Functional Interactions between Mldp (LSDP5) and Abhd5 in the Control of Intracellular Lipid Accumulation

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Cellular lipid metabolism is regulated in part by protein-protein interactions near the surface of intracellular lipid droplets. This work investigated functional interactions between Abhd5, a protein activator of the lipase Atgl, and Mldp, a lipid droplet scaffold protein that is highly expressed in oxidative tissues. Abhd5 was highly targeted to individual lipid droplets containing Mldp in microdissected cardiac muscle fibers. Mldp bound Abhd5 in transfected fibroblasts and directed it to lipid droplets in proportion to Mldp concentration. Analysis of protein-protein interactions in situ demonstrated that the interaction of Abhd5 and Mldp occurs mainly, if not exclusively, on the surface of lipid droplets. Oleic acid treatment rapidly increased the interaction between Abhd5 and Mldp, and this effect was suppressed by pharmacological inhibition of triglyceride synthesis. The functional role of the Abhd5-Mldp interaction was explored using a mutant of mouse Abhd5 (E262K) that has greatly reduced binding to Mldp. Mldp promoted the subcellular colocalization and interaction of Atgl with wild type, but not mutant, Abhd5. This differential interaction was reflected in cellular assays of Atgl activity. In the absence of Mldp, wild type and mutant Abhd5 were equally effective in reducing lipid droplet formation. In contrast, mutant Abhd5 was unable to prevent lipid droplet accumulation in cells expressing Mldp despite considerable targeting of Atgl to lipid droplets containing Mldp. These results indicate that the interaction between Abhd5 and Mldp is dynamic and essential for regulating the activity of Atgl at lipid droplets containing Mldp.

Growing evidence indicates that lipogenesis and lipolysis are regulated by protein-protein interactions that occur on the surface of specialized intracellular lipid droplets (1, 2). PAT1 ( perilipin, adipophilin, and TIP-47) proteins, are thought to be key regulators of these processes by serving as scaffolds that organize and regulate the protein trafficking at lipid droplet surfaces (1–3). Mldp (muscle lipid droplet protein; alternatively, OXPAT, LSDP5) is a PAT family member that is highly expressed in tissues, like muscle and liver, having high oxidative capacity (4–6). Expression of Mldp is up-regulated under conditions such as fasting and diabetes, in which the systemic supply of lipid to target tissues is increased, and in vitro studies suggest that Mldp plays a role in facilitating triglyceride storage as well as fatty acid oxidation (4–6). It is not presently known how Mldp is involved in these functions, but we hypothesize that it is likely to involve direct or indirect interactions with lipases and lipase co-activators (3, 7).

Abhd5 (α/β hydrolase domain-containing protein 5; alternatively CGI-58) is an evolutionarily conserved protein that acts as a potent activator of Atgl (adipose triglyceride lipase; alternatively, PNPLA2, desnutrin, TTS-2.1) (8). Both proteins are expressed in a variety of tissues, and rare homozygous mutations of either gene in humans produces a similar (but not identical) lipid storage disease that is characterized by ectopic lipid accumulation in skin, muscle, and liver (9–11). Regulation of lipid metabolism by Abhd5 is not fully understood. Abhd5 has been shown to bind perilipin (Plin) (12, 13), and it has been proposed that the phosphorylation-dependent release of Abhd5 is a means of initiating lipolysis via activation of Atgl (3, 7). Abhd5 is expressed in several tissues that lack Plin (12), raising the possibility that this co-activator might interact with additional PAT proteins.

In the experiments detailed below, we investigated the potential interaction of Mldp and Abhd5 in vivo and in vitro. Our results show that Mldp and Abhd5 interact in vivo and in vitro. This interaction occurs on the surface of intracellular lipid droplets and is promoted by triglyceride synthesis. Atgl and Mldp are targeted to the same lipid droplets, and the interaction of Abhd5 with Mldp appears to be critical for regulating Atgl activity at these droplets.

