Lipoic acid is a sulfur-containing cofactor indispensable for the function of several metabolic enzymes. In microorganisms, lipoic acid can be salvaged from the surroundings by lipoate protein ligase A (LpLA), an ATP-dependent enzyme. Alternatively, it can be synthesized by the sequential actions of lipoate protein ligase B (LipB) and lipoyl synthase (LipA). LipB takes up the octanoyl chain from C8-acetyl carrier protein (C8-ACP), a byproduct of the type II fatty acid synthesis pathway, and transfers it to a conserved lysine of the lipoyl domain of a dehydrogenase. However, the molecular basis of its substrate recognition is still not fully understood. Using *Escherichia coli* LipB as a model enzyme, we show here that the octanoyltransferase mainly recognizes the 4′-phosphopantetheine-tethered acyl-chain of its donor substrate and weakly binds the apo-acyl carrier protein. We demonstrate LipB can accept octanoate from its own ACP and noncognate ACPs, as well as C8-CoA. Furthermore, our 1H saturation transfer difference and 31P NMR studies demonstrate the binding of adenosine, as well as the phosphopantetheine arm of CoA to LipB, akin to binding to LpLA. Finally, we show a conserved RGG loop, analogous to the lipoate-binding loop of LpLA, is required for full LipB activity. Collectively, our studies highlight commonalities between LipB and LpLA in their mechanism of substrate recognition. This knowledge could be of significance in the treatment of mitochondrial fatty acid synthesis related disorders.

Lipoic acid is a hydrophobic cofactor, essential for the function of several metabolic enzyme complexes, *viz.* pyruvate dehydrogenase complex (PDC) required for pyruvate oxidation, 2-oxoglutarate dehydrogenase as a component of Krebs cycle (OGDHc), glycine cleavage system (GCS) involved in glycine degradation, branched-chain keto acid dehydrogenases (BCKDHc) necessary for the metabolism of branched chain amino acids, and 2-oxoadipate dehydrogenase in lysine metabolism (1–5). These are multimeric enzymes with a conserved lipoyl domain that serves as a substrate for lipoic acid modification. Lipoyl lysine acts as a “swinging arm,” helping to shuttle intermediates between the active site of multienzyme complexes (1, 4, 6–9). The flexibility of lipoic acid is crucial for substrate channeling and electron transfer during oxidation–reduction reactions (10).

Most of the knowledge with regard to lipoic acid metabolism comes from studies conducted on *Escherichia coli*, *Listeria monocytogenes*, *Bacillus subtilis*, and *Staphylococcus aureus* (11). In the presence of free lipoate or octanoate, lipoic acid is salvaged from the environment by lipoate protein ligase A (LpLA) (12–14). An activated lipoyl-5′-AMP intermediate is formed from ATP and lipoic acid on the surface of LpLA. Thereafter, the ε-amino group of a lipoyl lysine attacks the noncovalently bound lipoyl-AMP, forming an amide linkage with lipoyl-group and releasing AMP (3, 8, 9, 14, 15). All LpLA molecules comprise a large N-terminal catalytic domain (lipoic acid binding) and a small C-terminal domain (16). In lipoic acid–deficient environments, lipoate protein ligase B (LipB) also known as Lipooyl-octanoyl transferase catalyzes the first biosynthetic step of lipoic acid synthesis (4). Studies on *Mycobacterium tuberculosis* (Protein Data Bank [PDB] 1W66) and *Thermus thermophilus* (PDB 2QHS, 2QHT, 2QHU, and 2QHV) LipB suggest conservation of a structural scaffold, similar to the N-terminal catalytic domain of LpLA. LipB functions as a cysteine/lysine dyad acyltransferase, cysteine 176 and Lys 142 of *M. tuberculosis* LipB (Cys169 and Lys 135 of *E. coli* LipB) functioning as acid/base catalysts (17–19). A covalent octanoyl-LipB thioester intermediate is formed by the transfer of octanoyl-chain from C8-ACP (acyl carrier protein) to Cys176 of LipB (20). Subsequently, the thioester bond is attacked by the ε-amino group of the lipoyl lysine, resulting in C8- modification of the latter. In the following step, octanoyl group covalently tethered to the lipoyl domain is converted to lipoic acid in the presence of LipA (lipoyl synthase) by insertion of two sulfur atoms.

In metazoans, three different enzymes participate in lipoic acid synthesis: (a) lipoyl transferase 2 (LipT2) that transfers octanoyl chain from C8-ACP to the lipoyl subunit of GCS, (b) lipoic acid synthase (LIAS), a sulfur insertion enzyme that adds two sulfur atoms to the C8- chain to form lipoic acid, and (c)
C$_8$-phosphopantetheine is recognized by LipB

LipT1, an amidotransferase, that relays octanoyl-chain/lipoic acid from the H subunit of GCS to the lipoyl carrier protein subunit of other dehydrogenases, viz. PDH, BCKDHC, αKGDH. Malfunction of any of the gene products causes lipoic acid synthesis defects in neonatal patients, underscoring the need to understand the pathway at the molecular level. Disorders range from defective mitochondrial energy metabolism, toxic levels of certain amino acids, neurological problems, and even death, depending on which lipoic acid synthesis gene malfunctions (11, 21, 22). Similar clinical phenotypes are also observed if the upstream mitochondrial fatty acid synthesis (FAS) that does not function properly, for example, mitochondrial enoyl-CoA reductase associated disorder (MECR) (21, 23), malonyl-CoA synthase (ACS3), malonyl-CoA-acyl carrier protein transacylase (MCAT), etc. (24). Defects in GCS gene also causes elevated glycine levels in the brain, leading to analogous neurological problems (25, 26). Impaired dehydrogenase activity causes respiratory defects and muscle weakness due to the disruption of Krebs cycle (5, 12, 13).

Interestingly, all lipoic acid metabolism enzymes belong to the Pfam PF03099 family and share a similar scaffold, despite low sequence conservatism. They differ remarkably in their catalytic mechanism and interactions (20, 27). Reasonable biochemical/structural data exist for the free, as well as substrate-bound forms of LplA from *E. coli*, *Thermoplasma acidophilum*, *Enteroctococcus faecalis*, *Streptococcus pneumoniae*, and *Mycoplasma hypopneumoniae*. However, the biosynthetic enzyme LipB is relatively less explored. Structures are available from only two sources, *M. tuberculosis* and *T. thermophilus* for this enzyme. The mechanistic understanding of LipB interaction with its substrates is also sparse. Two recent studies on LipB have highlighted its ability to allosterically distinguish between different acyl-ACP’s, viz. C$_8$-, C$_9$-, C$_10$-, and C$_12$-ACP based on differences in their surface conformation (28, 29).

Our work extends their findings one step further and shows that the enzyme can differentiate between apo-, holo-, and C$_8$-ACP as well. The enzyme binds C$_8$- and holo-ACP with moderate affinity but weakly associates with apo-ACP. It can uptake octanoate from its cognate C$_8$-ACP, noncognate C$_8$-ACP’s, as well as octanoyl-CoA. Chemical shift perturbation studies support the binding of adenosine and phosphopantetheine arm of CoA with LipB. Interestingly, mice LipT2 also successfully transferred octanoate-CoA from C$_8$-CoA to *E. coli* GcvH in vitro, suggesting a common mechanism of substrate recognition by most octanoyl transferases.

