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Regulation of homocysteine metabolism by *Mycobacterium tuberculosis* S-adenosylhomocysteine hydrolase

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Mycobacterium tuberculosis modulates expression of various metabolism-related genes to adapt in the adverse host environment. The gene coding for M. tuberculosis S-adenosylhomocysteine hydrolase (Mtb-SahH) is essential for optimal growth and the protein product is involved in intermediary metabolism. However, the relevance of SahH in mycobacterial physiology is unknown. In this study, we analyze the role of Mtb-SahH in regulating homocysteine concentration in surrogate host Mycobacterium smegmatis. Mtb-SahH catalyzes reversible hydrolysis of S-adenosylhomocysteine to homocysteine and adenosine and we demonstrate that the conserved His363 residue is critical for bi-directional catalysis. Mtb-SahH is regulated by serine/threonine phosphorylation of multiple residues by M. tuberculosis PknB. Major phosphorylation events occur at contiguous residues Thr219, Thr220 and Thr221, which make pivotal contacts with cofactor NAD⁺. Consequently, phosphorylation negatively modulates affinity of enzyme towards NAD⁺ as well as SAH-synthesis. Thr219, Thr220 and Thr221 are essential for enzyme activity, and therefore, responsible for SahH-mediated regulation of homocysteine.

-adenosylhomocysteine hydrolase (SahH) catalyzes reversible hydrolysis of S-adenosylhomocysteine (SAH) and produces homocysteine and adenosine¹ (Fig. 1A). Homocysteine is either used in the transsulfuration pathway to synthesize cysteine or re-methylated to methionine for S-adenosylmethionine (SAM) regeneration². Adenosine is either deaminated to inosine or used in the synthesis of nucleotides by salvage pathway². Role of SahH in regulating the concentration of important metabolites such as SAH and homocysteine has been studied in great detail in eukaryotes³-5. In humans, genetic defect in SAHH gene (encoding SAHH/AdoHcy hydrolase) is accompanied by several impaired phenotypes including elevated levels of methionine and homocysteine³. SAH catabolism by SahH helps in increased tolerance to salt stress in plants by maintaining the synthesis of an osmolyte- glycine betaine⁴. Disruption of sahh in bacteria Rhodobacter capsulatus exhibits loss of survival in minimal media, altered SAM and SAH levels and decreased bacteriochlorophyll synthesis⁵. Similarly, SahH inhibitor neplanocin A increases SAH levels in Alcaligenes faecalis⁶. In Mycobacterium tuberculosis, the levels of SAH and homocysteine are modulated in response to different carbon sources and drugs respectively²,²,8. The mechanism underlying this regulation is still unknown. SahH may have an important role in the regulation of these metabolites in M. tuberculosis. sahh is essential for growth of M. tuberculosis²; however, the physiological role(s) of enzyme as well as its regulation is largely unexplored.

SahH consists of two domains- a substrate-binding catalytic domain and a cofactor binding domain¹⁰. The catalytic processes in both, SAH-hydrolytic and SAH-synthetic directions essentially require NAD⁺ as a cofactor¹¹. *M. tuberculosis* is the first prokaryote in which the structure of SahH has been elucidated¹². *M. tuberculosis* SahH (*Mtb*-SahH) forms a homotetramer with one NAD⁺ molecule bound to each subunit¹². Residues involved in catalysis as well as those involved in hydrogen-bonding with NAD⁺ are conserved in *Mtb*-SahH. Interestingly, a distinct feature was observed near the homocysteine-binding site in solvent access channel involving His363 of *Mtb*-SahH¹². In this study, we analyze the regulation of homocysteine concentration by *Mtb*-SahH. We show the role of the conserved residue His363 in enzyme activity. Subsequently, we study the regulation of SahH by serine/ threonine phosphorylation, as well as identify important phosphoracceptor residues which, interestingly, are also involved in NAD⁺-binding. This study indicates that phosphorylation-mediated regulation of SahH is conserved in mycobacteria and it can be a novel mechanism of regulating homocysteine metabolism.



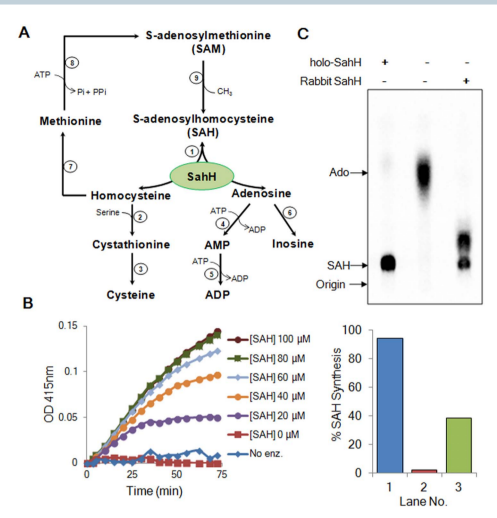


Figure 1 | Metabolic pathways and activity of *Mtb*-SahH. (A) Homocysteine metabolism and the role of SahH. SahH (green oval box) catalyzes reversible hydrolysis of SAH to homocysteine and adenosine as shown by bi-directional arrows. Numbers in circles denote the enzymes present in *M. tuberculosis* H37Rv as follows: 1- SahH (Rv3248c), 2- Cystathionine β-synthase (Rv1077), 3- Cystathionine γ-synthase (Rv1079), 4- Adenosine kinase (Rv2202c), 5- Adenylate kinase (Rv0733), 6- Adenosine deaminase (Rv3313c), 7- Methionine synthase (Rv2124c and Rv1133c), 8- S-adenosylmethionine synthetase (Rv1392) and 9- Methyltransferases, which transfer methyl group (CH₃) to substrate molecules. (B) SAH-hydrolytic activity of His₆-SahH purified from *E. coli.* 100 nM of holo-SahH was used with 0–100 μM of SAH. A reaction containing 100 μM SAH in the absence of enzyme (No enz.) served as a control for spontaneous SAH-hydrolysis. OD was measured at 415 nm and plotted as a function of time (minutes). SahH activity increased with increasing concentration of SAH. (C) SAH-synthetic activity of His₆-SahH purified from *E. coli.* Reaction was performed for 2 hours using 4 μM of holo-SahH and TLC image was analyzed by autoradiography (lane 1). SahH from Rabbit erythrocytes was used as a positive control (lane 3) and a negative control was setup without the (lane 2). Adenosine (Ado) and SAH spots are indicated by arrows. After the densitometric analysis of spots, percent SAH synthesis is plotted in lower panel. An extra spot above SAH-spot in lane 3 could be inosine due to slight contamination (<1%) of adenosine deaminase enzyme in SahH (from rabbit erythrocytes purchased from a commercial source, Sigma) that deaminates adenosine to inosine. Corresponding retention factors (Rf) were SAH (0.1), inosine (0.36) and adenosine (0.5).

Results

Reversible catalysis of SAH by Mtb-SahH. To validate reversible catalysis, Mtb-SahH was cloned in pProEx-HTc and purified as a recombinant His₆-tagged protein (HTc-SahH) from *Escherichia coli*. SAH-hydrolytic activity of purified Mtb-SahH was assessed by a colorimetric assay using Ellman's reagent (DTNB) that detects the product-homocysteine¹³. SahH requires NAD⁺ as a cofactor¹¹ and its NADH-bound form is inactive¹⁴. Since the purified protein contains NAD⁺ as well as NADH¹⁴, there was no detectable activity in purified protein in the absence of any added cofactor (data not shown). Therefore, we prepared apo-SahH, reconstituted it with NAD⁺ to obtain holo-SahH, which was used with increasing concentration of SAH (0–100 μ M) to analyze SAH-hydrolysis. An increase in SAH-hydrolytic activity was observed with increasing amounts of SAH (Fig. 1B), implying that holo-SahH was active. The reaction was found to attain saturation at 80 μ M of SAH after which the

enzyme activity did not increase further. The hydrolytic activity also increased with increasing amounts of holo-SahH (10–400 μ M) (Supplementary Fig. S1). These results confirm SAH-hydrolytic activity of Mtb-SahH.

