the data. *P < 0.05; n = 10.
transgenic mice generated in either a homogenous C57BL/6 background (23) or in an outbred strain background (Figure 1A) to yield ad-Lrp1+/− mice with either a homogenous C57BL/6 background or a mixed C57BL/6 and 129 background. In both colonies, the Lrp1 mRNA level was reduced by approximately 50% in white adipose tissue (WAT) and by approximately 85% in brown adipose tissue (BAT) of ad-Lrp1+/− mice compared with their respective ad-Lrp1+/+ littermates (Figure 1B). Western blot analysis of LRP1 protein levels in WAT and BAT was consistent with the mRNA expression levels (Figure 1C). When the WAT from subcutaneous (sWAT) and epididymal (eWAT) fat depots of ad-Lrp1+/+ and ad-Lrp1−/− mice were fractionated into mature adipocytes and stromal vascular cells, LRP1 expression in WAT of ad-Lrp1−/− mice was found to be due to its presence at normal levels in the stromal vascular cells (Figure 1D). In contrast, LRP1 expression in mature adipocytes was decreased by greater than 90% in subcutaneous fat depot and more than 70% in epididymal fat depot of ad-Lrp1−/− mice (Figure 1D). Expression of LRP1 in brain, liver, muscle, and macrophages was not affected in the ad-Lrp1−/− mice (Figure 1, C and E).

Comparison of ad-Lrp1+/+ and ad-Lrp1−/− mice revealed similar fasting plasma triglyceride and cholesterol levels when the animals were fed a standard low-fat chow diet (Figure 2A). However, postprandial lipid clearance from the circulation in response to a bolus fat load was markedly delayed in ad-Lrp1−/− mice compared with ad-Lrp1+/+ mice (Figure 2B). Since LRP1 binds apoE-containing lipoproteins as well as LpL, the delayed postprandial lipid clearance observed in ad-Lrp1−/− mice may be due to defective postprandial lipoprotein endocytosis or deficiency in cell-surface LpL that mediates triglyceride hydrolysis prior to cellular fatty acid uptake. These possibilities were explored initially by comparing total and heparin-extractable LpL activities in various tissues of ad-Lrp1+/+ and ad-Lrp1−/− mice. Results showed no significant difference in LpL activities in total homogenates of BAT and WAT from both subcutaneous and epididymal fat pads and in the muscle of ad-Lrp1+/+ and ad-Lrp1−/− mice (Figure 3). In contrast, cell-surface LpL, as measured by heparin-releasable LpL activity, was elevated significantly in both BAT and WAT (Figure 3). Although decreased heparin-releasable LpL activity was noted in muscle, the difference in heparin-releasable activities from muscle of ad-Lrp1+/+ and ad-Lrp1−/− mice did not reach statistical significance (Figure 3). Thus, the delayed postprandial triglyceride clearance in ad-Lrp1−/− mice was not due to defective expression of LpL.

The mechanism by which adipocyte LRP1 participates in postprandial lipid clearance were explored by experiments monitoring tissue distribution of dual radiolabels after infusing lipid emulsions containing [3H]cholesteryl oleate and [14C]triolein into ad-Lrp1+/+ and ad-Lrp1−/− mice. Results showed that uptake of 3H- and 14C-labeled lipids by adipose tissues in ad-Lrp1−/− mice was decreased to a similar extent, with the most dramatic decrease observed in BAT (Figure 4). Interestingly, despite the similar level of cholesteryl ester and triglyceride uptake by the liver of ad-Lrp1−/− mice.
and *ad-Lrp1*–/– mice, hepatic triglyceride levels were found to be lower in the *ad-Lrp1*–/– mice (Figure 4C). Triglyceride levels in muscle were similar in *ad-Lrp1*+/+ and *ad-Lrp1*–/– mice (Figure 4D).

