Customizing Functionalized Cofactor Mimics to Study the Human Pyridoxal 5’-Phosphate-Binding Proteome

Graphical Abstract

Highlights

- Enrichment of human vitamin B₆-binding proteins with cofactor-derived probes
- In situ target screening of vitamin B₆ antagonists
- Comparison of human cell lines suggests cell-type-dependent cofactor loading states

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In Brief

Fux et al. applied cofactor mimics to study human pyridoxal 5’-phosphate (PLP)-dependent proteins in situ, covering a large fraction of known vitamin B₆-binders. Moreover, cellular targets of anti-vitamin B₆ compounds were unraveled. Comparison of different human cell lines resulted in signature labeling profiles, suggesting different cofactor loading states of PLP-binding proteins.

Fux et al., 2019, Cell Chemical Biology 26, 1461–1468
October 17, 2019 © 2019 The Authors. Published by Elsevier Ltd.
https://doi.org/10.1016/j.chembiol.2019.08.003
Customizing Functionalized Cofactor Mimics to Study the Human Pyridoxal 5′-Phosphate-Binding Proteome

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https://doi.org/10.1016/j.chembiol.2019.08.003

SUMMARY

Pyridoxal 5′-phosphate (PLP) is a versatile cofactor that catalyzes a plethora of chemical transformations within a cell. Although many human PLP-dependent enzymes (PLP-DEs) with crucial physiological and pathological roles are known, a global method enabling their cellular profiling is lacking. Here, we demonstrate the utility of a cofactor probe for the identification of human PLP-binding proteins in living cells. Striking selectivity of human pyridoxal kinase led to a customized labeling strategy covering a large fraction of known PLP-binding proteins across various cancer-derived cell lines. Labeling intensities of some PLP-DEs varied depending on the cell type while the overall protein expression levels of these proteins remained constant. In addition, we applied the methodology for in situ screening of PLP-antagonists and unraveled known binders as well as unknown off-targets. Taken together, our proteome-wide method to study PLP-DEs in human cancer-derived cells enables global understanding of the interactome of this important cofactor.

INTRODUCTION

Vitamin B₆ refers to six interconvertible pyridine compounds: pyridoxine (PN), pyridoxamine (PM), pyridoxal (PL), and their 5′-phosphorylated versions (PNP, PMP, and PLP). The biologically active form is PLP, one of the most ubiquitous cofactors found in nature, that catalyzes around 238 different types of biochemical transformations, corresponding to 4% of all classified activities (Percudani and Peracchi, 2003). Although all living organisms rely on vitamin B₆, only plants and microorganisms are able to synthesize the vitamin de novo. All other organisms, including humans, acquire vitamin B₆ from nutrients and interconvert the vitamers to the cofactor PLP via formation of a Schiff base (the internal aldimine) between the PLP aldehyde group and an ε-amino group of the active-site lysine (Toney, 2005).

PLP-DEs catalyze a diverse set of chemical reactions mainly with amino acid substrates, including decarboxylation, transamination, racemization, α, β, γ-substitution or α, β, γ-elimination (Percudani and Peracchi, 2003). Although sequence similarity is low among PLP-DEs, they are categorized into five distinct fold types guided by structural similarity (El-Sayed and Shindia, 2012). Due to the large number of chemical reactions catalyzed by PLP-DEs, they play a crucial role in cellular metabolism, ranging from amino acid biosynthesis and turnover to synthesis of neurotransmitters or tetrapyrroles (Parra et al., 2018). Therefore, deregulated expression levels of PLP-DEs correlate with several diseases, which have sparked the development of inhibitors for these enzymes. For example, drugs targeting γ-amino-butyric acid (GABA) aminotransferase are used for the treatment of severe epilepsy (Amadasi et al., 2007). In addition, several human PLP-DEs, such as serine hydroxymethyltransferase (SHMT) (Renwick et al., 1998) or ornithine aminotransferase (OAT) (Zigmund et al., 2015), represent attractive targets for drug development due to their role in cell proliferation and tumor development. However, as all PLP-DEs share the same cofactor, the design of selective inhibitors is challenging.

We recently introduced a library of B₆ cofactor mimics composed of alkyn- and azide-functionalized PL-probes to study the Staphylococcus aureus PLP-binding proteome (Hoegl et al., 2018). With this strategy, we were able to detect 73% of all annotated PLP-DEs in S. aureus and assign functions to previously uncharacterized proteins. Here, we customize our methodology for eukaryotic cells and thereby cover a large fraction of the human PLP-ome. We validated our results exemplarily for two PLP-DEs. As an example of application, we demonstrated the utility of our labeling strategy to screen for in situ targets of vitamin B₆ antagonists, revealing insights into the selectivity of the examined compounds. Finally, we uncovered differences in labeling profiles among certain cell types that have the potential to inspire downstream applications for therapeutic purposes.

RESULTS

Bioactivation of PL-Probes by Human PLK

As PL phosphorylation is the limiting step for generating the biologically active form of vitamin B₆ (Figure 1A), we first monitored...
the turnover of PL-probes (Figure 1B) by human PLK (hPLK). PL-probes were synthesized as described previously (Hoegl et al., 2018) with slight modifications in the synthetic route. We purified hPLK as a dimer (Cao et al., 2006) (Figure S1A) and subsequently utilized the recombinant enzyme to record PL turnover at 392 nm, corresponding to the absorbance maxima of PLP (Figures 1C and S1B). The calculated $K_m$ value of about 2.16 mM for natural PL is in good agreement with previous observations (Figures 1D and S1C) (Kimura et al., 2014). The absorbance maxima of the probes were slightly redshifted, with PL1P ($\lambda_{\text{max}} = 416$ nm) exhibiting the most pronounced change compared with PLP. Although all PL-probes were phosphorylated by hPLK, the corresponding catalytic efficiencies varied significantly. For example, the $K_m$ value of PL1, bearing a short substitution, increased about 10-fold, while its catalytic efficiency was only about 3-fold reduced compared with PLP. Although all PL-probes were phosphorylated by hPLK, the corresponding catalytic efficiencies varied significantly. For example, the $K_m$ value of PL1, bearing a short substitution, increased about 10-fold, while its catalytic efficiency was only about 3-fold reduced compared with PLP. Although all PL-probes were phosphorylated by hPLK, the corresponding catalytic efficiencies varied significantly. For example, the $K_m$ value of PL1, bearing a short substitution, increased about 10-fold, while its catalytic efficiency was only about 3-fold reduced compared with PLP. Although all PL-probes were phosphorylated by hPLK, the corresponding catalytic efficiencies varied significantly. For example, the $K_m$ value of PL1, bearing a short substitution, increased about 10-fold, while its catalytic efficiency was only about 3-fold reduced compared with PLP.

