

1 ***Mycobacterium tuberculosis* SufR Responds to Nitric oxide via its 4Fe-4S cluster and**
2 **Regulates Fe-S cluster Biogenesis for Persistence in Mice**

3 Kushi Anand¹, Ashutosh Tripathi¹, Kaustubh Shukla², Nitish Malhotra³, Anil Kumar
4 Jamithireddy⁴, Rajiv Kumar Jha⁵, Susmit Narayan Chaudhury³, Raju S Rajmani¹, Arati Ramesh³,
5 Valakunja Nagaraja⁵, Balasubramanian Gopal⁴, Ganesh Nagaraju², Aswin Sai Narain Seshayee³,
6 and Amit Singh*^{1, 5}.

7 ¹Centre for Infectious Disease Research, Indian Institute of Science, Bangalore 560012, India.

8 ²Department of Biochemistry, Indian Institute of Science, Bangalore 560012, India. ³National
9 Centre for Biological Science, Bangalore 560065, India. ⁴Molecular Biophysics Unit, Indian
10 Institute of Science, Bangalore 560012, India. ⁵Department of Microbiology and Cell Biology,
11 Indian Institute of Science, Bangalore 560012, India.

12

13 Running Title: NO sensor SufR mediates *M. tuberculosis* persistence

14

15 **Correspondence**

16 *Amit Singh, Ph.D
17 Associate Professor
18 Wellcome Trust-India Alliance Senior Fellow
19 Department of Microbiology and Cell Biology (MCBL)
20 Centre for Infectious Disease Research (CIDR)
21 Indian Institute of Science (IISc)
22 Bangalore-12
23 Ph: +91 8022932604
24 asingh@iisc.ac.in
25

26

27 **Highlights**

28 (i) *Mycobacterium tuberculosis (Mtb)* induces the expression of *suf* operon for
29 Fe-S cluster biogenesis in response to nitric oxide (NO).

30 (ii) We found that a transcription factor SufR senses NO via its 4Fe-4S cluster
31 and regulates the expression of the *suf* operon for Fe-S cluster biogenesis.

32 (iii) SufR-regulated Fe-S cluster biogenesis confers respiratory and redox features
33 that promote recovery of *Mtb* from NO stress.

34 (iv) SufR activity is required to support the NO-dependent persistence of *Mtb* in
35 macrophages and mice.

36

37

38

39

40

41

42

43

44

45

46 **ABSTRACT**

47 The persistence of *Mycobacterium tuberculosis* (*Mtb*) is a major problem in managing
48 tuberculosis (TB). Host-generated nitric oxide (NO) is perceived as one of the signals by *Mtb* to
49 reprogram metabolism and respiration for persistence. However, the mechanisms involved in NO
50 sensing and reorganizing *Mtb*'s physiology are not fully understood. Since NO damages Fe-S
51 clusters of essential enzymes, the mechanism(s) involved in regulating iron-sulfur (Fe-S) cluster
52 biogenesis could help *Mtb* persist in host tissues. Here, we show that a transcription factor SufR
53 (Rv1460) senses NO via its 4Fe-4S cluster and promotes persistence of *Mtb* by mobilizing the
54 Fe-S cluster biogenesis system; *suf* operon (*Rv1460-Rv1466*). Analysis of anaerobically purified
55 SufR by UV-visible spectroscopy, circular dichroism, and iron-sulfide estimation confirms the
56 presence of a 4Fe-4S cluster. Atmospheric O₂ and H₂O₂ gradually degrade the 4Fe-4S cluster of
57 SufR. Furthermore, electron paramagnetic resonance (EPR) analysis demonstrates that NO
58 directly targets SufR 4Fe-4S cluster by forming a protein-bound dinitrosyl-iron-dithiol complex.
59 DNase I footprinting, gel-shift, and *in vitro* transcription assays confirm that SufR directly
60 regulates the expression of the *suf* operon in response to NO. Consistent with this, RNA-
61 sequencing of *MtbΔsufR* demonstrates deregulation of the *suf* operon under NO stress. Strikingly,
62 NO inflicted irreversible damage upon Fe-S clusters to exhaust respiratory and redox buffering
63 capacity of *MtbΔsufR*. Lastly, *MtbΔsufR* failed to recover from a NO-induced non-growing state
64 and displayed persistence defect inside immune-activated macrophages and murine lungs in a
65 NO-dependent manner. Data suggest that SufR is a sensor of NO that supports persistence by
66 reprogramming Fe-S cluster metabolism and bioenergetics.