The reduced lipid assimilation by adipocytes in the *ad-Lrp1*–/– mice resulted in lower body weight (BW) as compared with that observed in the *ad-Lrp1*+/+ mice in both inbred (Figure 5A) and mixed-strain backgrounds (Figure 5B). This BW difference was entirely due to smaller fat mass (Figure 5C), which was also consistent with lower plasma leptin levels observed in the mutant mice (526 ± 87 pg/ml vs. 3,126 ± 978 pg/ml in wild type; \( P < 0.05 \)). The reduction of fat mass in *ad-Lrp1*–/– mice was due to a remarkable and macroscopically obvious reduction of interscapular and epididymal fat pads observed in both C57BL/6 and mixed-strain background (Figure 6, A and B). Similarly, BAT mass was reduced by more than 65% in the *ad-Lrp1*–/– mice (Figure 6C). Histological analysis of fat tissues revealed smaller-sized and fewer lipid droplets in white and brown adipocytes in both lines of *ad-Lrp1*–/– mice compared with wild-type mice (Figure 6, D–F). The consis-
Figure 7
Increased food intake under normal chow–fed conditions and reduced fat mass after food deprivation in ad-Lrp1–/– mice. (A) Food intake data collected from inbred ad-Lrp1+/– (open squares) and ad-Lrp1–/– (filled circles) mice (n = 12). (B) Changes in fat as a percentage of BW in ad-Lrp1+/– (white bar) and ad-Lrp1–/– (black bar) mice on the mixed background after 24 hours of food deprivation (n = 16–18). *P < 0.05, **P < 0.005 compared with the ad-Lrp1+/– groups.

tency of these phenotypes in both inbred and outbred lines of ad-Lrp1–/– mice underscores the importance and highly penetrant role of LRP1 in the regulation of adipose tissue metabolism. Surprisingly, ad-Lrp1–/– mice were found to consume more food than the ad-Lrp1+/– mice despite their reduced BW and fat mass (Figure 7A). Moreover, when the animals on the mixed-strain background were fasted for 24 hours, an exaggerated loss of fat mass was observed in ad-Lrp1–/– mice in comparison with their ad-Lrp1+/– littermates (Figure 7B), which is indicative of an increase in the metabolic rate in the absence of adipocyte LRP1. Consolidating observations on the mixed background with additional studies conducted in the inbred population revealed that all inbred ad-Lrp1–/– mice exhibited difficulty in maintaining body temperature, with increased shivering and body surface reduction by cooling up that was visibly noticeable initially at the ambient environment of 22°C and was more pronounced at 4°C (see Supplemental Videos 1 and 2; supplemental material available online with this article; doi:10.1172/JCI31929DS1). This was further quantified using a multidimensional light beam system designed to analyze motor activity (Figure 8A). In contrast, ambulatory movement was not altered by LRP1 gene disruption (Figure 8B).

The potential impairment of BAT functions in ad-Lrp1–/– mice was further illustrated by results showing that ad-Lrp1–/– mice displayed similar body core temperature but significantly lower surface temperature than ad-Lrp1+/– mice when they were kept at an ambient temperature of 22°C (Figure 9). The difference in body surface temperature was further exaggerated when the animals were exposed to a 4°C environment (Figure 9). Interestingly, body core temperature also decreased much faster in ad-Lrp1–/– mice than ad-Lrp1+/– mice when the ambient temperature was lowered (Figure 9).

The increase in muscle shivering activity, decrease in body surface temperature, and the rapid rate of cold-induced core temperature decline may reflect a reduction in resting thermogenesis. Therefore, we examined age-matched ad-Lrp1+/– and ad-Lrp1–/– mice in an indirect calorimeter to test this hypothesis. While the respiratory quotient was found to be similar (Figure 8C), the mutant mice displayed substantially higher oxygen consumption and energy expenditure (Figure 8D). The increased energy expenditure in mice specifically lacking LRP1 in adipocytes was not due to differences in uncoupling protein–1 (UCP-1) expression in BAT or UCP-2 and UCP-3 expression in muscle (Figure 8E). Thus, the unexpected increased energy expenditure in ad-Lrp1–/– mice, which could explain their lower fat mass, may reflect at least in part a potential compensatory increase in muscular contractile activity due to the dramatic decrease of BAT.