### Table 1

| Substrate | $K_m$ [µM] | $V_{\text{max}}$ [µM/min] | $k_{\text{cat}}$ [min$^{-1}$] | $k_{\text{cat}}$ [min$^{-1}$·µM$^{-1}$] |
|-----------|------------|--------------------------|------------------|------------------------|
| PL        | 2.16 ± 0.38 | 0.79 ± 0.02              | 3.16 ± 0.09      | 1.46 ± 0.26            |
| PL1       | 28.0 ± 5.31 | 2.87 ± 0.13              | 11.5 ± 0.51      | 0.41 ± 0.08            |
| PL2       | 582 ± 43.8  | 4.32 ± 0.14              | 17.3 ± 0.55      | 0.03 ± 0.00            |
| PL3       | 516 ± 57.8  | 8.05 ± 0.36              | 32.2 ± 1.43      | 0.06 ± 0.01            |

Metabolic Labeling in HEK293 with PL1 Provides Access to a Large Fraction of the Human PLP-ome

As previous experiments in bacteria demonstrated a superior labeling efficiency upon supplementation of growth media with PL-probes (Hoegl et al., 2018), we cultivated human adenovirus-transfected embryonic kidney cells (HEK293) in the presence of PL1. We maintained cells for seven passages on PL1-containing media lacking natural B$_6$ vitamers to allow for complete turnover of the proteome (Ong, 2002) and therefore accumulation of PL1P in PLP-DEs. We included HEK293 cells passaged in defined complete media with natural PL as control. To account for the lower concentrations (5 µM) of natural PL in standard cell culture media and the increased $K_m$ value of the probe, we cultured cells in the presence of 10 µM PL1. Notably, the doubling rate and morphology of HEK293 was not affected by incubation with PL1 compared with the PL control, indicating that the probe supports essential functions of natural vitamin B$_6$. To exclude that remaining intracellular vitamin B$_6$ levels may impair the labeling results, we relatively quantified PL and PLP levels over the passaging time. Satisfyingly, concentrations of endogenous vitamers decreased markedly after three passages with no PL and PLP left at the time point of the proteomic experiments (seven passages in PL1-containing media; Figure 2A).

Cells were lysed in the presence of sodium borohydride (NaBH$_4$) to attach the probe irreversibly to target proteins by reductive amination. Subsequent bioorthogonal ligation of the alkyne tag to biotin-azide using copper-catalyzed azide-alkyne
1,3-dipolar cycloaddition (CuAAC, click chemistry) allowed for enrichment of probe-bound proteins on avidin beads (Rostovtsev et al., 2002; Speers and Cravatt, 2004; Tomoe et al., 2002). After tryptic digest, peptides were analyzed using Liquid chromatography combined with tandem mass spectrometry (LC-MS/MS) and compared with the PL-cultured control using label-free quantification (LFQ) (Figure 2B) (Cox et al., 2014). We detected 15 PLP-DEs (GO term 0030170: PLP binding; Ashburner et al., 2000) significantly enriched upon PL1 labeling in HEK293 compared with the control (Figure 2C and Table S1), which accounts for around 30% of the complete human pool of PLP-binding enzymes (53 reviewed proteins annotated with GO term 0030170: PLP binding within the human proteome [UniProt: 9606]; Ashburner et al., 2000), including members of fold types I, II, and IV (El-Sayed and Shindia, 2012). Several PLP-DEs related to cancer proliferation and tumor development (e.g., SHMT1; Renwick et al., 1998 or cytosolic aspartate aminotransferase, GOT1; Thornburg et al., 2008) or to Parkinson disease (Bras et al., 2008) and atherosclerosis (Park et al., 2008) were identified. These therapeutically relevant proteins are currently being targeted through the development of selective inhibitors (Ducker et al., 2017; Lowther et al., 2010). Interestingly, pyridoxal 5′-phosphate binding protein (PLPBP), a PLP-DE with an unknown cellular function, displayed the highest fold change by comparing LFQ intensities of PL1 labeling with the PL control.

PL1P Is the Species Bound to PLP-DE Active Sites

In order to validate binding of PL1P to PLP-DEs, we exemplarily selected the well-studied anti-cancer target SHMT1 as well as PLPBP, the strongest enriched protein upon PL1 labeling. Both enzymes are highly conserved throughout evolution and expressed in all tissues (Ikegawa et al., 1999; Renwick et al., 1998). SHMT1 catalyzes the conversion of serine and tetrahydrofolate (THF) into glycine and 5,10-methylene-THF (Ducker et al., 2017), which donate carbon for the de novo synthesis of purine and pyrimidine nucleotide bases for DNA replication (Snell et al., 1988). PLPBP was shown to bind PLP, but no enzymatic activity toward any of the 20 proteinogenic amino acids or their corresponding D-enantiomers was detected (ito et al., 2013).

Inspection of the active-site pocket of SHMT1 with bound PLP (PDB: 1BJ4; Renwick et al., 1998) indicates sufficient space for the additional alkyne group of PL1P (Figure 3A). To test PL1P binding, recombinant SHMT1 was purified as the holoenzyme and transformed to the apo-form by nucleophilic displacement of bound PLP using hydroxylamine (Demoss, 1962). Internal aldimine absorbance re-appeared upon reconstitution of apo-SHMT1 with PL1P and PLP as a control (SHMT1-PL1P and SHMT-PLP) (Figure 3B). Notably, the absorbance maximum of SHMT1-PL1P was redshifted compared with holo-SHMT1 and SHMT-PLP, which is in line with changes observed for unbound PL1P. Thermal denaturation studies revealed a selective increase in melting temperature of apo-SHMT1 upon incubation with increasing PLP amounts, whereas the un-phosphorylated vitamer did not show these characteristics (Figure 3C, left). PL1P also resulted in significant stabilization of the apo-enzyme, suggesting that probe and natural vitamin Bε behave similarly (Figure 3C, right). Finally, apo-SHMT1 was incubated with PL1
or PL1P, the corresponding internal aldimines were reduced with NaBH4, and the protein-bound alkynes conjugated to a Rhodamine-azide tag via CuAAC. SDS-PAGE with fluorescence readout revealed specific and concentration-dependent labeling of SHMT1 with PL1P and only weak background labeling when incubated with PL1 (Figure 3 D). This demonstrates that only the phosphorylated version of the probe can bind the enzyme.

Next, we expressed and purified poorly characterized recombinant human PLPBP. The protein is predominantly dimeric even under reductive conditions, indicating that dimerization is not caused by a disulfide bridge (Figure S2 A). To validate binding of PL1P to PLPBP, we also performed thermal stability assays and in vitro labeling. We observed a melting temperature shift of the dimer from 57°C to around 66°C upon incubation with increasing amounts of PLP, whereas no changes were observed upon incubation with PL (Figure S2B, left). As for SHMT1, similar results were obtained with PL1 and PL1P, where only PL1P led to stabilization (Figure S2B, right). In line with thermal stability assays, we detected a concentration-dependent fluorescence increase exclusively for PL1P, whereas only weak background labeling was observed for PL1 (Figure S2C). PLPBP monomer behaved similarly (Figures S2D and S2E). These validation results are in line with previous experiments using recombinant S. aureus alanine racemase (Hoegl et al., 2018).