**Results**

LipB (UNP60720) catalyzes a bisubstrate reaction, using C$_8$-ACP (molecular weight 9.4 kDa) as a donor substrate and lipoil domain of dehydrogenases, for example, GCS H protein (GcvH), PDC, or 2-oxoglutarate dehydrogenase complex (OGDH) as octanoate acceptors (25). Limited structural/ biochemical data exist for the LipB-ACP complex (25). Limited structural/ biochemical data exist for the LipB-ACP complex (25). Limited structural/ biochemical data exist for the LipB-ACP complex (25). Limited structural/ biochemical data exist for the LipB-ACP complex (25).

In the absence of an *E. coli* LipB structure, *M. tuberculosis* LipB 1.08 Å crystal structure (MtLipB, PDB 1W66) has been used as a structural model to interpret our findings.

*M. tuberculosis* LipB comprises five central β-strands 1β$_1$-1β$_2$-1β$_3$-1β$_4$-1β$_5$, enclosed by three long helices, α$_2$ and α$_3$ on one face and α$_1$ on the other side of β sheet, illustrated in **Figure 1A**. Some of the catalytically important residues of *M. tuberculosis* LipB are shown as sticks and labeled cyan. The corresponding *E. coli* LipB residues are also labeled black. A short stretch of beta strands comprising 1β$_2$-1β$_1$-1β$_3$, called the ‘capping sheet’, is also present in LipB, of unknown function. **Figure 1B** shows the molecular surface of MtLipB, colored based on coulombic charge, displaying the position of active site cavity. *E. coli* LipB amino acid sequence has been compared with other octanoyl transferases in **Fig. S1**. *E. coli* LipB (UNP 60720) shares 32% sequence identity with *M. tuberculosis* LipB (UNP P9WK83, PDB 1W66), 28% with human mitochondrial LipT2 (UNP A6NKi8), 29% with *Mus musculus* LipT2, and 26% with *Leishmania major* enzyme (UNP Q4A1B).

**LipB binds octanoyl-ACP with moderate affinity**

To gain insights into the LipB:C$_8$-ACP interaction, a catalytically inactive *E. coli* LipBK135A/C169A double mutant was prepared, and its interaction with $^{15}$N labeled *E. coli* C$_8$-ACP followed in a $^1$H$^{15}$N transverse relaxation optimized spectroscopy (TROSY)–heteronuclear single quantum coherence (HSQC) NMR experiment. C$_8$-ACP was stable at room temperature for approximately 7 days, as shown in **Fig. S2**. The protein did not display any hydrolysis during this time, confirming its suitability for NMR studies. ACP backbone was assigned based on the BMRB ID 27061, and standard 3D NMR experiments were acquired to assign the octanoyl-ACP spectra. To give equal weights to proton and nitrogen chemical shifts, changes upon LipB interaction have been reported as their weighted average, determined using Equation 1.

$E. coli$ ACP displays a four-helix bundle fold, consisting of helix I (Thr 2-Gly 16), helix II (Asp 35-Asp 51), helix III (Asp 43-Leu 46, Glu 49, Thr 52, Ile 54, Asp 56, Glu 60-Thr 64, Glu 65, and Ala 68-Tyr 71 (Fig. 1C). These residues are present in helix II, III, and IV of ACP. The peaks for Asp 56, Thr 64, Gly 46, Leu 43, Ala 42, Glu 41, Val 40, Glu 39, and Asp 38, Val 37, Ala 36, Gly 35, Asp 34, Val 33, Asp 32, Val 31, Thr 30, Ser 29, Ser 28, Tyr 27, and Tyr 26 were broadened beyond 1:1 ACP:LipBK135A/C169A ratio, shown by a (*) in the figure. Leu 37 and Glu 48 displayed chemical exchange line broadening right from the first titration point, shown as (●) in the figure. One SD has been used to demarcate significant chemical shift change. Perturbations have been mapped to the *E. coli* holo-ACP structure (PDB 5VCB, chain D) represented as a worm in **Figure 1D**. Thickness of the worm is directly proportional to the magnitude of chemical shift change. **Fig. S3A** shows the multiple overlaid $^1$H$^{15}$N TROSY-HSQC spectra for C$_8$-ACP:LipBK135A/C169A titration. Each titration point is shown in a different color: free C$_8$-ACP is colored red; C$_8$-ACP:LipBK135A/C169A 1:0.25 M ratio blue; 1:0.5 magenta, 1:0.75 cyan, 1:1 green, 1:1.5 orange, and 1:2 purple. Same color coding has been used throughout for all the titrations reported in this study.
The binding affinity of LipBK135A/C169A for C₈-ACP was determined by surface plasmon resonance (SPR) measurements, immobilizing LipBK135A/C169A on the surface of a sensor chip, and passing increasing concentrations of C₈-ACP over the chip surface. Fig. S4A shows the SPR sensogram and Fig. S4B binding curve for C₈-ACP:LipBK135A/C169A interaction. The two proteins interact in micromolar range, with a dissociation equilibrium constant ($K_D$) of 36.4 ± 0.19 μM.

**LipB binds holo-ACP with relatively lower affinity**

Holo-ACP contains a phosphopantetheine modification akin to C₈-ACP, but lacks the octanoate group. The contribution of C₈-chain in the interaction of C₈-ACP with LipB was also investigated. $^{15}$N-labeled holo-ACP (containing $^{15}$N-labeled 4'-phosphopantetheine arm) was titrated with unlabeled LipBK135A/C169A. Noticeable changes in the amide chemical shift were observed for several holo-ACP
C8-phosphopantetheine is recognized by LipB

residues. As shown in Figure 1E, Glu 14, Glu 21, Val 29, Ser 36, Leu 37, Thr 39, Val 40, Glu 41, Leu 42, Val 43, Met 44, Ala 45, Leu 46, Glu 47, Asp 51, Thr 52, Glu 53, Ile 54, Glu 57, Ala 59, and Glu 60 displayed significant perturbations. The aforementioned residues are present in helix I, loop I, helix II and helix III of ACP, shown as a worm in Figure 1F. Fig. S3B displays multiple overlaid 1H15N TROSY-HSQC spectra for holo-ACP upon titration with LipBK135A/C169A. The direction of chemical shift change with increasing LipBK135A/C169A concentration is shown by a forward pointing arrow. Interestingly, one of the two NH resonances of 4'-phosphopantetheine moiety (15N-labeled phosphopantetheine expressed in holo-form in E. coli) observed at a chemical shift of 8.1 ppm in the 1H15N TROSY-HSQC spectra also displayed significant change during the titration (Fig. S3D).

Fig. S4C shows the SPR sensorgram, and Fig. S4D shows binding curve for the interaction of LipBK135A/C169A with holo-ACP. The binding curve did not reach saturation, and therefore, the estimate of affinity has been reported as apparent binding constant. A KD app of 100 ± 0.43 μM was observed for the LipB:holo-ACP interaction.

apo-ACP weakly interacts with LipB

In the cell, ACP is expressed as an apo-protein and post-translationally modified to holo-ACP by the catalytic action of 4'-phosphopantetheinyl transferase. The importance of 4'-phosphopantetheine arm in the interaction of LipB with its substrate was also probed. 1H15N apo-ACP was titrated with unlabeled LipBK135A/C169A (Fig. 1G). Insignificant changes in the amide chemical shift were observed (<0.06 ppm) upon addition of up to 2 M equivalents of LipBK135A/C169A. In the superimposed spectra, most of the amide resonances of apo-ACP overlapped with the peaks in presence of LipBK135A/C169A (1:2 M ratio, Fig. S3C). A worm figure of apo-ACP is also shown, displaying insignificant changes upon interaction with LipBK135A/C169A (Fig. 1H).

