The phosphatidic acid–binding, polybasic domain is responsible for the differences in the phosphoregulation of lipins 1 and 3

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Lipins 1, 2, and 3 are Mg2+-dependent phosphatidic acid phosphatases and catalyze the penultimate step of triacylglycerol synthesis. We have previously investigated the biochemistry of lipins 1 and 2 and shown that di-anionic phosphatidic acid (PA) augments their activity and lipid binding and that lipin 1 activity is negatively regulated by phosphorylation. In the present study, we show that phosphorylation does not affect the catalytic activity of lipin 3 or its ability to associate with PA in vitro. The lipin proteins each contain a conserved polybasic domain (PBD) composed of nine lysine and arginine residues located between the conserved N- and C-terminal domains. In lipin 1, the PBD is the site of PA binding and sensing of the PA electrostatic charge. The specific arrangement and number of the lysines and arginines of the PBD vary among the lipins. We show that the different PBDs of lipins 1 and 3 are responsible for the presence of phosphoregulation on the former but not the latter enzyme. To do so, we generated lipin 1 that contained the PBD of lipin 3 and vice versa. The lipin 1 enzyme with the lipin 3 PBD lost its ability to be regulated by phosphorylation but remained downstream of phosphorylation by mammalian target of rapamycin. Conversely, the presence of the lipin 1 PBD in lipin 3 subjected the enzyme to negative intramolecular control by phosphorylation. These results indicate a mechanism for the observed differences in lipin phosphoregulation in vitro.

In vertebrates, the lipin family of Mg2+-dependent phosphatic acid (PA)3 phosphatases consists of three members (lipins 1–3), which form diacylglycerol from PA in neutral and phospholipid synthesis (1–6). Lipins are cytosolic enzymes that must associate with membranes to access substrate (1, 7–9).

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3 The abbreviations used are: PA, phosphatidic acid; mTOR, mammalian target of rapamycin; PE, phosphatidylethanolamine; PBD, polybasic domain; NEM, N-ethylmaleimide; βME, β-mercaptoethanol; SRD, serum-rich domain; PC, phosphatidylcholine.
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Additionally, we present evidence that the specific PBDs of lipins 1 and 3 are responsible for the observed differences in their in vitro phosphoregulation.

Results

Purification and enzymatic activity of recombinant lipin 3

Adenovirus was used to overexpress Mus musculus FLAG-lipin 3 in HeLa cells. After 2 days, lipin 3 protein was affinity-purified, eluted with FLAG peptide, and dialyzed. An aliquot of the purified lipin 3 was resolved by SDS-PAGE, and the gel was stained with Coomassie Blue. B, PAP enzymatic activity of 50 ng of purified lipin 3 was measured as a function of time in Triton X-100/PA mixed micelles. C, PAP activity of lipin 3 against PA at a constant concentration of 0.5 mM with an increasing concentration of Triton X-100. D, PAP activity of lipin 3 as a function of pH. E, PAP activity of lipin 3 as a function of MgCl2. F, PAP activity of lipin 3 against as a function of MnCl2. G, PAP activity of lipin 3 with increasing concentrations of NEM. H, PAP activity of lipin 3 with increasing concentrations of β-ME. Experiments shown in D–H were performed using Triton X-100/PA mixed micelles with a surface concentration and bulk concentration of PA at 10 mol % and 0.5 mM, respectively. Each data point is a mean of triplicate determinations ± S.E. (error bars).

Phosphorylation of lipin 3 and sensing of PA charge

Phosphorylation has been shown to play an important role in the regulation of lipin 1 PAP activity (10, 11, 14). To investigate the phosphorylation of lipin 3, the purified protein was trypsin-digested and subjected to LC-MS/MS. Phosphorylated residues were assigned a site localization probability describing the confidence that a particular site within the sequence region is phosphorylated. We found 15 novel phosphorylation sites with localization probabilities of 0.95 or greater throughout the length of the protein (Fig. 2A and Table 1). An important phosphorylation site for lipin 1 regulation is Ser106. This site was previously identified on lipins 1 and 2 (10, 15). A novel phosphorylation site for lipin 3 was identified at Ser186, which is conserved in lipins 1 and 2 (10, 15, 25). The addition of reducing agents stimulates the activity of lipins 1 and 2 (15, 25). Indeed, 20 mM β-mercaptoethanol (β-ME) maximally stimulates lipin 3 activity (Fig. 2B).
of this site on lipin 3, a confidently identified peptide with phosphorylation on Thr^{105} or Ser^{106} was observed in lipin 3 (see Table 5 (in red) and supplemental Table S1). An important lipin protein region, enriched in phosphorylation sites, is the serine-rich domain (SRD). The SRD contains several 14-3-3 consensus-binding motifs and is important for mediating lipin 1 subcellular localization (13). On lipin 3, the region corresponds to amino acids Ser^{185}–Ser^{230}, of which Ser^{198}, Ser^{218}, Ser^{220}, and Ser^{230} are phosphorylated with a localization probability of 0.95 or greater and Ser^{215} falls just short of statistically significant localization probability of 0.93 (Table 1 and supplemental Table S1). Phosphorylation on residue Ser^{218} is also present in lipins 1 and 2 (Ser^{285} in lipin 1 and Ser^{243} in lipin 2), and Ser^{287} is also phosphorylated in lipin 1 (homologous to Ser^{220} in lipin 3), albeit with a localization probability of 0.77 (Table 1 and supplemental Table S1) (10, 15).