Finally, to monitor whether the probe binds to the active-site lysine residues of PLP-DEs, we exemplarily incubated PL1P-bound PLPBP with increasing amounts of the natural cofactor (Figure S2F). Satisfyingly, labeling was outcompeted by a 10-fold molar excess of PLP over probe, validating its binding into the active-site pocket.

In Situ Target Screening of Vitamin B₆ Antagonists
As inhibitor development for PLP-DEs is often challenged by conserved cofactor binding sites, in situ target screening of vitamin B₆ antagonists is an important step to characterize their cellular selectivity. We previously applied our cofactor mimics to uncover new off-targets of D-cycloserine (DCS) in S. aureus (Hoegl et al., 2018). To adapt this selectivity screen for human cells, we exemplarily selected the two enantiomers of the PLP-DE inhibitor penicillamine (Pen), which display poorly characterized cellular specificities and thus represent excellent candidates for competition experiments in PL1-labeled HEK293.

Pen is a degradation product of penicillins (Weigert et al., 1975), which is therapeutically applied in the treatment of rheumatoid arthritis (Jaffe, 1964) and Wilson’s disease (Walshe, 1956). Although the D-enantiomer is the natural occurring isomer, both configurations of Pen display an anti-vitamin B₆ effect (Kuchinskas et al., 1957; Rumsby and Shepherd, 1981; Takahashi and Matsuda, 1976). Several PLP-DEs, including cystathionine γ-lyase (CTH) (Brancalione et al., 2016), SHMT1 (Sukanya et al., 2008), and GOT1 (du Vigneaud et al., 1957), were shown to be inhibited by Pen in vitro. However, our competitive
proteomics studies in HEK293 using l- and d-Pen revealed only a limited number of PLP-DE targets (Figures 4A and 4B, Table S2), suggesting a higher selectivity in situ. The cofactor is displaced by Pen via formation of a thiazolidine adduct (Figure 4C) (Heyl et al., 1948). Both enantiomers show a similar target scope, including PLPBP and SGPL1. Contrary to previous postulations, the primary target within the H2S pathway is cystathionine β-synthase (CBS) instead of CTH (Brancaleone et al., 2016). Interestingly, mitochondrial cysteine desulfurase (NFS1) competed exclusively with l-Pen, indicating an absolute configuration-dependent selectivity. To validate our competition results, we incubated recombinant PLPBP with increasing concentrations of l- and d-Pen. UV-vis spectra displayed a decrease in absorbance of the internal aldimine (Figures 4D and S2G), confirming that Pen indeed binds to PLPBP.

**Expanding Metabolic Labeling to Study Cell-type-Dependent Targeting Profiles**

With a validated PLP-DE profiling tool for human cells in hand, we determined if labeling of this enzyme class across various cell lines would reveal individual signature profiles. For this, we cultivated human cancer-derived cell lines K562 (chronic myelogenous leukemia), HeLa (cervix epithelioid carcinoma), as well as HCT116 (colorectal carcinoma) in the presence of PL1. Satisfyingly, we obtained significant enrichment of several PLP-DEs in all cell lines examined (Figure S3, Table S1). However, throughout all experiments, we observed a certain amount of non-PLP-DE background binders, which are not annotated as putative PLP-DEs. We selected glucosylceramidase (GABA) as common non-vitamin B6-dependent protein detected in all cell lines to evaluate putative PLP-dependency (Figure S4A). Neither a characteristic internal aldimine peak at around 430 nm (Figure S4B) nor substantial stabilization of GBA upon incubation with PLP and PL1P, which is a characteristic feature of PLP-DEs, was observed (Figure S4C), indicating that GBA is most likely bound to the probe in an unspecified manner. Indeed, PLP exhibits a highly reactive aldehyde, which can in principle bind to any e-amino group on reactive lysine residues (Phillips, 2015). In fact, several of the significantly enriched non-PLP-binders carry nucleophilic lysine residues as identified by an amine-reactive probe (Hacker et al., 2017).

Taken together, we were able to enrich 18 PLP-DEs, which account for 34% of the complete human PLP-ome. Interestingly, differences in the number and fold changes of significantly enriched PLP-DEs were observed dependent on the type of cell (Figure 5A, Table S1). Enzymes such as molybdenum cofactor sulfurase (MOCOS), sphingosine-1-phosphate lyase 1 (SGPL1), or PLPBP were enriched throughout all cell lines studied. Remarkably, other PLP-DEs were exclusively detected by our method in particular cell lines, such as mitochondrial 5-aminolevulinate synthase (ALAST) solely in HCT116 or mitochondrial 2-amino-3-keto-4-methylbutyrate dehydrogenase (GCAT) in HEK293. This is an intriguing finding as expression levels of PLP-DEs as well as vitamin B6 salvage enzymes PLK and PNPO were comparable throughout all cell lines (Figures 5B, 5C, and S5) (Schmidt et al., 2018), indicating that the level of cofactor binding is cell-type specific and does not necessarily correlate with protein abundance. This was corroborated by a PLP-ome analysis with an anti-PL antibody (Whittaker et al., 2015). First, we monitored the specificity of the antibody toward reduced PLP-DEs. Satisfyingly, NaBH₄-treated PLPBP and SHMT1 were recognized with only a slight signal resulting from the non-reduced PLP-DEs, whereas hPLK, which does not covalently bind the cofactor, was not detected at all (Figures 5D, left and S5G). We proceeded with the analysis of NaBH₄-treated and non-treated proteomes (Figures 5D, right and S5H). In line with the chemoproteomic data, we observed different cofactor loading...
states dependent on the type of cell for the reduced samples, whereas only slight labeling was observed for the non-reduced control, further verifying antibody specificity. Notably, for HeLa cells, we observed the lowest number of PLP-DEs in both the proteomic and the western blot analysis. This result further supports cell-type specific alterations in PLP-DE cofactor loading states independent of labeling with the probe, which might indicate a so far unrecognized means of enzyme regulation.

