Modulation of myocellular fat stores: lipid droplet dynamics in health and disease

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STORAGE OF FAT IN WHITE ADIPOSE tissue (WAT) is the most widely studied form of triacylglycerol (TAG) storage. In periods of caloric excess, adipose tissue mass expands by increasing TAG content. During starvation, hydrolysis of TAG results in mobilization of fatty acids to meet the organism’s energy requirement and TAG content in adipocytes declines. Stored TAG, hence, represents a dynamic pool of fatty acids. Whereas adipose tissue is by far the largest storage site of fatty acids as TAG, subcellular TAG-containing structures—referred to as lipid droplets (LD)—are also present in other tissues. Until recently, LD were considered inert storage sites of energy dense fats. Nowadays, however, LD are increasingly considered dynamic functional organelles involved in many intracellular processes like lipid metabolism, vesicle trafficking, and cell signaling. Next to TAG, LD also contain other neutral lipids such as diacylglycerol. Furthermore, LD are coated by a monolayer of phospholipids decorated with a variety of proteins regulating the delicate balance between LD synthesis, growth, and degradation. Disturbances in LD-coating proteins may result in disequilibrium of TAG synthesis and degradation, giving rise to insulin-desensitizing lipid intermediates, especially in insulin-responsive tissues like skeletal muscle. For a proper and detailed understanding, more information on processes and players involved in LD synthesis and degradation is necessary. This, however, is hampered by the fact that research on LD dynamics in (human) muscle is still in its infancy. A rapidly expanding body of knowledge on LD dynamics originates from studies in other tissues and other species. Here, we aim to review the involvement of LD-coating proteins in LD formation and degradation (LD dynamics) and to extrapolate this knowledge to human skeletal muscle and to explore the role of LD dynamics in myocellular insulin sensitivity.

adipose triglyceride lipase; hormone-sensitive lipase; perilipin adipocyte differentiation-related proteins; muscle vesicle trafficking and cell signaling (16, 92). The physiological role of lipid droplets varies with the subcellular localization, the morphology, and the (sub)proteome of the droplet. In adipocytes, the cytoplasm is almost entirely occupied by the LD, whereas in tissues with a high lipolytic rate like cardiac and skeletal muscle, numerous small LD are present. Size, and possibly function, is sensitive to diet, disease, nutritional status, and training status (92) and on the balance between LD synthesis and LD degradation.

Abundant storage of TAG in LD in metabolically dynamic tissues like cardiac and skeletal muscle, as well as the liver and pancreas, may interfere with subcellular processes like insulin signaling. Thus, a strong positive correlation between ectopic TAG storage, measured by 1H NMR spectroscopy as intramyocellular lipid content (IMCL), and insulin resistance (57, 69, 101) has been reported in insulin-resistant subjects. Interestingly, endurance training, an insulin-sensitizing intervention, also coincides with increased IMCL content (40). In fact, we have shown that an increase in IMCL is a very early response to training (119). In addition, it has been shown that a 4-mo-insulin sensitizing intervention of combined weight loss and physical activity significantly decreased LD size, without a decline in total myocellular triacylglycerol content (49). It is, therefore,
unlikely that the presence of LD per se impedes insulin signaling. Rather, it has been hypothesized that intermediates in LD synthesis and degradation, such as DAG, ceramides, and fatty acyl CoA esters might be responsible (56, 104, 112, 122). The balance between LD synthesis and degradation, here referred to as LD dynamics, partly determines the fraction of lipid-derived intermediates.

In adipocytes, the phospholipid monolayer coating the LD is decorated with several proteins involved in lipid droplet dynamics referred to as PAT proteins. PAT proteins [perilipin, ADRP (for adipocyte differentiation-related protein), and TIP47 (for tail-interacting protein of 47 kDa)], are characterized by a common N-terminal motif (PAT domain). Other recently discovered proteins, like S3-12 and OXPAT, possess a high-sequence homology with the PAT domain and are also considered PAT proteins. Throughout this review, we will use the term PAT proteins not only for perilipin, ADRP, and TIP-47, but also for S3-12 and OXPAT.