67

68

69 **Key words**

70 Dinitrosyl-iron-dithiol complex, Spare respiratory capacity, Redox potential, Gene regulation,

71 Inducible nitric oxide synthase, Transcriptomics

72

73

74

75

76

77

78

79

80

81

82

83

84

85

86

87 **Introduction**

88 About 90% people infected with *Mycobacterium tuberculosis* (*Mtb*) remain asymptomatic for
89 tuberculosis (TB). This indicates that the host immunity effectively suppresses the bacterial
90 replication without eradicating the pathogen. The ability of the host to produce nitric oxide (NO)
91 by an inducible nitric oxide synthase (iNOS) is known to modulate immunity [1] and microbial
92 physiology [2], thereby controlling diverse infections [3, 4] including TB [5]. Upon infection
93 with *Mtb*, lesional macrophages in humans and macaques express functional iNOS [6, 7], and
94 exhaled breath of TB patient contains NO [8]. Importantly, iNOS activity seems to control TB in
95 humans [9, 10]. Mechanistically, NO inhibits respiration and arrests growth of *Mtb* [2]. The
96 bacterial mechanisms responsible for sensing NO and mobilizing adaptation programs are poorly
97 understood.

98

99 Previous models suggested that *Mtb* exploits a three-component system, DosR/S/T, to induce
100 transcriptional changes, growth arrest, and a switch from aerobic to anaerobic respiration under
101 NO stress [11, 12]. However, the NO-mediated activation of DosR regulon is transient [13] and
102 other gases, such as carbon monoxide (CO) and oxygen (O₂)-limitation, similarly activate the
103 DosR pathway [14, 15]. Therefore, the mechanism that specifically coordinates changes in
104 metabolism, growth, and respiration of *Mtb* in response to NO is not fully understood. In this
105 context, a recent study demonstrated active degradation of several iron-sulfur (Fe-S) cluster
106 proteins coordinating respiration, central metabolism, and amino acid biosynthesis in NO-treated
107 *Mtb* [13]. Importantly, a seven-gene operon, the Suf system (Rv1460-Rv1466) that is likely
108 involved in Fe-S cluster biogenesis/repair [16], showed prolonged, elevated expression in
109 response to NO [13]. Interestingly, excluding Rv1460 (*sufR*), other genes of the *suf* operon are

110 essential [16, 17]. Since the reaction of NO with Fe-S clusters generates a lethal dinitrosyl-iron
111 dithiol complex (DNIC) [18], active degradation and calibrated regeneration of Fe-S clusters via
112 the *suf* operon represents a potential adaptive defense against NO. Despite these reports, how
113 *Mtb* senses NO and regulates Fe-S cluster biogenesis remains uncharacterized. Filling this
114 knowledge gap is crucial for understanding the molecular underpinning of *Mtb* persistence.

115

116 Recently, the first protein of the Suf system (SufR; *Rv1460*) has been shown to coordinate a 2Fe-
117 2S and function as a putative regulator of the *suf* operon [19]. However, the C-terminus of SufR
118 contains five cysteine residues (C₁₇₅-C₁₇₉-C₁₉₂-C₂₁₈-C₂₂₀), which suggests serving as a ligand for
119 a 4Fe-4S cluster [20, 21]. The presence of a 2Fe-2S cluster was demonstrated *in vitro* that
120 invariably results in poor Fe-S cluster incorporation [19, 22]. Also, without a functional assay
121 (*e.g.*, DNA binding), the authenticity of the 2Fe-2S form of SufR cannot be validated. Lastly,
122 phenotypic characterization of the *sufR* mutant remained ambiguous as one study showed the
123 requirement of SufR for growth under standard culture conditions but not under stress (*e.g.*, iron-
124 limitation) [19]. In contrast, other studies reported the opposite findings [22, 23]. Several
125 discrepancies thus exist in the previously reported characterization of SufR, and consequently, in
126 our understanding of its physiological function. In this study, we performed biochemical,
127 biophysical, and genetic characterization of SufR. We show that SufR contains a NO-sensitive
128 4Fe-4S and regulates Fe-S biogenesis, redox balance, and bioenergetics of *Mtb* to promote
129 persistence in response to NO *in vivo*.