To determine whether phosphorylation affects lipin 3 enzymatic activity, FLAG-lipin 3 was purified using affinity chromatography as described above. Before protein elution, the protein was incubated in phosphatase buffer with and without protein phosphatase, eluted, and dialyzed. This yielded phosphorylated (−α) and dephosphorylated (+α) lipin 3 (Fig. 2B). We veri-
Phosphorylation of lipin 3 in 3T3-L1 cells

Lipin 1 is reported to be phosphorylated and negatively regulated upon acute insulin treatment, in a rapamycin- and Torin 1-sensitive manner (10, 11, 13, 14). We sought to determine whether lipin 3 is similarly affected by phosphorylation. Lipins 1 and 3 were expressed in differentiated 3T3-L1 cells by adenoviral transduction. Two days post-infection, the cells were serum-starved in low-phosphate buffer for 2 h, incubated with 32P orthophosphate, and treated with vehicle or a pan-mTOR inhibitor, Torin 1, for 45 min plus 10 milliunits/ml insulin for the last 15 min or insulin alone, and phosphate incorporation into lipin 3.

The PAP activity of PBD mutants with liposomes

All three lipin family members contain homologous PBDs immediately C-terminal to the N-terminal domain at residues 153–161 in lipin 1 and 136–144 in lipin 3 (Fig. 4A) (23, 25). In lipin 1, deletion of this site excluded the enzyme from the nucleus (22). Mutation of the lipin 1 PBD to alanines confirmed that it is a nuclear localization sequence and further that it is critical for PA binding but not for PAP activity (13, 14, 22). Our laboratory has shown that PBD is important for sensing of the PA electrostatic charge, and phosphorylation sites within lipin 1 may hinder PA association with the PBD (14). Our data thus
Figure 3. Phosphorylation of lipin 3 in 3T3-L1 cells. A, 3T3-L1 cells infected with adenovirus expressing FLAG-lipin 3 were serum-starved in low-phosphate buffer for 2 h, incubated with 0.2 mCi/ml orthophosphate, and treated as indicated. The cells were harvested, incubated with anti-FLAG beads for 4 h, displaced from beads, and separated on an SDS-polyacrylamide gel and transferred to PVDF. Top, a phosphor image of $^{32}$P incorporation into lipin 3 (top; $^{32}$P) and an immunoblot image of total lipin 3 protein (bottom; lipin 3). Bottom, quantitation of three independent experiments. B, the experiment was performed as in A, but with FLAG-lipin 1. Shown are scatter plots with mean of the quantitation of three independent experiments ± S.E. (error bars). One-way ANOVA was used to analyze statistical significance followed by Sidak post hoc analysis. *, $p < 0.05$; N.S., no statistical significance.

Figure 4. Enzymatic activity of lipin PBD exchange mutants in liposomes. A, schematic of lipin PBD exchange mutants. B, PBD mutants were expressed, affinity-purified, incubated in phosphatase buffer with (+ $\lambda$) or without (− $\lambda$) $\lambda$ phosphatase, and eluted as described before. Proteins were separated on an SDS-polyacrylamide gel along with BSA standards and stained with Coomassie Blue dye. The PAP activities of purified phosphorylated (− $\lambda$) and dephosphorylated (+ $\lambda$) lipin 1 (3PBD) were measured with PC/PA (C) and PC/PE/PA (D) liposomes. Shown are the PAP activities of purified lipin 3 (1PBD) with PC/PA (E) and PC/PE/PA (F) liposome. Activities were assayed at pH 7.5. Each data point is a mean of triplicate determinations ± S.E. (error bars).
far contrast with those for lipin 1 and indicate that under the present conditions, lipin 3 phosphorylation does not prevent its recognition and binding to di-anionic PA (Fig. 2).

We wondered about the molecular basis for such differences in the phosphoregulation of lipin enzymes. One possibility is that the distinct PBDs of lipins 1 and 3 are at least partially responsible for conferring sensitivity to phosphorylation. To test this hypothesis, constructs exchanging the PBD of WT lipin 1 with that of WT lipin 3 (lipin 1 (3PBD)) and the PBD of WT lipin 3 with that of WT lipin 1 (lipin 3 (1PBD)) were expressed. The PBD mutant constructs (Fig. 4A) were purified from HeLa cells to yield phosphorylated (−λ) and dephosphorylated (+λ) proteins (Fig. 4B). The PAP activities of the PBD exchange mutants were tested in the context of PC/PA and PC/PE/PA liposomes. Insertion of the lipin 3 PBD in lipin 1 eliminated the ability of phosphorylation to hinder the recognition of the electrostatic state of PA, and PE significantly augmented the specific activity of both phosphorylated and dephosphorylated enzyme (Fig. 4, C and D). As seen in Table 3, the \( K_{\text{m, app}} \) did not significantly decrease with 30 mol % PE, but the catalytic efficiency and turnover number of the lipin 1 (3PBD) mutant increased independent of phosphorylation.

In contrast, only dephosphorylated lipin 3 (1PBD) activity was significantly augmented in the presence of 30 mol % PE, but the activity of phosphorylated lipin 3 (1PBD) was not (Fig. 4, E and F). The \( k_{\text{cat}} \) of dephosphorylated lipin 3 (1PBD) increased by about 2.5-fold, compared with 1.3-fold for the phosphorylated enzyme (Table 3). These data suggest that the phospho-

regulation of lipin 1 occurs, at least in part, because of its specific PBD.

**The PAP activity of PBD mutants within micelles**

In the context of Triton X-100/PA mixed micelles, the activity of WT lipin 1 is significantly higher than its specific activity in liposomes (14). In contrast, lipin 3 activity is closely matched between these two modes of substrate presentation. It is possible that lipin 3 enzymatic activity is sensitive to inhibition by detergent. To test whether the variation in lipin 1 and 3 PAP activity in micelles is due to the PBD, we measured the activity of the lipin PBD mutants under the described conditions. The lipin 1 (3PBD) displayed specific activity comparable with that of WT lipin 1 (Fig. 5A) (14). Lipin 3 (1PBD) activity was also similar to that of WT lipin 3 (Figs. 2E and 5B).