EXPERIMENTAL PROCEDURES

Double immunofluorescence Staining of Mouse Cardiomyocytes

Antisera to glutathione S-transferase fusions of mouse Mldp and Abhd5 were generated in rabbits and affinity purified by Proteintech. Antibodies to Adr (adipose differentiation-related protein, adipophilin) were from Research Diagnostics.

PBS, phosphate-buffered saline; BSA, bovine serum albumin; ECFP, enhanced cyan fluorescent protein; EYFP, enhanced yellow fluorescent protein; IB, intracellular buffer.
Alexa-fluor secondary antibodies were obtained from Molecular Probes, whereas Cy3-labeled and unlabeled Fab fragments were obtained from Jackson Immunochemicals.

Hearts were removed from C57/Bl6 mice, washed in PBS, and fixed in 2% paraformaldehyde for 7 h on ice. Tissues were washed in PBS, teased into single fibers, and preincubated in blocking buffer (50 mM glycine, 5% bovine serum albumin (BSA), 0.03% saponin, and 0.02% NaN₃ in PBS) for at least 1 h at 4 °C. The samples were then optically cleared and bleached as described by Dickie et al. (14). Briefly, samples were dehydrated in increasing concentrations of methanol, fixed overnight in Dent’s fix (1:4 v/v DMSO: methanol), bleached for 5 h in 7.5% H₂O₂ in Dent’s, then rehydrated with a descending methanol series.

Double staining of Mldp and Abhd5 was performed using the procedure of Negoescu et al. (15) for employing two primary antibodies from the same species (rabbit), as we have described previously (7, 16). Specimens were successively incubated with the following antibodies (diluted in PBS, 5% BSA, 0.03% saponin): 1) affinity-purified rabbit anti-Mldp (1:100) overnight at 4 °C, 2) an excess of Cy3-conjugated goat-anti-rabbit Fab (1:200) for 3 h at room temperature, 3) affinity-purified rabbit anti-Abhd5 (1:250) overnight at 4 °C, and 4) Alexa 488-conjugated goat-anti-rabbit (Fab)₂ (1:500) for 2 h at room temperature. The samples were washed extensively with PBS between antibody incubations. Under these conditions, there was no cross-reactivity between the two pairs of primary/secondary antibodies; control experiments demonstrated that omission of each primary antibody resulted in complete and specific elimination of fluorescent signals only in the corresponding channel.

For double labeling of Abhd5 and Adrp, the specimens were incubated successively with the following antibodies: 1) affinity-purified rabbit anti-Abhd5 (1:250) overnight at 4 °C, 2) Cy3-conjugated goat-anti-rabbit Fab (1:200) for 3 h at room temperature, 3) guinea pig anti-Adrp (1:200) for 2 h at room temperature, and 4) Alexa 488-conjugated goat-anti-guinea pig antibodies (1:500) for 2 h at room temperature.

**FIGURE 1.** Colocalization of Mldp, Adrp and Abhd5 in mouse cardiomyocytes. A. permeabilized microdissected cardiac muscle fibers were subjected to double-label immunofluorescence for the indicated antigens. Confocal images are shown on the left, and representative line scans of those images are shown on the right. Peaks in the line scan graphs represent individual lipid droplets. RFU, relative fluorescence units. B. average coefficients of determination (Pearson’s r²) indicating the magnitude of colocalization from 4–5 samples. Values are the means ± S.E.

**Generation of Fluorescent Fusion Proteins and Protein Complementation Constructs**

Mldp1 (GI:116292165) and Mldp2 (GI:27754108) were amplified by PCR from mouse heart cDNA and cloned in-frame into EYFP-N1 (Clontech) vector. Mldp1 (hereafter denoted Mldp) is full-length, whereas translation of Mldp2 starts at the second ATG in the full-length mRNA and, thus, lacks the first 15 amino acids of full-length Mldp. Except where noted, results presented below are for full-length Mldp; however, Mldp2 behaved identically to full-length Mldp in all assays tested. Adrp (GI:116235488) and Rab18 (GI:12292992) were cloned by PCR from mouse white fat cDNA into the ECFP-C1 (Clontech) vector. Wild type and E262K mutant Abhd5 constructs were generated by PCR from mouse white fat cDNA and cloned in-frame into EYFP-N1 (Clontech) vector. Wild type and E262K mutant Abhd5 constructs were generated by PCR and fluorescently tagged with ECFP, EYFP, or mCherry, as indicated.