Numerous mouse models with modest reduction in adiposity and lower leptin levels have been shown to be more insulin sensitive with improved glucose metabolism in vivo (22, 29–31). However, other mouse models with reduced adipose tissues such as PPARγ2-knockout mice have impaired insulin sensitivity (32). Therefore, we tested the effects of reduced adipose tissue mass and leptin levels in glucose homeostasis in the ad-Lrp1–/– mice. Results showed that the ad-Lrp1–/– mice have markedly lower fasting glucose and insulin levels compared with wild-type mice (Figure 10A). Lower fasting NEFA levels were also observed in inbred ad-Lrp1–/– mice (Figure 10A). Glucose tolerance tests showed that the mutant mice also displayed an improved response to glucose challenge (Figure 10B). Tissue distribution of 2-deoxy-[3H]glucose revealed that the skeletal muscle was responsible for the elevated plasma glucose clearance in ad-Lrp1–/– mice (Figure 10C). This observed increase in glucose uptake by skeletal muscle of ad-Lrp1–/– mice is consistent with the increased muscular activity in these animals.

The reduced adipose tissue mass along with increased metabolic rate and improved glucose tolerance in ad-Lrp1–/– mice suggested that adipose-specific LRP1 inactivation may be protective against diet-induced obesity and diabetes. This hypothesis was examined by placing ad-Lrp1+/– and ad-Lrp1–/– mice on a high-fat diet containing 35.8% fat. Results showed that both ad-Lrp1+/– and ad-Lrp1–/– mice displayed reduced respiratory quotients under the high-fat feeding conditions and that the difference in energy expenditure between ad-Lrp1+/– and ad-Lrp1–/– mice was exaggerated (Figure 11, A and B). It is likely that the relative decrease in dietary carbohydrates further impairs the thermogenic efficiency in ad-Lrp1+/– mice, with both BAT and muscle metabolism even more dependent on the ability to use lipids. Importantly, the substantially elevated energy expenditure in ad-Lrp1–/– mice prevented most of the high-fat diet–induced BW gain and adiposity observed in their ad-Lrp1+/– littermates (Figure 11, C and D). Finally, in accordance with their protection against diet-induced obesity — and in contrast to wild-type mice, which displayed elevated fasting plasma glucose and insulin levels upon high-fat feeding — plasma glucose and insulin levels in the ad-Lrp1+/– mice remained at the normal level after feeding the high-fat diet for 16 weeks (Figure 12, A and B).

Glucose tolerance in high-fat diet–fed ad-Lrp1–/– mice was also similar to that observed in chow-fed mice, while ad-Lrp1+/– mice displayed delayed glucose clearance (Figure 12C).

Discussion
Here we showed that in the absence of LRPI, TGRL assimilation is impaired in adipose tissue, resulting in decreased fat mass. The data also revealed that glucose metabolism and energy balance are enhanced in ad-Lrp1–/– mice. The metabolic alterations we have reported with male mice were also evident, albeit less robust, in female mice (data not shown). Furthermore, these phenotypic manifestations due to the loss of LRPI expression in adipose tissues were essentially identical in mice on a homogeneous C57BL/6 background as well as on a mixed-strain background, thus documenting the importance and highly penetrant role of adipocyte LRPI in the regulation of adipocyte tissue...
The aP2 gene promoter commonly used to generate fat tissue–specific knockout and transgenic mouse models (25, 26, 33) was used in the current study to drive Cre recombinase transgenic expression in \( Lrp1^{\text{flx/flx}} \) mice for the generation of adipose-specific LRP1–knockout mice. The aP2-Cre recombinase transgene had no effect on LRP1 expression in brain, liver, and macrophages. However, LRP1 expression levels were reduced by approximately 85% in BAT and by approximately 75%–90% in mature adipocytes of \( ad-Lrp1^{+/–} \) mice compared with those observed in \( ad-Lrp1^{+/+} \) mice. These findings are consistent with the high LRP1 expression levels reported in nondifferentiated 3T3-L1 preadipocytes and murine embryonic fibroblasts (34, 35) and the expected expression of the aP2 promoter–driven Cre recombinase transgene during the late phase of adipocyte differentiation (36).