**The binding of lipin 1 and 3 PBD mutants to PC/PA liposomes**

The lipin PBDs are the main site of substrate interaction (14, 22). It is likely that changes in activity caused by the PBD mutants are due to altered association of the mutant enzymes with PA. Using liposome flotation assays, we investigated the ability of phosphorylated and dephosphorylated Venus-lipin 1 (3PBD) and Venus-lipin 3 (1PBD) purified enzymes to associate with PA-containing liposomes. The binding of Lipin 1 (3PBD) increased by about 25–30% in the presence of 30 mol % PE regardless of phosphorylation state (Fig. 6, A and B). The binding of phosphorylated lipin 3 (1PBD) increased slightly from 0 to 30 mol % PE, whereas the dephosphorylated enzyme binding increased by ~40% from PC/PA to PC/PE/PA liposomes (Fig. 6, C and D). It should be pointed out that the PBD exchange mutants were catalytically active but displayed significantly reduced \( K_{\text{m, app}} \) for PA, compared with WT lipins 1 and 3 (Fig. 4 (C–F) and Table 3) (14), suggesting that the PBD exchange did not affect affinity to bulk substrate. However, \( K_{\text{m, app}} \) takes into account both bulk and surface concentration (24).

**Phosphorylation of Ser\(^{106}\) on lipin 1 (3PBD)**

There are two possibilities by which exchanging the PBDs alters the ability of phosphorylation to regulate enzymatic

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**Table 3**

*Steady-state kinetic data for phosphorylated (−λ) and dephosphorylated (+λ) lipin PBD mutants*

The data were generated from assays using PC/PA or PC/PE/PA liposomes containing 10 mol % PA, the indicated mol % PE, and additional PC to bring the combined concentration of phospholipids to 100 mol %. The specific activity was measured as \( V_{\text{max}}/\text{mg protein} \). Student’s \( t \) test was used to analyze statistical significance.

| PE | \( K_{\text{m, app}} \) mol % | \( K_{\text{cat}} \) s\(^{-1} \) | \( K_{\text{cat}}/K_{\text{m, app}} \) \( \mu \text{M}^{-1} \text{s}^{-1} \) |
|----|-----------------|-----------------|-----------------|
| Lipin 1 (3PBD) | | | |
| −λ | 0 | 211 ± 70 | 0.76 ± 0.23 | 0.003 |
| −λ | 0 | 191 ± 88 | 1.66 ± 0.17 | 0.008 |
| +λ | 0 | 170 ± 99 | 0.64 ± 0.06 | 0.004 |
| +λ | 0 | 146 ± 54 | 1.64 ± 0.34 | 0.011 |
| Lipin 3 (1PBD) | | | |
| −λ | 0 | 120 ± 69 | 0.67 ± 0.09 | 0.005 |
| −λ | 30 | 117 ± 31 | 0.92 ± 0.05 | 0.007 |
| +λ | 0 | 107 ± 40 | 0.78 ± 0.19 | 0.007 |
| +λ | 30 | 155 ± 38 | 2.02 ± 0.26 | 0.014 |

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**Figure 5. Enzymatic activity of lipin PBD exchange mutants in Triton X-100/PA micelles.** Phosphorylated (−λ) and dephosphorylated (+λ) lipin 1 (3PBD) (A) and lipin 3 (1PBD) (B) activities were assayed using Triton X-100/PA mixed micelles at pH 7.5. The surface concentration of PA was 10 mol %. Each data point is a mean of triplicate determinations ± S.E. (error bars).
activity. Either the phosphorylation sites themselves are altered by the generation of the PBD mutants or the existing phosphorylation sites can regulate the PBD of lipin 1 but not lipin 3. Serine 106 is a conserved site among the lipins and in lipin 1 is phosphorylated in response to insulin, downstream of mTOR (10, 12). Ser106 on lipin 2 is not responsive to insulin stimulation or Torin 1 treatment (15). We attempted to investigate the phosphorylation of this site on lipin 3 and the PBD mutants but could not detect phospho-Ser106 on lipin 3 or lipin 3 (1PBD) using Western blot analysis. Lipin amino acid alignments revealed that the region surrounding Ser106 that was used to generate phospho-specific antibodies is conserved within lipins 1 and 2 but not in lipin 3. The LC-MS/MS analysis of lipin 3 revealed that either Thr105 or Ser106 was phosphorylated, but the precise localization could not be resolved with statistical significance (supplemental Table S1). Therefore, we investigated the regulation of Ser106 phosphorylation in lipin 1 (3PBD). Wild-type lipin 1 and lipin 1 (3PBD) proteins were overexpressed in HeLa cells and treated with vehicle or 250 nM Torin 1. Changes in the phosphorylation of Ser106 were detected using the phospho-Ser106 antibody. Under basal conditions, the residue Ser106 on lipin 1 (3PBD) was phosphorylated (Fig. 7A). Treatment with Torin 1 decreased phosphorylation of Ser106 on WT lipin 1, as reported previously (Fig. 7, A and B) (12). Exchanging the lipin 1 PBD with the PBD of lipin 3 had no effect on the ability of Torin 1 to inhibit phosphorylation of this site (Fig. 7, A and B). In addition, gel mobility shifts between phosphorylated and dephosphorylated PBD mutants were similar to what has been seen with the WT enzymes (Fig. 7C). Together, immunoblotting data suggest that mutation of the lipin 1 PBD does not alter the phosphorylation of Ser106 and does not eliminate the lipin 1 (3PBD) as a downstream substrate of mTOR.