Bimolecular fluorescence complementation (BiFC) constructs were created by substituting split fragments of EYFP (17) for the full-length fluorescent proteins above (7). The N-terminal fragment, designated Yn, contained amino acids 1–158, whereas the C-terminal fragment, designated Yc, contained amino acids 155–239.
for the fluorescent proteins above. The N (amino acids 1–92)- and C (amino acids 93–187)-terminal fragments, designated Ln and Lc, were cloned in-frame onto the N and C termini of test proteins, as indicated in the legends to Figs. 4–6.

Colocalization of Fluorescent Fusion Proteins

**Intact Cell Experiments**—3T3-L1 or COS-7 cells were plated on 25-mm glass coverslips and cultured in Dulbecco’s modified Eagle’s medium containing 10% fetal bovine serum overnight. Cultures were transfected at about 60% confluence using Lipofectamine or Lipofectamine LTX (Invitrogen), as recommended by the manufacturer. After transfection, cells were incubated overnight in growth medium supplemented with 400 μM oleic acid complexed to BSA (0.5%) to facilitate formation of lipid droplets.

**Binding of Recombinant Abhd5 to Permeabilized Cells**—Recombinant Abhd5 tagged with ECFP or mCherry was prepared from lysates of transiently transfected 293T cells. Briefly, transfected 293T cells were washed in PBS, collected by centrifugation, and suspended in intracellular buffer (IB; 10 mM HEPES, pH 7.3, 140 mM KCl, 6 mM NaCl, 1 mM MgCl₂, 2 mM EGTA) at a concentration of 0.8 ml/10-cm plate. Cells were frozen and thawed, then passed 15 times through a 26-gauge needle and centrifuged for 10 min and 16,000 × g to obtain soluble extract. For experiments comparing wild type and E262K Abhd5-Cherry, probe concentrations were determined with a fluorescence plate reader and equalized. Supernatants were diluted 1:1 with IB containing 0.02% saponin and 1% BSA. 3T3-L1 or COS-7 cells were transfected with Mldp-EYFP or control vector as above and after 24 h lightly fixed in 1% fresh paraformaldehyde at 4 °C for 30 min. Cells were washed with PBS then permeabilized for 30 min at room temperature in IB containing 0.02% saponin and 1% BSA. Permeabilized cells were washed once in IB and incubated with ECFP-Abhd5 extracts for 45 min at room temperature. Binding reactions were washed 2–3 times with IB, then postfixed with 1% paraformaldehyde before confocal microscopic analysis. Capture parameters were equal for each fluorescent protein across experimental conditions. Binding of fluorescent Abhd5 to Mldp and Atgl was determined by line scan analysis of merged confocal images using IPLabs software and evaluated by linear regression (GraphPad).

G. princeps Luciferase Complementation

**In Vivo Complementation in HEK 293T Cells**—293T cells were grown in 48-well plates and transfected in quadruplicate with N- and C-luciferase fragments fused to droplet proteins and controls detailed above. In some experiments cells were cultured for 24 h in media containing 400 μM oleic acid, BSA complexes, whereas in other experiments oleic acid/BSA complexes or BSA (0.5%) alone were added 1 h before harvesting cells. To examine the effects of inhibiting triglyceride synthesis, cells were treated separately by coupled in vitro transcription/translation using the Active Pro kit (Ambion) as recommended by the manufacturer. Briefly, 1 μg of plasmid DNA was transfected and translated in a 50-μl reaction for 2 h at 37 °C in a Thermomixer set at 1200 rpm. Reactions were diluted 6-fold in IB vehicle (0.4% final concentration). Reagents were added as 20× stocks, and cells were incubated for 2 h at 37 °C.