To validate that Mtb-SahH is also active in reversible direction, we analyzed its ability to synthesize SAH from homocysteine and adenosine. Mtb-SahH was incubated with [8-\frac{14}{C}]-adenosine and homocysteine and the reactions were carried out for 1 or 2 hours. Reaction products were then resolved by thin layer chromatography (TLC) and analyzed by autoradiography. Apo-SahH showed negligible activity in the absence of NAD+ (Supplementary Fig. S2A), while holo-SahH could synthesize considerable amounts of SAH (Fig. 1C). Further, we observed that the activity increased with increasing concentration of holo-SahH (1-4 μ M) (Supplementary Fig. S2B). Taken together, these assays show that M. tuberculosis SahH is an active enzyme that can catalyze reversible hydrolysis of SAH.



Role of His363 in SahH activity. SahH is one of the highly conserved proteins in both eukaryotes and prokaryotes¹⁰. His363 is a conserved residue in Mtb-SahH, which based on the sequence (Supplementary Fig. S3) and structural analyses, is proposed to act as a switch, that opens to allow access to the substrate SAH11,12. His363 needs to flipout in order to accommodate homocysteine moiety of SAH12, although its role in enzyme activity has not been proven experimentally. To gain insight into this mechanism, His363 residue of Mtb-SahH was mutagenized to alanine and the SahH-H363A mutant was purified as a recombinant protein from E. coli. SAH-hydrolytic assays were performed with equal amounts of wild type SahH (SahH-WT) and SahH-H363A, where the mutant was found to be completely inactive (Fig. 2A). To confirm the role of His363 in catalysis we also checked the efficiency of H363A mutant to synthesize SAH. SahH-H363A mutant showed reduced activity to synthesize SAH, as compared to SahH-WT (Fig. 2B). These results confirm that His363 does act as a key residue for SahH enzyme activity.

Regulation of homocysteine concentration by SahH. SahH catalyzes a reversible reaction where *in vitro* equilibrium favors the SAH-synthetic direction, but under *in vivo* conditions, hydrolytic activity is preferred as the products of this reaction (homocysteine and adenosine) are constantly being used by downstream enzymes¹. In order to observe the favored direction of catalysis in mycobacteria, we examined the role of SahH in the regulation of intracellular

homocysteine concentration in the surrogate host *Mycobacterium smegmatis*. Any change in the expression or activity of SahH may lead to an alteration in the concentration of homocysteine. *Mtb*-SahH was over-expressed in *M. smegmatis* MC² 4517 using pYUBDuet shuttle vector. Over-expression of *Mtb*-SahH was confirmed by immunoblotting using antibodies generated against *Mtb*-SahH (Supplementary Fig. S4). Total intracellular homocysteine concentration was then measured in *M. smegmatis*. The homocysteine amounts were $1.411 \pm 0.25 \, \mu \text{moles}/10^{15}$ colony forming units (CFU) in *M. smegmatis* (empty) and $1.346 \pm 0.18 \, \mu \text{moles}/10^{15}$ CFU in *M. smegmatis* harboring only the vector used for over-expression of *Mtb*-SahH (Table 1). Over-expression of *Mtb*-SahH increased homocysteine level to $6.363 \pm 2.06 \, \mu \text{moles}/10^{15} \, \text{CFU}$ (Table 1 and Fig. 2C). Thus, increase in levels of SahH in mycobacteria causes increase in this critical metabolite.

Conservation of SahH in mycobacteria and its regulation by phosphorylation. Since SahH is a key enzyme involved in metabolism, understanding its regulation will be helpful in revealing the mechanisms underlying homocysteine metabolism. In a prior study, during the analysis of intracellularly expressed proteins of *M. tuberculosis*, multiple spots of *Mtb*-SahH were found with distinct isoelectric points in two-dimensional electrophoresis, suggesting post-translational modifications of this protein¹⁵. Interestingly, *Mtb*-SahH was found to be a phosphorylated protein in a large-scale phosphoproteome analysis of *M. tuberculosis*¹⁶. Based

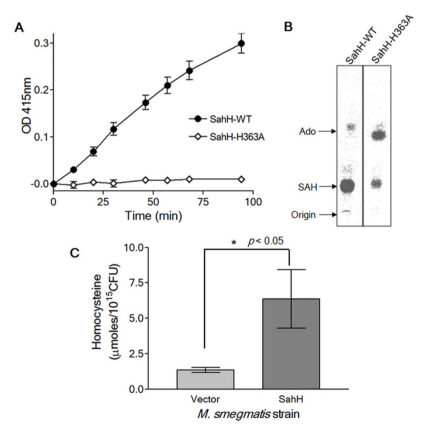


Figure 2 | Role of His363 in activity and effect of SahH on homocysteine concentration in M. smegmatis. (A) Role of His363 in SAH-hydrolytic activity. Activities of 1.74 μ M each of holo SahH-WT and holo SahH-H363A were compared. Graph shows increase in OD at 415 nm with time when SahH-WT was used. No increase in OD at 415 nm was observed with SahH-H363A mutant. Error bars represent standard deviation (SD). (B) Role of His363 in SAH-synthetic activity. Activities of 4 μ M each of holo SahH-WT and holo SahH-H363A were compared. Autoradiograph of TLC shows SAH synthesis in reactions containing SahH-WT while reduced product formation was seen in reactions containing SahH-H363A. Adenosine (Ado) and SAH spots are indicated by arrows. (C) Histogram representing intracellular homocysteine concentration in M. smegmatis MC² 4517 with and without the overexpression of Mtb-SahH. Homocysteine concentration was analyzed using UPLC and plotted as μ moles/ μ 0.15 CFU. Homocysteine levels increased to about six times in cells over-expressing SahH (see also Table 1). Error bars represent standard deviation of three independent experiments and * p < 0.05 as calculated by Student's t-test.



Table 1 Homocysteine concentration [(μmoles/10 ¹⁵ CFU) ± SD] in M. smegmatis MC ² 4517					
M. smegmatis strains	Empty (no vector)	pYUBDuet vector $^{\alpha}$	pYUBDuet:SahH [∞]	pYUBDuet:SahH-T219A/T220A/T221A	
Homocysteine concentration	1.411 ± 0.251	1.346 ± 0.176	6.363 ± 2.057	1.030 ± 0.258	
² p value is 0.0136 as assessed by Student's Hest.					

on these earlier findings, we analyzed the phosphorylation status of SahH in different *Mycobacterium* species. SahH showed high amino acid sequence conservation in ten different species of *Mycobacterium* genus including both pathogenic and non-pathogenic bacteria (Supplementary Fig. S3). In order to assess conserved phosphorylation mediated regulation of SahH across different mycobacterial species, we analyzed phosphorylation status of SahH in *Mycobacterium bovis* BCG and *M. smegmatis*. Native SahH from *M. bovis* BCG (*Mbo*-SahH) was immunoprecipitated and its phosphorylation status was analyzed. The identity of SahH in immunoprecipitated sample was first confirmed by immunoblotting with anti-SahH antibodies

(Supplementary Fig. S5A). Phosphorylation was then confirmed by immunoblotting with anti-phospho-threonine (anti-pThr) anti-bodies as the major phosphorylated amino acid in *M. tuberculosis* is Thr¹⁶. *Mbo*-SahH was found to be phosphorylated (Fig. 3A). PknB_c (catalytic domain of *Mtb*-PknB¹⁷, 1–331 amino acids) and GST were used as positive and negative controls, respectively. This result demonstrates that SahH is a potential substrate of serine/threonine protein kinases (STPKs) in *M. bovis* BCG.