To better understand the origin of insulin-desensitizing lipid moieties in skeletal muscle, a detailed understanding of the processes involved in LD formation and degradation is hence warranted. To date, however, data on LD dynamics in (human) muscle are still scarce. A rapidly expanding body of knowledge on the process of LD synthesis and degradation, such as DAG, ceramide and fatty acyl CoA, not only in adipose muscle are still scarce. A rapidly expanding body of knowledge on the process of LD synthesis and degradation, such as DAG, ceramide and fatty acyl CoA, not only in adipose muscle is similar to adipose tissue

ATGL. For the past 30 years, HSL was considered the only lipase involved in TAG degradation. Interestingly, in HSL knockout mice, total adipose tissue mass did not change (100, 104), while hydrolysis of TAG was still possible, but DAG levels were increased (48). This suggests that a lipase other than HSL must be involved in TAG hydrolysis. This protein has been identified and was named adipose triglyceride lipase (ATGL) (162). Independent studies identified the same protein at about the same time and called it iPLA2 (58) and desnutrin (142). ATGL is well preserved across species, and homologues have been reported in yeast [Tgl4 (7)], in plants [SUGAR-DEPENDENT1 lipase (31)], and in Drosophila [Brummer lipase (46)]. Here, we will use the term ATGL for this lipase.

ATGL mRNA is predominantly expressed in WAT and brown adipose tissue (BAT), and to a lesser extent in cardiac and skeletal muscle (47, 60, 63, 65, 142, 162). Cell studies revealed PPARγ-mediated control of ATGL gene expression (65, 142). Both thiazolidinedione-based PPARγ agonists and nonthiazolidinedione-based PPARγ agonists increased ATGL mRNA and ATGL protein in 3T3-L1 adipocytes (64). Knockdown of PPARγ reduced ATGL mRNA and protein expression in adipocytes (64). Treating mice with rosiglitazone increased ATGL mRNA and protein content (64, 77, 121).

During fasting, a state in which PPARγ is activated through elevated free fatty-acid levels, ATGL mRNA and ATGL protein expression is upregulated (63, 142). Refeeding results in a decreased transcription of ATGL in mice (63, 65). Decreases in transcription of the ATGL gene after refeeding can be a direct effect of reduced PPARγ activity in adipose tissue; it can also be mediated via insulin, transient hyperglycemia or reduced FFA levels by insulin-mediated suppression of adipose tissue lipolysis. Later studies revealed that transcriptional control of the ATGL gene in adipose tissue is under control of insulin rather than glucose (63, 65, 68). Thus, ATGL appears to be under control of PPARγ, as well as insulin.

In 3T3-L1 adipocytes and under basal conditions, the majority of ATGL protein was present in the cytoplasm, whereas a distinct fraction of ATGL was tightly associated with LD (162). ATGL hydrolyzes the first ester bond of TAG and is likely to limit lipolysis in smooth muscle, cardiac muscle, and possibly skeletal muscle (47, 48, 120). Although knock-down of ATGL (or its homologue Brummer lipase in the case of Drosophila) resulted in a decreased lipolysis and an increase in LD size in Drosophila, 3T3-L1 cells and in HeLa cells (35, 46, 63, 124), adipose tissue mass in mice lacking ATGL was...
increased, but not massively. Rather, fat storage in nonadipose tissue increased to supraphysiological levels in skeletal and cardiac muscle (47, 120). At 12 wk of age, a striking 20-fold rise in cardiac TAG content was observed. As a result of the increase in TAG accumulation, cardiac function was severely hampered, and the knockout of ATGL resulted in cardiac failure and premature death (47). In humans, carriers of an ATGL polymorphism, resulting in lower ATGL activity, possessed excess ectopic TAG accumulation along with myopathy (35).

Reduced TAG lipolysis in muscle of ATGL knockout mice was paralleled by a decrease in fat oxidation and a concomitant increase in glucose oxidation during fasting and an unanticipated increased glucose uptake (47, 54). Increased glucose uptake by the muscle, despite excessive TAG storage, may reflect substrate availability; in the absence of ATGL activity, fatty acids cannot be released from the LD and can hence not be used as an energy source. Thus glucose uptake and oxidation must increase to meet the energy demand.

The relatively modest (~2-fold) increase in adipose tissue mass and liver fat in ATGL knockout mice (47), suggests that in these tissues ATGL may not be the major TAG lipase. In cardiac muscle and to a lesser extent in skeletal muscle, however, TAG storage in ATGL knockout mice is more pronounced, suggesting an important role for ATGL-based lipolysis of TAG in muscle. Overexpression of ATGL in 3T3-L1 cells, COS-7 cells, and HeLa cells (46, 63, 124, 142), as well as overexpression of the Brummer lipase (homologue of human ATGL) in Drosophila (46, 63, 124, 142), resulted in increased TAG lipolysis and a concomitant decrease in LD size. Unilateral overexpression of ATGL in skeletal muscle increased oxidation of LD-derived fatty acids and increased insulin-desensitizing lipid intermediates like DAG and ceramide (148), suggesting that increased ATGL activity was not matched with increased HSL activity. Thus, disequilibrium between ATGL and HSL activity may increase DAG content and impede insulin signaling.