**In Vivo Complementation in COS-7 Cells**—COS-7 cells were grown in 24-well plates and transfected in quadruplicate with N- and C-luciferase fragments fused to Atgl or wild type and E262K mutant Abhd5 with EYFP control vector or Mldp-EYFP. After transfection, cells were cultured for 18–24 h in media containing oleic acid. Luciferase activity was determined as previously described (18). Briefly, cells were washed once and lysed in 120 μl of phenol red-free DMEM by freeze-thawing and brief trituration. Lysates (100 μl) were transferred to 96-well plates, and luminescence was read after the addition of 100 μl of 20 μM coelenterazine substrate (NanoLight).

**Cell-free Protein Complementation Assay**—Ln-Abhd5, Ln-Rab18, and Mldp-Lc were cloned into the AMB-CAT (Ambion) vector. Recombinant proteins were produced separately by coupled in vitro transcription/translation using the Active Pro kit (Ambion) as recommended by the manufacturer. Briefly, 1 μg of plasmid DNA was transcribed and translated in a 50-μl reaction for 2 h at 37 °C in a Thermomixer set at 1200 rpm. Reactions were diluted 6-fold in IB with Triacsin-C (10 μM final concentration, Biomol) or DMSO.
with protease inhibitors (Roche Applied Science), and 25 μl of each in vitro transcription/translation reaction was transferred alone and in combination into a white 96-well plate, incubated for 4 h at room temperature, and read for luminescence as above. Protein production was verified by Western blotting with a protein that detects both luciferase fragments. Protein production was verified by Western blotting with a protein antibody (Nanolight) that detects both luciferase fragments.

**Microscopy**

Images were acquired with an Olympus IX-81 microscope equipped with automated filter controls and a spinning disc confocal unit. Images were captured using a 60× 1.2 NA plan apo water immersion lens or 40× 0.9NA plan apo dry lens and a Hamamatsu ORCA cooled CCD camera. The following Chroma filter sets were used for the indicated fluorophores: Alexa 488, ECFP, mCherry, YFP, EYFP, ECFP, and nile red. Lipidtox Deep Red was imaged with the Semrock 4040A filter set. Microscope control and data acquisition were performed using IPlabs (Scanalytics, BD Biosciences) software.

**Image Analysis**

**Double-label Immunofluorescence**—Colocalization of fluorescence was determined by line scan analysis of merged confocal images using IPlabs software, with the analyst blind to the labeling conditions. For analysis of muscle immunofluorescence, at least 3 scans averaging >75 μm in length were made for each cell. Pearson’s linear regression analyses performed for each scan using GraphPad software, and coefficients of determination (r²) were averaged for each cell, with n = number of cells examined. Lipid droplet size was determined by measuring the cross-sectional length of respective immunofluorescence signal of >60 droplets from samples of 3 mice. FRET was performed using the three-filter method (22), and net FRET was calculated using the FRET extension of IPlabs software. For a given experimental dye, all acquisition parameters were equivalent for all cells and all channels. The adequacy of the FRET constants in eliminating bleed-through was verified with independent singly transfected cells. FRET signals were only observed at lipid droplets containing ECFP, and photobleaching of the acceptor eliminated the calculated net FRET signals.