Further, phosphorylation of *Mtb*-SahH was confirmed in *M. smegmatis*. *Mtb*-SahH was over-expressed as a His₆-tagged protein and purified from *M. smegmatis*. Purified fraction (pSD5-SahH) was

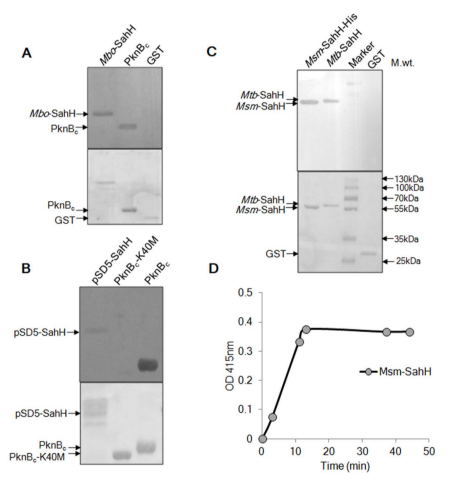


Figure 3 | *In vivo* phosphorylation of SahH and characterization of *Msm*-SahH. (A) *In vivo* phosphorylation of SahH in *M. bovis* BCG. Native SahH was immunoprecipitated from *M. bovis* BCG and its phosphorylation was assessed by immunoblotting using anti-pThr antibodies. Upper panel shows immunoblot and lower panel shows ponceau-stained membrane. PknB_c and GST were used as positive and negative controls, respectively. The protein band corresponding to SahH in *Mbo*-SahH was not visible on ponceau image but was found to be highly phosphorylated on Thr residues (See also Fig. S5A). (B) *In vivo* phosphorylation of SahH in *M. smegmatis*. *Mtb*-SahH was cloned and over-expressed in *M. smegmatis* as a His₆-tagged protein and phosphorylation was assessed in purified fraction (pSD5-SahH) by immunoblotting using anti-pThr antibodies. Upper panel shows immunoblot and lower panel shows ponceau-stained membrane. PknB_c and PknB_c-K40M were used as positive and negative controls, respectively (See also Fig. S5B). (C) Detection of *Msm*-SahH by immunoblotting. *Msm*-SahH was over-expressed and purified from *E. coli* and analyzed by immunoblotting using anti-SahH antibodies. Upper panel shows immunoblot and lower panel shows ponceau-stained membrane. *Mtb*-SahH and GST were used as positive and negative control respectively. (D) Activity analysis of *Msm*-SahH purified from *E. coli* as a His₆-tagged protein. 1.74 μM of holo *Msm*-SahH was used for the activity analysis and OD at 415 nm is plotted as a function of time. Graph shows increase in SAH-hydrolytic activity over time.



phospho-enriched and immunoblotted with anti-SahH and antipThr antibodies. The phospho-enriched sample contained multiple protein bands and presence of SahH was confirmed by immunoblotting with anti-SahH antibodies (Supplementary Fig. S5B). Protein identity was also confirmed by mass spectrometry (data not shown). Immunoblotting with anti-pThr antibodies showed that *Mtb*-SahH is phosphorylated in *M. smegmatis* by native STPKs (Fig. 3B). PknB_c and PknB_c-K40M¹⁸ (catalytically inactive mutant of PknB) were used as positive and negative controls, respectively. These results together confirm that SahH gets phosphorylated *in vivo* in different species such as *M. bovis* BCG and *M. smegmatis* suggesting that SahH is a conserved substrate of STPKs.

M. smegmatis sahh was initially thought to be a pseudogene with a frame-shift mutation, although later it was corrected and proved to be a sequencing error¹⁹, still there was no conclusive experimental proof of presence of active *M. smegmatis* SahH (*Msm*-SahH). We therefore, cloned the gene coding for *Msm*-SahH in pProEx-HTc and purified the protein from *E. coli*. In sequence analysis, this gene did not show any frame-shift mutation and the sequence coded an ORF of 485 amino acids (data not shown). Anti-SahH antibodies generated against *Mtb*-SahH were able to detect the purified *Msm*-SahH (Fig. 3C). Also, *Msm*-SahH was found to be enzymatically active (Fig. 3D) confirming conservation of SahH in *M. smegmatis*.

In vitro phosphorylation and dephosphorylation of *Mtb*-SahH. Ser/Thr phosphorylation is a vital component of signaling machinery of *M. tuberculosis* consisting of 11 STPKs (PknA to PknL) and one Ser/Thr phosphatase (PstP)²⁰. Out of 11 STPKs, only four (PknA, PknB, PknG and PknL) are conserved in *Mycobacterium leprae*, which has evolved after extensive gene decay and elimination,

leaving only the genes that are critical for survival21. pknA and pknB (the genes coding for PknA and PknB, respectively) are located in the same genomic region as pstP (the gene coding for PstP)20. The gene coding for PknB is essential for the growth and survival of M. tuberculosis²² and PknB regulates several fundamental processes like cell shape, cell division, metabolism, protein synthesis and response to stress 18,23-27. To assess the phosphorylation of SahH by PknB, in vitro kinase assays were performed with Mtb-SahH and PknB_c. As seen in the autoradiograph (Fig. 4A), Mtb-SahH was found to be phosphorylated by PknBc. Phosphorylation was specifically localized to the protein and not on the His6-tag as evident after the cleavage of tag by TEV protease (Supplementary Fig. S6). We also found that SahH is a target of multiple STPKs of *M*. tuberculosis (data not shown), which is consistent with earlier observation¹⁶. The reversible regulation of Mtb-SahH by phosphorylation was confirmed by the dephosphorylation assay with PstP_c (catalytic domain of PstP28, 1-300 amino acids). SahH phosphorylated by PknB_c was found to be dephosphorylated by PstP_c in a time-dependent manner (Fig. 4B). These experiments show that SahH is a substrate of Mtb-STPK PknB and Ser/Thr phosphatase PstP.

Phosphorylation of *Mtb*-SahH in the surrogate host *E. coli*. To further substantiate PknB-specific *in vitro* SahH phosphorylation, *Mtb*-SahH was co-expressed in *E. coli* with PknB in a dual expression vector pETDuet-1. As a control, catalytically inactive mutant PknB-K40M¹⁸ was co-expressed with *Mtb*-SahH instead of PknB. To discern the *in vivo* phosphorylation of SahH, *E. coli* cells containing the dual expression constructs were metabolically labeled with ³²P-orthophosphoric acid to label the phosphorylated proteins.

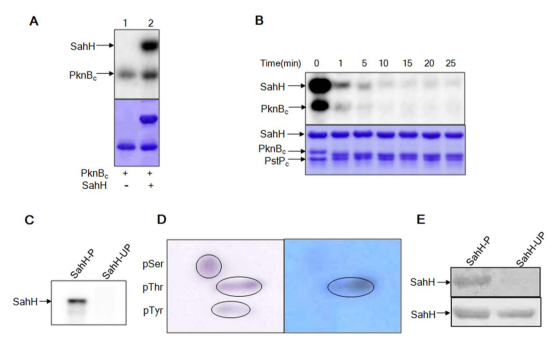


Figure 4 | Phosphorylation of *Mtb*-SahH by PknB_c. (A) *In vitro* kinase assay of *Mtb*-SahH with PknB_c. Upper panel shows autoradiograph with PknB_c alone (lane 1) and PknB_c with SahH (lane 2). Lower panel shows the corresponding coomassie-stained gel image. As evident, SahH is phosphorylated by PknB_c. (B) *In vitro* dephosphorylation of *Mtb*-SahH by PstP_c. SahH phosphorylated by PknB_c was subjected to dephosphorylation by PstP_c and samples were taken for analysis at the indicated time points, for a total of 25 minutes. Upper panel shows autoradiograph and lower panel shows coomassie-stained gel image. (C) Autoradiograph of metabolic labeling of SahH-P and SahH-UP. *E. coli* expressing SahH-P or SahH-UP was subjected to metabolic labeling and autoradiograph is shown. SahH co-expressed with PknB (SahH-P) was observed to be phosphorylated as opposed to SahH co-expressed with PknB-K40M (SahH-UP). (D) Phosphoamino acid analysis of SahH-P by 2D-TLE. Left panel shows ninhydrin-stained TLC plate and right panel shows the corresponding autoradiograph. Residues pThr, pSer and pTyr have been encircled. As observed in the autoradiograph, only pThr is detected in SahH phosphorylated by PknB. (E) Immunoblot analysis with anti-pThr antibodies. SahH-P and SahH-UP were purified and their phosphorylation status was analyzed. Upper panel shows the immunoblot and lower panel shows ponceau-stained membrane. SahH-P was found to be phosphorylated on Thr residues.



Ni+-NTA affinity pull-down was then used to extract SahH and purified proteins were run on SDS-PAGE followed by autoradiography. pETDuet-SahH:PknB (renamed SahH-P) was found to be phosphorylated while pETDuet-SahH:PknB-K40M (renamed SahH-UP) was not phosphorylated (Fig. 4C). To identify the phosphorylated amino acid residues of SahH, phosphoamino acid analysis was performed with purified SahH-P followed by twodimensional thin layer electrophoresis (2D-TLE). Phosphorylation was present specifically on Thr residues (pThr) while neither phospho-Ser (pSer) nor phospho-Tyr (pTyr) was detected on SahH-phosphorylated by PknB in E. coli (Fig. 4D). To validate Thr phosphorylation, SahH-P and SahH-UP were purified as His₆-tag proteins and analyzed for the phosphorylation by immunoblotting using anti-pThr antibodies after their interaction with PknB or PknB-K40M in E. coli cells. While, SahH-UP was found to be unphosphorylated, SahH-P was found to be phosphorylated on Thr residues (Fig. 4E). These experiments reaffirm that Mtb-SahH is phosphorylated by PknB.

