Research Article

SufB intein splicing in *Mycobacterium tuberculosis* is influenced by two remote conserved N-extein Histidines

**Running Title:** Conserved histidines regulate *Mtu* SufB cleavage

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**Abbreviations:**

*Mtu: Mycobacterium tuberculosis*

WT: Wild type

SI: Splicing inactive

DM: *Mtu* SufB Double mutant
IPTG: Isopropyl β- d-1-thiogalactopyranoside
TCEP: Tris (2-carboxyethyl) phosphine
HA: Hydroxylamine
DTT: Dithiothreitol
HRP: Horseradish peroxidase
ECL: Enhanced chemiluminescence
TBST: Tris-buffered saline, 0.1% tween 20
BSA: Bovine serum albumin
β-gal: Beta galactosidase
ABC: Ammonium bicarbonate
ACN: Acetonitrile
TFA: Tri-fluoro acetic acid
HPLC: High pressure (or performance) liquid chromatography
FPLC: First protein liquid chromatography
MALDI-TOF/TOF: Matrix assisted laser desorption ionization time-of-flight
SDS: Sodium dodecyl sulphate
PAGE: Polyacrylamide gel electrophoresis
RMSD: Root mean square deviation
SASA: Solvent accessible surface area
RMSF: Root mean square fluctuation
PME: Particle-mesh Ewald
HPCC: High performance computing cluster
MD Simulations: Molecular dynamic simulations
Abstract: Inteins are auto-processing domains that implement a multi-step biochemical reaction termed protein splicing, marked by cleavage and formation of peptide bonds. They excise from a precursor protein, generating a functional protein via covalent bonding of flanking exteins. We report the kinetic study of splicing and cleavage reaction in [Fe-S] cluster assembly protein SufB from *Mycobacterium tuberculosis*. Although it follows a canonical intein splicing pathway, distinct features are added by extein residues present in the active site. Sequence analysis identified two conserved histidines in the N-extein region; His-5 and His-38. Kinetic analyses of His-5Ala and His-38Ala SufB mutants exhibited significant reductions in splicing and cleavage rates relative to the SufB wild-type precursor protein. Structural analysis and molecular dynamics simulations suggested that *Mtu* SufB displays a unique mechanism where two remote histidines work concurrently to facilitate N-terminal cleavage reaction. His-38 is stabilized by the solvent-exposed His-5, and can impact N-S acyl shift by direct interaction with the catalytic Cys1. Development of inteins as biotechnological tools or as pathogen specific novel antimicrobial targets requires a more complete understanding of such unexpected roles of conserved extein residues in protein splicing.

Keywords: Intein splicing; Splicing regulation; conserved histidines; *Mycobacterium tuberculosis* SufB; SufB intein.

1. Introduction

Protein splicing is a self-catalyzed event that generates a continuous extein protein by ligating two intein separated extein regions with a peptide bond. This post-translational auto-excision of the intervening intein protein is critical for the formation of an active protein (1-4). Splicing and cleavage products (Figure 1C) are obtained through a series of nucleophilic displacement reactions mediated in a coordinated fashion by the catalytic residues (5-9). Control over this
reversible interruption of the functional form of the host exteins can play a regulatory role in protein activation (10, 11). The extein sequences upstream and downstream of N-terminal and C-terminal intein ends are termed ‘N-extein’ and ‘C-extein’ respectively. The intein folds into a horseshoe-shaped structure with a catalytic cleft that brings the conserved catalytic residues and the extein splice junction close enough for initiation of splicing reaction (12).

Inteins themselves have N-terminal and C-terminal regions with conserved sequence segments (termed as Blocks or motifs) that facilitates the splicing and cleavage reaction(s). N-terminal intein region comprises of A, N2, B, and N4 structural motifs or Blocks while F- and G-Blocks are part of the C-terminal intein region. Typically, A-Block contains Cys/Ser or Thr; B-Block includes His and Thr residues; F-Block usually has Asp and His and the G-Block bears two conserved residues; a penultimate His and a terminal Asn. Classical or Canonical (Class 1) intein splicing involves 4 sequential acyl rearrangements (Figure 1C), a nucleophilic attack by C1 or S1/T1 leads to an N-S/N-O acyl shift converting the peptide bond of N-terminal splice junction to a thioester linkage, a second nucleophilic attack by C+1 forms a branched intermediate at C-terminal splice junction through esterification, the branched intermediate is resolved by terminal Asn cyclization through cleavage of C-terminal splice junction, and finally an (S-N/O-N) acyl shift fully ligates the two extein segments by an amide bond formation.

These bond rearrangements at splice junctions during the cleavage and splicing reaction(s) are assisted by non-catalytic intein residues through stabilization of various intermediate structure(s)(13). The Block A residues Cys, Ser, or Thr participate in the first step of splicing with significant assistance from the Block B His and Thr residues. The highly conserved Block B His destabilizes the scissile peptide bond either by reducing the energy barrier or by loss of resonance or protonation of the Cys1 amide bond via His imidazole ring to catalyze the N-S acyl
shift (14-18). The Block F Asp residue drives the thioesterification through the tetrahedral intermediate by ground-state destabilization (8, 16, 19). It is also proposed that Block B histidine plays a dual catalytic role; being weakly basic it deprotonates the Cys1 to accelerate the N-S acyl shift and subsequently acts as an acid to stabilize the tetrahedral intermediate (15, 17, 18). The Block F Asp residue also deprotonates the C+1 residue to stabilize the net positive charge on Cys1 that drives the transesterification reaction. The F- and G-Block His residues are critical in the coordination of terminal Asn cyclization. The F-Block His increases nucleophilicity of Asn, and the G-Block His accelerates the Asn cyclization by increasing the electrophilicity of backbone peptide (20-23). The final acyl-shift is energetically favorable and does not require assistance from either intein or extein residues(24). Inteins do show polymorphisms in the catalytic residues leading to variation in the splicing mechanism as seen in Class 2 and Class 3 intein splicing (11, 25-29).

Interactions between extein residues and the catalytic intein core in the regulation of splicing reaction have been studied by modulating intein activity, changing the conformation of catalytic cleft, and restraining the activity of catalytic residues (6, 30-34). Previous studies on intein-extein partnership in intein splicing have suggested mediation by extein residues both near and remote to the N- and C-terminal splice junctions (7, 30, 31, 33, 35-37). The N-extein residue at the first position (-1) is important for the first thioester reaction and shows enhanced N-terminal cleavage rate (>4-fold) or attenuated cleavage by 1,000-fold by replacing the native residue to aspartate and proline respectively (30). Substitution of bulky amino acids at this position can cause local distortion and induce N-cleavage reaction (17). The participation of the first C-extein Cys+1 in the second and third steps of the splicing reaction has been demonstrated by its mutation dramatically augmenting or inhibiting splicing and generating off-pathway N-cleavage products.
Earlier, extein effects were assumed to be limited to residues proximal to the intein (12, 35, 38-40). Earlier, extein effects were assumed to be limited to residues proximal to the intein (12, 35, 38-40), but recent work has shown that distal exteins are implicated as environmental sensors with the role in regulating splicing depending on solution environment and temperature in *Pho* RadA precursors (6, 10, 34).

SufB is a critical component of [Fe-S] cluster assembly and repair machinery called SUF (mobilization of sulfur) complex. This is a stress response system that gets upregulated during periods of oxidative stress and Fe starvation (42, 43). Though there are multiple pathways for [Fe-S] cluster biogenesis among the three kingdoms of life such as NIF (nitrogen fixation) and ISC (Iron-sulfur cluster), the SUF complex is unique in mycobacteria. Fe-S cluster containing proteins execute a broad spectrum of cellular functions in organisms such as respiration, gene regulation, RNA modification, DNA repair, and replication (44-46). Although the SUF system has been well characterized in the *E. coli* system, (39, 42, 43, 47-49), the importance of SufB intein splicing in the formation of functional Suf complex has been shown in mycobacteria (49).

The present study reports on the splicing and cleavage reactions of a full-length *Mtu* SufB (FL-SufB) precursor protein. We delineated the different structural domains of *Mtu* SufB, analyzed whether it follows a canonical or non-canonical intein splicing pathway, identified intein and extein residues that participate in catalytic cleft formation, assessed both their conservation in different mycobacterial species and their role in regulating cleavage and/or splicing reactions, and analyzed distinctions from other intein precursor proteins. We found intein residues highly conserved in different mycobacterial species that favor a canonical splicing mechanism (Figure 1A and 1B). We detected two distal histidines in the N-extein region, His-5 and His-38, that are conserved in mycobacteria, archaea, and other microbes where SUF is the exclusive system for Fe-S cluster biogenesis (Figure 2A and 2B). Biochemical analyses of H5A and H38A SufB
mutants confirmed their influence on splicing and cleavage. Structural modeling of *Mtu* SufB and explicit-solvent molecular dynamics (MD) simulations of the model were used to analyze the SufB precursor splicing active site dynamics, and these simulations suggested that N-terminal cleavage could be supported by an interaction between H-38 and H-5. These observations suggest that the two distal H-5 and H-38 N-extein residues participate in SufB precursor stabilization and aid *Mtu* SufB intein splicing.