Results of the current study revealed the importance of adipocyte LRP1 in postprandial TGRL metabolism. Previously, the rate of plasma triglyceride accumulation after a high-fat meal was thought to be dictated by the rate of LpL-mediated triglyceride hydrolysis and the subsequent cellular uptake of the NEFAs (37), as well as receptor-mediated uptake of remnant lipoproteins (38, 39). Turnover studies have shown that the liver is the main site of chylomicron remnant clearance and that this process is mediated by the LDL receptor and LRP1 (38–40). Both receptors are also expressed almost ubiquitously in other tissues. Results of the current study showed that the absence of LRP1 in adipose tissues of \( ad-Lrp1^{+/–} \) mice resulted in delayed postprandial TGRL clearance as well as decreased uptake of both cholesterol esters and triglycerides from lipid emulsions injected into the circulation. Interestingly, adipose tissue uptake of both lipid moieties was
decreased to a similar extent even though heparin-releasable LpL was elevated in adipose tissues of ad-Lrp1−/− mice. Taken together, these results indicated that LRP1 is important in mediating both the lipolytic and the endocytic processes responsible for TGRL catabolism in adipocytes. This interpretation is consistent with previous in vitro data showing a role of LRP1 in mediating LpL binding and lipolysis of TGRL at the cell surface (41), as well as its well-documented role in binding and uptake of LpL- and/or apoE-enriched lipoproteins (15, 21, 42). It is noteworthy, however, that fasting triglycerides were maintained at the normal levels in ad-Lrp1−/− mice despite their delay in clearance of postprandial TGRL. These data are supportive of previous reports demonstrating that LRP1 is important for clearance of the intestinally derived apoB48-containing lipoproteins but not the catabolism of endogenously derived apoB100-containing lipoproteins (43).

In addition to the delay in postprandial TGRL clearance and alteration in lipid delivery to target tissues, the ad-LRP1−/− mice were also shown to have an overall decrease in fat mass and were protected from high-fat diet–induced glucose intolerance and obesity. Previous studies have clearly established the role of BAT in energy expenditure through thermogenesis in the form of heat dissipation (44, 45). Hence, the lower body-surface temperature (less heat dissipation) and increased sensitivity to cold-induced body core temperature decline observed in ad-Lrp1−/− mice are consistent with the reduced BAT mass observed in these animals. Although it has been shown that a complete absence of BAT function is associated with obesity (46), the ad-Lrp1−/− mice described here are more resistant to diet-induced obesity and more glucose tolerant than wild-type mice. We concluded that the different phenotype of the ad-Lrp1−/− mice is due to loss of a specific function in all fat depots in contrast to the effect of complete BAT ablation. Similarly, the metabolic phenotypes of the ad-Lrp1−/− mice are also different from those of the Cidea-deficient mice, which are resistant to diet-induced obesity and diabetes but are more resistant to cold-induced body core temperature drop (47).

The requirement for increased muscular activity in body core temperature maintenance in the ad-Lrp1−/− mice was supported by observations of their muscle shivering, especially during the light phase, when the animals rest and ambulatory activity is minimal. Such an alternative heating mechanism is substantially less effi-

Figure 9
Cold adaptation studies in ad-Lrp1+/+ and ad-Lrp1−/− mice. (A) Heat production visualized by a high-resolution infrared camera from 6 mice in each group. (B) Body core and surface temperature measurements from ad-Lrp1+/+ and ad-Lrp1−/− mice (n = 10). *P < 0.05, **P < 0.005, ***P < 0.001.
cient and requires increased caloric combustion. This interpretation of the data is also in line with the increased food intake of these mice. The increased oxygen consumption and energy expenditure displayed by ad-Lrp1−/− mice, with their reduced BAT mass and impaired heat dissipation mechanism, is consistent with this hypothesis. These results are also consistent with the concept of muscle shivering as the first-order compensatory mechanism for impaired BAT functions in body temperature maintenance (45).