**DISCUSSION**

PLP-DEs play crucial physiological and pathological roles in human cells. Here, we customize the application of cofactor mimics to the study of the human PLP-ome in living cells. We observed altered metabolic conversion of the probes by hPLK compared with previous studies with SaPLK, and only probe PL1, with a minimal alkyne substituent, could be efficiently phosphorylated. Despite this restricted scope and the intrinsic electrophilicity of the alkyne linker in PL1, which is prone to nucleophilic attack by cellular cysteines (Lehmann et al., 2016), the probe significantly enriched a large fraction of the human PLP-ome in HEK293, including several clinically relevant PLP-DEs such as SHMT1 and little characterized proteins such as PLBPB. Both proteins were subjected to in-depth studies validating that PL1P is the species bound to recombinant PLP-DEs. The methodology was further utilized for in situ target screening of both enantiomers of the PLP-DE inhibitor Pen, demonstrating an unexpected high cellular PLP-DE selectivity contrary to the results of previous in vitro studies and opening new perspectives for therapeutic applications. Finally, we compared PLP-DE labeling in cancer-derived cell lines K562, HCT116, and HeLa with labeling in HEK293. Interestingly, we observed signature profiles depending on the cell type although corresponding PLP-DEs were ubiquitously expressed. Such changes in cofactor loading are largely unknown and highlight the utility of the probe for a direct readout. This was also supported by an antibody-based PLP-ome analysis of unmodified cells, revealing differences in vitamer loading supporting chemoproteomic data, which might display a mechanism of cell-type-specific enzyme regulation.

**SIGNIFICANCE**

Many essential cellular processes such as glucose, lipid, and amino acid metabolism are driven by PLP-DEs. Thus, mining the human PLP-ome represents an attractive strategy for accessing a large and important enzyme class associated with multiple physiological and pathological functions. We show that probe design, including the chemical positioning and length of the bioorthogonal tag, is a crucial determinant for the success of labeling, which, contrary to bacteria, does not work efficiently for hPLK if the tag is too large. We figured out that a probe with a small modification is able to utilize native PL uptake systems to be incorporated across a significant portion of cellular PLP-DEs. Trapping the transient binding state within PLP-DE active sites permitted labeling and downstream identification of cancer-associated PLP-DEs. Interestingly, individual enzymes largely varied in the cofactor loading status across the panel of cells studied.
Furthermore, application of the probe in a competitive mode revealed targets and off-targets of human PLP-DE inhibitors, which highlight this methodology as a general approach to test for inhibitor selectivity in drug development. Thus, our studies not only provided unprecedented insights into the human PLP-ome but also allowed direct monitoring of cofactor loading as well as inhibitor off-targets.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.chembiol.2019.08.003.

**ACKNOWLEDGMENTS**

This project has received funding from the European Research Council (ERC) and the European Union’s Horizon 2020 research and innovation program (grant agreement no. 725985, CHEMMINE, ERC consolidator grant). Further financial support was provided by the Deutsche Forschungsgemeinschaft (DFG) SFB 749. M.P. acknowledges a PhD fellowship from the Studienstiftung des deutschen Volkes. We thank Mona Wolff, Katja Gliesche, and Katja Bäuml for technical assistance. We thank Stephan M. Hacker for critical evaluation of the manuscript.

**AUTHOR CONTRIBUTIONS**

A.F. designed, planned, and conducted all experiments, including kinetic studies, proteomics, and metabolomics sample preparation, as well as protein expression and analysis. A.F. performed proteomic and intact protein mass spectrometry measurements and statistical analysis of the data. V.C.K. contributed expertise to metabolomics sample preparation, measured metabolomics samples, and analyzed metabolomics data. M.P. and A.H. synthesized PL-probes. S.A.S. supervised experiments. A.F. and S.A.S. wrote the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Mouse monoclonal anti-PLPBP antibody (clone 1G2) | OriGene | Cat# TA505162; RRID: AB_2622886 |
| Rabbit polyclonal anti-SHMT2 antibody | abbexxa | Cat# abx128462 |
| Rabbit polyclonal anti-PNPO antibody | Sigma | Cat# HPA023204; RRID: AB_1855506 |
| Rabbit polyclonal anti-ALAS1 antibody | Thermo Fisher Scientific | Cat# PA5-57434; RRID: AB_2637832 |
| Rabbit polyclonal anti-SHMT1 antibody | abcam | Cat# ab55736; RRID: AB_2285970 |
| Goat anti-rabbit antibody conjugated to horseradish peroxidase | invitrogen | Cat# 32260 |
| Goat anti-mouse antibody conjugated to horseradish peroxidase | invitrogen | Cat# 32230 |
| Mouse monoclonal anti-PLK antibody | Santa Cruz Biotechnology, Inc. | Cat# sc-365173, RRID: AB_10708566 |
| Rabbit polyclonal anti-PL antibody | GeneTex | Cat# GTX12625 |
| **Bacterial and Virus Strains** |        |            |
| *Escherichia coli* Rosetta 2 (DE3) | Merck | Cat# 71400 |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| PL1                  | Hoegl et al., 2018 | N/A |
| PL2                  | Hoegl et al., 2018 | N/A |
| PL3                  | Hoegl et al., 2018 | N/A |
| Glycosylceramidase (GBA) | ProSpec | Cat# enz-908 |
| Lysyl Endopeptidase | Wako | Cat# 125-05061 |
| Trypsin, Sequencing grade | Promega | Cat# V5111 |
| L-Penicillamine | TCI | Cat# P1370 |
| β-Penicillamine | Acros Organics via Fisher Scientific | Cat# 10224750 |
| BTAA ligand | Jena Bioscience | Cat# CLK-067-25 |
| **Critical Commercial Assays** |        |            |
| Roti Quant Universal | Roth | Cat# 0120.1 |
| Pierce™ BCA Protein Assay Kit | Thermo Fisher Scientific | Cat# 23225 |
| ECI western blotting substrate solution | Pierce | Cat# PIER80196 |
| Ponceau S | Sigma | Cat# P3504 |
| **Deposited Data** |        |            |
| raw files, Fasta files, and MaxQuant analysis to PRIDE | This manuscript | https://www.ebi.ac.uk/pride/archive/; PXD014771 |
| **Experimental Models: Cell Lines** |        |            |
| Human HeLa cell line | ECACC via Sigma Aldrich | Cat# 93021013; RRID: CVCL_0030 |
| Human K562 cell line | ECACC via Sigma Aldrich | Cat# 89121407; RRID: CVCL_0004 |
| Human HCT116 cell line | ECACC via Sigma Aldrich | Cat# 91091005; RRID: CVCL_0291 |
| Human HEK293 cell line | ECACC via Sigma Aldrich | Cat# 85120602; RRID: CVCL_0045 |
| **Oligonucleotides** |        |            |
| hPLK forward primer (5’->3’): ggggacaaatgttcgaactaaaaa gcagggctttgagaatctttattttcagggcgaggaggagtgccgggtg | eurofins | custom made |
| hPLK reverse primer (5’->3’): ggggaccactttgtacagagaag gccttgggtcacacccggttgctg | eurofins | custom made |
| SHMT1 forward primer (5’->3’): ggggacaaatgttcgaactaaaaa agcagggctttgagaatctttattttcagggcgaggaggagtgccgggtg | eurofins | custom made |
| SHMT1 reverse primer (5’->3’): ggggaccactttgtacagagaag gccttgggtcacacccggttgctg | eurofins | custom made |

(Continued on next page)
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Stephan A. Sieber (stephan.sieber@tum.de). Plasmids generated in this study have been deposited to Addgene (http://www.addgene.org, Deposit No. 77114; IDs 131231, 131230, 131229).