To more thoroughly investigate whether the mutation of the lipin PBD domains altered the global phosphorylation of the rest of the protein, we analyzed phosphorylation sites of lipin 1 (3PBD) and lipin 3 (1PBD) using LC-MS/MS analysis and compared them with those of WT lipins 1 and 3. Previous work investigating lipin 1 phosphorylation was performed on HA-tagged lipin 1 purified from 3T3-L1 cells after acute insulin stimulation (10). To maintain consistency in protein acquisition, purification, and LC-MS/MS analysis, we purified FLAG-tagged WT lipin 1 from HeLa cells in an identical manner and analyzed phosphorylation sites for all four proteins.

Twenty-two phosphorylation sites were identified on WT lipin 1 and 16 on lipin 1 (3PBD) and localized with a probability of 0.95 or greater (Table 4). Many of the sites found on WT lipin 1 agreed with previous reports (10). Of the 16 phosphosite on lipin 1 (3PBD), 11 were in common with WT lipin 1, including Ser106 and sites within the SRD (Table 4 and Fig. 8). Fifteen phosphorylation sites were identified on lipin 3 and 16 on lipin 3 (1PBD).
Table 4

WT lipin 1 and lipin 1 (3PBD) phosphorylation analysis by LC-MS/MS

| Modified Sequence | Residue | WT Lipin 1 Loc Prob | Lipin 1 (3PBD) Loc Prob |
|-------------------|---------|---------------------|-------------------------|
| IIP(M)ox|YLATS(ph)PILSE | S106 | 0.99 | 1 |
| IIPMT(YLATS(ph)PILS(ph)E) | S110 | - | 0.99 |
| Tiph(LNPDYPPQQDDIPK | S124 | 0.99 | - |
| EWS(ph)PSSLVDCQR | T200 | 1 | - |
| WSPS(ph)PSSLVDCRTTPHCLAE | S237 | 0.99 | 0.97 |
| Tiph(PPLAEGVLS(VS)CPGLSCHFHAESPSGSR | S239 | - | 0.99 |
| TPSGSRPST(ph)PKSDSE | T249 | 1 | 1 |
| PSTPKS(ph)DSELVLSK | T262 | - | 0.82 |
| SDDs(ph)ELVSK | S287 | 0.98 | 0.98 |
| LVS,(ph)ADRLTPKNNLE | S293 | 0.77 | - |
| LVSDKADRL(ph)KNNLE | T296 | 0 | 0.99 |
| SPS(ph)MHKESPPLSR | S320 | 1 | 0.05 |
| ESS(ph)PLGSRK | S328 | 0.99 | 0.99 |
| Tiph(PDKMNQFAHSESSDTSDFQSTMAR | T335 | 1 | 0.99 |
| SSTDTSF(ph)JQOS(ph)PTMARGLHISSQKAOTE | S353 | 0 | 1 |
| MNFQAIHSESSDTSDFQ(ph)PTMAR | S356 | 0.99 | 0.99 |
| AQTEMOFVNEEDLSGLAAAAPPS(ph)PVAEELK | S392 | 1 | 0.05 |
| SRHLGADGVYVLDLT(ph)DMDPEVAYLFPK | T437 | 1 | - |
| SANQS(ph)PSVSGGGSDGVESTSDSLR | S472 | 0.99 | 1 |
| SANQS(ph)PSVSGGG(ph)SIDGVESTSDSLR | S479 | 0.76 | 0.99 |
| SANQS(ph)PSVSGGG(ph)GVESTDSLR | S483 | 0.38 | 0.69 |
| NAT(ph)KEEKSVEQCLTGK | T597 | 1 | - |
| GHGNGTEGQAPQGLAT(ph)R | T626 | 0.05 | - |
| Tiph(LRLTSEQLKL | T661 | 0.99 | - |
| SDLT(ph)GHLPHLPGKDQWQIAK | T772 | 0.99 | 1 |
| GTVLQGPSLLL(ph)PSSLFSALHR | S789 | 0.95 | - |
| SHS(ph)CDFFCSSDFSSTFTWFR | S802 | 1 | 0.83 |
| EPLPFPENQDMHAS(ph)A | S923 | 0.95 | 0.95 |

Discussion

The lipins are enzymes with complex and multifunctional roles in cellular lipid metabolism. Whereas lipins serve the same catalytic function to dephosphorylate PA, their roles on the physiological level are distinct. Deleterious mutations in Lpin1 gene result in hepatic and adipose irregularities in mice, whereas in humans they cause muscle rhabdomyolysis (23, 27, 28). Mice lacking lipin 2 appear to have age-dependent neuronal dysfunction, and human LPIN2 mutations have been linked to the rare immunological disorder Majeed syndrome (29). Lipin 3-deficient rodents do not show an obvious phenotype but do not show an obvious phenotype but may be critical for adipogenesis (30). Such findings are perhaps a reflection of the intricate and distinct means by which these enzymes are regulated on the molecular level. The in vitro control of lipin 1 localization and activity by phosphorylation and the electrostatic charge of PA is well understood, but in these contexts, lipin 3 is relatively understudied.
in vitro. These findings are similar to what was observed for lipin 2 but contrast with what is known of lipin 1 phosphoregulation (14, 15). The molecular basis for such differences among enzymes, which otherwise catalyze the same reaction, is unclear.