Identification and validation of phosphorylation sites. Purified SahH-P and SahH-UP were subjected to mass-spectrometric analysis to identify the phosphorylation sites. Five Thr and one Ser residues were found to be phosphorylated on SahH by PknB in *E.coli* (Fig. 5A). Among the phospho-sites, Thr216, Thr219, Thr220 and Thr221 are present in the domain I, which is the substrate binding

catalytic domain. Thr257 and Ser376 are present in the domain II which is the cofactor binding domain¹². No phosphorylated residues were identified on SahH-UP (data not shown).

To validate these phosphorylation sites of SahH, specific Ser and Thr residues were mutated to alanine to generate phospho-deficient variants of SahH which cannot accept a phosphate group. SahH-WT and mutant proteins co-expressed with PknB in E. coli were thus purified, analyzed for phosphorylation by immunoblotting and subsequent densitometric analysis was plotted. Single site mutants of Thr219, Thr220, Thr221 and Thr257 residues showed 25-50% loss in phosphorylation signal except SahH-T216A which showed hyperphosphorylation (Fig. 5B, Supplementary Fig. S7A). No considerable loss in phosphorylation was observed in SahH-S376A by immunoblotting using anti-pSer antibodies (data not shown). Multiple Thr mutants of SahH were then generated and analyzed in the same way. Maximum loss (approximately 90%) in the phosphorylation signal was obtained in the triple mutant of Thr219, Thr220 and Thr221 (SahH-T219A/T220A/T221A) (Fig. 5C, Supplementary Fig. S7B). To substantiate the phosphorylation of these three residues, E. coli containing SahH-WT or SahH-T219A/T220A/T221A, both coexpressed with PknB, were subjected to metabolic labeling as discussed in previous section and proteins were analyzed by autoradiography. The SahH-T219A/T220A/T221A mutant showed approximately 60% loss in phosphorylation intensity when compared to SahH-WT (Fig. 5D, Supplementary Fig. S8). Thus,

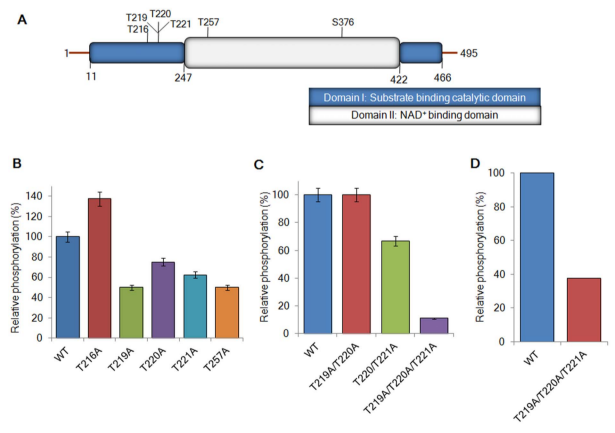


Figure 5 | Analysis of phosphorylation sites on *Mtb*-SahH. (A) Diagrammatic representation of phosphorylation sites at SahH-P identified by mass-spectrometry. Five Thr (Thr216, Thr219, Thr220, Thr221 and Thr257) and one Ser residues (Ser376) were identified and labeled above the respective domains of SahH. Domain I is shown in blue color and domain II is shown in white color. (B) and (C) Histograms depicting relative phosphorylation of SahH and its single or multiple Thr mutants, respectively. Purified SahH-WT and its mutants were analyzed by immunoblotting with anti-pThr antibodies. Percent relative phosphorylation was calculated considering the phosphorylation intensity of SahH-WT as 100%. Corresponding immunoblots are shown in Supplementary Fig. S7A and S7B. (D) Metabolic labeling of SahH-WT and SahH-T219A/T220A/T221A mutant. Histogram depicting densitometric analysis of autoradiograph of metabolically labeled proteins purified from *E. coli*. Percent relative phosphorylation is shown considering phosphorylation intensity of SahH-WT as 100%. Corresponding autoradiograph and coomassie-stained gel image is shown in Supplementary Fig. S8.



Thr219, Thr220 and Thr221 are the major sites of phosphorylation at SahH by PknB. A mutant of all the four Thr residues at sites 216, 219, 220 and 221 (SahH-T216A/T219A/T220A/T221A) was also generated and found to be hyperphosphorylated (Supplementary Fig. S7B). The mutation of Thr at the position 216 could have resulted in structural perturbations leading to opening of tertiary structure, which served as a non-specific substrate for the phosphorylation. As a result, both SahH-T216A and SahH-T216A/T219A/T220A/T221A mutants were hyperphosphorylated (Fig. 5B and Supplementary Fig. S7B).

Phosphorylation-mediated changes in SahH activity. The major phospho-sites at SahH (Thr219-Thr221) obtained in our study with PknB are also involved in NAD+ binding12. Therefore, we chose to analyze the phosphorylation-mediated changes in the affinity of SahH towards NAD+. To assess the kinetic differences in the activity of SahH due to its phosphorylation, SAH-hydrolytic activities were measured with the apo-forms of SahH-P and SahH-UP using increasing concentration of NAD⁺. Phosphorylation of Mtb-SahH by PknB in E. coli did not affect the Vmax of SAH-hydrolysis although the K_m for cofactor NAD⁺ increased significantly (Fig. 6A). While the K_m of unphosphorylated SahH was 0.31 \pm 0.042 μ M, phosphorylation significantly increased it to 1.08 \pm 0.122 μM (Table 2). Thus, phosphorylation decreased the affinity of SahH for NAD+, confirming that the Thr219-Thr221 residues play a decisive role in NAD+ binding. Similar negative effect of phosphorylation was observed when the enzyme activity of SahH was assessed in synthetic direction. SahH-P was less efficient in the synthesis of SAH using adenosine and homocysteine (Fig. 6B). Densitometric analysis indicated that SahH-P was only 50% efficient in the synthesis of SAH as compared to SahH-UP (Fig. 6C).

Table 2 Kinetic parameters of SahH				
$\text{Kinetic parameters}^{\alpha}$	SahH-P	SahH-UP		
Specific activity (pmoles/ mg/min) ± SE	11.42 ± 0.322	10.47 ± 0.215		
$K_{\rm m}$ (μ M) \pm SE $^{\beta}$	1.083 ± 0.1218	0.3152 ± 0.04249		
^a Values represent mean of three independent experiments with standard error (SE). ^β p value is 0.004 as assessed by Student's Hest.				

Since the major sites of phosphorylation were found to be Thr219–Thr221, we analyzed the enzyme activity of SahH-T219A/T220A/T221A mutant. We hypothesized that if the side-chains of these three Thr residues are involved in NAD⁺ binding, then the mutant should not be able to bind the cofactor. We found that SahH-T219A/T220A/T221A mutant had no SAH-hydrolytic activity when compared to SahH-WT protein (Fig. 6D), supporting our hypothesis. These results show the importance of Thr219–Thr221 residues and their phosphorylation in regulating SahH activity.