In diet-induced obese C57BL/6J mice and ob/ob mice low ATGL protein levels have been reported in muscle along with high muscle TAG storage (65, 142, 148). In contrast to ATGL protein levels, however, ATGL mRNA expression was higher in skeletal muscle of C57BL/6J high-fat mice and ob/ob mice compared with nonobese controls (148). Similar observations have been made in subcutaneous adipose tissue of obese subjects, in which low ATGL protein content was observed along with high levels of ATGL mRNA (127), suggesting that posttranscriptional processes profoundly affect ATGL protein content. Despite the dissociation between mRNA expression and protein content, the above-mentioned studies indicate an association between myocellular ATGL and obesity. In adipose tissue, however, diet-induced obesity in mice did not affect ATGL protein content (64, 121), and ATGL mRNA levels in ob/ob mice were comparable to lean littermates (71). Similarly, in adipose tissue of obese humans, ATGL gene expression (81) or protein content (114) was comparable to nonobese subjects. Furthermore, in insulin-resistant subjects, adipose tissue ATGL mRNA levels were lower compared with obese subjects with preserved insulin sensitivity irrespective of body fat mass or fat distribution (9).

In insulin-resistant subjects, weight loss resulted in a decrease in ATGL and HSL mRNA- and protein level (61), but in insulin-nonresistant subjects, this decrease was not detected (81). However, the insulin-resistant subjects were studied in negative energy balance, whereas the insulin-nonresistant subjects were studied after energy balance was restored (81). In normoglycemic young men, exercise training resulted in increased ATGL protein in muscle without changes in HSL protein level, but with a decline in muscle TAG content (4). Strikingly, exercise training in general improves insulin sensitivity, whereas the changes observed here could imply increased DAG levels, which is supposed to impede insulin sensitivity. Unfortunately data on DAG or insulin sensitivity have not been reported in this study.

ATGL activity requires coactivation by comparative gene identification-58 (CGI-58), also known as α/β-hydrolase domain containing 5 (ABHD5) (74, 120, 160). CGI-58 stimulates TAG hydrolysis in wild-type and HSL-deficient adipose tissue but fails to activate lipolysis in the absence of ATGL, suggesting that CGI-58 is not a lipase as such, but rather facilitates lipolysis by ATGL but not HSL (120). ATGL activity was enhanced 20-fold in the presence of CGI-58 (74). Furthermore, a mutation in CGI-58 is associated with Chanarin-Dorfman Syndrome, a rare genetic disease resulting in excessive ectopic TAG accumulation. Reintroduction of functional CGI-58 normalizes the TAG content in peripheral tissues of these patients (74). These results indicate that CGI-58 is required to maximize the rate of TAG hydrolysis. Regulation of ATGL activity by CGI-58 requires interaction with lipid droplet-coating proteins of the PAT family, which will be discussed in Lipid droplet-binding proteins not related to the PAT family.

Several studies have provided evidence that ATGL and HSL act coordinately in hydrolyzing triglycerides (73). The current hypothesis states that ATGL hydrolyzes the first ester bond of the TAG molecule, whereas HSL preferentially degrades DAG (36, 120). Data indicate that this is the case for TAG stored in adipose tissue, as well as for TAG stored ectopically. A disequilibrium in ATGL and HSL activation in insulin-responsive tissues like muscle may increase the content of the insulin-desensitizing lipid intermediate DAG. So far, however, this putative disequilibrium has not been reported in muscle of type 2 diabetic subjects.

MGL and other lipases. Information on the role and regulation of monoaoylglycerol lipase (MGL) is limited, especially for ectopically stored fat. Given the specific affinity of ATGL for TAG and of HSL for DAG, another lipase with a high affinity for MAG hydrolysis could be of importance in adipose tissue and skeletal muscle. In adipose tissue, it has, indeed, been shown that MGL is of importance for complete hydrolysis of MAG (37). Experiments in murine white adipose tissue in vitro to examine the relative contribution of ATGL and HSL to total lipolysis revealed that joint action of ATGL and HSL is responsible for 95% of total hydrolytic activity (120), suggesting a quantitatively minor role for other lipases, including MGL.

Triacylglycerol hydrolase (TGH) and triacylglycerol hydrolase homologue (TGH-2) are TAG-specific lipolytic enzymes highly expressed in liver and adipose tissue (98). In contrast to ATGL and HSL, TGH and TGH-2 efficiently hydrolyze triacylglycerol comprising short-chain fatty acids. Because of hepatic conversion of short-chain fatty acids into long-chain fatty acids by elongases and the absence in muscle of a short-chain fatty-acid specific synthase, the vast majority of the TAG
stored in the human body comprises long-chain fatty acids. Whereas the quantitative role of TGH and TGH-2 in hydrolysis of hepatic TAG may be highly relevant, its role in hydrolysis of myocellular TAG may be less prominent.