SPR measurements were also performed to understand the strength of this interaction. Relatively higher concentration of apo-ACP was required, compared to C8- and holo-ACP titrations. Fig. S4E shows the SPR sensorgram, and Fig. S4F shows binding curve for the interaction of LipBK135A/C169A with apo-ACP. A KD app of 2.1 ± 0.18 mM was obtained for the binding interaction.

Noncognate ACPs can serve as a donor substrate for LipB

Cross reactivity of LipB with noncognate C8-ACP substrates was also investigated. The enzyme assay comprised 5 μM E. coli LipB, an octanoyl donor (C8-ACP, 75 μM), and an octanoyl acceptor GcvH (25 μM). Three separate assays were performed using different octanoyl-donors: (a) E. coli C8-ACP (UNP 0A6A8), (b) C8-LmAACP (L. major ACP, UNP E9AD06), and (c) C8-PfACP (P. falciparum ACP, UNP Q7KWJ1). The product formed after the assay (C8-GcvH) in each case was qualitatively analyzed on a 12% SDS-PAGE gel (Fig. 2A).Lane 1 is pure apo-GcvH, lane 2: C8-EcACP, lane 3: C8-LmAACP, lane 4: C8-PfACP (used as a reference/control).

LipB can octanoylate GcvH using C8-CoA as a donor substrate

Next, we probed the importance of ACP scaffold in the substrate recognition by LipB. C8-CoA comprises a C8-chain and a phosphopantetheine arm, identical to the modifications present in C8-ACP. In addition, it contains an adenosine 3'-diphosphate group, not found in C8-ACP. To test whether LipB can use CoA analogs as a donor substrate, different lengths of acyl-CoA were used viz. C4-CoA, C8-CoA, C10-CoA, and C12-CoA, and the reaction progress was qualitatively monitored by following the modification of GcvH on an SDS-PAGE gel. Quantitatively, the reaction progress was followed by reverse phase HPLC chromatography. As shown in Figure 2E, LipB efficiently modified apo-GcvH using C8-CoA as acyl-chain donor (lane 5). No change in migration of the GcvH band was observed when C4-CoA, C10-CoA, or C12-CoA were used as donors, shown in lanes 4, 6, and 7, respectively in Figure 2E. MALDI-TOF mass spectrometry confirmed an increase in mass of GcvH by 128 Da after the enzyme assay, when C8-CoA was used as an acyl-chain donor. This change is mass is equivalent to the molecular mass of the C8-chain (127 Da, Fig. S5 C). The expected molecular mass of apo-GcvH is 15,834 Da (GcvH subunit containing an N-terminal His tag “GSSHHHHHHSSGLVPRSHGM”). The terminal methionine of His tag was cleaved by E. coli MAP1, endogenous methionine aminopeptidase MAP (31). Fig. S5A shows the mass spectra for apo-GcvH (used as a reference) and Fig. S5, B–E mass spectra of GcvH after the LipB assay using C4-CoA, C8-CoA, C10-CoA, and C12-CoA as acyl-chain donors. No change in mass was observed in the assays performed with C4-CoA, C10-CoA, and C12-CoA (Fig. S5, B, D and E).

Mammalian LipT2, an octanoyltransferase is associated with vast number of neurological disorders. The ability of mice MmLipT2 (Mus musculus, UNP Q9D009) to transfer octanoate from C8-CoA to E. coli GcvH was also investigated. As shown in Figure 2F, MmLipT2 also successfully converted E. coli GcvH using C8-CoA as a donor substrate, which was confirmed from the forward migration of the GcvH band in
Figure 2. LipB can use noncognate ACP as well as C₈-CoA as octanoate donor. A, a 12% SDS-PAGE gel displaying lane 1: Apo-GcvH, lanes 2 to 4: C₈-ACP, C₈-LmACP, and C₈-PfACP loaded for reference, lane 5: molecular weight marker, lane 6 to 8: LipB assay using C₈-ACP (E. coli), C₈-LmACP (L. major), and C₈-PfACP (P. falciparum), respectively, as substrates. A1, a standard curve prepared based on the migration of standards in the SDS-PAGE gel of (A). The migration of E. coli ACP corresponds to a molecular weight of 18 kDa, LmACP ~14 kDa, and PfACP ~12 kDa. C18-reversed phase chromatogram for the GcvH conversion after the LipB assay performed using (B) C₈-EcACP, (C) C₈-LmACP, and (D) C₈-PfACP, as octanoate donors. The chromatogram before the assay is shown in black and after the assay in blue. %GcvH conversion is shown at the bottom of each chromatogram. E, a 12% SDS-PAGE gel displaying the conversion of GcvH, using different acyl-CoA as octanoyl donors. Lane 1: molecular weight marker, Lane 2: apo-GcvH, Lane 3: assay in the absence of LipB, Lane 4 to 7: assay in presence of C₄-CoA, C₆-CoA, C₈-CoA, and C₁₀-CoA, respectively. F, a 12% SDS-PAGE gel, displaying the conversion of E. coli GcvH in presence of Mus musculus LipT2, using C₈-CoA as octanoyl donor. Lane 1: molecular weight marker, lane 2: apo-GcvH, lane 3: assay in the presence of mice LipT2. % conversion determined using HPLC is shown at the bottom of the lane.

C₈-phosphopantetheine is recognized by LipB
C₈-phosphopantetheine is recognized by LipB

C₈-CoA is preferred over C₈-ACP as a LipB substrate. It is noteworthy that in lane 5, a major pro-substrate and poorly utilizes C₈-CoA (32). Does PAGE. Lane 5 shows the assay performed in presence of equimolar proteins used as reference), respectively, on a 12% native-PAGE gel. In Figure 3 of apo-GcvH into its holo-form on a 12% SDS-PAGE. ACP plays bands for the assay carried out using C₈-ACP alone as a donor substrate. In a previous study, using C₈-ACP as a substrate, a GcvH displayed complete conversion. On the contrary, an band was low compared to lane 6, suggesting less consumption of C₈-CoA + C₈-ACP, while lane 7 shows the bands for the assay with C₈-ACP alone. Notably, both lanes 6 and 7 displayed complete conversion of apo-GcvH to C₈-GcvH. However, based on native-PAGE (Fig. 3B), in lane 5, the intensity of the holo-ACP (used as reference) in lane 2, and apo-GcvH (used as reference) in lane 2, and apo-GcvH (used as reference). The sample displays three peaks in the 6.0 to 9.0 ppm range, corresponding to 1, 2, and 1’ resonances of the adenosine component of CoA (atom numbering shown in Fig. 4A). In the 0 to 5 ppm range, 4’ and 5’ peaks from the adenosine component (colored black) and 1”, 2”, 3”, 4”, 5”, 6”, 7”, and 8” peaks arising from the phosphopantetheine arm were also observed. Figure 4B2 displays the NMR spectra for CoA in buffer (50 mM Tris–HCl, 200 mM NaCl, pH 7.8). Figure 4B3 shows the ¹H NMR spectra for CoA in the same Tris–HCl buffer pH 7.8, in presence of 1 mM WT LipB (for reference). Figure 4B4 illustrates the saturation transfer difference (STD) spectra for CoA in buffer to ensure no intramolecular NOE transfer occurs. Figure 4, B5–B8 displays the STD spectra for CoA in the presence of WT LipB, LipBK135A/C169A (catalytically inactive double mutant), LipBG72A/G73A, and LipBR71A, respectively. All these STD experiments were acquired using a CoA:LipB molar ratio of 10:1. NOEs corresponding to position 1, 2, and 1’ were observed in Figure 4, B5–B8, suggesting binding of the adenine ring. Similarly, in the 0 to 5 ppm range, peaks 2”, 3”, and 8” corresponding to the phosphopantetheine arm displayed NOEs in the STD spectra in presence of WT and the mutant LipB. The on resonance (red), off resonance (blue), and STD spectra (green) for the binding of LipBK135A/C169A with CoA in 50 mM Tris, 200 mM NaCl, pH 7.8 is illustrated in Fig. S6B. Fig. S6A shows the chemical structure of CoA.