Kinetic analyses of H-5A SufB mutant demonstrated 3-fold and 1.4-fold reductions in splicing and cleavage rates respectively, relative to wild-type *Mtu* SufB precursor. Likewise, a 3.4-fold and 3-fold reduction in splicing and cleavage rates were observed in the H-38A SufB mutant. A side-by-side *Mtu* SufB structure prediction was done using homology (chimera) modeling, secondary structure prediction through consensus with other protein sequences. Subsequently, molecular dynamics (MD) simulations in aqueous media were carried out to find the equilibrated structure. Furthermore, MD simulations clarified the structural features of the SufB intein active site and indicated that N-terminal cleavage reaction is catalyzed by H-38 with the assistance of H-5. Taken together, our study substantiates a distinct mechanism for N-terminal cleavage reaction shown by *Mtu* SufB. Although H-38 is relatively distal to the N-terminal splice junction when supported by His-5, it can efficiently activate the first step of splicing reaction. Finally, we have proposed a novel mechanism for the N-cleavage reaction mediated via the concerted actions of these conserved histidines in the N-extein region of *Mtu* FL-SufB precursor. Interestingly, attempts to express the SufB double mutant (H-5A/H-38A) protein resulted in a truncated protein. These observations suggest that H-5 and H-38 might have important biological role(s) in the SufB precursor stabilization and perhaps the functionality of *Mtu* SufB protein.
2. MATERIALS AND METHODS

2.1. Genetic constructs

The full-length Fe-S cluster assembly protein SufB from the *Mycobacterium tuberculosis* H37Rv strain (*Mtu*-SufB-FL) and its isolated intein (*Mtu*-SufB-I) genes were PCR-amplified using Pfu Ultra High-Fidelity DNA Polymerase (Agilent Technologies) from heat-killed *Mtu* genomic DNA. DNA purification by gel electrophoresis was followed by EcoRI and HindIII restriction digestion and cohesive end ligation (T4 DNA Ligase, NEB Cat. No. M0202S) for cloning. The genes were inserted into the multiple cloning site 1 (MCS1) of the low copy expression vector pACYCDuetTM-1 (Novagen), which was driven by a T7 promoter/lac operator with a chloramphenicol resistance gene for selection. The constructs were screened via colony PCR and confirmed by sequencing (Sequencing Core Facility, SUNY, Albany, and Eaton Bioscience Inc. sequencing service) using the ACYCDuetUP1 (Novagen Cat. No.71178-3) and DuetDOWN1 primers (Novagen Cat. No. 71179-3), as well as the original primers. *Mtu*-SufB-FL mutants H5A, H38A, C1A, N359A were generated by substituting respective key catalytic residues to alanine via phosphorylated inverse PCR primers. Splicing inactive (SI) *Mtu*-SufB-FL double mutant (C1A/N359A) was created via inverse PCR to add the N359A mutation into the C1A cleavage mutant. Similarly, H5A/H38A *Mtu*-SufB-FL double mutant (DM) was created by adding H38A mutation into the H5A mutant by inverse PCR. The cloned H38A and H5A/H38A *Mtu* SufB mutant genes were also confirmed separately by sequencing (Agrogenomics, Odisha) using ACYCDuetUP1 (IDT Cat. No. 103948189) and DuetDOWN1 (IDT Cat. No. 103948190) along with the SufB Primers. All the above primers are listed in Table S1.
2.2. Sequence analysis

Protein sequences for *Mtu* SufB-FL (Accession number YP_006514844.1, GI: 397673309) and the 477 amino acid intein-less SufB protein from *Mycobacterium smegmatis* strain MC2 155 (Msm-SufB-FL, accession number YP_887437.1, GI: 118472504) were pairwise sequence aligned in ClustalW to distinguish extein and intein regions in *Mtu* SufB-FL (50). The identified *Mtu* SufB intein sequence (*Mtu* SufB-I) was confirmed by sequence comparison using Blast with the sequences deposited in Inbase (The Intein Database, www.inteins.com) (51). Different structural domains of the intein-like homing endonuclease, and the N- and C-terminal inteins were clearly demarcated by sequence alignment and structural analysis of SufB intein with homing endonuclease domain (I-CreI) and intein homing endonuclease Ii (PDB: 2CW7). The *Mtu* SufB-FL and *Msm*-SufB-FL sequences, combined with one archaeal and other bacterial SufB proteins collected from the NCBI and intein databases (51), were aligned using Dialign2 software (52). Conservation of different intein and extein residues was edited and color-coded manually. Phylogenetic tree analysis was performed using the Maximum likelihood method in the MEGA X program (53, 54) for both SufB inteins [Figure 2C (ii)] and SufB precursor sequences [Figure 2C (i)] from different organisms.

2.3. Protein overexpression and purification

Full-length (FL) un-spliced precursor and mutant SufB proteins carrying an N-terminal 6X-His-tag were over-expressed in BL21 (DE3) *E.coli* cells via IPTG (Sigma 367-93-1) induction. Cells were resuspended in lysis buffer (20mM sodium phosphate, 0.5M NaCl, pH 7.4) and lysed via tip sonicator (Sonics vibra cell VCX-130). Proteins were over-expressed and isolated from inclusion bodies (IB) via centrifugation. The IB materials were solubilized by 8M urea (Merck, 1084870500)
buffer (lysis buffer, 8M urea, 20 mM of imidazole (MP–biochemicals-288-32-4) and centrifuged at 16,500g for 20 min to collect the supernatant. Then 6X-His-tagged full-length (FL) precursor and mutant proteins were purified by Ni-NTA affinity column (Ni-NTA His trap, HP GE Healthcare Life Sciences-17524802). Prior to sample application, columns were equilibrated with binding buffer (20mM sodium phosphate, 0.5M NaCl, 40mM imidazole). After sample loading, columns were washed several times (15 CV) in binding buffer. Finally, proteins were eluted as purified fractions in elution buffer (20mM sodium phosphate, 0.5M NaCl, 500mM imidazole) followed by quantification via Bradford’s assay.

2.4. In-vitro splicing and cleavage assays

2.5 µM of purified proteins were allowed to refold in 1 ml of renaturation buffer (20 mM sodium phosphate, 0.5 M NaCl, 0.5 M Arginine, 1 mM EDTA, pH 7.4) in presence of 2mM TCEP-HCl (sigma-51805-45-9) at 20˚C for 24 h. The 0hr sample was retrieved before renaturation and splicing was quenched by addition of loading dye (0.1% bromophenol blue, 50% glycerol, β-mercaptoethanol, 10% SDS, tris 6.8) followed by rapid freezing at -200 C. Our controls, splicing inactive(SI) SufB double mutant and empty expression vector pACYC Duet-1, were treated similarly for the in-vitro assays. For the N-cleavage assay, proteins were refolded in presence of reducing agents and nucleophiles such as 2mM TCEP-HCl, 50 mM DTT (Roche-10708984001), and 0.5 M 250 Hydroxylamine (SRL-66164) with 1mM TCEP in renaturation buffer (55). For splicing and cleavage analysis, sample extraction at each time interval was followed by the addition of loading dye to stop the reactions and then boiling at 95˚C for 5 min. Resultant products from various refolding reactions were resolved through 4~10% gradient SDS PAGE. Protein bands were stained with Coomassie blue R-250 and densitometric analysis was performed by using GelQuant.Net biochemical solutions. Percentage(s) of splicing and cleavage products were
measured by taking the percentage(s) of the ratio of the total splicing product (LE and I) over total proteins (LE+I+P) and total N-cleavage product (NE+NC) over total proteins (NE+NC+P). The 0hr splicing value(s) were subtracted at each time point for baseline correction.

2.5. Kinetic analyses

Since Mtu 6X (His)-tagged WT SufB and SufB mutants (H-5A, H-38A, H-5A/H-38A, and 263 C1A/N359A) were purified and renatured at different temperatures, after normalizing splicing and cleavage values at different time intervals, the plot was generated by taking the percentage of splicing or cleavage product with respect to time (in min). Next, the curve was fitted in pseudo-first-order kinetics, with an equation $Y = Y_0 + (\text{Plateau} - Y_0) \times (1 - \exp(-K \times X))$ [Where $X$=time, $Y_0$ =Y value when time $(X)= $ time0, Plateau= max Y value at time t, $K$ = rate constant, expressed in reciprocal of the X-axis (time units)] in graph pad prism software. The fitted curve was generated by automatic outlier elimination fitting in a nonlinear regression equation. The rate constant ($K$) and Vmax were generated by the software. Half-life (t1/2) was calculated by graph pad prism using the formula $(\ln(2)/K)$.