Jointly, ATGL and HSL can hydrolyze the vast majority of the TAG in adipose tissue and skeletal muscle; at present, there is little reason, suggesting that defects in hydrolysis of MAG in skeletal muscle can play a major role in modulating lipid-desensitizing intermediates in skeletal muscle.

Lipid Droplet-Coating Proteins

The LD is coated by a phospholipid monolayer decorated with a family of proteins, referred to as PAT proteins, characterized by a common NH2-terminal motif. PAT refers to Lipid Droplet-Coating Proteins desensitizing intermediates in skeletal muscle.

In adipose tissue, activation of perilipin by phosphorylation is required for full activation of HSL-related hydrolysis. Perilipin is the predominant PAT protein in adipose tissue involved in control of storage and degradation of lipid stored in the LD (1, 39, 125, 126). Expression of perilipin is under control of the transcription factor PPAR (1, 6, 26), and perilipin is predominantly found on the LD of adipose tissue (10, 43, 103). In the fed state and in the presence of insulin, perilipin is not phosphorylated on PKA site serines and coats the LD in adipose tissue (32, 89), thereby limiting lipolytic activity of HSL to lipid stored in the LD (15, 88). In contrast, under lipidic conditions, perilipin gets phosphorylated by PKA (15, 43, 44), allowing HSL to access the LD (89, 91, 126, 132, 133). In line with this, TAG levels in adipose tissue of perilipin knockout mice (83, 133) and in cells lacking perilipin is reduced (15, 83, 88), suggesting a pivotal role for perilipin in regulating HSL activity. Despite higher lean body mass and an increased metabolic rate, perilipin-null mice tend to develop glucose intolerance and peripheral insulin resistance (115). Interestingly, perilipin levels in adipose tissue from obese Zucker rats and obese humans appear lower than in lean controls if matched for fat cell mass (26, 91, 144). It is important to note that most of the data on regulation of lipolysis of lipid stored in LD and of interaction of perilipin with HSL has been published in the pre-ATGL era, and hence, one cannot exclude the possibility that some of these observations have been confounded by lipase activity of ATGL, which by then, was not yet discovered.

Although hydrolysis of TAG in myocellular LD involves HSL activity, skeletal muscle is devoid of perilipin. Rather, adrenaline and contractile activity-associated increases in TAG hydrolysis in myocellular LD goes along with recruitment of HSL to ADRP and TIP-47-coated LD (105), suggesting that these PAT proteins may be myocellular analogs of perilipin that operate in muscle to control basal and stimulated lipolysis.

ADRP. Shortly after the discovery of perilipin, adipose differentiation-related protein (ADRP) was identified (34, 59).

ADRP is expressed in a wider range of tissues than perilipin (13, 50), including rodent and human skeletal muscle (103, 105). ADRP is involved in the uptake of long-chain fatty acids from the cytoplasm and incorporation of fatty acids in LD (20, 38, 76, 80, 85). In livers from ob/ob mice (116) and etomoxir (i.e., CPT-1 inhibitor)-treated rats (128) and in specific diseases states involving ectopic fat accumulation (50, 90), ADRP expression is increased.

Like perilipin, ADRP decorates the LD membrane, thus affecting lipolytic activity of ATGL and recruitment of another PAT protein, TIP47, to the lipid droplet (76, 97). In ADRP-expressing tissues, hydrolysis of TAG in LD might indeed be under control of ADRP (105, 150). Tissue-specific and whole body knockout studies of ADRP resulted in resistance to diet-induced hepatic steatosis, while adipocyte differentiation and lipolysis were unaffected (20). The absence of changes in adipogenesis may originate from a compensatory induction of TIP47. In ADRP knockout cells with concomitant siRNA-mediated downregulation of TIP47, LD formation was attenuated, and fatty acids added to the system incorporated into phospholipids in cellular membranes other than LD (131). Downregulation of hepatic ADRP using antisense oligonucleotides reduced hepatic steatosis, hypertriglyceridemia, and insulin resistance in obese mice (55) and rescued diet-induced obese mice from developing insulin resistance. These favorable effects were attributed to lower levels of the insulin-desensitizing lipid intermediate DAG in the livers of the treated animals (140).