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Cell Lines**

All cell lines were obtained from ECACC via Sigma Aldrich. K562 (female, Cat# 89121407; RRID: CVCL_0004), HEK293 (female, Cat# 85120602; RRID: CVCL_0045), and HCT116 (male, Cat# 91091005; RRID: CVCL_0291) were cultivated in RPMI-1640, HeLa (female, Cat# 93021013; RRID: CVCL_0030) in DMEM media supplemented with 10% L-glutamine and 10% FCS at 37°C and humidified 5% CO₂ atmosphere. The cells were routinely tested for mycoplasma contamination.
**Microbe Strains**

For protein expression, *Escherichia coli* Rosetta2 (DE3) (Merck, Cat# 71400) were grown in lysogeny broth (LB)-media containing ampicillin (100 μg/mL) and chloramphenicol (34 μg/mL) at 37 °C and with shaking at 200 rpm until reaching the exponential phase (OD600nm 0.6-0.8).

**METHOD DETAILS**

**Chemical Synthesis**

Synthesis of PL2 and PL3 was performed as described previously (Hoegl et al., 2018). Synthesis of PL1 was performed as described with slight modifications. PN hydrochloride was protected with TBS-Cl. Meta-chloro perbenzoic acid (m-CPBA) (77%, 3.86 g, 17.2 mmol, 1.2 eq) was added to a solution of TBS-protected PN (8.00 g, 15.7 mmol, 1.0 eq) in anhydrous dichloromethane (DCM, 150 mL) at 0 °C in five portions. After addition, the reaction was stirred vigorously at r.t for 2 h. Then, the reaction mixture was washed with 20% sodium sulfite solution (1 × 80 mL) and saturated sodium bicarbonate solution (1 × 80 mL), dried over sodium sulfate and concentrated to yield crude TBS-protected N-oxide quantitatively as a white solid, which was used directly in the next step.

Trifluoroacetic anhydride (TFAA, 10.3 mL, 15.5 g, 73.9 mmol, 5.0 eq) was slowly added to a solution of crude TBS-protected N-oxide (7.80 g, 14.8 mmol, 1.0 eq) in anhydrous DCM (120 mL) via syringe at 0 °C. The ice bath was removed and the reaction was stirred overnight. Then, anhydrous MeOH (30 mL) was slowly added at 0 °C and the reaction was stirred for 20 min and subsequently allowed to warm to r.t. while stirring for another 20 min. The mixture was diluted with DCM (200 mL) and neutralized with saturated sodium bicarbonate solution (1 × 200 mL). After separation, the organic phase was dried over sodium sulfate and concentrated under reduced pressure. The residue obtained was purified by flash chromatography (EtOAc/hexanes, 10-50%) to yield TBS-protected diol (3.64 g, 8.80 mmol, 60%) as a pale yellow solid. All further steps were carried out as described (Hoegl et al., 2018).

**Biological Methods**

**Cloning and Overexpression of Proteins**

GBA was purchased from ProSpec (Cat# enz-908). N-terminally Strep-II tagged hPLK, SHMT1, and PLPBP (Addgene IDs No.: 131231, 131230, 131229) were cloned and expressed using the primers and conditions listed in Table S3. Cloning was performed using the Invitrogen Gateway cloning system with pDONR201<sup>Kan</sup> as the donor vector and pDest007<sup>Ampr</sup> as the destination vector. E. coli Rosetta2 (DE3) (Merck, Cat# 71400) carrying the expression plasmids were cultured as described and expression was induced through the addition of 0.2 μg/mL anhydrotetracycline (ATET). Bacteria were harvested and washed with PBS (6,000 x g, 4 °C) prior to cell lysis and protein purification.

**Purification and Analytics of Proteins**

Buffer compositions, as well as columns used for purification are listed in Table S4. The bacterial overexpression culture was resuspended in strep binding buffer and lysed by sonication. The lysate was clarified by centrifugation (36,000 × g, 37 °C) prior to the next step.

**hPLK Kinetic Assay**

PL-probes were phosphorylated using SaPLK as described previously (Hoegl et al., 2018). UV/Vis spectra of PL and PL-probes, as well as their phosphorylated counterparts were recorded in assay buffer (20 mM HEPES pH 7.0, 100 mM KCl, 1 mM MgCl₂) at 37 °C using an InfiniteM200 PRO reader (TECAN, Cat# IN-MNANO) and an extinction coefficient of 40,000 M⁻¹·cm⁻¹ at 280 nm (calculated with ProtParam). Concentrations of holo- and apo-SHMT1 were determined with the Bradford assay (Roti Quant Universal, Roth, Cat# 0120.1) and of PLPBP with the bicinchoninic acid (BCA) assay (Thermo Fisher Scientific, Cat# 23225). Molecular weights of the proteins were confirmed by intact-protein MS (Table S4). Proteins were stored at -80 °C in small aliquots. UV/Vis spectra of protein samples (100 μM) were recorded in corresponding SEC buffers or in phosphate buffered saline (PBS) in the case of GBA at 37 °C on an InfiniteM200 PRO reader (TECAN, Cat# IN-MNANO) (300-600 nm, 2 nm increments).

**Reconstitution of apo-SHMT1 with PLP and PL1P**

For reconstitution with PLP and PL1P, 50 μM apo-SHMT1 were incubated with 5-fold molar excess of cofactor in 10 mM 4-([2-hydroxyethyl]-1-piperazineethanesulfonic acid (HEPES) pH 7.5, 100 mM NaCl, 1 mM dithiothreitol (DTT), 0.5 mM ethylenediaminetetra-acetic acid (EDTA), and 10% glycerol overnight at 4 °C under rotation. Residual PLP or PL1P was removed by desalting the proteins.
using a HiTrap Desalting column (5 mL, GE Healthcare, Cat# 17-1408-01) and an Äktá purification system (GE Healthcare) simultaneously placing them into 10 mM HEPES pH 7.5, 100 mM NaCl, 1 mM DTT, and 0.5 mM EDTA.

**Sample Preparation for Intact-Protein MS**

10 μM protein samples (25 μL, SHMT1 and PLPBP in corresponding SEC buffer) were treated with 10 mM NaBH₄ (2 μL of 250 mM stock prepared fresh in 0.1 M NaOH) at r.t. for 30 min. Residual NaBH₄ was quenched by acidification to pH 5-6 with HCl (5-10 μL of 0.5% FA) and neutralized to pH 7 with NaOH (5-10 μL of 0.1 M NaOH). Samples were diluted to 50 μL with PBS (5 mM final enzyme concentration). 2 μM hPLK was prepared in SEC-buffer for measurement by intact-protein MS.