2.6. Western blot

Western blot analysis was performed using an anti-His antibody (Invitrogen, LOT 1902132) to confirm the identity of splicing and cleavage products. Following resolution through SDS PAGE, test proteins were transferred to a nitrocellulose membrane; at 50v, 2h. After a successful transfer, blocking was done with 5% skim milk for 2h at room temperature. Then the blot was incubated with HRP conjugated anti-His antibody (Invitrogen, LOT 1902132) at 1:5000 dilutions for 16h at 40 C. Then blot was washed with 1X TBST and developed using 13 ECL as the substrate. N-extein detection was done with 1:2500 antibody dilution.
2.7 Mass spectrometry and chromatography

After renaturation, proteins were resolved through 4~10% SDS PAGE. Protein identification by mass spectrometry was performed at Central Proteomics Facility, Institute of Life Sciences, Bhubaneswar using the following standardized protocol. Well resolved protein gel bands were destained with 25mM ammonium bicarbonate (ABC). Cleared gel bands were reduced with 20 mM DTT and then alkylated by 100 mM iodoacetamide solution. The gel plugs were dehydrated sequentially with 50%, 100% acetonitrile (ACN) and then digested with 0.3 μg of trypsin (Absciex 4352157) in 25 mM ABC solution at 37°C for 16 hours. Peptides were extracted sequentially using extraction buffer containing 0.1%, 0.5%, and 5% tri-fluoro acetic acid in 50% ACN from gel bands. Extracted peptides were desalted by using C18 Zip-Tip (Millipore) and then dried in a centrifugal vacuum concentrator. Peptide samples were reconstituted with 0.1% TFA in 50% ACN solution mixed 1:1 ratio with α-cyano-4-hydroxycinnamic acid solution (C8982; 10 mg/ml) and then spotted onto metal target plate in triplicates. MS and MS/MS spectra were acquired using MALDI TOF/TOF (AbSciex TOF/TOF 5800). Acquired spectra were searched against *Mycobacterium tuberculosis* from NCBI nr database using mascot search engine in Protein Pilot Software for protein identification. Searches were performed allowing trypsin mis-cleavage up to Carbamido-methylation of cysteine, oxidation of methionine was also included as variable and fixed modification respectively. The peptide mass tolerance was 300 set as 100 ppm for precursor ion and 0.8 Da for fragment ion with +1 charge. Since all the splicing and cleavage products were identified as *Mtu* FL-SufB, the acquired spectra were researched against a customized database containing probable splicing products and cleavage products of SufB protein of *Mycobacterium tuberculosis* using mascot search engine in Protein Pilot Software with the same parameter. β-gal and BSA were run as internal calibration. The protein score, percent coverage, theoretical molecular weight, and Iso-electric pH value were obtained. The mass spectrometry proteomics
data have been deposited to the ProteomeXchange Consortium via the PRIDE [1] partner repository with the dataset identifier PXD015199.

Purified and renatured $Mtu$ FL-SufB protein was examined by analytical HPLC carried out on an Agilent 1200 series instrument equipped with a Zorbax gf-450 column (6µm, 9.4x250mm) with a flow rate of 1ml/min. All HPLC runs used the following solvent: 1:1 water and isopropanol (solvent 1) and 0.1% TFA (Trifluoroacetic acid) in water (solvent 2). The column was equilibrated using solvent 1 and solvent 2. A protein sample was inserted into the column and ran for 30mins. The retention time vs protein intensity measured at 280 nm was noted. For reference, we ran 6µl of Precision plus protein ladder (Biorad1610374) diluted to 30µl using Sodium phosphate buffer. The retention time vs molecular weight of the known protein standards was measured and plotted to make a standard curve. The unknown protein peaks from the test samples were compared with the standard curve to find out the relative molecular weights. The molecular weights of the expected fragments from the MALDI-TOF/TOF MS (Table S3) data were compared with the standard curve and molecular weights of the unknown peaks were determined.

Molecular weight of the SufB DM (H5A/H38A) was determined by FPLC. H215A/H248A SufB mutant protein (DM) was isolated and purified under denaturing conditions. The relative molecular weight of DM was confirmed by Gel filtration chromatography using Hi prep 16/60 Sephacryl S200 GPC column (HiPrep 16/60 GE Healthcare, Cat. no. 10255928). 8M Urea buffer was removed from the protein sample by dialysis in 20mM sodium phosphate and 0.15M sodium chloride buffer with pH 7.4. Then the sample was concentrated using Protein concentrator (Thermo Scientific 88515). For FPLC analysis, the column was equilibrated with 2 column volume (cv) of sodium phosphate buffer (pH 7.4) followed by loading of molecular weight standard Bovine Serum Albumin (BSA) at a concentration of 1mg/ml. Then 2mg of purified DM protein was loaded
onto the GPC column. The run was performed at a flow rate of 1ml/min while keeping the temperature at 20° C. The fractions eluted at an absorbance of 280nm were collected and analyzed by SDS-PAGE under reducing and non-reducing conditions.

2.8 Molecular Dynamics (MD) Simulations

2.8.1. Homology modeling of Mtu SufB precursor

A homology model of the Mtu FL-SufB precursor was built using the SufB chain of the E. coli SufB-SufC-SufD complex (PDB ID: 5AWF, chain A), the S. cerevisiae intein homing endonuclease (PDB ID: 1VDE, chain A), and the T. kodakarensis homing endonuclease (PDB ID: 2CW7, chain A) using Bioluminate 2.7 from Schrödinger. The model was assessed by Ramachandran plot analysis and PROSA(Z-score) (56).

2.8.2. MD simulation analysis

Explicit-solvent MD simulations were performed using GROMACS (V5.1.4) (57) with the OPLS- AA force field (58) and the SPC/E water model (59, 60). The Mtu SufB model (846 aa) was solvated in a cubic box with at least 1 nm distance between the protein and the edge of the box, and neutralized with NaCl. The system was minimized with target Fmax of no greater than 1000 kJ mol-1 349 nm-1 with steepest descent minimization with a spherical cut-off at 1 nm was imposed on all intermolecular interactions with verlet cut-off scheme (61, 62). The leap-frog algorithm with a timestep of 2fs (63) and a canonical NVT ensemble was used to run the simulation for 100 ns, with temperature maintained at 300K through velocity-rescale coupling and no temperature coupling. H-bonds were constrained using lincs with the order of 4 (64). The particle-mesh Ewald (PME) algorithm was used for implementing long-range electrostatic interactions with the grid dimension of 0.16nm and interpolation order of 4 (65). Histidine at positions 5 and
were mutated to alanine by homology modelling, and MD simulations were performed on these mutant proteins in an identical fashion. The MD simulation trajectories were analyzed with GROMACS and Pymol, and plotted using Origin 8.0.

3. RESULTS

3.1 Structural domains of *Mt u* full-length (FL) SufB precursor

We obtained the protein sequences for Fe-S cluster assembly protein SufB [*Mycobacterium tuberculosis* H37Rv], an 846 amino acid protein (Accession number WP_003407484.1, GI: 397673309) and the intein-less SufB protein [*Mycobacterium smegmatis* MC2 155], a 477 amino acid protein (Accession number YP_887437.1, GI: 118472504) from NCBI protein database. A pair-wise sequence alignment using the ClustalW server very clearly delineated the intein sequence as well as identified ~95% sequence similarities between the extein sequences. A blast search against the sequences deposited in Inbase also identified the intein sequence as 359 amino acid *Mt u* SufB (*Mt u PpsI*) intein From *M. tuberculosis* strains H37Rv. Based on sequence alignment and structural analyses of SufB intein and homing endonuclease domain I-CreI and intein homing endonuclease Ii (PDB: 2CW7), the final demarcation of different structural domains of *M. tuberculosis* and their respective sequences appeared to be as shown in Table S2.