ADRP may also be involved in regulating subcellular localization of the LD. In preadipocytes, ADRP associates with the intermediate filament protein vimentin (10, 43, 103), suggesting that intracellular trafficking of the lipid droplets may occur. This notion was substantiated by the observation that the microtubular motor protein dynein—allowing the droplet to move within the cell along the microtubular system—cooperates with ADRP (12). Extending these observations to skeletal muscle is intriguing; in mature skeletal muscle, LD can be found in a highly organized manner along myofibrils and adjacent to Z-discs (Fig. 1) in direct vicinity of the Z-disc interconnecting intermediate filament desmin (102). The highly organized subcellular structure of skeletal muscle is initiated during differentiation when vimentin expression decreases and desmin expression increases, allowing alignment of myofibrils in a regular array of Z-discs interconnected by desmin (139). In this respect, it is interesting to note that in fully differentiated skeletal muscle, the motor protein dynein and a microtubular network machinery both are present to traffic nascent LD from the ER toward a location in vicinity of the Z-disc and in association with mitochondria.

In 3T3-L1 preadipocytes and in early stages of differentiating 3T3-L1 adipocytes, the gene encoding for ADRP is induced, and ADRP protein content increases (13, 59). In fully differentiated adipocytes, however, ADRP protein levels drop, despite the persistence of high levels of ADRP mRNA. At the protein level, it has been shown that maintenance of ADRP content highly depends on the amount of neutral lipids present in the cell. ADRP, which is not bound to neutral lipids, is posttranslationally targeted for proteasomal degradation (84, 157).

Like perilipin, the ADRP promoter contains a PPAR-responsive element. Likewise, ADRP gene expression can be stimu-
lated by PPARα- (25, 27, 134) and PPARβ/δ ligands (21, 118). It remains uncertain whether PPARγ is also involved, since several investigations present conflicting results (reviewed in Ref. 27).

**TIP47.** Another member of the PAT protein family is the 47 kDa protein TIP47 [also known as pp17 (136)]. TIP47 is ubiquitously expressed (26) and shares 43% homology with ADRP (8). TIP47 gene expression is not regulated by PPARγ (26, 27). Originally, TIP47 was described to be a mannose-6-phosphate receptor sorting device (8, 29). However, knockdown of TIP47 had no effect on mannose-6-phosphate receptors but did affect LD biogenesis, lipid incorporation, and lipid liberation (17). Knocking-out ADRP in human hepatocytes (Huh7) cells or in immortalized fibroblasts derived from ADRP-ablated embryonic cells, resulted in a compensatory increase in TIP47 (97, 131).

In contrast to ADRP, cytosolic TIP47 that is not associated with neutral lipids is not degraded but may translocate to nascent LD, allowing LD growth (87, 97, 153). Studies using freeze fracture followed by Immunogold labeling revealed that TIP47 and ADRP both are indeed extensively found in association with the LD membrane but may also invade the LD (109–111). Thus, TIP47 may have a role in the packaging of neutral lipids into droplets as well (17, 153). Exactly how TIP47 exerts its role, especially in skeletal muscle lipid droplet packaging, however, remains to be established.

**S3-12.** The PAT member family with the lowest sequence homology to the other PAT proteins is S3-12 (117). S3-12 shares a 33-amino acid motif with ADRP and has a protein sequence identity of 50% with both TIP47 and ADRP (79) but does not have a PAT domain. S3-12 mRNA is induced in early stages of adipocyte differentiation (117) and coats nascent lipid droplets during TAG synthesis in 3T3-L1 adipocytes (151, 154). S3-12 is highly expressed in WAT and to a lesser, but detectable, extent in BAT, cardiac and skeletal muscle (117).

Transcriptional regulation of S3-12 has been shown to be PPARγ dependent (26). So far, little information is available on the exact role of S3-12 in muscle lipid metabolism.

**OXPAT/MLDP/LSDP5.** Three independent laboratories recently identified a new PAT domain containing a LD-coating protein expressed mainly in tissues with a high capacity for fat oxidation, like (type 1) muscle fibers, cardiac muscle, brown adipose tissue, and liver. This tissue expression profile resulted in the protein being named OXPAT (152). The same protein has also been identified in the myocardium and was named myocardial lipid droplet protein (MLDP) (158), while a third research group named the protein lipid storage droplet protein (LSDP5) (25). Throughout this review, we will use the name OXPAT. The primary structure of OXPAT shares a high degree of homology to ADRP and TIP47 (25). Importantly, OXPAT has the same PAT domain as the other PAT proteins but has its own unique tissue expression pattern. Via mechanism(s) not yet identified, OXPAT promotes both long-chain fatty acids esterification into TAG, as well as long-chain fatty acid oxidation (152). Because there is a significant overlap in tissues expressing OXPAT and S3-12, it has been suggested that OXPAT and S3-12 are coordinately and reciprocally regulated (152). However, detailed information on the concordant regulation of S3-12 and OXPAT is currently lacking.