One possibility is that the protein-wide phosphorylation of lipins 1 and 3 is intrinsically unique. Both proteins are highly phosphorylated, as shown previously for lipin 1 and demonstrated here by LC-MS/MS analysis and the incorporation of $^{32}$P (Figs. 2 and Table 1) (10). However, subtle differences in phosphorylation may alter enzyme activity and regulation. In lipin 1, the SRD is a region enriched in serines and involved in the nucleocytoplasmic trafficking of this protein (10, 13). Lipin 3 contains a homologous domain that contains nine serines and one threonine. Of these, we found that four serines are phosphorylated (Fig. 2A and Table 1). One serine in this region, Ser226, is phosphorylated in lipin 1 (as determined by Harris et al. (10)) but not in lipin 3. Although this may play a role in the regulatory differences between lipins 1 and 3, it is probably not the only factor.

PA effector proteins, including lipins, do not contain a recognized lipid association motif, but their recruitment and ability to interact with membranes is important for enzymatic activity (14, 17). Kooijman et al. (18) showed that small peptides composed of lysines and arginines can alter the electrostatic charge of PA by lowering $pK_a$ below the physiological pH and causing proton dissociation. Basic residues in native proteins demonstrate these principles. For example, the FRBP12 rapamycin-binding domain of mTOR contains a single arginine, crucial for the interaction of the FRBP12 rapamycin-binding domain with PA (31). Other examples include Raf-1, which binds PA at a non-conserved sequence of 35 amino acids that includes a cluster of four positively charged residues responsible for the initial electrostatic attraction between this enzyme and lipid (32).

In yeast, the best-described example of protein-PA association is with Opi1, a gene repressor (33–35). The activity of Opi1 is regulated by its sequestration to the endoplasmic reticulum membrane, mediated by binding to PA (36). Young et al. (37) showed that a basic domain in Opi1 is responsible for direct interaction with PA, and this interaction was diminished when cells were acidified. Further, they identified two lysines and one arginine as being critical for the pH-dependent electrostatic interactions with this lipid. A methyl-PA lacking the $pK_a$ displayed weak and pH-independent binding to Opi1 (37, 38). These studies eloquently demonstrate the dynamic nature of PA ionization state and the importance of the interactions between basic residues and PA for proper protein function.

Due to the significance of basic residues in protein-PA association, we hypothesized that the unique PBD of lipin 1 plays an important role in the ability of phosphorylation to regulate its activity. In the present work, we show that replacing the lipin 1 PBD with that of lipin 3 eliminates the ability of phosphorylation to negatively regulate lipin 1 activity (Fig. 4, C and D). Conversely, the specific activity of lipin 3 (1PBD) increased significantly only after treatment with protein phosphatase and

| Modified Sequence | Residue | WT Lipin 3 Loc Prob | Lipin 3 (1PBD) Loc Prob |
|-------------------|---------|---------------------|-------------------------|
| LGVLRS(ph)REK     | S60     | 1                   | -                       |
| LODSGEAFFVQELDS(ph)DEEDVPR | S94 | 1 | 1 |
| LCTIS(ph)IPWGLGSLGPS(ph)DSQIGTASEPEGLVTGK | S106 | 0.50 | 0.54 |
| LTLEKPT(ph)PES(ph)P5AQE | T175 | 1 | 1 |
| PTPES(ph)P5AQEEAEPPSQPK | S178 | 0.96 | 0.99 |
| PTPES(ph)P5AQEEAEPPSQPK | S180 | 0.99 | 0.72 |
| PTPES(ph)P5AQEEAEPPSQPK | S180 | 0.99 | 0.72 |
| AEEPPSQ(ph)QPKDIPYSQDGE | S189 | - | 0.99 |
| DHIPYS(ph)DGECPTQANLSSQGDMSPK | S198 | 1 | 1 |
| DHIPYSQGECT(ph)PGQANLSSQGDMSPK | T203 | 0.52 | 0.78 |
| DHIPYSQGECT(ph)PGQANLSSQGDMSPK | S209 | 0.61 | 0.83 |
| DHIPYSQGECT(ph)PGQANLSSQGDMSPK | S210 | 0.56 | 0.87 |
| DHIPYSQGECT(ph)PGQANLSSQGDMSPK | S215 | 0.93 | 1 |
| SipR(ph)SEKLRSLSLEPSLPLAESHMQVWVGRGLPK | S218 | 0.96 | 0.97 |
| SDS(ph)LELIR | S230 | 0.95 | 1 |
| SipR(ph)LEPS(ph)PLR | S226 | - | 1 |
| SLEPS(ph)PLAESHLM(oxQWVGWGR | S230 | 1 | 1 |
| TQNS(ph)RGAAHPPTK | S332 | 1 | - |
| SWSh(ph)WTTPESHTPSHPQVSR | S345 | 0.99 | 1 |
| YGMPARRWS(ph)PEPNQKLE | S408 | 0.99 | 0.99 |
| LLES(ph)PNPEHIAETLSVDK | S418 | 0.99 | - |
| FTOHMSD(ph)YEDLTK | S460 | 0.99 | - |
| VLS(ph)SDDDVPDVSPVLE | S559 | 0.9 | 0.96 |
| TEVLS(ph)SDDDVPDVSPVLEPVLPSSTPGYVPTK | S567 | 0.99 | 0.99 |
| GPSTDSLAS(ph)PEY6NLSYWR | S520 | 0.99 | 0.99 |

The polybasic domain determines lipin phosphoregulation

In the present work, we show that replacing the lipin 1 PBD with that of lipin 3 eliminates the ability of phosphorylation to negatively regulate lipin 1 activity (Fig. 4, C and D). Conversely, the specific activity of lipin 3 (1PBD) increased significantly only after treatment with protein phosphatase and
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only in the presence of PE, indicating that phosphorylation hinders its recognition of di-anionic PA (Fig. 4, E and F). In accordance with this finding, the \( k_{\text{cat}} \) of dephosphorylated lipin 3 (1PBD) increased to a much greater extent, from 0 to 30% PE, than its phosphorylated counterpart (Table 3).