**Intact-Protein MS**

Full-length protein measurements were performed as described previously (Hoegl et al., 2018). Proteins were measured on a MassPREP On-Line Desalting Cartridge (Waters) on an Ultimate 3000 HPLC system (Dionex) coupled to a Finnigan LTQ-FT Ultra mass spectrometer (Thermo Fisher Scientific) with electrospray ionization (spray voltage 4.0 kV, tube lens 110 V, capillary voltage 48 V, sheath gas 60 arb, aux gas 10 arb, sweep gas 0.2 arb). Protein Deconvolution Software (Thermo Fisher Scientific, Cat# IQLAAEGABSFANOMBAQ) was used for data analysis and deconvolution.

**Gel-Based Labeling of Recombinant Proteins**

Gel-based labeling of recombinant proteins was performed as described previously (Hoegl et al., 2018). In brief, proteins were labeled with PL1 or PL1P and reduced with NaBH₄ previous to CuAAC to attach rhodamine azide. For competitive labeling, PLPBP dimer at 10 μM was incubated with five equivalents of PL1P (added from a 2 mM aqueous stock) in SEC buffer for 30 min at 37°C. Probe-labeled protein was further incubated with indicated molar excess of PLP (added from 2 and 100 mM stocks, respectively) for 30 min at 37°C previous to reduction and CuAAC. Samples were analyzed by SDS-PAGE and fluorescence scanning.

**Thermal Stability Assay**

2 μM of protein in SEC buffer or GBA in PBS were incubated with varying excess of PL(P) and PL1(P) (2 μM of corresponding stocks) and SYPRO orange protein gel stain (Thermo Fisher Scientific, Cat# S6650) was added to a final concentration of 1 x from a 5,000 x dilution. After 30 min at 37°C, the samples were separated on a 12.5% SDS-gel (80 μg of lysate per cell line) and plotted on a PVDF membrane (Roti-PVDF, 0.2 μm, Roth, Cat# 8989.1) using a semi-dry blotting station (Trans-Blot SD Semi-Dry Transfer Cell, Bio-Rad, Cat# 1703940). For PLPBP, SHMT2, PNPO, ALAS1, and SHMT1 blocking was performed using 5% milk-powder (w/v) in PBS-T (PBS supplemented with 0.5% Tween-20) for 1 h. Antibodies (mouse monoclonal anti-PLPBP antibody (clone 1G2, OriGene, Cat# TA505162, RRID: AB_2622886); rabbit polyclonal anti-SHMT2 antibody (abbeza, Cat# abx1284262); rabbit polyclonal anti-PNPPO antibody (Sigma, Cat# HPA023204, RRID: AB_1855506); rabbit polyclonal anti-ALAS1 antibody (Thermo Fisher Scientific, Cat# P5-57434, RRID:AB_2637832); rabbit polyclonal anti-SHMT1 antibody (abcam, Cat#ab55736, RRID: AB_2285970)) were diluted according to manufacturers protocol in 5% milk-powder in PBS-T and added to the membranes. Immunobinding carried out overnight at 4°C. After extensive washing the membranes were incubated with secondary antibodies (goat anti-rabbit antibody conjugated to horseradish peroxidase (0.5 mg/mL, Invitrogen, Cat# 32260) or goat anti-mouse antibody conjugated to horseradish peroxidase (0.5 mg/mL, Invitrogen, Cat# 32230) diluted 1:10,000 in 5% milk-powder in PBS-T and added to the membranes. Immunobinding carried out overnight at 4°C. After extensive washing the membranes were incubated with the secondary antibody (goat anti-mouse antibody conjugated to horseradish peroxidase (0.5 mg/mL, Invitrogen)) diluted 1:2,500 in 3% BSA in PBS-T for 1 h at r.t. The membranes were washed and chemoluminescence was detected after incubation with freshly prepared ECI western blotting substrate solution (Pierce, Cat# PIER80196) in a Luminescent LAS 4000 image analyzer (Fujifilm, ordered via GE Healthcare, Cat# 28955810). Membranes were stained with ponceau S (Sigma, Cat# P3504) to inspect for equal protein loading amounts.

**PLP-one Analysis with the anti-PL Antibody**

Cell lines were grown as described and incubated for 15 min at 4°C in lysis buffer (1% (w/v) sodium deoxycholate, 1% (v/v) NP-40 in PBS pH 7.4). Supernatant was clarified at 21,000 x g for 20 min at 4°C. Protein concentration was adjusted with a BCA-assay. Samples were separated on a 12.5% SDS-gel (70 μg of lysate per cell line and 8 μL of the 15 μM protein samples) and plotted on a PVDF membrane (Roti-PVDF, 0.2 μm, Roth, Cat# 8989.1) using a semi-dry blotting station (Trans-Blot SD, Bio-Rad, Cat# 1703940). For PLK, blocking was performed using 3% BSA (w/v) in PBS-T at r.t. For PLPBP, SHMT2, PNPO, ALAS1, and SHMT1 blocking was performed using 5% milk-powder (w/v) in PBS-T and added to the membranes. Immunobinding carried out overnight at 4°C. After extensive washing the membranes were incubated with secondary antibody (goat anti-rabbit antibody conjugated to horseradish peroxidase (0.5 mg/mL, Invitrogen, Cat# 32260) or goat anti-mouse antibody conjugated to horseradish peroxidase (0.5 mg/mL, Invitrogen)) diluted 1:2,500 in 3% BSA in PBS-T for 1 h at r.t. The membranes were washed and chemoluminescence was detected after incubation with freshly prepared ECI western blotting substrate solution (Pierce, Cat# PIER80196) in a Luminescent LAS 4000 image analyzer (Fujifilm, ordered via GE Healthcare, Cat# 28955810). Membranes were stained with ponceau S (Sigma, Cat# P3504) to inspect for equal protein loading amounts.
32260) diluted 1:5,000 in 5% milk-powder in PBS-T for 2 h at r.t. The membranes were washed and chemo-luminescence was detected after incubation with freshly prepared ECL western blotting substrate solution (Pierce, Cat# PIER80196) in a Luminescent LAS 4000 image analyzer (Fujifilm, ordered via GE Healthcare, Cat# 28955810). Membranes were stained with ponceau S (Sigma, Cat# P38504) to inspect for equal protein loading amounts.