Twelve hours of fasting increased OXPAT mRNA, and 24 h of fasting increased OXPAT protein expression in liver and to a lesser extent in other tissues like the heart. After 12 h of refeeding, OXPAT mRNA levels had returned to baseline levels (25). Interestingly, in the highly oxidative soleus muscle, the already abundant levels of OXPAT did not respond to fasting or refeeding. Activation of PPARα induced OXPAT expression in wild-type mice but not in PPARα knockout mice, suggesting that the presence of PPARα is required for induction of the OXPAT gene. Fasting increased OXPAT mRNA levels in wild-type mice, but for reasons not yet understood,
also in the PPARα knockout animals. It should be noted, though, that because of a considerably lower baseline expression of OXPAT in the PPARα knockout mice, OXPAT mRNA content, upon fasting, did not reach the levels observed in fed wild-type mice. Thus, it can be concluded that fasting-induced increases in OXPAT expression do not require a functional PPARα gene, whereas functional PPARα is important for basal expression of OXPAT in the liver and heart (25, 152, 158). Interestingly, mice and humans treated with pioglitazone (a PPARγ agonist with PPARα affinity) increased gene expression of OXPAT in adipose tissue but not in skeletal muscle (152). Recently, we showed that in skeletal muscle from humans that OXPAT protein content in obese control subjects was comparable to levels found in body mass index (BMI)-matched type 2 diabetic patients. Interestingly, we also showed that upon rosiglitazone-induced insulin sensitization, OXPAT levels declined significantly without changes in myocellular LD content (86).

Lipid droplet-binding proteins not related to the PAT family. Full activation of ATGL requires interaction of a coactivator known as CGI-58, which, in turn, interacts with perilipin and ADRP on the surface of LD in 3T3-L1 cells (41, 130, 159–161). It has been hypothesized that physical association of ATGL with CGI-58 requires release of CGI-58 from the (adipocyte) droplet to exert full lipolytic activity of ATGL (130, 160). Experimental data in 3T3-L1 adipocytes indicate that in the basal condition, CGI-58 strongly interacts with perilipin. Upon phosphorylation of perilipin, CGI-58 detaches from perilipin and is released from the droplet to translocate and interact with ATGL at the surface of small LD at sites devoid of perilipin (41).

Given the ubiquitous expression of CGI-58 and the multiple tissues expressing ATGL, it is likely that activation of TG hydrolysis by ATGL and its coactivation by CGI-58 is not restricted to adipose tissue, but it is also operational in, for example, cardiac and skeletal muscle. The notion that CGI-58 might also be operational in skeletal muscle is substantiated by the observations that patients suffering from mutations in either the CGI-58 gene or in the ATGL gene present with severe muscle weakness and myopathies (3, 66). As perilipin is not present in muscle tissue, coregulation of ATGL by CGI-58 in skeletal muscle most likely involves other (PAT) protein(s). Indeed, it has recently been shown in mouse cardiac myocytes that in virtually all OXPAT-containing LD also, CGI-58 was detected. In LD, mainly coated by ADRP, no such interaction was observed. Loading cardiac myocytes with oleate promoted interaction of OXPAT with CGI-58 and appeared critical in regulating ATGL lipolytic activity in OXPAT containing LD (42).

By detailed examination of multiple proteins undergoing major changes in expression during adipogenesis, two members of the Cide domain-coating proteins, CideA and CideC (also known as fat-specific protein 27, FSP27), have recently been identified as LD-binding proteins (62, 95, 106, 107, 138). These proteins appear to be involved in the negative regulation of lipolysis and, hence, in LD morphology and LD dynamics in a perilipin-like manner (106). Ectopically overexpressed CideC colocalizes to perilipin (106), suggesting interaction of both proteins in facilitating LD lipolysis.

CideA is predominantly expressed in BAT, with small amounts of mRNA detected in among others cardiac and skeletal muscle. CideC is more widespread, with high levels in WAT and moderate levels in BAT and skeletal muscle (155). Overexpression of CideA and CideC blunts lipid droplet degradation and results in increased ectopic fat storage (106). Downregulation of CideA using RNAi markedly elevates lipolysis in human adipocytes (107). Transcriptional rate of both CideA and CideC is controlled by PPARγ. Hence, treating mice with rosiglitazone induced CideA gene expression and lipid deposition in WAT (107). In humans, expression of CideA and CideC correlates positively with insulin sensitivity. A third member of this family, CideB appears to be involved in hepatic fat storage. Indeed, Cide B knockout mice possessed reduced high-fat diet-induced hepatic steatosis and lower circulating plasma TGs, and they were insulin sensitive (75).