What remains unclear is how minor changes in the amino acid arrangement of the lipin PBD can drastically alter the ability of phosphorylation to control lipin activity. Kooijman et al. (21) demonstrated that the increase in the PA charge induced by lysine residues is greater than the increase induced by arginines. It is possible that lysines more readily hydrogen-bond with the phosphorylated amino acids within the protein, which weakens their interaction with PA. Interestingly, lipin 1 contains lysines on three sites within its PBD where lipin 3 contains arginines (Fig. 4A). Perhaps this underlies the mechanism whereby lipin 1 but not lipin 3 is negatively regulated by its own phosphorylation. In addition, it may be that not the number of lysines but a very precise arrangement of basic residues, both lysines and arginines, is necessary for phosphorylation to hinder recognition of PA charge and substrate association.

Unfortunately, there are no structural data available for lipins, and the region between the C-terminal and N-terminal domains does not conform to any known protein fold (Fig. 2A) (5). As such, the precise mechanisms whereby the lipin 1 PBD allows for phosphoregulation are speculative. The elimination of phosphorylation of lipin 1 (3PBD) could alternatively be explained by inherent changes in the overall phosphorylation of the mutant enzyme. However, the observed changes in gel mobility upon \( \lambda \) phosphatase treatment of the purified PBD mutants suggest that this is not the case (Fig. 7C). Additionally, Ser\(_{106}\) of lipin 1 (3PBD) is phosphorylated under basal conditions and remains downstream of mTOR in \( \lambda \) vitro, as demonstrated by reduction of phospho-Ser\(_{106}\) with Torin 1 treatment (Fig. 7B). Furthermore, LC-MS/MS analysis of lipin PBD mutants revealed that they shared many of the same phosphorylation sites as their WT counterparts (Tables 4 and 5 and Fig. 8). Based on these data, it appears that the exchange of the PBDs between lipins 1 and 3 does not significantly affect their phosphorylation. However, it is possible that the phosphorylated residues not found in the PBD mutants play crucial role in the regulation of PAP activity.

It is of interest whether the lipin PBD was always subject to this intermolecular regulatory mechanism rather than gaining this function later, during the emergence of more complex organisms. The PBD does not exist in yeast, and the recruitment of the yeast Pah1 is dependent on an N-terminal amphipathic helix rather than a PBD-like domain (39). The PBD domain appeared before the evolution of vertebrate species and the three distinct lipins. Whether the single lipin PBD was regulated by phosphorylation is unknown. However, although the lipin 1 domain notably changed from its first appearance to mammals, it is highly conserved, hinting at its important role in lipin function. Conversely, the lipin 3 PBD sequence is more variable in its evolution, at least from reptiles to humans. These observations suggest that perhaps lipin 3 PBD evolved to function under different cellular conditions and that its recognition of substrate charge was fine-tuned to be unresponsive to control by the enzyme’s phosphorylation sites.

Another feature of the lipin PBDs is their unusual length. The regions implicated in PA association in most PA effector proteins are composed of no more than 2–5 basic residues, and often just one lysine or arginine is critical for binding (16). Lipins, however, have a nine-residue PBD, and the differences between lipin 1 and 3 PBDs are in the first five residues. Interestingly, mutation of the first four amino acids in this domain (KKRR) perturbed the binding of lipin 1 to PA. Mutations of the remaining five amino acids did not negatively affect binding to any further extent but abolished nuclear localization (22). Lipins are known to be multifunctional; for example, they are PA phosphatases but also transcriptional co-regulators (1, 3–5, 40). In the context of the literature, our data suggest that whereas the PBDs are critical for substrate recognition and association, they are probably involved in other lipin functions. These and maybe yet unknown roles for lipins are perhaps reasons for the evolution of a relatively longer PBD. The nuances surrounding the differences in the molecular regulation of lipins may be indicative of the specialization of their activities or perhaps speak to their tissue-specific functions and necessity for either more lax or alternative regulatory mechanisms (6, 30, 41). Our findings elucidate a novel intramolecular mode of lipin 1 regulation, implicating the PBD as the determinant factor in the ability of lipin activity to be controlled by phosphorylation.

Experimental procedures

Materials

\([\text{\[^{32}P\]ATP}}\) was from PerkinElmer Life Sciences. The 1,2-dioleoyl-sn-glycerol, 1,2-dioleoyl-sn-glycero-3-phosphate, 1,2-sn-glycero-3-phosphocholine, 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine, and all materials for the preparation of liposomes were from Avanti Polar Lipids (Alabaster, AL). The Escherichia coli DGK, FLAG beads, FLAG peptide, primary FLAG antibody, and alkaline-conjugated secondary anti-rabbit and anti-mouse antibodies were from Sigma-Aldrich. The generation of anti-rabbit polyclonal antibody against phosphoserine 106 of lipin 1 was described by Harris et al. (10). All other commonly used reagents were also from Sigma-Aldrich unless otherwise indicated. All cell lines were obtained from ATCC (Manassas, VA).

Cell culture

Human cervical cancer cells (HeLa) were cultured in DMEM supplemented with 5% FBS (VMR Life Science Seradigm, Radvor, PA) and 1% penicillin/streptomycin (Life Technologies, Inc.). Fibroblasts were cultured in DMEM with 10% FBS and antibiotics.