These metabolic phenotypes observed in the ad-Lrp1−/− mice, with alterations in the use of fat versus muscle tissues for body core temperature maintenance, closely parallel the metabolic phenotypes of Plin−/− mice with targeted inactivation of the perilipin gene (48). The Plin−/− mice displayed constitutively active hormone-sensitive lipase activity in adipocytes, resulting in less fat storage and smaller adipocytes. Interestingly, the Plin−/− mice also consumed more food, displayed increased sensitivity to the cold environment, and were more resistant to diet-induced obesity (48). Importantly, indirect calorimetry measurements revealed increased oxygen consumption by Plin−/− compared with wild-type mice. Moreover, muscle mass was also increased in the Plin−/− mice (48). Thus, the Plin−/− mice with defective lipid storage potential in both WAT and BAT also required increased energy utilization by the muscle to compensate for their ineffective BAT thermogenic mechanism in maintenance of body core temperature. Taken together, the metabolic phenotypes of ad-Lrp1−/− and Plin−/− mice indicate that poor lipid storage in brown and white adipocytes, due to defective lipid uptake and storage mechanisms, respectively, results in shifting from BAT thermogenesis to the use of a less energy-efficient muscular activity mechanism to maintain body core temperature. This shift of tissue energy metabolism is protective against diet-induced obesity.

In addition to the reduction in BAT and WAT, the ad-Lrp1−/− mice also displayed a marked improvement in whole-body glucose disposal compared with the ad-Lrp1+/+ mice. The determination of glucose uptake by individual tissues revealed that the skeletal muscle is the major tissue responsible for the elevated plasma glucose clearance. This difference in glucose uptake between skeletal muscle of ad-Lrp1+/+ and ad-Lrp1−/− mice may reflect the increase in energy demand for muscle thermogenesis and the overall decrease in adipose fat storage in ad-Lrp1−/− mice. Thus, this study showed that inactivation of adipose LRP1 activity not only reduces fat storage in both WAT and BAT and protects against diet-induced obesity, but that it also protects against glucose intolerance and diabetes due to increased energy expenditure shifting from brown adipose thermogenesis to muscular activity for maintenance of body temperature.

In summary, this study documents the prominent role of adipocyte LRP1 in modulating energy metabolism and sensitivity to diet-induced obesity and diabetes. These results add an impor-
tant perspective to the currently established models in showing the unique importance of various LDL receptor family proteins in these metabolic processes. Previously, VLDL receptor expression in adipocytes was shown to play a role in adiposity by modulating LpL-mediated lipid uptake (14). Additionally, LRP5 was shown to participate in glucose-induced insulin secretion by islets, with its defective expression causing impaired glucose tolerance (49). More recently, LRP6 mutations in humans have been implicated in the metabolic syndrome consisting of hyperlipidemia, hypertension, and diabetes via impairment of the Wnt signaling cascade (50). These continued investigations aimed at identifying the functional roles of each of the LDL receptor family proteins expressed in various tissues will add further insights into lipid transport–dependent and –independent functions of these receptor proteins in physiology and pathophysiology of metabolic diseases.