**BiFC**—3T3-L1 preadipocytes were grown on 25-mm coverslips and transfected with 500 ng each of the complementary Yn and Yc fusion constructs along with 130 ng of an ECFP tracer to identify transfected cells. Cells were cultured for 24 h at 32 °C in growth media supplemented with oleic acid, then fixed in PBS containing 1% paraformaldehyde. Fixed cells were examined using a 40× 0.9NA air objective by an observer that was blind to the labeling conditions. For analysis of muscle immunofluorescence, at least 3 scans averaging >75 μm in length were made for each cell. Pearson’s linear regression analyses performed for each scan using GraphPad software, and coefficients of determination (r²) were averaged for each cell, with n = number of cells examined. Lipid droplet size was determined by measuring the cross-sectional length of respective immunofluorescence signal of >60 droplets from samples of 3 mice. FRET was performed using the three-filter method (22), and net FRET was calculated using the FRET extension of IPlabs software. For a given experimental dye, all acquisition parameters were equivalent for all cells and all channels. The adequacy of the FRET constants in eliminating bleed-through was verified with independent singly transfected cells. FRET signals were only observed at lipid droplets containing ECFP, and photobleaching of the acceptor eliminated the calculated net FRET signals.
Lipid Droplet Formation and Accumulation of Neutral Lipids—COS-7 cells were transfected with Mldp-EYFP (or control), ECFP-Atgl (or lipase-dead S47A mutant), and wild type or E262K mutant Abhd5-Cherry at a plasmid ratio of 2.5:1:1. After transfection, cells were incubated with media supplemented with 400 μM oleic acid complexed to BSA and fixed in 1% paraformaldehyde after 18 h. In the first experiment transfected cells were localized using ECFP-Atgl and scored as to the presence or absence of lipid droplet clusters using differential interference contrast by an analyst who was blind to the transfection conditions (see supplemental Fig. 3 for representative images). Three coverslips were evaluated per condition per experiment, and the results of 5–6 independent experiments were combined for presentation. In the second experiment cells were transfected and treated with oleic acid as above. After fixation, cellular neutral lipids were stained with Lipidtox Deep Red (1:1000, Molecular Probes) and imaged in a wide-field using a 40× objective. Five non-overlapping fields were captured for each transfection condition using identical capture parameters. Cellular Lipidtox fluorescence was quantified by using the segmentation tool of IPlabs software to identify specific staining of lipid droplets and eliminate nonspecific cytosolic staining (threshold set at 380 or 400 grayscale units of 12 bit images). Identical segmentation parameters were used for all transfection conditions of a given experiment. Total fluorescence of the segmented areas was summed for each cell in a field and averaged for each experiment. Normalized data from three independent experiments were combined for presentation and statistical analysis.

Statistical analyses

Differences in colocalization coefficients, total fluorescence, and luciferase activity were evaluated by one-way analysis of variance. Group means were compared by Bonferroni’s test for multiple comparisons. BiFC efficiency was assessed by χ².

RESULTS

Abhd5 and Mldp Are Colocalized on Cardiomyocyte Lipid Droplets—Mldp is highly expressed in tissues like heart with high oxidative capacity, whereas Adrp is more widely expressed (5). We examined the subcellular localization of these endogenous PAT proteins and Abhd5 in intact microdissected mouse cardiac muscle fibers (Fig. 1). Mldp and Adrp were targeted to numerous small lipid droplets that were aligned between myofibrils. Mldp and Adrp were partially colocalized on individual droplets; however, numerous lipid droplets could be readily found that contained one or the other droplet protein. Although there was considerable overlap in the sizes of droplets containing these PAT proteins, droplets identified by Mldp immunofluorescence were on average larger than those identified by Adrp immunofluorescence (0.57 ± 0.09 versus 0.47 ± 0.08 μm, mean ± S.D., p < 0.0001). These data indicate that cardiac lipid droplets are heterogeneous with respect to size and protein content.

The degree of colocalization between Abhd5 and the droplet scaffold proteins was evaluated by double label immunofluorescence. Confocal imaging demonstrated that virtually every individual lipid droplets that contained Mldp immunofluorescence also contained proportional levels of Abhd5 (Fig. 1A). In contrast, numerous droplets containing Adrp lacked detectable Abhd5 and vice versa (Fig. 1B). Averaging the results of line-scan analysis from several cells demonstrated that Mldp and Abhd5 were significantly more colocalized (p < 0.0001) than were Adrp and Abhd5, suggesting a functional relationship between Mldp and Abhd5.

Expression of Mldp Targets Abhd5 to Lipid Droplets—Mldp-EYFP expression in 3T3-L1 fibroblasts led to the formation of small, highly uniform lipid droplet clusters (Fig. 2, A and B). Such
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uniform lipid droplet clusters were not observed in nontransfected cells or in cells transfected with control vector or Abhd5 alone. By contrast, ECFP-Abhd5 appeared to be targeted to perinuclear membranes (Fig. 2C) with minimal targeting to randomly distributed intracellular droplets (Fig. 2D). When cotransfected, Abhd5 and Mldp were highly targeted to lipid droplets (Fig. 2, E and F). The subcellular targeting of ECFP-Abhd5 at individual lipid droplets of any given cell was precisely predicted by subcellular localization of Mldp-EYFP.