Recombinant expression plasmids

The M. musculus lipin 3 cDNA was subcloned in frame with FLAG tag in the pCMV-TAG2B vector. The PBD exchange
mutants (lipin 1 (3PBD) and lipin 3 (1PBD)) were generated by PCR mutagenesis. Venus-tagged lipin 3, lipin 1 (3PBD), and lipin 3 (1PBD) were generated by subcloning the Venus cDNA in frame with the N terminus of lipin 3 downstream from the FLAG epitope tag. Adenoviruses were generated using the pAdEASY system (42). All amino acid numbering conforms to M. musculus phosphatidylinositol phosphatase LPIN3 (accession number NP_075021.1) or, when appropriate, to LPIN1 isoform b (accession number NP_056578).

**Purification of recombinant lipin proteins**

Cells from 40 15-cm plates were cultured in DMEM with 5% FBS and infected with an adenoviral vector expressing the indicated FLAG-lipin constructs for 48 h. The cells were fed daily during this time, harvested, and homogenized using a 22-gauge needle in buffer A (20 mM HEPES, 150 mM NaCl, 0.1% Brij, pH 7.2). The cell homogenates were centrifuged at 16,000 rpm for 10 min, and the cleared lysates were incubated with 15 μl/plate of FLAG (M2) beads for 4 h at 4 °C. Following the incubation, beads were separated into two fractions and incubated in phosphatase buffer (50 mM HEPES, 100 mM NaCl, 1 mM MnCl2, 2 mM DTT, pH 7.0) with (+) and without (−) 2000 units of λ protein phosphatase for 30 min at 30 °C. The beads were then loaded on an affinity column and washed in 1 ml of buffer A 10 times. The lipin proteins were eluted with five successive additions of an equal volume of 0.5 mg/ml FLAG peptide and dialyzed against buffer A without detergent. The purified lipins were loaded on an 8.5% acrylamide SDS-polyacrylamide gel along with BSA standards and stained with Coomassie Blue dye for quantification of protein yield.

**Preparation of [32P]PA**

1,2-Dioleoyl-sn-glycerol was phosphorylated by E. coli dia-cylglycerol kinase using [γ-32P]ATP. The labeled PA was purified by thin-layer chromatography as described by Han and Carman (43).

**Substrate preparation**

Briefly, 90 mol % PC and 10 mol % PA or 60 mol % PC, 30 mol % PE and 10 mol % PA were dissolved in chloroform and combined. To this was added [32P]PA (10,000 cpm/nmol). The lipids were dried in vacuo. The dried lipids were hydrated in buffer B (50 mM Tris-HCl, 0.5 mM MgCl2, and 10 mM β-ME, pH 7.4) to 10 ml and subjected to five freeze-thaw cycles, followed by extrusion 11 times with a mini-extruder through a 100-nm polycarbonate filter (44). The preparation of Triton X-100/PA mixed micelles followed a protocol described previously (14). Briefly, PA and [32P]PA were combined in chloroform and dried in vacuo. Triton X-100 was added to solubilize the lipids to the indicated concentrations. Micelles were generated with 10 mol % PA and were calculated using the formula, mol % = 100 × [lipid (m)]/[([lipid (m)] [Triton X-100 (m)])].

**Measurement of PAP activity**

The basic principles for measuring PAP activities were derived from previous work by Han and Carman (43) and further elaborated by Eaton et al. (14). Briefly, purified lipin proteins, radioactive liposomes, and buffer B were combined to a final volume of 100 μl. The final concentrations were as follows: 50 mM Tris-HCl, 0.5 mM MgCl2, 10 mM β-ME, and the indicated concentrations of PA. The reactions were allowed to proceed for 20 min at 30 °C with gentle agitation and were terminated with the addition of 500 μl of acidic methanol (methanol + 0.1 N HCl). To this was added 1 ml of chloroform followed by 1 ml of 1 M NaCl. The organic extraction was vortexed, and 500 μl of the aqueous phase was placed in scintillation vials to measure the removal of 32P from PA by a scintillation counter. The activity from assays containing lipins was normalized to activity in assays without enzyme.

**Measurement of PA binding**

Binding of lipin to liposome vesicles was measured following a slightly modified version of Hofer et al. (45). In short, PC/PA (80 mol % PC, 20 mol % PA) and PC/PE/PA (50 mol % PC, 30 mol % PE, 20 mol % PA) were hydrated to 10 mM in buffer C (50 mM Tris-HCl, 1 mM EDTA, pH 7.2). Liposomes were prepared as described above. The reaction volume was 100 μl and contained Venus-FLAG-lipins, 2 μM PA, and buffer C. The reactions were allowed to proceed for 30 min at 30 °C with gentle agitation and were terminated by the addition of an equal volume of 80% sucrose (w/v). This mixture was layered over 270 μl of 80% sucrose (w/v) and was diluted in 4 equal volumes of 50 mM Tris-HCl, 0.5 mM MgCl2, and 10 mM β-ME, pH 8.2; reduced using 5 mM dithiothreitol at 55 °C for 30 min; and digested overnight using trypsin (Promega Corp., Madison, WI), Lys-C (Wako Chemicals, Tokyo, Japan), or Glu-C (Worthington). The resulting peptides were desalted and analyzed in duplicate by LC-MS/MS on both Thermo LTQ-Velos-Orbitrap and Q-Exactive mass spectrometers using a data-dependent acquisition strategy. Raw mass spectra were searched using Maxquant (version 1.5.5.1) (46) against the UniProt human protein sequence database concatenated to sequences of the WT recombinant mouse lipins including the lipin 1 and 3 PBD-switch mutants, allowing for phosphorylated serine, threonine, or tyrosine. The “match between runs” feature of Maxquant was disabled, and peptide-spectral match false discovery rate was filtered to <1%. Phosphorylation sites were mapped if the...
corresponding peptide had a posterior error probability of \( \leq 0.05 \) and if the site localization probability was 0.95 or greater. The posterior error probability score is a measure of confidence in the identity of the peptide. The site localization probability is a measure of confidence in the precise localization of the phosphorylation to a residue within the peptide.