Table 1

| Enzyme kinetic parameters for E. coli WT LipB and the mutants |
|---------------------------------------------------------------|
| Enzyme parameters | LipBWt | LipBG72A/G73A | LipBR71A | LipBK135A |
| Vₘₐₓ (μmol/min) | 1.08 ± 0.01 | 0.32 ± 0.01 | 0.26 ± 0.02 | 0.20 ± 0.01 |
| Kₐ₀(μM) | 12.24 ± 0.79 | 6.71 ± 0.98 | 34.39 ± 8.8 | 42.37 ± 4.58 |
| kₘ (min⁻¹) | 0.22 | 0.06 | 0.05 | 0.04 |

The assay was performed using C₈-CoA as octanoate donor and E. coli GcvH as octanoate acceptor.
shows the $^{31}$P NMR spectra for 1 mM LipB in 50 mM Tris, 200 mM NaCl, pH 7.8 at 298 K. No peaks were observed, suggesting protein as such does not contain any phosphate group. Figure 4C2 displays the $^{31}$P spectra for CoA in 50 mM Tris, 200 mM NaCl, pH 7.8. Three prominent peaks were visible, corresponding to 3'-phosphate, 1'-phosphate, and 5'-phosphate. Figure 4, C3–C6 show the $^{31}$P spectra for CoA in presence of LipBK135A/C169A, WT, LipBG72A/G73A, and LipBC169A, respectively. The peak corresponding to 3'-phosphate displayed an upfield change in chemical shift upon addition of WT LipB (Fig. 4C4), LipBG72A/G73A (Fig. 4C5), and LipBC169A (Fig. 4C6). The new peak arising as a result of this interaction is indicated by a downward pointing arrow in the figures. No change in peak position was observed in LipBK135A/C169A sample (Fig. 4C3), suggesting interaction of Lys135 with 3'-phosphate of CoA, that is lost in the double mutant. A CoA:LipB ratio of 8:1 was used for these experiments.

Figure 3. LipB prefers C$_8$-CoA over C$_8$-ACP as a donor substrate. A, kinetic studies using - LipB, - LipBR71A, - LipBG72A/G73A, and - LipBK135A as octanoyl-transferases, C$_8$-CoA as a donor substrate, and GcvH as an acceptor substrate. Reported values are the average of two independent measurements done on a Symmetry C18 reversed phase HPLC column. The figure was drawn using GraphPad prism version 7.0. B, a 12% native-PAGE gel displaying lane 1 to 4: apo-ACP, C$_8$-ACP, holo-ACP, and apo-GcvH, respectively, used as controls. Lane 5 was loaded with LipB assay performed in presence of equimolar concentration of C$_8$-ACP and C$_8$-CoA used as donor substrates. Lane 6: assay done using C$_8$-ACP alone as a donor substrate. C, the same samples were loaded on a 12% SDS-PAGE gel. Lane 1: Marker, lane 2 to 5: apo-ACP, C$_8$-ACP, holo-ACP, and GcvH samples loaded as a reference. Lane 6: assay carried out using equimolar concentration of C$_8$-CoA and C$_8$-ACP, while lane 7 was loaded with the assay carried out in presence of C$_8$-ACP alone. In both the assays, GcvH was used as an acceptor substrate for the bisubstrate reaction, and its conversion to C$_8$-GcvH was followed by forward migration. C18-reversed phase chromatogram for (D) Apo-GcvH used as a reference, LipB assay in presence of (E) equimolar concentration of C$_8$-ACP: C$_8$-CoA, and (F) C$_8$-ACP alone. Samples in Figure (D–F) were the same samples that were loaded in lanes 4 to 6 of the Native-PAGE and lanes 5 to 7 of SDS-PAGE gels in figures (B) and (C).
Are adenine- and phosphate-binding sites also conserved in LipB?

$^{15}$N-labeled *E. coli* LipBK135A/C169A (catalytically inactive double mutant) was titrated with C$_8$-CoA, and the titration was followed by NMR. Several amides in the $^{15}$N TROSY-HSQC experiment displayed significant chemical shift change (Fig. S7). In the figure, LipBK135A/C169A 1:0 M ratio (red), 1:0.2 (cyan), 1:0.4 (yellow), 1:0.8 (green), 1:1 (magenta), and 1:2 (orange) are shown as multiple overlaid spectra. The direction of chemical shift change is shown by a *black arrow*. Most of the peaks displaying perturbations appeared partially broadened due to chemical exchange. Unfortunately, owing to the low

Figure 4. LipB interacts with adenosine 3′,5′-diphosphate and phosphopantetheine arm of CoA. A, the chemical structure of CoA is shown, displaying its two components phosphopantetheine (labeled brown) and ADP (labeled black). B1, $^1$H NMR spectra of CoA in D$_2$O. B2 $^1$H spectra of CoA in Tris–HCl. B3 $^1$H spectra of CoA in Tris–HCl in presence of 1 mM WT LipB. Saturation transfer difference (STD) spectra for B4 CoA alone in buffer, B5 CoA: wild LipB, B6 CoA:LipBK135A/C169A, B7 CoA:LipBG72A/G73A, and B8 CoA:LipBR71A. C, $^{31}$P NMR spectra of C1 WT LipB in Tris–HCl, C2 CoA in Tris–HCl buffer, C3 CoA: LipBK135A/C169A, C4 CoA: WT LipB, C5 CoA:LipBG72A/G73A, C6 CoA:LipBC169A. The molar ratio of CoA:LipB was maintained at 10:1 in all STD experiments and 8:1 in the $^{31}$P NMR experiments.
stability of the enzyme at high concentrations (necessary for backbone assignments), we were unable to assign the backbone amides of *E. coli* LipB. The pattern of amide perturbations though confirmed that *E. coli* LipB binds C₇-CoA.

Notably, the lipoate scavenging enzyme LplA has adenine- and phosphate-binding sites that favor lipoic acid and ATP binding. We asked whether adenine and phosphate binding sites are also conserved in LipB, given the evolutionary relationship of the two proteins. Thus, *M. tuberculosis* MtLipB structure (PDB 1W66) was superimposed on the catalytic domain of *E. coli* LplA complexed with C₇-AMP (LplA, PDB 3A7A), using the matching option of Chimera, shown in Fig. S8A (33). In most cases, atoms N2, N6, C2, and C8 of adenine, present at the Watson–Crick edge interact with the backbone and side chain atoms of adenine-binding proteins (34). In the *E. coli* LplA structure (PDB 3A7A), N1 and N6 of adenine form a hydrogen bond with the side chain of Asn 83 OD1, Thr 151, and the backbone O of Phe 78. Also, N7 forms a hydrogen bond with the backbone N of Phe 78. In *M. tuberculosis* LipB (MtLipB, 1W66), remarkably similar residues are present at some of these sites, Gln 87 and Trp 82 (instead of Asn 83 and Phe 78 in LplA). In *E. coli* LipB, Gln 82 and Tyr 77 are present, while the equivalent residue is Lys 142 in *E. coli* LplA. Arg 71 and Gly 72 are hydrogen bond with the carbonyls of Gly73, Gly75, and side chain of Ser72 in the LplA structure (Fig. S8B, colored brown). These interactions have been speculated to stabilize the lipoate-binding loop of LplA. In MtLipB as well, arginine 76 side chain forms identical hydrogen bonds with the backbone carbonyls of Gly 77 and Lys 79, colored cyan in Fig. S8D. The structural similarities between LipB and LplA hint at the possibility of conservation of adenine- and phosphate-binding sites in LipB, like LplA.