The co-expression data demonstrates that subcellular targeting of Abhd5 is directed by Mldp, possibly through direct binding. To explore this possibility further, we examined whether recombinant ECFP-Abhd5, generated from 293T cell lysates, would bind to Mldp in permeabilized 3T3-L1 cells. As shown in Fig. 3, ECFP-Abhd5 bound to cells expressing Mldp-EYFP but not to nontransfected cells or to cells expressing a control EYFP-tagged protein. Of several hundred cells examined, all cells expressing Mldp bound Abhd5, whereas control cells did not exhibit binding above the levels seen in nontransfected controls. Importantly, the binding of ECFP-Abhd5 was directly proportional to Mldp-EYFP concentration (supplemental Fig. 1A) and unrelated to cellular lipid content or the presence of lipid droplets, as detected by nile red staining.

FRET measurements demonstrated that bound ECFP-Abhd5 was an effective FRET donor with Mldp-EYFP, strongly indicating that these proteins interact directly (supplemental Fig. 1B). Interestingly, although minor colocalization of Mldp-EYFP and ECFP-Abhd5 could be observed in the cytoplasm away from lipid droplets, FRET signals were only observed on the surface of droplets.

The Interaction of Abhd5 and Mldp Mediates Protein Complementation in Cells and in Cell-free Assays—Protein complementation analysis is a technique for assessing the interaction of proteins in live cells (23) and is based on the ability of interacting proteins to reconstitute functional activity of a reporter protein (e.g. EYFP and luciferase) that has been split in two and fused to the test proteins of interest.

Complementary split luciferase fragments fused to Abhd5 and Mldp generated robust luciferase activity when co-expressed in lipid-loaded 293T cells (Fig. 4A) or 3T3-L1 cells (not shown). The luciferase activity was comparable with that generated by the interaction of Plin with Abhd5, which served as a positive reference (12, 13). By contrast, minimal activity was observed when Abhd5 or Mldp constructs were cotransfected with Rab18 or Adrp (see below), which are targeted to lipid droplets (24–26). Ln-Abhd5 and Mldp-Lc that were synthesized in vitro using a bacterial transcription/translation system (Fig. 4B) also reconstituted luciferase activity, demonstrating that the interaction occurs without additional mammalian proteins.

BiFC analysis was used to localize the intracellular sites of Abhd5–Mldp interactions. Strong BiFC was observed in >95% of cells expressing complementary EYFP fragments fused to Mldp and Abhd5 (supplemental Fig. 2, p < 0.0001 versus control). Importantly, BiFC signals were observed only on the surface in lipid droplets.

Lipid Loading Promotes Abhd5–Mldp Interactions—The above data demonstrated that Abhd5 and Mldp interact on lipid droplet surfaces and suggested that lipid droplet formation might facilitate this interaction. To test this possibility, 293T cells were transfected with combinations of tagged Mldp and Abhd5 or control vectors and exposed to 400 μM oleic acid (OA) complexed to BSA or BSA alone for 1 h (Fig. 5A). OA treatment increased Mldp and Abhd5 complementation by greater than 2.5-fold (p < 0.001). Interestingly, cotransfection of Mldp fused to complementary luciferase fragments also reconstituted strong luciferase activity, indi-

FIGURE 6. E262K mutation of Abhd5 disrupts its interaction with Mldp. A, COS-7 cells were transfected with Mldp-EYFP and treated with oleic acid overnight. Cells were fixed and permeabilized and incubated with 293T cell lysates containing equal concentrations of wild type or E262K Abhd5-Cherry. Binding of wild type Abhd5 was performed three times with similar results.