Structural analysis of NAD⁺ **binding.** To further understand the phosphorylation-mediated changes in the cofactor binding, we analyzed the available crystal structure of *Mtb*-SahH (PDB ID: 2ZIZ) with respect to the NAD⁺ binding. NAD⁺ binds to the domain II of SahH (Fig. 7A) but still makes close contacts with the three Thr residues at positions 219, 220 and 221 of domain I (Fig. 7B). As shown in the figure, the side-chain oxygen atoms (OG1) of all the three Thr residues were within 5 Å reach of the oxygen atom (O2D) at the C-2' of the ribose moiety containing the nicotinamide of NAD⁺. The region encompassing the Thr219, Thr220 and Thr221 residues of SahH was then enlarged to show their interaction with

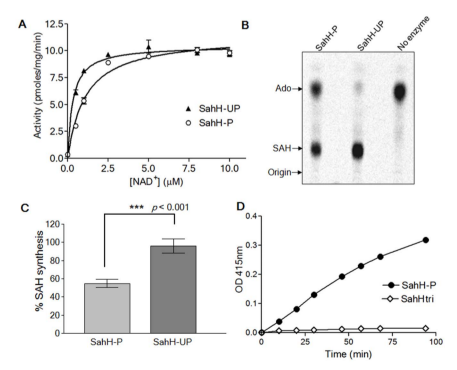


Figure 6 | Effect of phosphorylation on enzyme activity of Mtb-SahH. (A) Effect of phosphorylation on SAH-hydrolytic activity. Michaelis-Menten plot of SahH-P and SahH-UP is shown with concentration of NAD⁺ at horizontal axis. Phosphorylation of SahH is accompanied by increase in K_m and no change in V_{max} (see also Table 2). (B) Effect of phosphorylation on SAH-synthetic activity. Autoradiograph of TLC is shown with reactions containing 4 μ M each of SahH-P, SahH-UP and no enzyme. Intensity of SAH spot decreases when SahH-P is used as compared to SahH-UP. (C) Histogram showing percent SAH synthesis by SahH-P and SahH-UP as observed in (B). SahH-P is only 50% active in synthesis of SAH. Error bars represent SD of three independent results. *** p value equal to 0.001 as calculated by Student's t-test. (D) Role of Thr219, Thr220 and Thr221 in SAH-hydrolytic activity. Activities of 1.74 μ M each of holo SahH-WT (SahH-P) and holo SahH-T219A/T220A/T221A (marked as SahHtri) purified from E. coli were compared. Graph shows increase in OD at 415 nm of SahH-P with time while SahHtri does not show any detectable activity.



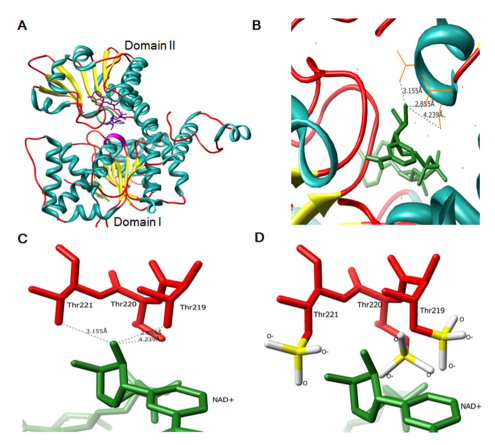


Figure 7 | Structural analysis of SahH and its major phosphorylation sites. (A) Cartoon representation of *Mtb*-SahH bound to NAD⁺ cofactor. Both the domains of SahH have been labeled. Thr219–Thr221 residues have been colored (magenta) to mark their proximity with NAD⁺ (purple). Other color representations are: helix (cyan), strand (yellow) and turn (red). (B) Region depicting interaction of NAD⁺ (green) with Thr219–Thr221. Side chains of Thr residues are shown in orange color. Distances of side-chain oxygen of Thr residues from oxygen of NAD⁺ are: 2.855 Å (Thr219), 4.239 Å (Thr220) and 3.155 Å (Thr221). Other color representations are according to that in (A). (C) and (D) Stick diagram representation of Thr219–Thr221 residues with NAD⁺. Side-chain oxygen molecules of Thr residues were modified to attach a phosphate group with two negative charges in (D). Color representations are as follows: Thr219–Thr221 (red), NAD⁺ (green), Phosphorus (yellow) and Oxygen (white).

NAD⁺ (Fig. 7C). We next used UCSF Chimera to simulate the presence of pThr at the positions 219, 220 and 221. Phosphorylation imparts two negative charges to the side-chain of each Thr by adding a phosphate group containing three oxygen atoms (Fig. 7D). This charge possibly disrupts the electrostatic interaction of the protein with NAD⁺. Although NAD⁺ may still bind to SahH, the efficiency with which it binds may greatly diminish. Also, to accommodate the phosphate group, additional structural perturbation may occur leading to interference with NAD⁺ binding. This analysis explains the possible reasons for the decrease in affinity of SahH-P for NAD⁺ and loss in activity of SahH-T219A/T220A/T221A mutant.

Validation of SahH phosphorylation sites in *M. smegmatis*. To further validate the role of Thr219–Thr221 residues in SahH phosphorylation, we mutated these residues simultaneously in pYUBDuet-SahH and over-expressed SahH-T219A/T220A/T221A in *M. smegmatis* MC² 4517. Over-expression was confirmed by immunoblotting using anti-SahH antibodies (Supplementary Fig. S4). We used two-dimensional gel electrophoresis to resolve the isoforms of SahH after the mutation of three contiguous Thr residues. After this, the different isoforms of SahH were analyzed by immunoblotting using anti-SahH antibodies. We identified four isoforms of SahH-WT, separated according to their isoelectric points (Fig. 8A). Interestingly, one isoform was lost when SahH-T219A/T220A/T221A mutant was used indicating loss of phosphorylation. This shows that the Thr219–Thr221 residues are important phosphorylation sites of *Mtb*-SahH in *M. smegmatis*.

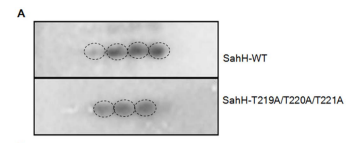
Importance of Thr219-Thr221 in regulating homocysteine levels.

We next used SahH-T219A/T220A/T221A as a phospho-deficient as well as enzymatically inefficient derivative to study the role of SahH in regulating homocysteine concentration. Homocysteine amounts were measured in *M. smegmatis* MC² 4517 with over-expression of SahH-T219A/T220A/T221A. No measurable alteration in homocysteine levels were observed in cells over-expressing SahH-T219A/T220A/T221A mutant as compared to vector-alone (Table 1 and Fig. 8B), while SahH-WT increased homocysteine concentration (Table 1 and Fig. 2C). These results validate the role of Thr219–Thr221 residues in SahH-mediated regulation of homocysteine concentration *in vivo*.

Discussion

Homocysteine is a key intermediate in the synthesis of metabolites such as methionine, cysteine, SAM, SAH and adenosine (Fig. 1A). Sulfur-containing metabolites- methionine and cysteine are required for the protein synthesis while SAM is essential for the one-carbon metabolism. In *M. tuberculosis*, transsulfuration pathway helps in conversion of methionine to cysteine where homocysteine is an intermediate²⁹. Adenosine is metabolized via the purine salvage pathway and proteins involved in this pathway are considered to be drug targets in *M. tuberculosis*³⁰. Thus, homocysteine stands at an important metabolic junction where it helps in regulating the levels of crucial metabolites. Although the genes required in these metabolic reactions have been identified in *M. tuberculosis*, there is no information on regulation of the homocysteine metabolism. Here, we present





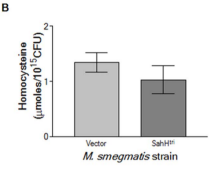


Figure 8 | Validation of Thr219–Thr221 as phosphorylation sites and role in homocysteine regulation. (A) Validation of Thr219–Thr221 as phosphorylation sites in *M. smegmatis*. SahH-WT and SahH-T219A/ T220A/T221A were purified from *M. smegmatis*, separated by two-dimensional PAGE and immunoblotted with anti-SahH antibodies. For clarity only the region containing SahH isoforms has been shown and different isoforms have been encircled. As seen, one isoform is lost in SahH-T219A/T220A/T221A. (B) Role of Thr219–Thr221 residues in regulating homocysteine concentration. Histogram shows intracellular homocysteine concentration in *M. smegmatis* MC² 4517 with and without the over-expression of SahH-T219A/T220A/T221A (marked as SahH^{tri}). Homocysteine concentration was analyzed using UPLC and plotted as μmoles/10¹⁵ CFU. Homocysteine levels were found to be unaffected by SahH-T219A/T220A/T221A (see also Table 1). Error bars represent SD of three independent results.

the evidence for the role of SahH in homocysteine metabolism, its reversible enzyme mechanism and role of phosphorylation in its regulation.

Our first objective was to characterize reversible enzyme activity of *Mtb*-SahH. NAD⁺ was found to be a requisite for both SAH-hydrolytic and SAH-synthetic activities. NAD⁺-bound SahH was active and did not require any other cofactor or metal ion. Further, loss in SAH-hydrolytic and decrease in SAH-synthetic activity by H363A mutation strengthens the earlier structural observation, where His363 needs to flip out in order to accommodate ethylthioadenosine (ETA, a partial analog of SAH)¹². His363 is thus a critical residue for enzyme activity.