**C₇-phosphopantetheine is recognized by LipB**

The lipoate-binding loop 69RRXXGGG₇₅ of LplA (PDB 3A7A) is also somewhat conserved in MtLipB and has a very similar conformation, shown in Fig. S8, C and D (35). MtLipB backbone and side chain are colored cyan, while LplA molecule is colored brown. Arg 70 side chain and Gly 73 backbone of LplA (PDB 3A7A) structurally overlaps with Arg 76 and Gly 78 of *M. tuberculosis* LipB (PDB 1W66). Arg 71 and Gly 72 are the equivalent residues in *E. coli* LipB. Interestingly, LplA Arg 70 side chain forms unique side chain to backbone hydrogen bonds with the carbonyls of Gly73, Gly75, and side chain of Ser72 in the LplA structure (Fig. S8D, colored brown). These interactions have been speculated to stabilize the lipoate-binding loop of LplA. In MtLipB as well, arginine 76 side chain forms identical hydrogen bonds with the backbone carbonyls of Gly 77 and Lys 79, colored cyan in Fig. S8D. The structural similarities between LipB and LplA hint at the possibility of conservation of adenine- and phosphate-binding sites in LipB, like LplA.

**‘RGG’ loop is necessary for full LipB activity**

The importance of RGG loop in LipB function was investigated by generating LipBR71A and LipBG72A/G73A mutants and performing enzyme assays using C₇-CoA as octanoyl donor and GcvH as acceptor substrate. Figure 5A shows a 12% SDS-PAGE gel, displaying molecular weight marker in lane 1, purified apo-EcGcvH in lane 2 (used as a reference), and assays with WT LipB, LipBG72A/G73A, LipBR71A, LipBK135A, and LipBK135A/C169A, respectively, in lanes 3 to 7. Apo-GcvH to C₇-GcvH conversion was ascertained from the forward migration of the GcvH band in the gel. Compared to WT LipB, GcvH conversion was remarkably reduced in lanes 4, 5, 6, and 7, suggesting loss of activity of the mutants we tested. LipBG72A/G73A displayed 39%, while LipBR71A displayed 21% activity compared to WT LipB, the values determined using C18 HPLC reversed-phase chromatography.

Similar LipB assays were also performed using C₇-ACP as an octanoate donor and GcvH as acceptor. A marked reduction in enzyme activity was observed in LipBG72A/G73A and LipBR71A mutants compared to WT LipB (Fig. 5B). In the 12% SDS-PAGE gel, lane 1 is molecular weight marker, lane 2 apo-GcvH (used as a reference), and lanes 3 to 5 assay performed with WT LipB, LipBG72A/G73A and LipBR71A mutants, respectively. C₇-GcvH band was most intense in lane 3 and relatively faint in lanes 4 and 5.

Kinetic measurements were performed on LipBR71A (filled triangles) and LipBG72A/G73A mutants (filled squares), using C₇-CoA as an octanoate donor and GcvH as acceptor. A marked reduction in enzyme activity was observed in LipBG72A/G73A and LipBR71A mutants compared to WT LipB (Fig. 5B). In the 12% SDS-PAGE gel, lane 1 is molecular weight marker, lane 2 apo-GcvH (used as a reference), and lanes 3 to 5 assay performed with WT LipB, LipBG72A/G73A and LipBR71A mutants, respectively. C₇-GcvH band was most intense in lane 3 and relatively faint in lanes 4 and 5.

STD experiments were also performed to monitor the binding of CoA with LipB. STD spectra were also performed using C₇-ACP and C₇-phosphopantetheine were visible in both the spectra, suggesting binding of CoA with the mutants. Figure 4C5 shows the 3¹P NMR spectra for the interaction of CoA with LipBG72A/G73A mutant. The 3¹-phosphate peak displayed a chemical shift change similar to WT protein, suggesting phosphate binding.

**Molecular dynamics simulations suggest RGG sequence imparts conformational flexibility to the LipB active site loops**

The lipoate-binding 69RRXXGGG₇₅ loop of LplA experiences large scale conformational changes that are necessary for binding its substrate (35). Therefore, molecular dynamics (MD) simulations were performed on WT LipB and its RGG mutants using GROningen Machine for Chemical Simulations (Gromacs 2021.5; University of Groningen; Royal Institute of Technology; Uppsala University) to understand the role of the loop in backbone dynamics (36). PDB 1W66 was edited using the swapaa command of Chimera in order to generate...
LipBR76A and LipBG77A/G78A mutant structures, and the resulting PDB files were saved and stored for GROMACS simulations. Figure 6, A–C display the conformation of the capping sheet, 'RGG' loop, and cysteine containing loop in the 21 models generated using 20 ns MD simulations on PDB 1W66 (WT), MtLipBG77A/G78A, and MtLipBR76A mutants. In both WT and MtLipBG77A/G78A simulations, Arg 76 side chain was observed to flip out of the active site cavity as a function of time (Fig. 6, A and B). During the course of simulations, capping sheet and the RGG loop were observed moving upward, away from the active site in WT LipB (Fig. 6A), widening the mouth of the cavity. The starting structure used for simulations is shown as a green ribbon. The direction of capping sheet and loop movement is shown by black arrows. In MtLipBG77A/G78A mutant, the capping sheet displayed restricted movement, but the loop containing Cys176 displayed reasonable change. In MtLipBR76A mutant, motions were restricted in both Cys176 containing loop as well as the capping sheet, with all the loops in the structures converged (Fig. 6C). Figure 6D shows the molecular surface of WT LipB, displaying the active site cavity tunnel, and the RGG loop masking it. We speculate that the movement of the RGG loop away would widen the mouth of the active site. Figure 6E displays root mean square fluctuation (RMSF) values as a function of residue number for WT MtLipB (PDB 1W66, black lines), MtLipBR76A (green lines), and MtLipBG77A/G78A mutant (red lines) simulations. In the WT protein, there were large magnitude RMSF fluctuations in the capping sheet and the RGG loop. The motions in the capping sheet were subdued in MtLipBG77A/G78A and MtLipBR76A mutants. Figure 6F shows the rmsd values for the three proteins as a function of time. All the structures converged within 20 ns simulation time, though MtLipBR76A converged relatively faster and at a much lower rmsd value. Overall, our results highlight the possible role of RGG loop in regulating the movement of the capping sheet and maintaining the conformation of the Cys176 loop, thus, making active site accessible to the substrate and releasing the product.