**PLPBP Incubation with Penicillamine (Pen)**

130 μM of PLPBP were incubated with 10, 25, or 50 mM L-Pen (TCL, Cat# P1370) and α-Pen (Acros Organics via Fisher Scientific, Cat# 10224750) (added from 10x stocks in water, pH adjusted to neutral), respectively, for 20 min at 25°C in a total volume of 10 μL. Afterwards, UV/Vis-spectra were recorded as described.

**Proteomics**

**Metabolic Labeling with PL1 and Competition**

All cell lines were cultivated as described. HeLa were transferred into in-house prepared DMEM media (Table S5) lacking pyridoxine, supplemented with 10 μM **PL1** (added fresh from a 100 mM stock in DMSO) and 10% dialyzed FCS (Sigma, Cat# F0392). K562, HEK293, and HCT116 were transferred into in-house prepared RPMI media (Table S5) lacking pyridoxine, supplemented with 10 μM **PL1** (added fresh from a 100 mM stock in DMSO) and 10% dialyzed FCS. All cell lines were additionally transferred into corresponding in-house prepared media containing 5 μM **PL** added fresh from a 50 mM stock in DMSO to ensure for equal amounts of organic solvent.

Cells were passaged seven times in media with **PL1** or **PL**, respectively, and afterwards seeded onto plates (148 cm²) and grown until 90% confluence. One separate plate was seeded for biological replicate (HEK293, HCT116, K562: n = 6; HeLa: n = 4). For competition experiments L- and α-Pen were diluted in in-house prepared media lacking dialyzed FCS to a final concentration of 10 mM (L- and α-Pen) and pH values were adjusted. **PL1**-labeled HEK293 cells were incubated with B6-antagonists for 2 h at 37°C. Cells were washed once with cold PBS (4°C) and then detached in cold PBS. After harvesting at 600 x g at r.t., cells were re-suspended in 1 mL lysis buffer (1% (w/v) sodium deoxycholate, 1% (v/v) NP-40 in PBS pH 7.4) containing 20 mM NaN3H4 (added from a 500 mM stock prepared fresh in 0.1 M NaOH) and incubated for 30 min at 4°C under rotation. Cell debris was removed at 4°C for 20 min (21,000 x g).

Precipitated proteins were precipitated by adding 4 x volume of cold acetone (-80°C) overnight at -20°C. Precipitated proteins were pelleted by centrifugation (21,000 x g, 15 min, 4°C) and washed with ice-cold methanol (MeOH, 2 x 1 mL), using sonication to resuspend the pellets between washes. Proteins were solubilized in 0.5 mL PBS containing 0.4% (w/v) SDS and protein concentrations were adjusted to 2 mg/mL (= 1 mg total per sample). Samples were subjected to CuAAC by adding 0.1 mM biotin-azide (5 μL of 10 mM stock in DMSO), 0.5 mM 2-((bis(1-(tert-butyl)-1H-1,2,3-triazol-4-yl)methyl)amino)methyl)-1H-1,2,3-triazol-1-yl)acetic acid (BTAA, Jena Bioscience, Cat# CLK-067-25) ligand (25 μL of 10 mM stock in ddH2O), 1 mM CuSO4 (10 μL of 50 mM stock in ddH2O) and 2 mM sodium ascorbate (10 μL of 100 mM prepared fresh in ddH2O) to each sample and incubating for 1 h at r. t. in the dark. Upon precipitation and washing (as described above), the protein pellets were re-solubilized in 0.5 mL PBS containing 0.4% SDS (w/v) and centrifuged (18,000 x g, 5 min, r.t.) prior to avidin bead enrichment. Enrichment, reduction, digestion, desalting, and sample preparation for MS was performed as described previously (Hoege et al., 2018) with the exception that enrichment was performed for 1.5 h. In brief, proteins were added to pre-equilibrated beads, enriched under rotation and washed extensively to remove background binders. Samples were reduced and alkylated prior to pre-digest with Lys-C (Wako, Cat# 125-05061) for 2 h at 37°C. Trypsin (Promega, Cat# V5111) digestion took place overnight at 37°C and samples were desalted on Sep-Pak C18 columns (Waters, Cat# WAT04960) afterwards. Peptides were evaporated in vacuo and re-dissolved in water containing 1% FA previous to LC-MS/MS analysis.

**LC-MS/MS Analysis of Proteomics Samples**

Samples were analyzed via LC-MS/MS using a UltiMate 3000 nano HPLC system (Thermo Fisher Scientific) equipped with an Acclaim C18 PepMap100 75 μm ID x 2 cm trap (Thermo Fisher Scientific, Cat# ES803A) and an Acclaim PepMap RSLC C18 separation column (75 μm ID x 50 cm, Thermo Fisher Scientific, Cat# 164535) coupled to an EASY-source equipped Thermo Fisher LTQ Orbitrap Fusion mass spectrometer (Thermo Fisher Scientific). Samples were loaded onto the trap column at a flow rate of 5 μL/min with aqueous 0.1% trifluoroacetic acid (TFA) and then transferred onto the separation column at 0.3 μL/min. Buffers for the nano-chromatography pump were aqueous 0.1% FA (buffer A) and 0.1% FA in acetonitrile (ACN, buffer B). Samples were separated using a gradient raising buffer B from 5 to 22% in 112 min, followed by a buffer B increase to 32% within 10 min. Buffer B content was further raised to 90% within the next 10 min and held another 10 min at 90%. Subsequently buffer B was decreased to 5% and held until end of the run (total: 152 min). During sample separation MS full scans were performed at 120,000 resolution in the orbitrap with quadrupole isolation. The MS instrument was operated in a 3 s top speed data dependent mode. The scan range was set from 300 to 1,500 m/z with 60% RF lens amplitude. The automatic gain control (AGC) target was set to 2.0e5, the maximum ion injection time was 50 ms and internal calibration was performed using the lock mass option. Peptides with intensity higher than 5.0e3 and charge state 2-7 were fragmented with higher-energy collisional dissociation (HCD) (30%). MS² scans were recorded in the ion trap operating in rapid mode. The isolation window was set to 1.6 m/z and the AGC target to 1.0e4 with maximum injection time of 100 ms. Ions were injected for all available parallelizable time. Dynamic exclusion time was 60 s with 10 ppm low and high mass tolerance. Samples for HCT116 **PL1** labeling were analyzed on a UltiMate 3000 nano HPLC system (Thermo Fisher Scientific) equipped with an Acclaim C18 PepMap100 75 μm ID x 2 cm trap (Thermo Fisher Scientific, Cat# ES803A) and an Acclaim PepMap RSLC C18 separation column (75 μm ID x 50 cm, Thermo Fisher Scientific, Cat# 164535) coupled to an EASY-source equipped Q Exactive Plus mass spectrometer (Thermo Fisher Scientific). Sample loading and separation was performed as for measurements on the Fusion instrument. Full scans.
were performed at 140,000 resolution over a scan range of 300–1,500 m/z using an AGC target of 3e6 and a maximum ion injection time of 80 ms. For data-dependent MS2 scans the AGC target was set to 1e5 at 17,500 resolution and 1.6 m/z isolation window. Fragmentation was performed at a normalized collision energy of 30%. The maximum injection time was set to 100 ms. Q Exactive Plus was operating in a TopN = 10 mode. Dynamic exclusion time was set to 60 s.