We further analyzed functional conservation of SahH by measuring its effect on homocysteine concentration. In earlier studies, attempts to measure homocysteine-thiolactone in absence of exogenous homocysteine supplements could detect only very low levels in M. smegmatis³¹. Even using mass spectrometry based techniques, the sensitivity of detecting endogenous homocysteine was very low and it is only the upregulation that was reported in the presence of drug8. In our study, we used Ultra-high performance liquid chromatography (UPLC) to separate the thiol-containing molecules in combination with a sensitive method of fluorescence-based detection. We were able to detect homocysteine in concentrated cell-free extracts of M. smegmatis. We found that concentration of homocysteine in M. smegmatis was indeed very low (Table 1) and overexpression of SahH increases homocysteine levels to about six times. This increase was in accordance with our in vitro results where SAH hydrolysis augmented when SahH amounts were increased.

Over-expression of SahH in *M. smegmatis* led to increased SAH-hydrolysis resulting in elevated homocysteine levels. This also indicates that the probable favorable direction of catalysis *in vivo* is SAH-hydrolysis; however, further investigations are required to confirm this hypothesis. SahH may regulate concentration of metabolites like homocysteine, adenosine and SAH in the cell and any change in its activity may perturb their concentration in the cell. As discussed in later sections, our study suggests that SahH activity can be regulated by phosphorylation of critical residues involved in NAD⁺ binding.

We investigated the phosphorylation-mediated regulation of SahH using M. tuberculosis PknB as a representative of the M. tuberculosis STPKs. We observed that Mtb-SahH is a substrate of PknB among other kinases. Since reversible regulation is a defining characteristic of STPK-mediated phosphorylation, we checked dephosphorylation of SahH using PstP_c. SahH was found to be dephosphorylated by PstP_c in a time-dependent manner. Phosphorylation of SahH was also evaluated in the E. coli dual expression system co-expressing SahH and PknB. A convincing way to look at in vivo phosphorylation is metabolic labeling using 32P-orthophosphoric acid. In this experiment SahH-P (phosphorylated by PknB) was phosphorylated while SahH-UP (co-expressed with PknB-K40M) was unphosphorylated. Phosphoamino acid analysis helped us to identify the phospho-residues in SahH. In this analysis, Thr residues were found to be phosphorylated. SahH was then purified (SahH-P and SahH-UP) from E. coli where SahH-P was found to be phosphorylated at Thr residues while SahH-UP was unphosphorylated. These experiments confirmed that SahH is modified by reversible Ser/Thr phosphorylation mediated by PknB and PstP.

Mass spectrometry analysis revealed that PknB phosphorylates SahH on multiple Ser/Thr residues. Interestingly, three Thr residues (Thr219, Thr220 and Thr221) were also identified by Prisic et al. in M. tuberculosis phosphoproteome¹⁶. This indicates that PknB is likely to be the kinase responsible for SahH phosphorylation in M. tuberculosis. Also, four different isoforms of Mtb-SahH were identified when it is purified from M. smegmatis. This suggests that M. smegmatis STPKs phosphorylate Mtb-SahH on multiple residues reaffirming our mass spectrometry results. Multiple phosphorylation events at a single protein are a common phenomenon in M. tuberculosis signaling. In M. tuberculosis phosphoproteome 301 phosphoproteins were identified with 516 sites, implying that there are several proteins that are phosphorylated on multiple sites¹⁶. The multiple sites of phosphorylation individually contributed almost equally to total phosphorylation at SahH as loss in any single residue does not completely abrogate the phosphorylation intensity. But, when multiple Thr residues were mutated to alanine, a triple mutant SahH-T219A/T220A/T221A showed maximum loss in phosphorylation. Importance of these three Thr residues in phosphorylation was also seen in vivo by metabolic labeling of E. coli expressing SahH-T219A/ T220A/T221A and PknB. Here again, SahH-T219A/T220A/T221A mutant showed significant loss in phosphorylation.

SahH was found to be a conserved protein with high sequence identity among mycobacterial species and the conservation of Thr219–Thr221 residues is noticeable. In light of this observation, we reveal that phosphorylation of SahH is also a conserved phenomenon in *Mycobacterium* species. Native SahH from *M. bovis* BCG was found to be phosphorylated. Also, recombinant *Mtb*-SahH was found to be phosphorylated in *M. smegmatis* suggesting a common phosphorylation-mediated regulatory mechanism of SahH. Since Thr219–Thr221 residues were proposed to be involved in NAD⁺ binding¹², we suspected that multiple phosphorylated residues on SahH may affect its enzymatic activity. Therefore, we measured kinetic parameters of SahH-P and SahH-UP in SAH-hydrolytic direction. Phosphorylation accompanied increase in K_m indicating decrease in affinity towards NAD⁺, the essential cofactor of SahH. Since NAD⁺ is required for activity in both the directions, phosphorylated SahH was found to be



only 50% efficient in SAH-synthesis as compared to unphosphorylated SahH. Loss in activity of SahH-T219A/T220A/T221A mutant reinforced the role of Thr219-Thr221 residues in NAD+ binding. NAD+ binds in a pocket constituted by domain II, which is a Rossmann fold, but also makes electrostatic interactions with residues in domain I12. We analyzed this interaction and observed that the distance between side chains of Thr219-Thr221 residues and ribose oxygen of NAD⁺ is ≤ 5 Å. Phosphorylation of Thr219-Thr221 modifies its interaction with NAD+ and thus modulates activity of SahH. We also simulated the replacement of Thr219-Thr221 residues with pThr by modifying the atoms in already available structure of SahH. It was clear that there would be hindrance in the binding of NAD⁺ once the residues at these positions are phosphorylated. Interestingly, Thr219-Thr221 residues are also important for SahHmediated regulation of homocysteine concentration as SahH-T219A/ T220A/T221A mutant does not upregulate homocysteine concentration in M. smegmatis.

In *E. coli*, acetate has been proposed to inhibit re-methylation of homocysteine to methionine leading to elevated intracellular homocysteine concentration which is toxic to the cell^{33,34}. Although such inhibitory effect of homocysteine needs to be elucidated in mycobacteria, our results indicate that exogenous homocysteine retards the growth of *M. smegmatis* (data not shown). Phosphorylation of SahH may be used in certain conditions to lower its activity and consequently prevent upsurge of homocysteine concentration. Interestingly, SahH phosphorylation was found to be induced in the presence of acetate¹⁶. However, further investigations are required to assess the role of SahH and its phosphorylation in homocysteine-mediated toxicity.

Loss of NAD+ binding either by mutation of critical amino acid residues (Thr219-Thr221) or by phosphorylation leads to decreased SahH activity and the latter mechanism may be used by mycobacteria to fine tune the homocysteine concentration in the cell. Recently, another report had shown that phosphorylation of M. tuberculosis SahH by PknA affects V_{max} of SAH-hydrolytic activity³². Thr2 and Thr221 were identified as phosphorylation sites but the mechanism by which phosphorylation at these sites decreases V_{max} remains obscure. Here we not only reveal the effect of phosphorylation on SahH activity but also provide the evidence that phosphorylation affects NAD+ binding and thus has an overall negative effect on catalysis. We find a novel regulator of homocysteine concentration in mycobacteria and suggest a mechanism that governs its modulation. SahH, a highly conserved protein, is regulated by phosphorylation and the phenomenon is conserved across Mycobacterium species. The regulation of SahH by phosphorylation is an important phenomenon as it governs concentration of key metabolites.

Methods

Bacterial strains and gene manipulations. *E. coli* cells were grown and maintained in LB medium as described³⁵. *M. smegmatis* MC² 155, *M. smegmatis* MC² 4517³⁶ (kindly provided by Dr. Rajesh S. Gokhale) and *M. bovis* BCG were grown in Middlebrook

7H9 broth supplemented with 0.5% glycerol, 10% ADC, 0.05% Tween-80. The gene coding for SahH (*rv3248c* and *msmeg_1843*) was PCR amplified from *M. tuberculosis* H37Rv and *M. smegmatis* MC² 155 genomic DNA, respectively and cloned in PProEx-HTc, pETDuet-1 (MCS-1), pSD5³⁷ and pYUBDuet³⁸ vectors. Gene coding for PknB or PknB-K40M (full-length, 1–626 aa) was inserted in MCS-2 of pETDuet-1^{27,39}. Cloning of PknB_c, PknB_c-K40M and PstP_c in pProEx-HTc was performed as explained earlier⁴⁰. pETDuet-SahH or pYUBDuet-SahH was used as template to generate mutations using QuikChange XL-site directed mutagenesis kit, according to manufacturer's instructions (Stratagene). Tables 3 and 4 provide the detailed information about plasmid constructs and primers used throughout this study.