Discussion

In prokaryotes, two different pathways for lipoic acid modification coexist: (a) an endogenous lipoic acid synthesis pathway, catalyzed by the consecutive action of LipB and lipoyl synthase (LipA) or (b) a salvage pathway carried out by LplA. Yeast and higher eukaryotes lack the scavenging pathway and derive lipoic acid exclusively from the biosynthetic pathway by the sequential action of LipT2, LIAS, and LipT1 (5). The three enzymes work in concert, performing nonredundant functions of the mitochondria. Genetic mutation of any one of these or the upstream type II fatty acid synthesis enzyme causes metabolic dysfunction (37, 38). The disorders cannot be treated by supplementation with exogenous lipoic acid, as all three enzymes cannot uptake free lipoic acid and the cofactor has to be synthesized on-site. Thus, there is urgent need to understand lipoic acid synthesis at the molecular level (11, 21, 22).

LipT2, LIAS, and LipT1 are mammalian homologs of E. coli LipB, LipA, and LplA. LipB (the biosynthetic enzyme) and the salvage enzyme LplA differ from each other in several respects: (a) C8-ACP, an intermediate of the fatty acid synthesis pathway is the only known substrate of LipB, while LplA is a lipoic acid/ATP-dependent enzyme; (b) a cysteine-lysine dyad serve as catalytic residues of LipB, whereas a lysine performs lipoylate adenylation as well as transfer reactions of LplA (35); (c) reaction proceeds via different intermediates in the two enzymes; LplA forms a noncovalent lipoyl-5’-AMP intermediate from lipoic acid and ATP, while a C8-LipB covalent adduct is formed using mitochondrial fatty acid synthase intermediate (C8-ACP). Despite these dissimilarities, unique parallels appear...
to exist between the two enzymes with regard to substrate recognition, as revealed by our studies. STD experiments and $^{31}$P NMR studies suggest binding of adenosine and phosphopantetheine moiety of CoA to LipB, similar to LplA (39). Stronger affinity of holo-ACP for LipB compared to apo-ACP and perturbations in one of the NH resonances of its phosphopantetheine moiety upon LipB titration support the conservation of a phosphate-binding site in LipB, like LplA. Structural overlay of MtLipB (1W66) on the catalytic domain of E. coli LplA (3A7A) also hints toward conservation of residues in LipB at sites that interact with adenine and phosphate groups of AMP in LplA structures (PDB 3A7A, 2ART, 2ARU). Interestingly, the two sites are conserved in the sequence of lipoate transferase 2 (LipT2, UNP A6NK58) as well, and the latter enzyme also converts GcvH using $C_8$-CoA, divulging evolutionary relationships (32).

The salvage enzyme LplA interacts with lipoic acid via a well-defined lipoate-binding loop. An $^{76}$RGG$^{78}$ loop present in the active site of (M. tuberculosis) MtLipB (1W66) has sequence and conformation fairly similar to the lipoate-binding loop $^{69}$RRSSGGGAV$^{77}$ of LplA. In the LplA structure (PDB 3A7A), Arg 69 forms unique side chain to backbone hydrogen bonds with the glycine and Ala of the loop, in a manner similar to Arg 76 of LipB (M. tuberculosis). This lipoate-binding loop is crucial for lipoylation in LplA and experiences large scale conformational changes upon lipoic acid binding (35). MD simulations of M. tuberculosis LipB (PDB 1W66) suggest similar motions in the capping sheet and the RGG loop that are quenched upon mutation of Arg 76 in the sequence to alanine. Conversely, mutation of the conserved glycines in the RGG sequence to Ala increased the overall dynamics of the Cys176 (catalytic cysteine) loop, while the
C₈-phosphopantetheine is recognized by LipB
capping sheet displayed reduced motions, inducing major changes in the conformation of the active site. Interestingly, ‘RGG’ sequences are frequently found in RRM domains and are known to bind phosphates of nucleotides/quadruplexes (40). Electrostatic attraction/hydrogen bonds between positively charged guanidinium of arginine and the negatively charged phosphates contribute to this interaction (41, 42). Interaction between phosphates and phosphotransacetylase enzyme from Methanosarcina thermophila has also been attributed to arginines (43). We surmise that the ‘RGG’ loop of LipB regulates the active site opening of LipB and may be transiently involved in interaction with the phosphate group of its donor substrate, owing to the side chain motions of Arg 76 during simulations.

Taken together, the present study highlights the importance of ACP modifications in substrate recognition by LipB. Apo-ACP weakly binds LipB, while the presence of a phosphopantetheine moiety (in holo-ACP) allows ACP to bind LipB with several-fold higher affinity. Addition of a C₈-chain further increases the binding affinity. These are new observations, shedding light on the unique mechanism of substrate recognition by LipB. We show that the phosphopantetheine moiety, a component of the substrate C₈-ACP not just serves as a swinging arm to transfer acyl chain to the active site cavity of FAS enzymes but also plays a crucial role in substrate recognition by LipB. In the absence of the phosphopantetheine group, binding of ACP and LipB is remarkably weak. The present work complements a previous study on LipB that emphasizes the importance of surface complementarity between C₈-ACP and LipB in substrate recognition by LipB. Based on that study, LipB can differentiate between different acyl intermediates allosterically, without even flipping the acyl chain (28, 29).

Above all, in the present work, we have identified a new substrate analog of LipB, that is C₈-CoA, which can also serve as a substrate for LipT2. This information might be of assistance while treating patients with human mitochondrial fatty acid synthase mutations; genes encoding mitochondrial trans-2-enoyl-coenzyme A-reductase (MECR) (21, 44), malonyl-CoA synthetase (ACS3), malonyl CoA-acetyl carrier protein transacylase (MCAT), etc. characterized by neurological manifestations due to the loss of lipoylation (21). The symptoms overlap with CoPAN and PKAN disorders (defects in CoA biosynthesis), as all these conditions have the same consequence, that is, reduction of lipoylation (25). Two candidates have been proposed as a therapy so far, lipoic acid and octanoic acid. Studies with radioactive lipoic acid supplements show that exogenous lipoic acid enters the blood stream and tissues of mammals but is rapidly degraded by β-oxidation with no noticeable lipoic acid modification (5). Likewise, in LipT2, LIAS, or LipT1 KOs, lipoic acid supplements do not provide any respite (5). We surmise that octanoic acid analogs might serve as a better substitute for C₈-ACP, owing to their ability to pass through the mitochondrial membrane, convert into octanoyl-CoA inside the mitochondria, and act as a donor substrate for LipT2. Support for this proposition also comes from studies on individuals with medium chain acyl-CoA dehydrogenase deficiency, a fatty acid oxidation defect characterized by elevated C₈-carnitine concentration, and lack of acyl-CoAs >C₁₀-CoA. These individuals display increased lipoylated PDC levels compared to healthy controls, associating medium chain CoA to lipoic acid synthesis (45). Another recent study on yeast with deleted 3-hydroxyacyl thioester dehydrogenase, an important enzyme of mitochondrial FAS (Ahtd2) but harboring a mitochondrially mislocalized fatty acyl-CoA ligase (Fam1-1 suppressor allele) was able to rescue the cells by growth on C₈-supplemented media. Fam1-1 has long back been shown to activate fatty acids by attachment to CoA or ACP (46). Authors attributed the suppression of respiratory growth defects in cells lacking mtFAS enzymes by Fam1-1 to a LipB-dependent mechanism, as the cell growth could not be achieved in the absence of lipoylating enzymes or C₈-supplements (47). Our in vitro studies validate their findings, offering proof of concept that LipT2 can indeed efficiently use C₈-CoA as a substrate, besides C₈-ACP. Further studies are warranted, to test the efficacy of octanoate in rescuing FAS-deficient mutants.