Metabolomics
Metabolomics Sample Preparation
HEK293 cells were cultured as described. For time-point zero measurements, cells were seeded into 148 cm² dishes in chemically defined media complemented with 5 μM PL (added fresh from a 50 mM stock in DMSO). For later time points, cells were transferred into chemically defined media complemented with 10 μM PL1 (added fresh from a 100 mM stock in DMSO) and cultured till the desired passaging numbers previous to seeding them onto 148 cm² dishes. Cells were grown to 90% confluence. After that, media was removed and cells were scraped to detach in 10 mL cold (4°C) 0.9% NaCl (m/v) in water per plate. Cells were harvested at 600 x g for 2 min. Supernatant was removed and cells were lysed in 1 mL ice-cold MeOH (-80°C, containing 1 μM trimethoprim as internal standard) overnight at -20°C. Samples were centrifuged for 20 min at 21,000 x g and 4°C. Supernatant was dried in vacuo and metabolites were re-dissolved in 90 μL 50/50 Vol-% MeOH/H₂O containing 1% FA for LC-MS/MS measurement.

LC-MS/MS Analysis of Metabolomics Samples
Metabolic profiling and MS/MS analysis was carried out on an UltiMate 3000 RSLC system (Thermo Fisher Scientific) coupled to an LTQ Orbitrap XL mass spectrometer (Thermo Fisher Scientific). Chromatographic separation was performed using an Accucore HILIC column (150 x 2.1 mm, 2.6 μm, 80 Å) (Thermo Fisher Scientific) at 40°C and 80 μm NH₄OAc in 100% H₂O pH = 6.8 (A) and 100% ACN (B). After 5 min pre-equilibration with 95% B, samples were eluted with a linear gradient from 95% to 0% B over 30 min at a flow rate of 250 μL/min followed by 8 min re-equilibration with 95% B. Mass spectrometric measurements were accomplished in positive ion mode (HESI-II source, Thermo Fisher Scientific) with following source parameters: capillary voltage 4.5 kV, tube lens 40 V, vaporizer temperature 43°C, sheath gas 50 L/h, aux gas 10 L/h, capillary voltage 12 V and capillary temperature 320°C. Full scan measurements were recorded between 50 – 1,000 m/z in profile mode at 30,000 resolution in the orbitrap. Parent ions of interest (PL m/z 168.06 and PLP m/z 248.03) were isolated (isolation width 1.0 m/z) subjected to collision-induced dissociation (normalized collision energy 35 V) in the MRM mode and most intense daughter ions (PL/PLP m/z 150.05) were used for quantification (SIM width 10 m/z) at a resolution of 7,500 in the orbitrap. In order to take metabolic degradation over time into account, sample order was randomized. Overall instrument performance was monitored using trimethoprim.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical Analysis of hPLK Kinetic Data
Data were processed using Prism 6 (GraphPad, RRID: SCR_002798). Absorbance values were transformed into concentrations using molar extinction coefficients of PLP and PLP-probes at corresponding absorbance maxima derived from the recorded UV/Vis-spectra (ε₃92nm (PLP) = 6,054.42 M⁻¹cm⁻¹, ε₄16nm (PL1P) = 9,857.49 M⁻¹cm⁻¹, ε₃96nm (PL2P) = 9,374.44 M⁻¹cm⁻¹, ε₄400nm (PL3P) = 6,240.76 M⁻¹cm⁻¹). Initial slopes from linear regions were determined from three technical replicates (with corresponding error) and plotted against concentrations of PL or PL-probe, respectively. Kinetic values (kcat, Km, Vmax) and corresponding errors (SEM) were derived from non-linear regression analysis (kcat and Michaelis-Menten) using default parameters. Means of kinetic values and corresponding errors from the three biological replicates were calculated using error propagation. The number of replicates together with mean values and corresponding SEM are reported in the figure and legend, as well as Supplemental figure and legend for clarity.

Statistical Analysis of Thermal Stability Data
Denaturation curves were fitted according to a sigmoidal trace. Melting temperatures were calculated as mean value with corresponding error (SEM). The number of replicates together with mean values and corresponding SEM are reported in the figure and legend, as well as Supplemental figure and legend for clarity.

Statistical Analysis of Proteomics Data
MS raw data were searched with MaxQuant software (version 1.6.0.1) (Cox and Mann, 2008) and default settings (with the exceptions that label-free quantification and match between runs were activated). All replicates from one condition (e.g. PL1 treated samples) were defined as one fraction. Searches were performed against a UniProt database of Homo sapiens proteome (taxon identifier: 9606, reference reviewed and unreviewed, downloaded on 25.12.2018). Resulting data were further analyzed using Perseus software version 1.6.0.0 (Tyanova et al., 2016). The rows were filtered (only identified by site, potential contaminant, reverse) and log₂ transformed. Biological replicates were grouped, filtered for 100% valid values in at least one group, and missing values were imputed for total matrix using default settings. A both sided, two-sample Student’s t-test was performed and derived p-values were corrected for multiple testing by the method of Benjamin and Hochberg with a significance level of p = 0.05. Volcano plots were generated by plotting log₂fold change of different conditions) against –log₁₀ (p-value). PLP-dependent proteins (gene ontology
(GO-term: 0030170; PLP-binding) were identified with the help of a GO annotation file for H. sapiens submitted on 26.06.2018 (Ashburner et al., 2000). The number of replicates, fold-change and p-value cut-offs are reported in the figures and legends, Supplemental figures and legends, as well as Supplemental tables for clarity.

**Statistical Analysis of Metabolomics Data**
Raw data were processed with Xcalibur Quan Browser (Thermo Fisher Scientific) using Genesis algorithm and manual integration mode. Intensities of the corresponding daughter ions were extracted and further processed using Excel. Finally, daughter ion intensities were plotted for PL and PLP for respective time points using Prism 6 (GraphPad, RRID: SCR_002798). The number of replicates is reported in the figure legend.

**DATA AND CODE AVAILABILITY**
All mass-spectrometric data and corresponding analysis generated during this study have been deposited at the ProteomeXchange Consortium (https://www.ebi.ac.uk/pride/archive/) via the PRIDE partner repository (Vizcaino et al., 2016) with the dataset identifier PXD014771. Plasmids generated in this study have been deposited to Addgene (Deposit No. 77114; IDs 131231, 131230, 131229). The authors declare that all other data supporting the findings of this study are available within the article and its Supplemental Information or from the corresponding author upon request.