Protein expression and purification. For expression of His₆-tagged proteins from pProEx-HTc or pETDuet-1, BL21 (DE3) cells were transformed with recombinant plasmids. Proteins were expressed and purified as described earlier⁴⁰. The purified proteins were analyzed by SDS-PAGE and concentrations were estimated by Bradford assay (Bio-Rad).

Enzymatic assay of SahH. *E. coli*-purified His₆-SahH and its mutants were subjected to ammonium sulfate precipitation to remove bound nucleotides, using the protocol of Gomi *et al.*⁴¹. Detailed protocol is described in Supplementary methods. The 'apo' forms of protein thus obtained were free of NAD⁺ cofactor. To reconstitute the active 'holo' form, enzymes were incubated with 1 mM β -NAD⁺ for 30 minutes at 4°C in buffer A (50 mM potassium phosphate buffer [pH 7.0] and 1 mM EDTA).

Enzyme activity was measured in hydrolytic direction using a colorimetric assay with 0–100 $\,\mu M$ SAH. Reaction buffer contained 100 $\,\mu M$ Ellman's reagent (or DTNB, Sigma) and 1 unit of adenosine deaminase (Sigma) in buffer A. Reaction was started by adding 100 nM of holo-form of SahH, unless otherwise indicated. For the determination of K_m and V_{max} 1.74 $\,\mu M$ of apo-forms of SahH-P and SahH-UP were used with 0–10 $\,\mu M$ $\,\beta$ –NAD $^+$. The reaction was allowed for 25 min at 37°C and OD was measured at 415 nm (microplate reader 680 XR, Bio-Rad). Absorbance units were converted to activity units (pmoles/mg/min). K_m and V_{max} values were obtained by non-linear regression analysis using GraphPad Prism software. Values from three independent experiments were presented as mean \pm standard error.

Enzymatic activity in synthetic direction was measured as described 42 . Briefly, 1–4 μM of holo-SahH was incubated with 25 μCi [8– $^{14}C]$ adenosine (American Radiolabeled Chemicals, St. Louis, MO; specific activity 53.5 mCi/mmol) and 5 mM DL-homocysteine (Sigma) in 20 mM potassium phosphate buffer [pH 7.0] (with 1 mM EDTA and 1 mM DTT). Reaction samples were incubated at 37 $^{\circ}C$ for one and two hours followed by boiling at 100 $^{\circ}C$ for 5 minutes. 10 μ l sample from the total reaction volume of 50 μ l was spotted on cellulose thin-layer chromatography (TLC) plates (Merck) and developed with butanol:methanol:water:NH $_4$ OH (60:20:20:1, v/v). Plates were dried and analyzed by autoradiography using Personal Molecular Imager (PMI, Bio-Rad). Densitometric analysis was done using Quantity One analysis software (Bio-Rad). Percent SAH synthesis was calculated as fraction of substrate adenosine converted to SAH.

Homocysteine measurement. M. smegmatis MC2 4517 cells were grown to OD600 of 0.6-0.8 and induced with 1 mM IPTG and 0.2% acetamide for 48 hours. Cells were harvested and washed three times with sterile water. Harvested cells were dissolved in sterile water and lysed by bead beating using 0.1 mm Zirconium beads. Cell debris was removed by centrifugation and cell-free lysate was processed for homocysteine measurement as described^{43,44}. Briefly, 100 µl of cell lysate was treated with 35 µl of 1.43 M Sodium borohydride dissolved in 0.1 N NaOH (to reduce the disulfide bonds). Then, 10 µl of n-amyl alcohol was added followed by addition of 35 µl of 1 N HCl. To this, 50 µl of 7 mM bromobimane (B4380, Sigma) dissolved in 5 mM sodium EDTA [pH 7.0] was added (to conjugate the reduced thiols with fluorophore). The solution was incubated at 42°C for 12-15 minutes followed by incubation at room temperature for 30-45 minutes. 50 µl of 1.5 M perchloric acid was used to precipitate proteins. Supernatant was neutralized with 6 µl of 2 M tris base. 4 µl of derivatized sample was injected into a 100 × 4.6 mm, 1.8 micron Eclipse plus C18 column using Agilent-1260 Ultra-high performance liquid chromatography (UPLC). Column was equilibrated with 90% buffer 1 (5% methanol and 0.86% acetic acid) and 10% Buffer 2 (100% methanol). Thiols were eluted from the column with linear

Table 3 List of plasmid constructs used in this study			
Plasmid construct	Description	Reference or source	
pProEx-HTc	E. coli expression vector with N-terminal His ₆ -tag	Invitrogen	
pProEx-HTc-SahH	Expression of His ₆ -SahH in <i>E. coli</i>	This study	
pProEx-HTc-PknB _c	Expression of His ₆ -PknB _c (cytosolic domain) in <i>E. coli</i>	40	
pProEx-HTc-PstP _c	Expression of His ₆ -PstP _c (cytosolic domain) in <i>E. coli</i>	40	
pETDuet-1	E.coli dual expression vector containing His6-tag in MCS-1 and S-tag in MCS-2	Novagen	
pETDuet-SahH	Expression of His ₆ -SahH in MCS-1 in <i>E. coli</i>	This study	
pETDuet-PknB	Expression of MBP-PknB in MCS-2 in <i>E. coli</i>	27, 39 [*]	
pSD5	Mycobacterial expression vector with Kanamycin resistance	37	
pSD5-SahH	Expression of His ₆ -SahH in <i>M. smegmatis</i> MC ² 155	This study	
oYUBDuet	Inducible shuttle vector for expression of proteins in M. smegmatis MC ² 4517	38 ′	
pYUBDuet-SahH	Expression of SahH under IPTG-inducible promoter in M. smegmatis MC ² 4517	This study	