Experimental procedures
Cloning and mutagenesis
Genes encoding ACP, LipB, and GcvH were PCR amplified from the E. coli genome and cloned in a pET28a expression vector between NdeI and HindIII restriction sites. The corresponding plasmids have been named as follows: pET28a(+)-EcACP, pET28a(+)-EcLipB, pET28a(+)-EcGcvH. Mice LipT2 gene (synthesized after codon optimization) was also cloned similarly (pET28a(+)-MmLipT2). Mutagenesis studies were performed using site-directed mutagenesis approach using Pfu turbo as the polymerase and pET28a(+)-EcLipB as a template. The PCR product was subjected to DpnI digestion overnight at 25 °C, in order to digest the methylated circular DNA. Approximately 15 μl of this product was transformed in XL-1 blue competent cells. Colonies obtained were inoculated in 5 ml LB media, followed by overnight culture, pelleting, and finally plasmid isolation. For the double mutant, LipB plasmid containing lysine mutation was used as a template. All the mutations (EcLipBK135A, EcLipBC169A, EcLipBK135A/C169A, EcLipBR76A, and EcLipBG72A/G73A) were confirmed by DNA sequencing.

Protein expression
pET28a(+)-EcACP and pET28a(+)-EcLipB plasmids were transformed separately into E. coli BL21(DE3) cells and plated on LB agar supplemented with 50 μg/ml of kanamycin. One colony of each was inoculated in 5 ml of LB media supplemented with kanamycin (50 μg/ml). The primary culture was incubated at 37 °C for 14 to 16 h, accompanied with shaking at 200 rpm. For large scale expression, 1 L LB medium (secondary culture supplemented with 50 μg/ml kanamycin) was inoculated with 1% of the overnight grown primary culture. The secondary culture was kept at 37 °C with shaking until its A₆₀₀ reached a value of 0.6 to 0.8. For ACP and LipB expression, secondary culture was induced with 0.5 mM IPTG and
incubated at 25 °C for 14 to 16 h, with shaking at 200 rpm. Post induction, cells were harvested by centrifugation at 4000 rpm at 4 °C for 30 min, and the pellet was used for protein purification.

\[ ^{15}\text{N}^{13}\text{C}-\text{labeled protein expression} \]

The protocol for transformation and primary culture were exactly the same as described for the unlabeled protein. However, for the secondary culture, 500 ml M9 media was prepared by dissolving Na2HPO4 (3.4 g), KH2PO4 (1.5 g), NaCl (0.25 g), and NH4Cl (0.5 g) in distilled water. The media was autoclaved and cooled. Prior to culture inoculation, 1 g of unlabeled/labeled glucose (40%), 1 ml of 1 M MgSO4, 10% glycerol, and Mn2+ ions were used to activate the natural phosphodiesterase, thereby converting holo-ACP to apo-ACP, enriching the apo-ACP concentration in the lysate (48).

For ACP, culture was incubated at 25 °C for 14 to 16 h, with shaking at 200 rpm. Post induction, cells were harvested at 4000 rpm at 4 °C for 30 min, and the pellet was used for protein purification.

**Purification of ACP**

Pellets were suspended in 30 ml of 20 mM sodium phosphate buffer (pH 7.8) and 500 mM NaCl. EDTA free protease inhibitor cocktail (Sigma) was added to the suspended pellet and subjected to sonication (5 s ON, 5 s OFF, and 30% amplitude). As EcACP is expressed in both apo- and holo-forms, an additional phosphodiesterase activation step was introduced. The sonicated cells were incubated with 10 mM MgCl2 and 2 mM MnCl2 at 35 °C, 120 rpm for 2 h. Both, Mg2+ and Mn2+ ions were used to activate the natural *E. coli* phosphodiesterase, thereby converting holo-ACP to apo-ACP, and enriching the apo-ACP concentration in the lysate (48).

Sonicated cells were centrifuged at 10,000 rpm, and the supernatant was passed through a Ni2+-NTA (Sigma) column, equilibrated with 20 mM sodium phosphate buffer, pH 7.8, containing 500 mM NaCl (washing buffer). The protein-bound resin was washed with 100 ml of washing buffer, followed by 100 ml of 20 mM sodium phosphate, pH 6.0, 500 mM NaCl wash (elution buffer) to remove the nonspecifically bound proteins. Next, the protein was eluted with imidazole in elution buffer and concentrated by 3 kDa cutoff centricons and 10% glycerol, and 300 mM NaCl (resuspension buffer). The concentrated protein was passed through a Superdex 75 FPLC column (size-exclusion chromatography) equilibrated with 50 mM Tris–HCl, pH 7.8, 200 mM NaCl, and 10% glycerol. Pure fractions based on SDS-PAGE were pooled, concentrated, and stored in 50 mM Tris–HCl, pH 7.8, 200 mM NaCl, and 50% glycerol at ~20 °C. For NMR titration studies, glycerol was removed completely prior to the experiments.

Most of the proteins were purified within 1 day starting from pellet, specially *EclipB* WT enzyme as well as the mutants. In case of EcACP, purification took nearly 2 days at 4 °C.

**NMR data acquisition**

NMR samples comprised uniformly labeled [*1H,15N,13C*] protein, in 50 mM Tris–HCl, 200 mM NaCl, pH 8.0, 90% H2O, and 10% D2O, 0.5% sodium azide. All the spectra were acquired on a Bruker Avance III 700 MHz NMR spectrometer, installed at the National Institute of Immunology, equipped with triple resonance TXI probe. Triple resonance NMR experiments, viz. HNCA, HNCAcB, CBCAcOH, etc., were used to assign apo-, holo-, and C8-ACP. *1H15N* TROSY-HSQC spectra were acquired using 1024 data points in the t2 dimension and 256 to 512 data points in the t1 dimension. NMR data were processed on a workstation running Red Hat Enterprise Linux 5.0, using NMRDraw/NMRPipe, and analyzed using Sparky (51). Experiments were performed at 298 K or otherwise as specified. *15N13C* spectra were indirectly referenced using a chemical shift standard, sodium 2,2-dimethyl-2-silapentane-5-sulfonate (52).

**STD experiments**

Saturation transfer difference experiments were acquired at 283 K using Bruker parameter sets for STDDIFFESGP3, a pseudo-2D experiment, and analyzed using Topspin 2.1pl3

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\[ C_8^{\text{phosphopantetheine}} \text{is recognized by LipB} \]
C₈-phosphopantetheine is recognized by LipB

(Bruker’s standard NMR software). A train of 50 ms Gaussian-shaped radio frequency pulses, attenuation level 55 dB, separated by intervals of 2 ms were used. Excitation sculpting was employed to suppress the water signal. The frequency for selective radiation for on resonance was placed at –3 ppm (for upfield shifted methyl protons) and off resonance at 35.5 ppm, respectively. A relaxation delay of 5 s was used. The acquisition time was 0.73 s. Subtracting on from off signal gave a positive STD signal from the bound CoA in the difference spectra. The ligand peaks observed in the 1D STD experiment were assigned based on ¹H NMR spectra of CoA.

³¹P NMR measurements

³¹P NMR measurements were performed on a 5-mm broad band in reverse probe, at 298 K, using Bruker ZG30 parameter set. A spectral width of 400 ppm, 65,536 data points, 2.0 s recycle delay, 14,336 scans, and 0.288 s acquisition time were used. A 30° flip angle was used in the experiment, with no proton decoupling. All the spectra were processed with a line broadening of 1.0 Hz, and phase correction using Topspin 2.1pl3 software.