Residues encompassing N- & C-terminal extein and intein sequences from *Mt u* SufB precursor are illustrated in Figure 1A. Multiple sequence alignment of *Mt u* SufB precursor and other intein bearing proteins highlighted conservation of catalytic cysteines (C1 and C+1), Block B His67, Block F Asp77, penultimate His358, and terminal Asn359 in the intein and extein regions (Figure 1B). Hence, SufB precursor protein (846 aa) is likely to follow a canonical intein splicing pathway to generate the following products (Figure 1C); ligated N-(252aa) and C-(235 aa) exteins
to form native SufB protein (487aa), SufB intein (359aa) with splicing domain (N- and C-terminal inteins; 155aa) and endonuclease domain (204aa). Off-pathway products may give rise to N-terminal cleavage products [NC(594aa) and N-extein (NE)] and C-terminal cleavage products [CC(611aa) and C-extein (CE)] (51, 66).
Figure 1. (A) Schematic representation of different structural domains of *Mtu* full-length SufB precursor. The conserved and the catalytic residues in the FL-SufB protein sequence are shown at the top. Further, numbering for different structural domains are also displayed; intein residues (1, 2…….n), N-extein residues (-1, -2…….-n) and C-extein residues (+1, +2…….+n). Catalytic cysteines responsible for sequential nucleophilic displacement reactions are C1 and C+1, terminal intein residue responsible for branched intermediate resolution is N359, conserved histidines in the N-extein region are H-5 (248, full-length protein) and H-38 (215, full-length protein). G+234 is the last residues of *Mtu* SufB. (B) Multiple sequence alignment of *Mtu* SufB with other intein bearing species shows conservation of critical residues (highlighted in different colors) that participate in a canonical splicing pathway. (C) Schematic diagram showing the mechanism of intein splicing. (i) There are four major reaction steps; first two steps of sequential nucleophilic attack by C1 and C+1 generate branched intermediate by trans-esterification. Next, cyclization of terminal Asn resolves the branched intermediate, ligates exteins and cleaves off SufB intein, (ii) and (iii) represent off-pathway products from protein splicing; N- and C-terminal cleavage products respectively.

3.2 Conservation of His-5 and His-38 in different bacterial species

Our sequence analysis results indicate that SufB from different mycobacterial and archael species exhibits a high degree of sequence similarities (Figure 2A and B). This is further supported by phylogenetic analysis to show that these mycobacterial proteins possibly have a common ancestral origin (67). Analysis cladogram of SufB intein sequences from different mycobacterial species shows significant similarities, except in *M. leprae, M. lepromatosis, M. triplex* and *M.*
Xenopi, possibly due to different intein insertion sites (39). Divergence is also noticed in Ferroplasma for intein [Figure 2C (ii)] and precursor sequence [Figure 2C (i)] perhaps due to differences in the kingdoms. The discrepancies in the phylogenetic tree analyses of FL-SufB precursor proteins suggest that there are variations in the extein sequences and intein insertion sites. This suggests intein evolution and independent intein transfer in different species and kingdoms.
Figure 2. Histidine conservation in the N-extein region of *Mtu* SufB protein (A) Multiple sequence alignment of SufB, focusing on N-extein~intein junction in mycobacterial and archaean species. Conserved histidine residues at -5 and -38 positions in the N-extein region are highlighted in yellow and green respectively. Catalytic C1 of inteins are marked in red. N-extein and intein residues are delineated by shading in grey and cyan blue. (B) Multiple sequence alignment of SufB, focusing on N-extein~intein junction in *Mtu* and other intein-less mycobacteria and in organisms where SUF is the exclusive system for Fe-S cluster assembly, (C) (i) & (ii) phylogenetic analyses of full-length (FL) SufB protein and SufB intein sequences in mycobacteria and *Ferroplasma acidarmanus* respectively.

Further, two conserved histidines were identified in the N-extein region of different mycobacterial and archaean SufB precursor proteins, His-5 and His-38 (Figure 2A). His-5 is also conserved in intein-less SufB proteins from *Staphylococcus aureus*, *Bacillus subtilis*, and certain *mycobacteria* that use SUF complex as the sole pathway for [Fe-S] cluster generation (Figure 2B) (68, 69). Although His-38 is conserved in both intein-bearing and intein-less mycobacterial SufB proteins, conservation is missing in intein-less bacterial systems like *Staphylococcus aureus* and *Bacillus subtilis* where SUF system is essential to synthesize [Fe-S] clusters (Figure 2B).

Although His-5 is proximal to the N-terminal splice junction in the protein sequence our MD simulations results (explained later in this manuscript) detected the proximity of H-38 to catalytic Cys1 in the predicted structural model for *Mtu* SufB. Identification of two highly conserved metal-chelating residues proximal to N-terminal extein~intein junction raises the possibility of
their regulatory roles on cleavage and/or splicing as well as in the functionality of [Fe-S] cluster assembly protein SufB. Despite their remote positions these two histidines possibly come closer to the active site and participate in protein splicing either via direct or indirect interaction with the catalytic residues near cleavage site(s). A comprehensive kinetic study of in-vitro splicing, and N-cleavage reactions are presented in the next section. Further mechanistic understanding of the roles of highly conserved histidines was explored via molecular dynamics (MD) simulations as shown later in this manuscript.

3.3 Kinetic study to evaluate the roles of conserved His-5 and His-38 on Mtu SufB intein splicing

Plasmid constructs for the current work were engineered as mentioned in the materials and methods section. Detection of two highly conserved His-5 and His-38 in the N-extein region raises the possibility of their regulatory effect on Mtu SufB intein cleavage and/or splicing either by interaction with catalytic residues directly or indirectly via other active site residues. To test this hypothesis, alanine substitution was made in place of His-5 and His-38 to generate single mutants (H-5A, H-38A) and double mutant (H-5A/H-38A, DM) (Materials and Methods). Alanine substitutions for active site residues have resulted in a complete blockage of cleavage and splicing reactions as shown by earlier studies (30). The negative control used for the above study was splicing inactive (SI) SufB double mutant. Substitution of both C1 and N359 to alanine (C1A/N359A) is expected to abolish intein splicing completely (70).

The above proteins were purified under denaturing conditions and refolded at 20°C as described in the materials and methods. FL-SufB precursor, H-5A and H-38A SufB mutants gave rise to the precursor (P, 95.98KDa), N-terminal cleavage (NC, 66 KDa) product, N-extein (NE,
29.9KDa), ligated exteins (LE, 55.7 KDa), and intein (I, 40.2 KDa) upon \textit{in vitro} refolding (Figure 3). As predicted none of these products were seen in the case of SI double mutant [Figure S2E (i)] and transformants expressing empty vector pACYC Duet-1 [Figure S2E(ii)]. C-terminal cleavage product (CC, 70.12 KDa) and C-extein (CE, 25.76 KDa) were not detected by SDS PAGE possibly due to protein degradation. Consequently, we were unable to analyze the effects of His-5Ala and His-38Ala mutation on the C-terminal cleavage reaction of \textit{Mtu} SufB. However, we did confirm the presence of P, LE, NE, and CC products via western blot (Figure 3B).
**Figure 3. Effect of H-5 and H-38 mutation on *Mtu* SufB splicing.** (A) Products from *in-vitro* refolding reactions were resolved through 4–10% gradient SDS PAGE as shown in, (i) FL-SufB precursor, (ii) H-5A and (iii) H-38A SufB mutant proteins exhibiting splicing over different time periods, (B) Western blot to confirm the identity of splicing and cleavage products for (i) FL-SufB precursor (ii) H-5A iii) H-38A SufB mutants and iv) controls; SI (splicing inactive double mutant SufB), H-I[6x(His)-tagged SufB intein], pACYC Duet-1 (cell lysates expressing empty expression vector). Anti-His antibodies detected presence of 6x(His)-tagged P, CC, LE and NE. NE is blotted separately with higher concentration of primary antibody. (C) Kinetic study of protein splicing in *Mtu* FL-SufB precursor, H-5A and H-38A SufB mutant proteins. Splicing products were quantified and plotted over different time periods and the curve was fitted to a pseudo first order reaction; $Y=Y_0 + (Plateau-Y_0) \times (1-e^{-K\times t})$. All the experiments were performed in triplicates and error bars represents (±1) SEM from 3 independent sets of experiments. A comparative analysis demonstrates statistically significant difference in splicing efficiency (p<0.0001) in FL-SufB precursor, H-5A and H-38A mutant SufB. P=Precursor, CC=C-cleavage, NC=N-cleavage, LE=Ligated Extein, I=Intein, NE=N-Extein, M=Protein marker.
3.3.1 His-5Ala and His-38Ala SufB mutants exhibit attenuated splicing reaction

Splicing efficiency was determined following in vitro renaturation of FL- and mutant SufB proteins (H-5A and H-38A) over a period for 24h. Precursor (P) proteins detected at time 0h were quite comparable for FL- and mutant proteins (Figure 3A). Ligated extein fractions were visualized at 0h as a possible soluble contaminant and accumulated product of protein purification. Densitometric analysis of protein products was performed via Gelquant.NET biochemical solutions. Splicing efficiency was calculated as a percentage of splicing 

\[ \frac{[I+LE/P+I+LE]}{100} \]