**Methods**

*Generation of ad-Lrp1–/– mice.* Two lines of adipose tissue–specific LRPI-knockout mice were generated independently at the University of Cincinnati College of Medicine and the University of Texas Southwestern Medical Center. For the generation of the ad-Lrp1–/– mouse strain in Cincinnati, adipose-specific Cre recombinase transgenic mice in a C57BL/6 background were obtained from Ronald M. Evans (Salk Institute, La Jolla, California, USA) (23) and crossbred with C57BL/6 Lrp1floxflox mice (28). Mouse genotype was confirmed by PCR using previously described primers and thermocycler conditions (23, 28). The ad-Lrp1–/– mouse colony in the Dallas laboratory was obtained by crossing aP2-Cre transgenic mice with Lrp1floxflox mice. These ad-Lrp1–/– mice were maintained on a mixed B6/129 background. Genotyping was performed by PCR using 3 primers with the following sequences: sense: 5′-GGAAATGATCTGGCCCCCATTGGTCACTCC-3′; 2 antisense: 5′-CACTGAGACAAGCTTCCAGGTTCCCACAA-3′ in aP2 gene and 5′-CGCGAACATCTTCAGGTTCTGCGGGAAACC-3′ in Cre transgene. PCR amplification began with a 1-minute denaturation step at 94°C and then 30 cycles of denaturation at 94°C for 15 seconds, annealing at 60°C for 1 minute, and extension at 72°C for 1 minute. The products were further analyzed by agarose gel electrophoresis to confirm the size.

Animals were housed in specific pathogen–free facilities with a 12-hour light/12-hour dark cycle. Animals were fed basal rodent chow 5058 PicoLab Mouse Diet 20 (9% fat by weight; LabDiet) or a high-fat diet containing 35.8% fat by weight (D12330; Research Diets), and age-matched male littersmates were used for all experiments. All experimental protocols were approved by the Institutional Animal Care and Use Committees of the University of Cincinnati and University of Texas Southwestern Medical Center.

*Preparation of mature adipocytes and stromal-vascular cell fractions from fat tissue.* Epididymal and subcutaneous fat pads were isolated from mice by dissection (n = 3) and washed in Krebs-Ringer-bicarbonate buffer (pH 7.4). The tissues were incubated with 0.1% collagenase (Worthington) on a shak-
B

A

FBS prior to homogenization in protease inhibitor–containing buffer for the differences observed in the presence or absence of 1 M NaCl. in tissue homogenates and in the heparin-released supernatants were determined based on the hydrolysis of the fluorogenic substrate 4-methylumbelliferone (Sigma-Aldrich) in the presence or absence of 1 M NaCl, as indicated before and after injection. At the 5-minute time point, mice were perfused with cold saline, tissues removed, and plasma decay and tissue counts measured.

LpL activity determinations. Total cell extracts were prepared by homogenization 90 mg of tissues in buffer containing 20 mM Tris-HCl, pH 8.0, 1 mM EDTA, and 300 μM sodium taurocholate. Cell-surface LpL was isolated from the minced tissues by collecting supernatants from centrifugation of samples for 10 minutes at 20,000 g in a benchtop microfuge after incubation for 1 hour at 37°C in buffer containing 10 mM Tris-HCl, pH 7.4, 10 U/ml heparin, and 0.5% heat-inactivated mouse serum. Total lipolytic activities in tissue homogenates and in the heparin-released supernatants were determined based on the hydrolysis of the fluorogenic substrate 4-methylumbelliferone (Sigma-Aldrich) in the presence or absence of 1 M NaCl, as described previously (52). LpL-specific lipolytic activity was calculated from the differences observed in the presence or absence of 1 M NaCl.

Body fat measurements. Body fat mass was measured in conscious mice (n = 16) using 1H magnetic resonance spectroscopy (EchoMRI-100; Echomagnetic Systems). BW measurements were performed monthly for animals maintained on chow diet and weekly for animals maintained on the high-fat diet. After 1 week of adaptation, food intake (in grams of food) of individual mice was recorded daily for a week. For the food deprivation experiment, mice were fasted for 24 hours with water ad libitum. The ad-Lrp1+/+ control group consisted of 16 mice, and the ad-Lrp1−/− experimental group had 18 mice. A BRUKER Minispec Time Domain Nuclear Magnetic Resonance (TD-NMR) analyzer was adopted to analyze fat/lean components of these live mice. The percentage of body fat was measured automatically by the TD-NMR analyzer. The reduction in the percentage of body fat after 24 hours of fasting was used as an indicator of fat loss.