In search for binding proteins to pigment epithelium-derived factors (PEDF), Notari et al. (96) serendipitously identified ATGL as a PEDF binding protein, where ATGL exerts lipase activity to phospholipids (96). Interestingly, PEDF coprecipitates with ATGL in hepatocytes (22), and the lack of PEDF with subsequent reintroduction was associated with significant increases in steatosis and normalization toward control values, respectively (22). Thus, via binding to ATGL, PEDF may have a role in hepatic triglyceride homeostasis, and, hence, insulin sensitivity. Indeed, it has recently been shown that administrating PEDF to mice stimulated adipose tissue lipolysis, augmented ectopic lipid deposition, and reduced insulin sensitivity. On the other hand, antibody-based neutralization of PEDF in obese mice enhanced insulin sensitivity (24). Overall, these results identify a causal role for PEDF in obesity-induced insulin resistance.

At this stage, it seems safe to conclude that next to the PAT proteins, other proteins with a role in LD dynamics start to emerge. It is, however, not yet fully understood how these proteins may interact to exert their role in modulation of LD dynamics.

Lipid Droplet Formation

Information on formation, maturation, and potential trafficking of LD in different cell types or tissues is still fragmentary and not always consistent. As reviewed by several authors (16, 82, 92, 93, 150), it has been hypothesized that neutral lipids are deposited between the leaflets of the ER membrane and then budded into the cytoplasm to form a primordial LD. LD-coating proteins might be either integral ER membrane proteins or may be added to the growing LD from the cytosol. Proteins coating the LD may originate from proteins resident in the cytoplasmic leaflet of ER membranes (16, 99). In contrast it has been shown that the fatty acid composition of LD phospholipids differed from that of rough ER phospholipids (135), reducing the likelihood that these droplets indeed originally budded from the ER. Alternatively, LD may be formed at specialized ADRP-enriched domains on the cytoplasmic surface of the ER (108).

At present, none of the existing models fully explains how TAG is incorporated into ER-associated LD.

Lipid droplet growth. Upon formation, LD continue to grow until they reach their mature size and the amount of TAG in the LD increases. The rate-limiting enzyme in TAG synthesis is diacylglycerol acyl-transferase (DGAT), which converts DAG into TAG. Hence, the involvement of DGAT in TAG accretion...
in the LD and increased LD size is anticipated (see Fig. 2). DGAT is present in two isoforms. DGAT1 and DGAT2 both have been located to the ER, albeit at distinct regions (18, 19). Fluorescence microscopy of polyene-labeled lipids revealed the presence of DAG to the outside of LD in COS7 cells (70). It has been shown that DGAT2 localizes to the ER under basal conditions and was targeted to the surface of LD in COS7 cells upon oleate loading (129) and that endogenous DGAT2 was localized to the vicinity of LD in 3T3-L1 adipocytes (70). This model is compatible with the origin of LD from the ER and demonstrates that LD can grow in size by inflow of TAG synthesized by DGAT2. Whether DGAT2 and DAG also localize to the LD in other cell types than fibroblasts and adipocytes and how newly formed TAG enters the lipid droplet is not yet known.

Recently, Olofsson et al. (11) hypothesized that primordial droplets are trafficking in the cytosol and fuse in vicinity of other LD (11). Fusion of LD requires membrane fusion of the lipid droplet-coating phospholipid monolayers, in a process facilitated by SNAP23 (11). Physical interaction of myocellular LD under TAG-storing conditions (DGAT 1 overexpression combined with a high-fat diet) is shown in Fig. 2B. Interestingly, fusion of GLUT4-containing vesicles with the sarcolemma upon insulin stimulation also requires SNAP23 (for review, see Huang and Czech (53)). Increasing the amount of SNAP23 to the membrane of LD upon oleate loading resulted in reduced GLUT4 fusion with the sarcolemma, possibly due to the low levels of SNAP remaining for GLUT4 membrane fusion (11). In line with this, overexpressing SNAP23 rescued oleate-mediated insulin resistance (11).

Involvement of PAT proteins in trafficking of LD has been hypothesized and partly been proven by Wolins et al. (150). In his model, PAT proteins are classified on the basis of their stability if not bound to neutral lipids (150). Perilipin and ADRP can be classified as proteins requiring neutral lipids to maintain stability in the absence of neutral lipids. Perilipin and ADRP are degraded via the ubiquitin/proteasome (156, 157). Tip47, S3-12, and OXPAT, on the other hand, are situated in the cytosolic pool under nonlipid-loading conditions and may translocate to nascent LD upon lipid loading (150). In line with this, in 3T3-L1 cells incubated with long-chain fatty acids, TAG-filled droplets coated with TIP47, S3-12, and ADRP appear. Subsequently, S3-12 and TIP47 coat the smallest and most peripheral LD, whereas ADRP coats a more medial population of larger LD (151, 154), and the largest droplets are coated with perilipin. Although part of this hypothesis has been confirmed in adipocytes, it should be noted that only adipocytes express all five members of the PAT family at significant levels. Most other cell types express two, or at most three PAT proteins.