Splicing efficiency was plotted over time as pseudo-first-order reaction kinetics in GraphPad Prism version 9.0 or 8.0 (Figure 3C). At 20°C, WT FL-SufB shows a splicing efficiency (Y_max) of (53.7±1.31) % with a rate constant (K) value of (9.92±0.001 x 10^{-3} \text{ min}^{-1}) and a half-life (t_{1/2}) of 1.1h (Table 1). Under the same experimental conditions, H-5A SufB mutant splices out with maximum efficiency (Y_max) of (37.8±0.68%), a rate constant (K) value of (3.3±0.0001 x10^{-3} \text{ min}^{-1}) and a half-life (t_{1/2}) of 3.4 h. H-38A SufB mutant splices out with an efficiency (Y_max) of (43.35±1.7%), the rate constant (K) (2.9±0.0003x10^{-3}), and half-life (t_{1/2}) 3.8h. This clearly demonstrates a 3fold and 3.4-fold reduction (p<0.0001; one-way ANOVA) in splicing efficiency for H-5A and H-38A mutants, respectively, relative to FL-SufB precursor. These observations further strengthen our hypothesis that histidines at -5 and -38 positions may have regulatory role on Mtu SufB intein splicing.
Table 1. Comparative analysis of different kinetic parameters for splicing in FL-SufB precursor, H-5A and H-38A SufB mutant proteins, at 20°C temperature. These data were extracted from Figure 3C.

| Kinetic parameters | FL-SufB       | H-5A         | H-38A        |
|--------------------|---------------|--------------|--------------|
| \( Y_{max} \)     | 53.81±1.03    | 37.82±1.68   | 43.35±1.7    |
| Rate constant (K)  | 9.92±0.007x10^{-3} | 3.3±0.0003x10^{-3} | 2.9±0.0003x10^{-3} |
| (min^{-1})         |               |              |              |
| Half-life(\( t_{1/2} \)) | 1.1h         | 3.4h         | 3.8h         |

Further analysis was done by the kinetic study of H-5A/H-38A SufB DM under similar experimental conditions. Surprisingly, H-5A/H-38A DM expression gave rise to a faint precursor (P) band and a truncated product of 28KDa size detected by SDS PAGE analysis whereas western blot detected both the P and the truncated protein (Figure 4 and Figure S2). The 28KDa product was confirmed by FPLC as well (Figure S3). The possibility of a frameshift during cloning was ruled out by repeat sequencing. A comparative analysis of splicing and cleavage products for FL-SufB, N359A SufB (cleavage mutant with isolated NC products), C+1A SufB (cleavage mutant with isolated NC products), and H-5A/H-38A SufB DM provided further clarification (Figure 4). The 28KDa (H-5A/H-38A SufB DM) truncated protein was identified as a possible splicing intermediate or a product of protein degradation, but different from the N-extein (29.9KDa).
Figure 4. Identification of *Mtu* H-5A/H-38A SufB double mutant (DM): (A) SDS Gel showing splicing and N-cleavage reactions in *Mtu* FL-SufB precursor, C+1A, N359A, DM (H-5A/H-38A) and SI (C1A/N359A) SufB mutant (B) Confirmation of protein products for FL-SufB, C+1A, N359A, DM (H-5A/H-38A) and SI (C1A/N359A) SufB mutant proteins via immuno-blotting assay. Anti-His antibodies detected presence of 6X(His) tagged P and LE. NE and SufB DM truncated protein product are blotted separately with higher concentration of primary antibody. P=Precursor, NC=N-cleavage, LE=Ligated Extein, I=Intein, NE=N-Extein.

3.4 Kinetic study to assess the roles of His-5 and H-38 in the first amide and thioester equilibrium

Next, we examined TCEP [Tris(2-carboxyethyl) phosphine], DTT (Dithiothreitol), and HA (Hydroxylamine) induced N-cleavage reactions in FL- and mutant SufB proteins. DTT can act both as a nucleophile and reducing agent. Hydroxylamine acts as a nucleophile and like DTT targets the linear thioester intermediate in the first step of splicing (71). TCEP, a reducing agent was used as a control for DTT and HA-induced N-cleavage reactions (72).
Since the optimum temperature for FL-SufB splicing is found to be 20°C, N-cleavage reactions for the test proteins were also conducted at the same temperature. Purified proteins were renatured in presence of different reducing agents, DTT (50mM), HA (0.5M), and TCEP(2mM) for different durations of time (materials and methods). Products of N-cleavage reactions were visualized by 4%~10% gradient SDS PAGE (Figure 5, 6, and Figure S4) and densitometric analysis of the cleavage products was performed by Gelquant.NET biochemical solutions. The percentage of N-cleavage reaction product was calculated as [(NE+NC/P+NE+NC) x 100]

### 3.4.1 H-5A and H-38A SufB mutants display reduced TCEP mediated N-cleavage reaction

TCEP mediated reaction was analyzed for 4h in FL-SufB and mutant SufB proteins. The mutant proteins clearly exhibited diminished production of N-cleavage (NC) products relative to FL-SufB [Figure 5, Table 2(i)]. Reaction mediated by TCEP did not give a proper curve that could be fitted in linear or non-linear regression. In FL-SufB, % N-cleavage was accelerated until 40 min followed by a decline as splicing became distinct. H-5A and H-38A SufB mutants presented a sluggish course with the N-cleavage product peak at 60-80 min and then tapered off. At 40 min and 60 min, H-5A displayed 2-fold and 1.5-fold diminution in % N-cleavage respectively, relative to FL-SufB. Likewise, at 40 min and 60 min, H-38A mutant displayed a 3.8-fold and 3-fold reduction in % of N-cleavage respectively relative to FL-SufB (Figure 5).
Figure 5. Effect of H-5A and H-38A mutation on TCEP mediated N-cleavage reaction in *Mtu SufB*: (A) SDS Gel image of (i) FL-SufB precursor (ii) H-5A and (iii) H-38A SufB mutant proteins displaying N-cleavage reactions over different time periods; B) (i) Comparative analysis of TCEP mediated reaction in FL-SufB, H-5A and H-38A SufB mutant proteins, indicates statistically significant differences (P=0.0011, one way ANOVA) in % of N cleavage over different time periods. Reaction mediated by TCEP did not give a proper curve which could be fitted in linear or non-linear regression. All the experiments were performed in triplicates and error bars represent (±1) SEM. P=Precursor, NC=N-cleavage, LE=Ligated Extein, I=Intein, NE=N-Extein, CC=C-Cleavage.

3.4.2 H-5A and H-38A SufB mutants exhibit reduced Hydroxylamine (HA) induced N-cleavage reaction

HA-induced N-cleavage reaction was monitored over a 24h period and then kinetics analysis was performed. The extracted data for HA-induced reactions were plotted over different time frames.
and the curve was fitted to a pseudo-first-order reaction with the equation, \( Y = Y_0 + (\text{Plateau} - Y_0) \times (1 - \exp(-K \times x)) \), (Figure 6). H-5A and H-38A mutants exhibited about 1.4-fold and 1.5-fold reduction (\( p = 0.0025 \); one-way ANOVA) in % of N-cleavage, respectively, relative to FL-SufB although the \( K \) and \( t_{1/2} \) values were comparable [Figure 6C, Table 2 (ii)]. HA is an alpha nucleophile that intercepts between amide-ester equilibrium and induces N-cleavage. It enhances the nucleophilicity of residues that leads to N-cleavage (73, 74). The mutation effect on HA-induced N-cleavage reaction was similar to what is observed in presence of TCEP. However, no significant differences in N-cleavage efficiency were noticed between H-5A, H-38A mutants, and FL-SufB in presence of DTT (Figure S4 and Table S4). Similar to HA, DTT is also a nucleophile that enhances the nucleophilicity of residues and facilitates N-terminal cleavage reaction (72, 75). Besides, DTT interacts directly with the residues that may increase the flexibility of the active site structure (76). Possibly, this abolishes the effect of H-5A and H-38A mutation on N-terminal cleavage. Supported by our current results we hypothesize that histidines at -5 and -38 positions in the N-extein region may have a role in Cys1 activation directly or via structural alignment of other active site residues. This likely explains why catalytic cysteines Cys1 and Cys+1 are interacting differently in His mutants, hindering the N-cleavage reaction. Affirmation of the above hypothesis was done by MD simulations (upcoming section).
Figure 6. Effect of H-5A and H-38A mutation on HA mediated N-cleavage reaction in Mtu SufB: (A) SDS Gel showing HA induced N-cleavage reactions in (i) FL-SufB precursor (ii) H-5A and (iii) H-38A SufB mutant; (B) Confirmation of splicing and cleavage products for HA induced N-cleavage reaction in (i) FL-SufB precursor, (ii) H-5A and (iii) H-38A mutant proteins via western blot. Anti-His antibodies detected presence of 6X (His) tagged P, CC and LE. NE is blotted separately with higher concentration of primary antibody; (C) (ii) Kinetic analysis of HA induced N-cleavage in WT and H-5A mutant proteins suggests statistically significant differences (P= 0.0025; one way ANOVA) in % of N-cleavage over different time periods. These curves were fitted into a pseudo first order reaction with equation Y=Y₀ + (Plateau-Y₀) *[1-exp(-K*x)]. All the experiments were performed in triplicates and error bars represent (±1) SEM. P=Precursor, NC=N-cleavage, LE=Ligated Extein, I=Intein, NE=N-Extein, CC= C-Cleavage.
Table 2. (i) Comparative analysis of TCEP mediated N-terminal cleavage reaction at 20°C over different time up to 240 mins. The data were extracted from Figure 5B. (ii) Different kinetic parameters (Y\text{max}, rate constant, and half-life) for HA-induced N-terminal cleavage in FL-SufB precursor, H-5A, and H-38A SufB mutants. These data were extracted from Figure 6C.