B, luciferase complementation assays. E262K mutation disrupted binding of Mldp but not to Atgl. Shown is a representative experiment performed in quadruplicate. The experiment was performed three times with similar results.
cating that Mldp can form oligomers in vivo. In contrast to the Abhd5-Mldp, however, the homotypic Mldp interaction was not significantly affected by OA. Intact Gaussia luciferase was not affected by OA (not shown), whereas control transfections exhibited low activity that was not significantly affected by OA. No specific binding of wild type Abhd5 was observed in cells expressing Atgl alone (middle row). Mutant Abhd5 failed to bind lipid droplets containing Mldp and Atgl (bottom row). AFU, arbitrary fluorescence units, $r^2$, coefficient of determination for the linear association. $b$, linear regression slope indicating amount of Abhd5 bound per unit Atgl. $B$, expression of Mldp-ECFP increases the interaction of wild type, but not mutant Abhd5 with Atgl in protein complementation assay. PC, protein complementation. Values are the means ± S.E. for eight independent experiments.

FIGURE 7. Expression of Mldp increases the colocalization and interaction of wild type, but not mutant Abhd5 and Atgl. A, COS-7 cells were transfected with Atgl-EYFP with Mldp-ECFP or Atgl alone, permeabilized, and incubated with equal concentrations of wild type or E262K Abhd5-Cherry. Colocalization of bound Abhd5 with Atgl was determined by line-scan analysis (right column) of individual cells, identified by the asterisk in Atgl images. Wild type Abhd5-bound lipid droplets containing Mldp and Atgl (top row). No specific binding of wild type Abhd5 was observed in cells expressing Atgl alone (middle row). Mutant Abhd5 failed to bind lipid droplets containing Mldp and Atgl (bottom row). AFU, arbitrary fluorescence units. $r^2$, coefficient of determination for the linear association. $b$, linear regression slope indicating amount of Abhd5 bound per unit Atgl. $B$, expression of Mldp-ECFP increases the interaction of wild type, but not mutant Abhd5 with Atgl in protein complementation assay. PC, protein complementation. Values are the means ± S.E. for eight independent experiments.

The effect of lipid OA could be observed within 15 min and was maximal after 2–4 h (not shown), suggesting that the effect likely involves lipid metabolism rather than a direct effect of the free fatty acid. To explore this further, we examined the effects of Triacsin-C, a potent acyl-CoA synthase inhibitor that blocks the synthesis of triglyceride from oleic acid (19–21). Triacsin-C nearly eliminated the ability of OA to increase the interaction of Abhd5 and Mldp, as determined by luciferase complementation (Fig. 5B).

**E262K Mutation of Mouse Abhd5 Disrupts Its Interaction with Mldp**—Certain point mutations of ABHD5 produce a neutral lipid storage disease in humans (9, 27), and one such mutation (E260K) was found to disrupt the interaction of ABHD5 with perilipin (13). We examined the effects of the homologous mutation in mouse Abhd5 (E262K) on interactions with Mldp. Wild type Abhd5-Cherry prepared from 293T cells bound Mldp-EYFP in permeabilized COS-7 cells, whereas binding of mutant Abhd5 was reduced by >90% (Fig. 6A). Additionally, E262K mutation of Abhd5 reduced its interaction with Mldp by >85% in luciferase complementation assays (Fig. 6B). Wild type Abhd5 interacted with Atgl in the luciferase complementation assay, although the mag-
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Abhd5-Mldp Interaction Regulates Atgl Activity at Lipid Droplets Containing Mldp—Abhd5 is a critical regulator of neutral lipid accumulation because of its ability to regulate Atgl activity. The functional significance of the Abhd5-Mldp interaction on neutral lipid balance was examined in transiently transfected COS-7 cells. In the absence of Mldp, ~55% of cells expressing ECFP-Atgl accumulated lipid droplets when exposed to oleic acid (Fig. 8A). Co-expression of wild type ECFP-Abhd5 or ECFP-E262K Abhd5 virtually eliminated lipid droplet accumulation in cells expressing Atgl. These effects were absent in cells expressing lipase-dead (S47A) Atgl, demonstrating that the effects of wild type and mutant Abhd5 were mediated via Atgl activity. In the presence of Mldp-EYFP, nearly all cells expressing ECFP-Atgl exhibited well defined clusters of lipid droplets. Co-expression of wild type Abhd5 eliminated formation of lipid droplets, whereas co-expression of E262K was without effect.