Energy balance measurements. Animals (n = 16) were allowed to acclimate to respiratory chambers for a period of 2 days. Subsequently, energy expenditure (EE), respiratory quotient (RQ), food intake, and locomotive activity were measured simultaneously over 24-hour periods for 4 consecutive days using a customized 32-cage Indirect Calorimetry System combined with a Drinking and Feeding Monitor and TSE ActiMot system (TSE-Systems). The volume of oxygen consumed (VO2 [ml/(kg × h)]) and carbon dioxide produced (VCO2 [ml/(kg × h)]) were measured to calculate RQ (ratio of CO2 produced to VO2 consumed) and EE (EE = [3.815 + 1.232 × RQ] × VO2 × 0.001 [kcal/(kg × h)]).

Fine and locomotive activities. The TSE Indirect Calorimetry System was customized to include a multidimensional light beam and light sensor system, which records and localizes light beam breaks within the entire home cage environment of each mouse. The parallel beams were set up close enough so that no mouse could be in the calorimeter cage without at least 1 light beam being broken by its body. The motor activity-monitoring software of this system was programmed to interpret breaking of consecutive beams of the lower level of the beam grid to indicate ambulatory movement. Recorded breaks of light beams within a second beam grid located in the upper half of the home cage would be interpreted as rearing behavior, which often correlates with feeding from hanging feeders or climbing activities. Finally, rapid consecutive breaks and reconnection of the same light beams were interpreted as stationary motor activity or fidgeting. The recordings of such events were used to indicate shivering or fidgeting behavior.

Cold adaptation studies. Ten ad-Lrp1+/+ and 10 ad-Lrp1−/− mice were placed in a cold-temperature environment of 4°C and body core temperature measured using a rectal thermometer (Physitemp). Heat production in a subset of 6 mice from each group was visualized using a high-resolution infrared camera (FLIR PM280; FLIR Systems) that applies radiometry technology to convert infrared energy (radiant heat) into 12-bit dynamic range color images. The images were captured from the backs of the mice before and 1 hour after exposure to the 4°C environment. Infrared thermography images were taken from the upper half of the body to specifically visualize heat production from the BAT.

Glucose tolerance and tissue uptake. Blood samples were collected immediately before and at 15, 30, 60, and 120 minutes after intraperitoneal injection of glucose (2 g/kg BW) into a 6-hour-fasted animal. Tissue uptake of glucose was monitored by intravenous injection of glucose (1 g/kg BW) along with 2-deoxy-[1H]glucose (0.5 μCi/g BW; NEN Life Science Products) after a 6-hour fast. Animals were euthanized after 15 minutes, extensively perfused with buffered saline, and tissues were excised, weighed, and solubilized with NCS-R II Tissue Solubilizer (Amersham).

Figure 12

Improved glucose metabolism in inbred ad-Lrp1+/+ mice fed a high-fat diet for 16 weeks. Fasting plasma glucose (A) and insulin levels (B) in high fat diet–fed ad-Lrp1+/+ (white bars) and ad-Lrp1−/− (black bars) mice. (C) Plasma glucose levels in inbred ad-Lrp1+/+ (open squares) and ad-Lrp1−/− (filled circles) mice after 6 hours fasting followed by intraperitoneal injection of glucose (2 g/kg BW). Integration of the data (AUC) is shown in the inset in C. Data represent mean ± SEM from 12 mice in each group. *P < 0.05, **P < 0.005.
Radioactivity in each sample was quantified by liquid scintillation counting. Data are reported as total 2-deoxy[3H]glucose radiolabel incorporation per milligram of tissue weight.

**Blood chemistry.** Animals were fasted for 6 hours prior to blood collection from the tail vein. Plasma triglyceride and cholesterol levels were determined using the Infinity Triglyceride and Cholesterol Kits (Thermo Electron Corp.). Blood glucose was determined with an Accu Check Active Glucometer (Roche Applied Science). Plasma insulin and leptin levels were measured in individual samples with multiplexed biomarker immunoassays for Luminox xMAP (Linco).