### (i)

| Time (min) | FL-SufB (% of N-cleavage) | H-5A (% of N-cleavage) | H-38A (% of N-cleavage) |
|------------|---------------------------|------------------------|-------------------------|
| 20         | 28.2±5.5                  | 15.1±0.97              | 7.42±2                  |
| 40         | 36±4.3                    | 17.89±2.7              | 9.3±2.7                 |
| 60         | 30.63±5.1                 | 20.76±3.7              | 10.2±1.7                |
| 80         | 25.7±5.7                  | 14.9±2.5               | 12.42±1.9               |
| 120        | 19.8±1.6                  | 8.4±0.7                | 10.5±2.7                |
| 240        | 16.78±1.5                 | 6.2±0.89               | 7.07±1.9                |

### (ii)

| Kinetic parameters | FL-SufB  | H-5A     | H-38A    |
|--------------------|----------|----------|----------|
| Y\text{max}       | 50.67±1.6| 35.72±1.4| 32.45±0.94|
| Rate Constant(K)   | 3.7±0.01 x10^{-2} | 3.9±0.01x10^{-2} | 3.7±0.009x10^{-2} |
| Half-life(t_{1/2}) | 18.5 min | 17.3 min | 18.2 min |
3.5 Confirmation of protein products

The identity of splicing and cleavage products (P, CC, LE, and NE) for the N-terminal 6X (His) tagged test proteins were confirmed via western blot using anti-His antibodies (Figure 3B, 4B, 6B and S4B). SI (C1A/N359A) double mutant SufB was used as a control for the study and gave rise to unspliced precursor protein (P) [Figure 3B (iv)]. Interestingly, one additional protein band was noticed above LE and below CC for FL- SufB, H-5A, and H-38A mutants possibly due to the generation of a splicing intermediate as an off-pathway product. We also observed a product just below the P protein for FL-SufB, SI-SufB, H-5A, and H-38A mutants (Figure3B, 6B, and S4B). However, this protein was missing in cells expressing SufB intein or empty vector pACYC Duet-1 [Figure 3B (iv)]. Hereby, we conclude that we are getting a degradation product of SufB precursor in the cells over-expressing active and inactive Mtu full-length (FL) SufB precursor protein.

Next, MALDI-TOF/TOF mass spectrometry was performed for further confirmation of different splicing and cleavage products (Table S3). Different protein products were cut from SDS PAGE gel and subjected to protein identification by mass spectrometry. First, acquired MS and MS/MS spectra were searched against the taxonomy Mycobacterium tuberculosis from NCBInr (77) database using mascot search engine in Protein Pilot Software. It is found that all the splicing products were identified as SufB protein of Mycobacterium tuberculosis complex with a statistically significant score. As all the splicing products were derivatives of a single protein, the protein identification results confirmed the same (Table S3). Further, acquired MS and MS/MS spectra were checked against a customized database containing probable splicing and cleavage products of SufB protein of Mycobacterium tuberculosis using mascot search engine in Protein Pilot Software with same parameters.
The individual protein band spectra resulted in the identification of protein splicing products matching to it. Individual protein bands identification result is represented with protein score, percent coverage, and theoretical molecular weight and Isoelectric pH value in Table S3. Significance is measured from the expectancy-value (with a p-value $\leq 0.05$).

We performed HPLC analysis to find out fragment weight against a known protein ladder as mentioned in materials and methods. Protein fragments of _Mtu_ FL-SufB were found to be within the expected mass range as per retention time (RT) from HPLC (Figure S1).

The molecular weight of SufB DM (H-5A/H-38A) protein was determined by FPLC, as mentioned in the experimental section. The eluted fractions of both, BSA standard and DM protein were analyzed by SDS gel electrophoresis under both reducing and non-reducing conditions, yielding molecular weights of 66KDa and 28KDa, respectively (Figure S3). No significant difference was observed in their molecular weights under reducing and non-reducing conditions, thereby concluding the existence of both the proteins in their monomeric form.

### 3.6 Proposed mechanism for the cleavage of N-terminal intein-extein peptide bond

The current work investigates the mechanism of _Mtu_ SufB intein cleavage at the N-terminal cleavage site between Gly252 (G-1) and Cys253 (C1). We have identified two highly conserved histidines at -5 (248, full-length protein) and -38 (215, full-length protein) positions of the N-extein sequence from different intein-carrying and intein-less bacterial and archeal SufB proteins. Our *in vitro* experiments suggested that His-5 (248) and His-38 (215) are likely to exert a regulatory role on SufB splicing and/or N-cleavage reactions.

To ascertain this hypothesis, we performed MD simulations to predict a 3-dimensional (3D) model for _Mtu_ full-length (FL)-SufB precursor protein (Figure 7A). This also included
identification of possible critical residues at the SufB intein active site (N- and C-terminal splice
junctions) (Figure 7B and Figure S8), given their proximity to catalytic residues Cys253 (C1),
Cys612 (C+1), and Asn611 (N359).

A

B
C

Gly 252

Asp 217

His 215

Gly 252

Cys 253

Gly 252

Cys 253

His 215

Gly 252

Cys 253

Gly 252

Cys 253

Gly 252
**Figure 7. Structural model and proposed mechanism for N-cleavage reaction in Mtu FL-SufB precursor.** (A) The 3D model of Mtu FL-SufB precursor obtained in presence of water and NaCl. Different domains and conserved catalytic residues are color coded as shown in the legend, (B) Spatial arrangement of the N-terminal active site residues; Cys253 (C1) and Gly252 (G-1) along with His248 (H-5), His215 (H-38) and Asp217 (D-36) at the N-terminal cleavage site, (C) Proposed mechanism for N-terminal cleavage reaction at intein-exetin junction, (D) Free energy change, and (E) Energy and entropy of the system during QM calculation of different reaction states of N-cleavage junction from peptide bond to thio-ester bond of Mtu FL-SufB precursor protein. gs = ground state, gs+ = conformational transition state, ts* = transition state of proton transfer, ts = tetrahedral intermediate, ts = transition state of proton transfer, p = product.
A chimera structure of Mtu FL-SufB protein was built in Schrödinger Bioluminate 2.7 using different templates based on their sequence identity with Mtu FL-SufB (Figure 7A). The sequence identity of Mtu FL-SufB to SufBCD complex from *Escherichia coli* (5AWF_A), PI-SceI; a homing endonuclease with protein splicing activity (1VDE_A) and intein homing endonuclease II (2CW7_A) was found to be 40%, 45%, and 22% respectively. The resultant structure of Mtu FL-SufB was found to have many loops in the intein region because of low sequence identity to the templates. In addition, nearly 13% of amino acids are found to be present in the disallowed region in the Ramachandran plot (Figure S6A). Therefore, we carried out molecular dynamics (MD) simulations of the aforementioned sub-optimized Mtu SufB structure in presence of water and NaCl to mimic *in-vivo* conditions in GROMACS for proper folding of the protein. The structure was solvated in water using the SPC/E water model (59, 60) and interactions were described through the OPLS-AA force field (58). The system was minimized by steepest descent minimization with a target maximum force not greater than 1000 kJ mol⁻¹nm. The simulations were done using GROMACS with time steps of 2fs for a duration of 100ns.

After the 100ns simulation various parameters such as solvent accessible surface area (SASA) (Figure S5B), root mean square deviation (RMSD) was computed and the structure was found to be stabilized after 80ns, suggesting that the dynamics have reached equilibrium (Figure S5A). Models of Mtu FL-SufB and H-5A SufB mutant were simulated for 20ns. Their RMSD and Root Mean Square Fluctuation (RMSF) were compared (Figure S7A and B). The equilibrated Mtu SufB structure obtained from the above MD simulations was found to have only ~1% amino acids in the disallowed region of the Ramachandran plot (Figure S6B). We found that the long side chains of the amino acids that were present in the disallowed region of the Ramachandran plot
were exposed outside. The model was further validated by ProSA web before and after 100ns of MD simulation (Figure S6C, D).