The above experiment was repeated, and the cellular neutral lipid content was quantified by Lipidtox fluorescence, as detailed under “Experimental Procedures” (Fig. 8B; see supplemental Fig. 3 for representative images). In the absence of Mldp, both wild type and mutant Abhd5 significantly reduced neutral lipid accumulation. Expression of Mldp-EYFP increased accumulation of neutral lipid in the presence of Atgl by nearly 6-fold ($p < 0.001$). Co-expression of wild-type Abhd5 eliminated accumulation of neutral lipid ($p < 0.0001$), whereas co-expression of the E262K mutant was without effect.

DISCUSSION

The present study confirms that Mldp is highly expressed in cardiac myocytes and further shows that Abhd5 and Mldp are highly co-localized on individual lipid droplets. In contrast, Abhd5 immunofluorescence of myocyte lipid droplets was largely unrelated to Adrp content, even though numerous droplets were highly enriched in this scaffold protein. These results demonstrated that cardiomyocyte lipid droplets are heterogeneous with respect to PAT protein composition and suggested there is a functional relationship between Mldp and Abhd5.

The functional interactions between Mldp and Abhd5 were explored in transfected cells using fluorescently tagged proteins and protein complementation assays. As previously reported, ECFP-Abhd5 was largely cytosolic when expressed in 3T3-L1 fibroblasts (7, 12, 13), whereas Mldp-EYFP expression led to the generation of clustered lipid droplets, with Mldp targeted to the droplet surface. Co-expression recruited ECFP-Abhd5 to lipid droplets containing Mldp-EYFP, and the concentration of Abhd5 on droplets of individual cells was precisely predicted by the concentration of Mldp. ECFP-Abhd5 prepared from 293T cell extracts specifically bound to Mldp-containing lipid droplets in permeabilized 3T3-L1 cells and not to endogenous droplets that contain Adrp. Abhd5 binding was proportional to Mldp concentration and supported FRET between the ECFP-Abhd5 donor and Mldp-EYFP acceptor, indicating a close (~10 nm), if not direct interaction.

Although Mldp and Abhd5 can be cytosolic, FRET and BiFC experiments indicate the interaction of these proteins occurs mainly, if not exclusively, on lipid droplets. Oleic acid treatment rapidly promoted the interaction in a manner dependent on triglyceride synthesis. These observations...
suggest that the interaction of Mldp with droplet surface increases its affinity for Abhd5.

Growing evidence indicates that proper trafficking of lipolytic effectors to lipid droplets is critical in the regulation of cellular lipolysis (3). For example, mutations that mistarget human ATGL disrupt lipolysis in the context of the living cell yet do not disrupt lipase activity against artificial substrates (28). The present experiments support this general conclusion by demonstrating that Mldp coordinates the physical and functional interaction of Abhd5 and Atgl.

Imaging of tagged proteins demonstrated that Abhd5 and Atgl are largely cytosolic proteins and that Mldp expression leads to the formation of intracellular lipid droplets containing each of these proteins. Furthermore, binding and complementation experiments demonstrate that Mldp expression greatly increases the subcellular targeting of Abhd5 to droplets containing Atgl and thereby augments their interaction.

The significance of the interaction of Abhd5 with Mldp was investigated using wild type and mutant Abhd5. Previous investigators have reported that E260K mutation of human ABHD5 blocks its interaction with perilipin, a related PAT protein (1), but blocks its interaction with perilipin, a related PAT protein (1), and that E262K mutation of human ABHD5 leads to the formation of intracellular lipid droplets containing Atgl and thereby augments their interaction.

In summary, the present study shows that Mldp directs Abhd5 to lipid droplets through, if not direct interactions. The interaction of Abhd5 and Mldp is promoted by lipid loading, and this interaction is critical for regulating Atgl activity at Mldp-containing droplets.

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