NMR data analysis

Chemical shift perturbations have been reported as weighted average of the nitrogen and ¹H chemical shifts (ΔAvgHN), derived from Equation 1 (53)

\[ \Delta \text{Avg}_{\text{HN}} = \left[ (\Delta \text{HN})^2 + (\Delta N/5)^2 \right]^{1/2} \]  

where ΔHN and ΔN are the changes in the proton and nitrogen dimensions, respectively. One SD has been used as a cutoff to mark significant change.

NMR titration studies

¹⁵N-labeled C₈-, apo-, or holo-ACP in 50 mM Tris and 200 mM NaCl, pH 7.8 was titrated with LipBK135A/C169A, and chemical shift perturbations were followed in a ¹⁵N TROSY-HSQC experiment. ¹⁵N-labeled holo-ACP and octanoyl-ACP were first prepared by converting apo-ACP to C₈-ACP using B. subtilis phosphopantetheinyl transferase and respective CoA. For each titration point, a ¹H¹⁵N TROSY-HSQC spectra were acquired. ¹⁵N TROSY spectra of apo-/holo-/C₈-EcACP were acquired at a concentration of 0.3 to 0.4 mM and then increasing concentrations of the unlabeled protein EcLipBK135A/C169A was added starting from 0.25 M equivalents to 2 M equivalents sequentially and the spectra recorded after each addition. After each addition, the concentration of apo-/holo-/C₈-EcACP was recalculated taking into account the dilution factor due to EcLipBK135A/C169A addition. Thus, after each titration point, the amount of EcACP decreased and EcLipBK135A/C169A increased. The concentration of ACP, as well as the enzyme stocks used for NMR studies and enzyme assays, were calculated using Bradford reagent in triplicates, and every time a new standard concentration of ACP, as well as the enzyme stocks used for NMR studies.

Enzyme assay

Enzyme activity was performed using 5 μM LipB, 75 μM C₈-CoA/octanoyl-ACP as the donor substrate, and 25 μM GcvH as the acceptor substrate in 50 mM Tris–HCl, 200 mM NaCl, pH 8.0, at 30 °C. The reaction components were incubated for 2 h at 30 °C and then quenched by adding the 1× SDS loading dye and stored at −20 °C overnight. The samples were loaded on a 12% SDS-PAGE gel for qualitative estimation of the conversion of GcvH or native-PAGE gel to follow breakdown of C₈-ACP. For quantitative estimation, reverse phase Symmetry C18-5 μM Waters HPLC column was used. Twenty microliter of the enzyme assay was quenching using 0.1% TFA. The sample was injected onto the C18 Waters column. Proteins were eluted by linear gradient of 0 to 100% acetonitrile, containing 0.1% TFA. For all the assays, absorption was observed at 220 nm. The apo/unmodified protein eluted prior to holo/C₈-bound form. The area under the peak was integrated to determine the concentration of apo-form and holo-form. Kinetic parameters were estimated by GraphPad Prism version 7.0.

For LipT2 assay, the reaction mixture was incubated at 37 °C, and the buffer comprised 50 mM Tris–HCl, 500 mM NaCl, pH 8.0. The octanoyl acceptor substrate was GcvH in all the assays.

All assays were done in duplicates, and in some cases, the error bars were even smaller than the symbols.

Mass spectrometry

The assay samples were passed through C₈ Zip Tips to remove salt, and the final sample was eluted in 50% acetonitrile and 0.1% TFA for analysis in Applied Biosystems 4800 MALDI-TOF spectrometer. Mass Spectrometry was used only to confirm full conversion.
GROMACS simulations

All MD simulations were performed using GROMACS 2021.5 software, installed on a Mac, using OPLS-All Atom force field (36). M. tuberculosis LipB PDB file 1W66 (without the ligand or water) was used for WT protein simulations. For the LipBG77A/G78A and LipBR76A mutants, Chimera (33) was used to mutate the side chain in the PDB file, and the generated PDB file was used for simulations. The protein was solvated in a cubic box of TIP3P water molecules, based on single point charge water model (54). One Na+ in WT, as well as LipBG77A/G78A, and two Na+ in LipBR76A were used by randomly replacing solvent molecules to neutralize the protein. This resulted in a total of 41,015, 41,018, and 41,005 atoms for the WT, LipBG77A/G78A, and LipBR76A, respectively. A second energy minimization step was performed to prevent any steric clashes employing the steepest descent algorithm, until the maximum force in the system was less than 1000.0 kJ mol⁻¹ nm⁻¹ and potential energy was negative ~10⁸. The minimized structure was equilibrated with solvent and ions under constant number of particles (N), volume (V), and temperature (T) (NVT) and constant number of particles (N), system pressure (P) and temperature (T) (NPT) ensembles applying a positional restraining force on the heavy atoms. NVT thermal equilibration was performed using velocity-rescaling temperature coupling for 100 ps at 300 K using a modified Berendsen thermostat with a time constant of 0.1 ps. The temperature of the system reached the target value of 300 K and was stable for the rest of the equilibration. Equilibration of pressure (1 bar) was done for 100 ps under an NPT ensemble, using the Parrinello–Rahman temperature coupling, with a time constant of 2 ps. In NVT and NPT simulations, bonds to hydrogen atom were constrained using LINCS algorithm (55). Next, MD production simulation was done, for 20 ns at constant temperature (300 K) and pressure (1.0 bar) using 10,000,000 steps and integration time step of 2 fs. Electrostatic interactions were computed using a particle mesh Ewald summation algorithm (56), coulomb radius was set to 1 nm, and Van der Waals radius cut off to 1.0 nm. The command trjconv was used to center the protein in the unit cell, and a corrected trajectory was generated. RMSD/RMSF values as well as PDB files were generated from this corrected trajectory file. In all three proteins, RMSD relative to the structure present in the minimized, equilibrated system was found to level off within 20 ns.

Data availability

Data are available from the authors upon request. Please send request to Monica Sundd, monicasundd@nii.ac.in.

Supporting information—This article contains supporting information.

Acknowledgments—The authors thank the Department of Biotechnology, Govt. of India, and SERB (Science and Engineering Research Board) Sanction order No. “EMR/2017/002093” for financial support. Authors also thank Ms Shanta Sen, Mass Spectrometry Facility, NII, for help with Mass Spectrometry data, and the Advanced Instrumentation Research Facility, Jawaharlal Nehru University, India for SPR measurements.

Author contributions—S. K. and M. S. conceptualization; C. D., U. Y., and M. S. methodology; C. D. and U. Y. investigation; S. K. and M. S. writing—original draft; M. S. supervision.

Funding and additional information—Fellowship to C. D. from the Lady Tata Foundation, India, and U. Y. from the University Grants Commission, India is gratefully acknowledged.

Conflict of interest—The authors declare that they have no conflicts of interest with the contents of this article.

Abbreviations—The abbreviations used are: FAS, fatty acid synthesis; GCS, glycine cleavage system; MD, molecular dynamics; NPT, constant number of particles (N), system pressure (P) and temperature (T); NVT, constant number of particles (N), volume (V), and temperature (T); PDB, Protein Data Bank; PDC, pyruvate dehydrogenase complex; RMSF, root mean square fluctuation; SPR, surface plasmon resonance; STD, saturation transfer difference.

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C₈-phosphopantetheine is recognized by LipB

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