Next, the trajectory analysis of the *Mtu* FL-SufB structure was done to find out the possible role of critical residues including His248 (-5) present near the N- and C- cleavage sites. In particular, we followed the dynamics of His248 (-5) with respect to Gly252 (-1). Our MD simulations showed an interesting dynamical result. It was observed that His248 (-5) pushes another N-extein residue His215 (-38) towards the active site Gly252 (-1). In other words, the MD result suggests that His248 (-5) and His215 (-38) act in a concerted manner to facilitate *Mtu* SufB N-cleavage reaction. We should point out here, although His248 (-5) is closer to the active site compared to His215 (-38) in the primary sequence, our 3-dimensional structure finds the opposite i. e. His215 (-38) is closer to the active site relative to His248 (-5). Specifically, the shortest dynamical distance between His215 (-38) and Gly252 (-1) was observed to be 6.83Å, similar to catalytic distances observed between active residues in other intein systems (12, 34). Therefore, we propose that His248 (-5) and His215 (-38) act concordantly to facilitate the *Mtu* SufB N-cleavage reaction as explained in the discussion section.

Based on the above MD results, a schematic cleavage reaction is shown in (Figure 7C) along the line of canonical intein splicing mechanism (78). Thus, we propose that conjoint effects of active site (N-terminal intein–extein splice junction) residues such as Gly252 (-1), Cys253 (1), and His215 (-38) supported by structural and conformational changes due to His248 (-5) contribute towards *Mtu* SufB N-terminal cleavage reaction. To provide further confidence to our structural and dynamical data, quantum mechanical (QM) calculations were required to obtain energetics of N-cleavage reaction and compared with earlier results. All the quantum chemical calculations were performed with the Gaussian16 package (32, 79, 80). All geometry optimization of the
molecules was computed using the B3LYP method of DFT theory at 6-31G (d) level with charge-neutral singlet \( (32, 79, 81) \). Similar techniques were used in the study of intein cleavage of \( Mtu \) Rec Intein \( (17) \). After the optimized structures of the reactants and the products for each step were obtained, they could be used as the initial and final states respectively in the TS module to find the corresponding energy barriers. Frequency analyses were performed at the same level to identify all the stationary points of the intermediates (only real frequencies) and the saddle points of the transition states (one imaginary frequency). The reaction energy barrier was obtained from the energy difference between the transition state and reactant of the reaction with the zero-point correction. All stationary states were confirmed by means of vibrational frequency analysis, and the Gibbs free energy at physiological temperature was calculated for all stationary points, including the zero-point energy, the entropy, and the thermal energy (all at 298.15K) at the B3LYP/6-31G (d) \( (\text{Figure 7D, 7E}) \). After exploring the reaction energy landscape and obtaining the optimized geometries for intermediates, transition states, and products, we hereby propose a detailed reaction mechanism for \( Mtu \) full-length SufB N-cleavage reaction as elucidated in the discussion section \( (\text{Figure 7C}) \).

Further, the amino acids near the C-cleavage site in the 3D model of \( Mtu \) SufB are identified as His610 (358), Asn 611 (359), and Cys612 (+1) \( (\text{Figure S8}) \). The distance between Asn611 (N359) and Cys612 (C+1) was found to be \( 5.52 \pm 0.11\text{Å} \) and the distance between Asn611 (N359) and His610 (H358) was \( 4.33 \pm 0.25\text{Å} \) \( (\text{Table S5}) \). Due to their proximity, the C-cleavage reaction may be facilitated by interactions between these three amino acids. Future research could shed light on the possible roles of these residues in the mechanism of C-cleavage reaction for \( Mtu \) FL- SufB.
4. DISCUSSION

4.1. Mtu SufB harbors a classic intein splicing system with two conserved histidine residues in the N-extein region

SufB is the central and essential component of the mycobacterial SUF system. This work reports that SufB protein is well conserved in mycobacteria, bacteria, and archaea during the evolutionary course. Intein insertion points may vary in some species, but the conservation of critical residues in intein and extein regions does exist. Intein-carrying SufB proteins are found in different mycobacterial species (Figure 2) (51). Simple sequence analysis depicted the conservation of SufB intein in different organisms and phylogenetic analysis of intein and whole sequence inferred the possibility of common ancestor through evolution (Figure 2). Cladogram analysis suggested a trivial diversion in *M. leprae, M. lepromatosis, M. xenopi, M. triplex* and *Ferroplasma*. The discrepancies in the phylogenetic tree analyses of un-spliced SufB precursors indicated that there are diversity in the extein sequences and intein insertion sites. This suggests intein evolution and independent intein transfer in different species and kingdoms.

Domain analysis of *Mtu* SufB precursor; an 846 aa residue protein clearly demarcated the intein and extein structural regions (Figure 1, Table S2). We found that *Mtu* SufB intein spanning over 359 residues has an intact endonuclease domain. Catalytic residues critical for classic intein splicing pathways like Cys1 (Block A), Cys+1 (Block G), penultimate His (Block G), and terminal Asn (Block G) are conserved in different mycobacterial species including *Mtu*. Besides, *Mtu* SufB intein also contains conserved Block B His67 that is known to catalyze the first N-S acyl shift by destabilizing the scissile peptide bond due to loss of resonance and reduction in energy barrier. *In vitro* refolding of *Mtu* SufB precursor gave rise to splicing and cleavage.
products confirming the catalytic roles of the aforementioned conserved residues. Thus, we conclude that Mtu SufB carries a canonical (classic) cis-splicing intein system (Figure 1).

Furthermore, sequence analysis also detected two highly conserved His residues in the N-extein sequence of Mtu SufB protein; His-38 and His-5 located in all SufB proteins where SUF constitutes the exclusive pathway for [Fe-S] cluster generation irrespective of genus and kingdoms (Figure 2A, B). Later, these histidines were identified to play key regulatory roles during Mtu SufB N-cleavage reaction.

4.2. A distinct extein–intein partnership mechanism guided by highly conserved H-5 and H-38 regulates Mtu SufB N-cleavage reaction

Catalytic residues such as Cys1, Cys+1, and terminal Asn directly participate in intein splicing via promoting sequential nucleophilic displacement reactions or by rearrangement of bonds near splice junctions. Non-catalytic residues assist indirectly via activation of active site residues and stabilization of various intermediate structure(s). Conserved His residues within intein sequence are known to play important roles during protein splicing as well. Block B His accelerates N-S acyl shift and cleavage of N-terminal intein–extein peptide bond whereas F- and G-Block His are crucial in the coordination of terminal Asn cyclization and cleavage of C-terminal splice site. We found that Mtu SufB displays a unique mechanism where two remote histidines located in the N-extein sequence work together to facilitate N-cleavage reaction via activation of catalytic Cys1.

To ascertain the roles of H-5 and H-38, a detailed kinetic analysis was performed on Mtu SufB full-length (FL) precursor and mutant proteins (H-5A and H-38A) [Figure 3, 4, and 5]. We found that both H-5A and H-38A mutants exhibit a sluggish splicing reaction relative to FL- SufB protein under optimum experimental conditions (Table 1). N-cleavage reaction kinetics of the
aforesaid histidine mutants were examined in presence of TCEP and nucleophiles such as Hydroxylamine and DTT. H-5A and H-38A mutants demonstrated a significant reduction in N-cleavage efficiency in comparison to FL- SufB protein in presence of TCEP and nucleophile HA (Table 2 (i) and (ii), Figure5 and 6). But DTT-induced thiolysis reversed the effect of alanine mutation in the histidine mutants. DTT is a thiol protectant reducing agent and exhibits a more efficient thiolysis and di-sulfide reduction. A strong nucleophile such as DTT abolishes the inhibitory effect of alanine mutation on SufB N-cleavage reaction. Herein, we may conclude that H-5 and H-38 possibly have an assistive role in Mtu SufB N-cleavage reaction along with other active site residues. Consequently, the effect of H-5A and H-38A mutation can be reversed in presence of strong nucleophile(s) like DTT (Figure S4).