**Analysis of phosphate removal by λ protein phosphatase**

HeLa cells were cultured on 4 15-cm plates and infected with adenovirus overexpressing FLAG-lipin 3 for 48 h. Following the infection period, cells were incubated in low-phosphate buffer (145 mM NaCl, 5.4 mM KCl, 1.4 mM CaCl\(_2\), 1.4 mM MgSO\(_4\), 25 mM NaHCO\(_3\), 0.2 mM sodium phosphate, 5 mM glucose, 10 mM HEPES, 0.1% insulin-free BSA, pH 7.4), radiolabeled with 0.04 Ci/mmol. Following the incubation, the cells were harvested, homogenized as described above, and then incubated with 15 μl/plate of anti-FLAG beads. Following the incubation, the beads were separated into two fractions and were treated with (+λ) or without (−λ) 400 units of λ protein phosphatase. The lipin 3 was detached from the beads by the addition of 1× Laemmli buffer containing β-ME. The proteins were separated by an 8.75% SDS-PAGE gel under reducing conditions and transferred onto a PVDF membrane (Immobilon, Darmstadt, Germany). The radiolabeled protein and removal of phosphate were visualized using autoradiography.

**Phosphorylation of lipin 3 in 3T3-L1 cells**

7–10 days post-differentiation, 3T3-L1 cells (2 10-cm plates/condition) were infected with an adenoviral vector expressing FLAG-lipin 3 or FLAG-lipin 1 for 48 h. In the last 2 h of the infection, cells were serum-starved in low-phosphate buffer supplemented with 0.2 mCi/ml \([^{32}P]\)orthophosphate. During this time, the cells were treated with vehicle or Torin 1 (Tocris Bioscience, Bristol, UK) for the last 45 min, followed by 10 million/ml of insulin for the last 15 min. The cells were harvested and homogenized in buffer A with protease inhibitors and cleared by centrifugation at 16,000 x g for 10 min. The supernatant was incubated with 15 μl/plate of anti-FLAG (M2) beads for 4 h at 4°C. Proteins were displaced from the beads with the addition of Laemmli buffer containing β-ME and were separated on an 8.75% SDS-polyacrylamide gel. Phosphate incorporation into lipins 1 and 3 was visualized using autoradiography, and total protein expression was detected as described using Western blot analysis. The incorporation of \(^{32}P\) was normalized to total protein.

**Western immunoblot analysis**

To verify the expression of the indicated proteins and phosphorylation sites, the PVDF membrane containing proteins was blocked by incubation in Tris-buffered saline with detergent Tween 20 (TBST) containing 10% (v/v) dried milk for 1 h at 25°C. The TBST contained the following: 50 mM Tris, 150 mM NaCl, and 0.05% (w/v) Tween 20, pH 7.4. After the block, the membrane was incubated with the indicated antibodies (1:1000) in TBST at 25°C for 1 h with gentle agitation. The membrane was then incubated with alkaline phosphatase–conjugated mouse or rabbit secondary antibody (1:10,000) diluted in TBST with 2% (w/v) dried milk for 1 h at 25°C. After three 15-min washes with TBST, the membrane was briefly incubated in chemiluminescent alkaline phosphatase substrate (Applied Biosystems, Foster City, CA). The immunoreactivity was detected using a Fuji LAS 4000.

**Statistical analysis**

The statistical analysis and determination of all kinetic constants were done using GraphPad Prism software. For calculation of \( K_{\text{m}}^{\text{app}} \) values, non-linear regression of the Michaelis-Menten plots was used. The \( k_{\text{cat}} \) values were calculated using the equation, \( k_{\text{cat}} = V_{\text{max}}/E_{p} \), where \( E_{p} \) is the amount of enzyme catalytic sites in nmol and \( V_{\text{max}} \) is the maximal rate of the reaction in nmol/min. Statistical significance between two groups was determined using Student’s \( t \) test. Statistical significance between more than two groups was determined using one-way analysis of variance (ANOVA) followed by the indicated post hoc analysis. The statistical tests used for each figure are indicated. All values are expressed as means ± S.E. Unless otherwise noted, data show one of at least three independent experiments. When appropriate, all experiments were repeated with at least two separate protein preparations. Significance was set to \( p < 0.05 \).

**Author contributions**—S. B. and T. E. H. conceived, coordinated, and designed the study. T. E. H. also significantly contributed to the editing of the manuscript. S. B. designed, performed, and analyzed the experiments shown in Figs. 1 (C, D, and H), 2 (B–E), 3, 4, 5, 6, and 7 and Tables 2 and 3 and wrote the manuscript. S. T. designed, performed, and analyzed the experiments shown in Fig. 2F, and R. T. L., S. W. E., and J. V. designed, performed, and analyzed the experiments shown in Figs. 2A and 8 and Tables 1, 4, and 5. S. B., T. E. H., and M. E. G. analyzed and displayed the data seen in Figs. 2A and 8 and Tables 1 and 4. M. E. G. also assisted in the experiments performed in Fig. 7A. J. M. P. performed and analyzed the experiments shown in Fig. 1 (A, B, E, and F). J. M. E. designed, performed, and analyzed the experiments shown in Fig. 1A. G. R. M. assisted in the experiments performed in Fig. 2B and made significant contributions to the editing of the manuscript. All authors reviewed the results and approved the final version of the manuscript.

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