**Histology.** Tissues were fixed by immersion in neutral buffered formalin (10%), dehydrated in ethanol and then transferred to xylene solution for embedding in paraffin. Five- and 8-μm sections were stained with H&E.

**Quantitative real-time PCR.** Total RNA from WAT (epididymal) and BAT was obtained using RNA STAT-60 (TEL-TEST Inc.). The extracted RNA was treated with DNase (Ambion) before reverse transcription (Applied Biosystems). SYBR Green PCR Master Mix (Applied Biosystems) was used for quantitative real-time PCR. The sequences of 2 primers for LRP1 were: sense, 5′-GACCCAGGTGTTGACACAGATG-3′; antisense, 5′-AGTCGTTGTCCTCGTACACTTC-3′. GAPDH was used as a control to normalize the amount of RNA sample. The sequence of its sense primer was 5′-TGTGTTGCTGTTGAGATCGA-3′, and the antisense one was 5′-GCCTGACACCACCTTCTGTAG-3′.

**Immunoblot analysis.** The expression of LRP1 in mice from Dallas was analyzed by preparing membrane fractions from isolated white (epididymal) and brown fat pads. Briefly, the adipose pads were homogenized in buffer A (20 mM Tris-HCl, pH 8.0, 1 mM CaCl2 and 150 mM NaCl) plus fresh protease inhibitors (P8430, Sigma-Aldrich) followed by centrifugation at 300 g for 5 minutes at 4°C. The supernatant was then centrifuged at 150,000 g for 30 minutes at 4°C. The membrane pellet was resuspended in buffer B (50 mM Tris-HCl, pH 8.0, 2 mM CaCl2, 80 mM NaCl, 1% Triton X-100). After another centrifugation at 100,000 g for 30 minutes at 4°C, the supernatant was collected for immunoblotting. The membrane extract was resolved on SDS–polyacrylamide gel and transferred to nitrocellulose membranes (Hybond-C Extra; Amersham Biosciences). The blot was blocked with 5% skim milk, probed with rabbit anti-LRP1 antibody, and incubated with HRP-conjugated goat anti-rabbit secondary antibody (Amersham Biosciences). Immunoreactive bands were visualized using a chemiluminescence detection kit (Amersham Biosciences).

Immunoblot analysis conducted in Cincinnati was performed by enriching LRPI in tissue extracts by precipitation with recombinant receptor-associated protein (RAP) conjugated to glutathione-S-transferase in the presence of glutathione beads. The precipitated proteins were separated by SDS-PAGE, transferred to polyvinylidene fluoride paper, and blotted with antibodies against the β subunit of LRPI (1H4; ATCC). Expression of UCP-1, -2, and -3 in tissues was analyzed by immunoblotting tissue homogenates with UCP-1 and UCP-2 antibodies obtained from Calbiochem and UCP-3 antibodies from Alpha Diagnostics. Immunoreactive proteins were detected by incubating the blots with fluorescently labeled species-specific secondary antibodies (Molecular Probes; Invitrogen) and visualized by the Odyssey Infrared Imaging System (Li-Cor Biosciences).

**Statistics.** Numerical data are expressed as mean ± SEM. An unpaired 2-tailed Student’s t test was used for analysis of 1-group variables. A P value of less than 0.05 between the groups was considered significant.

**Acknowledgments.** We are grateful to Ronald Evans for providing the aP2-Cre recombinase transgenic mice for this study. This research was supported by grants DK74932 (to D.Y. Hui), DK069987 and DK056863 (to M.H. Tschoj), and HL20948 and HL63762 (to J. Herz) from the NIH. S.M. Hofmann is the recipient of a Scientist Development Award (0635079N) from the National Center of the American Heart Association. D. Gilham is the recipient of a Post-Doctoral Fellowship (0625329B) from the Ohio Valley Affiliate of the American Heart Association. J. Herz is an Investigator of the American Heart Association.

Received for publication February 22, 2007, and accepted in revised form August 1, 2007.

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