To further ascertain the roles of conserved histidines, a 3D model of the Mtu FL-SufB protein was built by homology modeling. Molecular dynamics (MD) simulations were performed to equilibrate the aforementioned sub-optimized Mtu SufB structure. We obtained an interesting result during the trajectory analysis of the dynamical data. Although H-38 is relatively distal to the N-terminal splice junction in the protein sequence, in the 3-dimensional structure, it comes closer to the active site. MD results suggest that His248 (-5) and His215 (-38) act in a concerted manner to facilitate Mtu SufB N-cleavage reaction. Thus, we propose that concomitant effects of N-terminal active site residues such as Gly252 (-1), Cys253 (1), and His215 (-38) supported by His248 (-5) contribute towards Mtu SufB N-terminal cleavage reaction. Next, QM/MM calculations were done to determine the energetics of the proposed N-cleavage reaction. Taken together, we hereby propose a detailed mechanism for Mtu SufB N-cleavage reaction. A schematic for the same is depicted in (Figure 7C) along the line of canonical intein splicing mechanism.
In the process to form thioester intermediate from peptide bond of Gly-Cys dipeptide capped with a methyl group, the steps are as follows. The thiol group of Cys253 (1) comes closer to peptidyl C=O by a conformational change to a rotational transition state. This state is enthalpy favored. Then the thiol H is polarized towards the peptidyl O atom. The C=O and S-H experiences change in charge density in the system. The C, S, H atoms in the system experience decrease in electron density, and the O atom experiences an increase in electron density. In the next transition state, C and S come closer as at this state the imaginary vibrational frequency displacement vector of C=O is towards thiol. Thus, a C-S bond is formed. The proton migration from S to O is highly favored energetically and entropically with a free energy change of -9.2 kcal/mol. The computed value agrees well with earlier results (82, 83). After this step sp² hybridized peptide carbon becomes sp³ hybridized forming a tetrahedral transition state, a 2-hydroxy thiazolidine ring. This state annihilates the peptide resonance, therefore free energy change of +3.7 kcal/mol in contrast to the ground state of Gly252 (-1) and Cys253 (1) system. To again regain the sp² hybridization and decrease the energy of the system from resonance stabilization, the 2-hydroxy group expels proton to the environment. This process is energetically advantageous to the system elucidating negative free energy. In a tetrahedral state, S and N atoms have a Mulliken charge of +0.05 and -0.55. Thus, the N atom of the thiazolidine ring accepts a proton from the environment to the system for charge neutrality. This process is also entropically preferred to form thioester intermediate (82, 84). The conversion of a peptide bond to a thioester bond is energetically equivalent at the start as well as the end of the reaction which suggests this process is entropy driven. Further thioester is hydrolyzed by the imidazole side chain of His215 (-38) mediated by water. Base catalyzed thioester hydrolysis in an aqueous environment is well known (85). It is also important to point out that the least distance between His215 (-38) and His248 (-5) obtained
in MD simulations after equilibrium is about 6.83 Å suggesting catalytic distance for an interaction between these two residues (12, 13). Asp217 (-36) prepares protonated histidine His215 (-38) for catalyzing reactions by neutralizing charge through proton transfer. Furthermore, His248 (-5) which is exposed to the solvent, experiences random collisions from water molecules. This action forces His215 (-38) towards the N-cleavage site. This is corroborated by the dynamical distance pattern between His215 (-38) and His248 (-5). Thus, a congruent effort by His248 (-5) facilitates the thioester hydrolysis for cleavage by attacking species i.e., hydroxide ion generation catalyzed by His215 (-38). These results provide confidence in the proposed role of His248 (-5) and His215 (-38). Since histidines are known as metal chelating centers for various metalloproteins, conserved histidines at -5 and -38 positions might coordinate to [Fe-S] clusters contributing towards Mtu SufB functionality as well.

4.3. Biological significance of conserved histidines in the N-extein sequence of Mtu SufB

[Fe-S] cluster-bearing proteins have important physiological roles in electron transfer, redox regulation, metabolic pathways, cellular responses to external stimuli, and as regulators of gene expression (42). The role of the [Fe-S] cluster is closely associated with the functionality of their bound protein framework. In mycobacteria, the SUF system is the sole pathway for [Fe-S] cluster assembly and repair, especially in response to oxidative stress and iron limiting conditions inside macrophages (43, 49). During such an event, the intracellular Fe supply is either from siderophore chelation or via the metabolism of [Fe-S] clusters associated with specific proteins (42). Thus, iron homeostasis plays a major role in mycobacterial survival and virulence. SufB is a [Fe-S] cluster scaffold protein and a vital component of the functional SUF system, and indirectly promotes mycobacterial persistence under stress (49). Further, SufB has been implicated in mycobacterial iron metabolism (42).
Earlier studies have specified the bonding of Fe in [Fe-S] clusters to mostly cysteines from the protein backbone although there is increasing evidence for other ligands such as histidine, aspartate, arginine, threonine, and tyrosine. The most common alternative ligand for [Fe-S] cluster coordination is histidine that is highly conserved with a role in redox tuning and proton-coupled electron transfer (86). His433 (SufB) and His360 (SufD) are identified as key protein-ligands for the de novo [Fe-S] cluster assembly in *E. coli* (48). It has been shown that these non-cysteine ligands can influence the stability and reactivity of [Fe-S] clusters.

We have identified two highly conserved His-5 and His-38 residues in the N-extein sequence of *Mtu* SufB. These were shown to interact concordantly to initiate N-cleavage reaction via catalytic Cys1 activation. Mutational analysis of alanine-substituted His-5 and His-38 residues revealed significantly attenuated cleavage and splicing rates. Attempts to express and analyze the *Mtu* SufB double mutant (His-5A/ His-38A) resulted in a truncated protein product. One of the possible explanations could be the concomitant role of His-5 and His-38 in SufB precursor stabilization. Lack of the histidine residues at the native position perhaps leads to loss of protein stability forming off-pathway splicing intermediate and/or protein degradation product. In summary, the two histidines (His-5 and His-38) work concurrently to activate/initiate N-cleavage reaction leading to the generation of functional SufB protein via protein splicing. As proposed by earlier studies, apart from Cys, His is also an important ligand for metal coordination (48, 86). Thus, it is plausible that the highly conserved His-5 and His-38 possibly coordinate to Fe$^{2+}$/ Fe$^{3+}$ during [Fe-S] cluster biogenesis. Lastly, His-5 and His-38 together might also play a role in *Mtu* SufB precursor stabilization prior to cleavage and splicing as we were unable to express a full-length SufB double mutant (His-5A and His-38A), unlike the single mutants. However, whether
the SufB precursor stabilization is executed by coordination of conserved histidines to [Fe-S] cluster or bonding to Fe\(^{2+}/Fe^{3+}\), it needs extensive experimental and computational confirmation.

Suf is an exclusive system for biogenesis of Fe-S clusters in many pathogenic organisms such as *Staphylococcus aureus*, *Mycobacterium tuberculosis*, *Plasmodium* sp. and *Toxoplasma*, making it an attractive drug target. For instance, D-Cycloserine is a clinical second-line drug currently used against *M. tuberculosis* and inhibits SufS. Investigation on *S. aureus* revealed that SUF is the target system for a polycyclic molecule 882 that has direct interaction with SufC (69). Likewise, the current work may have an implication towards the development of novel anti-TB drugs targeting SufB destabilization, to fight tuberculosis.

**Data Availability:** All the data are available in the manuscript and supplemental data.

The accession code for the *Mycobacterium tuberculosis* SufB protein used in this study is *P9WFP7 (UniProtKB/Swiss-Prot)* and *WP_003407484.1*, GI: 397673309 (NCBI protein database).

**Confirmation of the identity of different splicing and cleavage products of Mtu FL-SufB protein by mass spectrometric analysis:** The data has been deposited to the ProteomeXchange Consortium via PRIDE partner repository with data set identifier PXD015199. The details of the submission are given below.

**Project Name:** Identification of full length, splicing and cleavage products of SufB protein of SUF-complex of *Mycobacterium tuberculosis*.

**Project accession:** PXD015199

**Reviewer account details:**

- **Username:** reviewer73867@ebi.ac.uk
- **Password:** bRxRcbvP

**Confirmation of the identity of H-38A (215)/H-5A (248) Mtu SufB double mutant product by mass spectrometric analysis:** The data has been deposited to the ProteomeXchange Consortium.
via PRIDE partner repository with data set identifier PXD023785. The details of the project submission are given below.

**Project Name:** Identification of splicing and cleavage products of H-38A/H-5A SufB double mutant protein of SUF-complex of *Mycobacterium tuberculosis.*

**Project accession:** PXD023785

**Reviewer account details:**

**Username:** reviewer_pxd023785@ebi.ac.uk

**Password:** 1i6COWkr

*The homology model for the full length Mtu SufB protein has been submitted to Model Archive [Project: ma-x807d]:*

Full length *Mtu* SufB protein of SUF complex of *Mycobacterium tuberculosis*; [https://www.modelarchive.org/doi/10.5452/ma-x807d](https://www.modelarchive.org/doi/10.5452/ma-x807d) with the access code: 6pmXRNkvwR.

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