Table 4	list of primers	used in	this	study
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Name of the primer $^{\alpha}$	Primer sequence $(5' \rightarrow 3')^{\beta}$
SahH pETDuet F	TGAGCCCCAAAGGCGGCAGCCCCTATGACCGGAA (Notl)
SahH pETDuet R	GTCTGCTCGCGGG GCGGCCGC ACTCAGTAGCG (Notl)
SahH HTc F	GAGCCCCAAAGAAGGATGAGGATCCCTATGACCGG (BamHI)
SahH HTc R	CTACTGAGTGCGGCCGCCCG CGAGC AG (Notl)
SahH pSD5 F	GGATGAAAGCCCATATGACCGGAAATTTGG (Ndel)
SahH pSD5 R ^y	ATTTTGCGTCTGCTCACGCGTGGGAGGCACTCAATGGTGATGGTGGTGGTAGCGGTAGTGG (Mlul)
SahH pYUBDuet F	AAGAAGGAT <u>GGATCCCCT</u> ATGACCGGAAA (BamHI)
SahH pYUBDuet R	TGGGCGATTITGCGTAAGCTTGCGGGTGGGA (Hindlll)
MsmSahH HTc F	GAAAAAAGGGG <u>GAATTCCA</u> TGACCGAACTCAAGGCCG (EcoRI)
MsmSahH HTc R	GGGTTCCGGGGACTTAAGCTTGTCCGGGCTACCGGG (HindIII)
SahHT216A F	GCCGAGTCGGTCAAG <mark>GGCG</mark> TC <u>GCC</u> GAGGAGACCACCGGCG
SahHT216A R	CGCCGGTGGTCGCCTC <u>GGC</u> GACGCCCTTGACCGACTCGGC
SahHT219A F	CGGTCAAGGGCGTCACCG AGG AG <u>GCC</u> ACCACCGGCGTGCTGCGG
SahHT219A R	CCGCAGCACGCCGGTGGT <u>GGC</u> CT <u>CCTC</u> GGTGACGCCCTTGACCG
SahHT220A F	CAAGGGCGTCACCGAGGAGACCGCCACCGGCGTGCTGCGGCTCTAC
SahHT220A R	GTAGAGCCGCAGCACGCCGGT <u>GGC</u> GGTCTCCTCGGTGACGCCCTTG
SahHT221A F	GGCGTCACCGAGGAGACCACC <u>GCC</u> GGCGTGCTGCGGCTCTACC
SahHT221A R	GGTAGAGCCGCAGCACGCC <u>GGC</u> GTGGTCTCCTCGGTGACGCC
SahHT219/220A F	CAAGGGCGTCACCGAGGAG <u>GCCGCC</u> ACCGGCGTGCTGCGGCTCTAC
SahHT219/220A R	GTAGAGCCGCAGCACGCCGGT <u>GGCGGC</u> CTCCTCGGTGACGCCCTTG
SahHT219/220/221A F	CAAGGGCGTCACCGAGGAG <u>GCCGCCGC</u> CGGCGTGCTGCGGCTCTAC
SahHT219/220/221A R	GTAGAGCCGCAGCACGCC <u>GGCGGCG</u> TCCTCGGTGACGCCCTTG
SahHT220/221A F	CAAGGGCGTCACCGAGGAGACC <u>GCCGC</u> GGCGTGCTGCGGCTCTAC
SahHT220/221A R	GTAGAGCCGCAGCACGCC <u>GGCGGC</u> GGTCTCCTCGGTGACGCCCTTG
SahHT216/219/220/221A F	TCGGTCAAGGGCGTCGCCGAGGAGGCCGCCGCCGCCGCGTGCTGCGGCTC
SahHT216/219/220/221A R	GAGCCGCAGCACGCCGGCGGCGCCTCCTCGGCGACGCCCTTGACCGA
SahHT257A F	AAATTCGACAACAAGTACGGCGCTCGGCACTCCCTGATCGACGGC
SahHT257A R	GCCGTCGATCAGGGAGTGCC G GGCGCCGTACTTGTTGTCGAATTT
SahHS376A F	ACATGGCCGGGCTGGAGCGC <u>GCG</u> GGGCGACACGGGTCAAC
SahHS376A R	GTTGACCCGTGTCGCCCCGGCGCGCTCCAGCCCGGCCATGT
SahHH363A F	GATCCTGGGAAATATCGGCGCCTTCGACAACGAGATCGAC
SahHH363A R	GTCGATCTCGTTGTCGAA <u>GGCG</u> CCGATATTTCCCAGGATC

x'F' denotes forward primer and 'R' denotes reverse primer.

This primer was used for addition of C-terminal His₆-tag to the protein.

gradient of both buffers (from 90% Buffer 1:10% Buffer 2 to 0% Buffer 1:100% buffer 2 in 7 minutes), with flow rate of 0.7 ml/min. Standard curve was generated with known concentrations of homocysteine (0.195-3.125 μM). Results were quantified by taking the area for the homocysteine-bromobimane peak and calculating its concentration using a regression equation derived from the standard curve. Homocysteine concentration was represented as μmoles/1015 CFU and results from three independent experiments were plotted as mean \pm standard deviation.

Generation of polyclonal antibodies for SahH in mouse and immunoblotting.

Affinity purified His6-SahH was used for antibody generation (see Supplementary methods). Standard protocols for western blotting were followed as described earlier40. The antibodies and dilutions used were- anti-pThr antibody (Monoclonal, Cell Signaling) 1:20,000 dilution, anti-pSer antibody (Abcam) 1:20,000 dilution, anti-SahH antibody 1:20,000 dilution and HRP-conjugated anti-mouse IgG antibody (Bangalore Genei) 1:20,000 dilution. The blots were developed using $Immobilon^{\text{TM}} \ western \ chemiluminescent \ HRP \ substrate \ kit \ (Millipore) \ according \ to$ manufacturer's instructions. During analysis of Thr mutants for loss in phosphorylation, densitometric analysis of bands was done using alphaimager 3400 (Alpha Innotech Corporation, San Leandro, California, USA) and normalized to their corresponding ponceau images. The relative loss in phosphorylation was calculated by considering the phosphorylation of SahH-WT as 100%.

Immunoprecipitation of SahH. To assess the phosphorylation status of native SahH, M. bovis BCG was grown till the $OD_{600} = 1$. Cells grown in 10 ml media were harvested and resuspended in 1X PBS containing protease inhibitor cocktail followed by lysis using bead-beating. Native SahH was immunoprecipitated from 2 mg whole cell lysate using anti-SahH antibodies following the protocol as described by the manufacturer (Pierce® Crosslink Immunoprecipitation Kit). The immunoprecipitated sample was run on SDS-PAGE, analyzed by immunoblotting using anti-pThr and anti-SahH antibodies.

Purification of SahH from M. smegmatis. For purification of His₆-SahH from M. smegmatis electroporated with pSD5-SahH, similar protocol was followed as described earlier²⁷. The purified protein was phosphoenriched, concentrated and utilized for immunoblotting. Identity of SahH was also confirmed by massspectrometric analysis (TCGA, New Delhi). For purification of pYUBDuet-SahH, plasmid constructs were electroporated in M. smegmatis MC² 4517 and transformants were selected on 7H10 agar plates containing Kan and Hyg. Purification procedures were the same as with pSD5-SahH except that the cultures were induced with 1 mM IPTG.

In vitro kinase assays and phosphatase assays. In vitro phosphorylation of 4–5 μg SahH by PknB_c (1-2 μg) was carried out as described⁴⁵ using 2 μCi [γ-³²P] ATP. For the visualization of phosphorylation signal on cleaved proteins, removal of recombinant tag was achieved by a previous protocol²⁷. In vitro dephosphorylation assays were carried out by adding PstP_c (1 µg) as described by Sajid et al. 46. In all the assays proteins were separated by 10-12% SDS-PAGE and analyzed by PMI.

Metabolic labeling in E. coli. E. coli (BL21-DE3) transformants harboring either pETDuet-SahH:PknB, pETDuet-SahH:PknB-K40M or pETDuet-SahH-T219A/ T220A/T221A:PknB were used for metabolic labeling as described by Sajid et al46. Extracted samples were analyzed by autoradiography using PMI. Densitometric analysis of autoradiograph-signals was done using alphaimager 3400 and normalized to amount of protein as seen in their respective coomassie-stained images.

Phosphoamino acid analysis. For phosphoamino acid analysis, metabolically labeled SahH-P was used and similar protocol was followed as previously described 45,47. Detailed protocol is described in Supplementary methods.

Identification of phosphorylation sites at SahH-P. For identification of phosphorylation sites, SahH-P and SahH-UP (3 µg) was run on 12% SDS-PAGE, stained with coomassie, de-stained and band corresponding to SahH-P was excised from the gel and washed with MilliQ water. The samples were processed for identification of phosphorylation sites by using Thermo-Finnagen LTQ electrospray instrument (Proteomics Core Facility, Children's Hospital, Boston, USA). Detailed sample processing protocol has been described earlier46.

Sequence and structural analysis of proteins. Amino acid sequence analysis of SahH was done using clustalW248, using ten different species of Mycobacterium. Available structure of Mtb-SahH was downloaded from PDB website (PDB ID: 2ZIZ) and

BRestriction sites/mutations are underlined and Restriction enzymes are mentioned in parenthesis



analyzed in UCSF chimera ^{12,49}. Atoms were labeled using 2D labels in chimera tools and Phosphate group was added by modifying atoms using chimera tools.

Analysis of proteins by two-dimensional PAGE. Recombinant proteins purified from M. smegmatis MC² 4517 were desalted using chloroform/methanol precipitation and 1 μ g of both the proteins were resolved by two-dimensional PAGE as described previously²⁷. Isoelectric focusing was done using 7 cm IPG strips (Bio-Rad) with 4–7 pH range, followed by second dimensional SDS-PAGE. After this, proteins were transferred to nitrocellulose membrane and immunoblotted using anti-SahH antibodies.

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Author contributions

A. Singhal and G. A. conceived and designed the experiments. A. Singhal, G. A., A. Sajid, A. M., A. B., R. V. and S. U. performed the experiments. A. Singhal, G. A. and A. Sajid wrote the paper and prepared the figures. S. S., V. K. N. and Y. S. contributed reagents/materials/analysis tools. All authors read and approved the manuscript.

Additional information

Supplementary information accompanies this paper at http://www.nature.com/scientificreports

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