Lipid Droplet Dynamics and Insulin Resistance

Obviously, tight interaction of PAT proteins and well-balanced activity of lipases is required to effectively store fatty acids in LD in times of lipid overflow and for controlled release of fatty acids in times of enhanced demand. Dysregulation of any of these processes might lead to disturbed LD synthesis and/or LD degradation, resulting in a rise in insulin-desensitizing intermediates. It is, therefore, of importance to improve our understanding of regulation of lipid synthesis and degradation in ectopically stored LD and the proteins involved. This type of information in humans, and especially in patient populations, is largely lacking and mainly associative. For example, lower perilipin protein levels have been reported in adipose tissue from obese vs. lean women (91, 144). Another study investigated the relation between ADRP and insulin sensitivity in skeletal muscle after a weight loss intervention program and after a pharmacological intervention program with troglitazone (103). Although a correlation between ADRP and insulin sensitivity was not detectable at baseline, ADRP in skeletal muscle was significantly increased after treatment and correlated with improved insulin sensitivity (103). Furthermore, a population of 85 nondiabetic subjects with a wide range of BMI and insulin sensitivity was investigated, and a negative correlation between OXPAT gene expression in adi-
pose tissue and BMI was found (152). From these data, cause and consequence cannot be separated, and other processes, which were not examined, may also contribute to, or even cause, the observed correlation between OXPAT and BMI (152).

**Perspective and Significance**

For decades, intracellular lipid droplets (LD) in nonadipose tissue have been considered to be relatively inert sites of fat storage, which are well preserved across species and can be found in a wide variety of sizes and quantities in various tissues. In lipid bodies in adipocytes, however, it has been recognized that synthesis and degradation of lipid in LD are fairly dynamic processes. Predominantly arising from reports in adipocytes and progress in the imaging of lipid droplets in live cells, the body of knowledge on LD dynamics and the proteins involved have rapidly expanded recently. Defects in LD dynamics have been linked to multiple disorders in lipid and lipoprotein metabolism and may contribute to the insulin-desensitizing effects of lipid intermediates (partly originating from LD), in insulin-responsive tissues like adipose tissue, liver, and cardiac and skeletal muscle.

Hydrolysis of lipid stored in LD is catalyzed by the concerted action of distinct lipases, which differ in their affinity for triacyl-, diacyl- or monoacylglycerol. In muscle, the two most abundant and best described lipases identified so far are HSL and ATGL. These lipases regulate—both positively and negatively—the rate of lipolysis by tight cooperation with a family of structurally related lipid droplet-coating proteins (perilipin, ADRP, TIP47, S3-12, and OXPAT, collectively referred to as PAT proteins) and with coactivators like CGI-58. While expression of perilipin appears to be restricted to adipose tissue, the other PAT proteins can be found in other cell types as well. Adrenergic stimulated lipolysis in adipose tissue results in PKA-mediated phosphorylation of perilipin, which recruits HSL to the LD surface to stimulate lipolysis. In addition, PKA activation results in dissociation of CGI-58 from the LD in a perilipin-dependent manner, allowing ATGL-based lipolysis. Prolonged PKA-mediated lipolysis results in smaller LD decorated with ADRP, TIP47, and S3-12.

While interaction of CGI-58 with ADRP has been described as well, it remains unclear whether and in which tissues this interaction affects lipolytic rate. In muscle, interaction of OXPAT with CGI-58 appears to be critical in regulating lipolytic activity of ATGL, but more work is needed to test to what extent information derived from other tissues or cell types can be translated toward (human) skeletal muscle.

Our knowledge on involvement of PAT proteins and their putative interaction with lipases in the regulation of lipolysis of lipid droplets has rapidly expanded. In addition, the emerging body of knowledge from cell studies toward LD growth and fusion expands the field to further explore the mechanisms involved in (dys)regulation of LD dynamics in ectopically stored fat in insulin-responsive tissues like liver and (skeletal) muscle. With this area of research developing, targets to modulate ectopic fat storage, in terms of synthesis, degradation, and lipid turnover, may be identified in the near future that may shed new light on routes to rescue myocardial insulin resistance, the hallmark in the development of overt type 2 diabetes.

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