130

131

132 **Material and Methods**

133 **Culture Conditions**

134 The *Mtb* H37Rv, *Mtb* Δ *sufR*, and *sufR*-comp were grown in Middlebrook 7H9 broth (Becton,
135 Dickinson and Company (BD), USA) medium supplemented with 0.2% glycerol, 0.5% BSA,
136 0.2% dextrose, and 0.085% NaCl (ADS) with 0.05% Tween 80 as described previously [23]. For
137 culturing on solid medium, *Mtb* strains were cultured on 7H10/7H11 agar medium (Becton,
138 Dickinson and Company (BD), USA) supplemented with 1x OADC (Becton, Dickinson and
139 Company (BD), USA) and 0.2% glycerol. *E. coli* cultures were grown in LB medium
140 (HIMEDIA, India). Whenever required, antibiotics were added to the culture medium (for *E.*
141 *coli*, 100 μ g/ml kanamycin (Amresco, USA) and 150 μ g/ml hygromycin (Sigma-Aldrich, India);
142 for *Mtb* strains, 50 μ g/ml hygromycin).

143

144 **Generation of the *sufR*-complemented strain**

145 Primer pair: 5'-ATCGAAGCTTGTCCGTCCCTGCCGATCTCAC-3' and 5'-
146 ATGCGGTACCAACGCTCTGCTGGCCTCTG-3' having restriction site HindIII and KpnI
147 were used to amplify *sufR* from wild type *Mtb*. PCR was performed and the product contained
148 the wild-type *sufR* gene, encoded by Rv1460 with the promoter region (567 bp upstream), was
149 cloned into pCV125 (integrated plasmid). Sequence was verified and the constructed plasmids
150 were transformed into the *Mtb* Δ *sufR* strain. Expression of complemented strain was confirmed
151 by qRT-PCR.

152

153 **RNA Sequencing experiments**

154 The *Mtb* strains were grown to an O.D.₆₀₀ of 0.4 and exposed to 0.5 mM diethylenetriamine-
155 nitric oxide (DETA/NO [Sigma-Aldrich, India]) for 4 h at 37°C. The experiment was carried out
156 with three independent biological replicates. Total RNA extraction was conducted using the
157 FastRNA® Pro Blue Kit (MP Biomedicals, USA) in accordance with the manufacturer's
158 instruction and further purified using RNeasy spin columns (Qiagen, USA) as described [23].
159 Following purification, the RNA was quantified and assessed for purity by a 2100 Bioanalyzer
160 (Agilent Technologies, Waldbronn, Germany). RNA samples with an RIN (RNA Integrity
161 Number) value >8 were processed further for sequencing. Ribosomal RNA (16s and 23s rRNA)
162 was removed by hybridization with magnetic beads-coupled oligonucleotide (MICROBExpress
163 Kit, Life Technologies, USA) and concentration of enriched mRNA was quantified by Qubit
164 RNA HS Assay Kit (Life Technologies, USA). RNA-seq was performed as described [16]. In
165 brief, libraries were prepared using NEB Next Ultra Directional RNA Library Prep Kit for
166 Illumina (New England Biolabs, USA), according to manufacturer's instructions. The library
167 size distribution and quality were assessed using a high sensitivity DNA Chip (Agilent
168 Technologies, USA) and sequenced in HiSeq 2500 platform (Illumina, USA) sequencer using
169 1X50 bp single-end reads with 1% PhiX spike-in control.

170

171 **Differential gene expression and statistical analysis for RNA-Seq**

172 Raw reads were obtained for *Mtb* H37Rv strain as fastq files. The reference genome sequence
173 (.fna) and annotation (.gff) files for the same strain (accession number: NC_000962.3) were
174 downloaded from the ncbi ftp website ("ftp.ncbi.nlm.nih.gov"). The annotation file was
175 customized with the addition of annotations for non-coding RNAs [62]. The format of the
176 annotation file (.gff) was changed to .bed format using an in-house python script. The raw read

177 quality was checked using the Fast QC software (version v0.11.5;
178 <http://www.bioinformatics.babraham.ac.uk/projects/fastqc>). BWA (version 0.7.12-r1039)[63]
179 was used to index the reference genome. Reads with raw read quality ≥ 20 were aligned using
180 BWA aln -q option. SAMTOOLS (version 0.1.19-96b5f2294a) [64] was used to filter out the
181 multiply mapped reads. BEDTOOLS (version 2.25.0)[65] was used to calculate the reads count
182 per gene using the annotation file (.bed). The normalization and differential gene expression
183 analysis for the conditions were carried out using edgeR as mentioned previously [66]. Genes
184 with at least 10 reads were selected for each comparative analysis. DGE analysis was done in
185 RStudio (1.1.447) with R version 3.4.4. (<http://www.rstudio.com/>)

186

187 **Aconitase assay**

188 The activity of aconitase (Acn) was measured by monitoring the disappearance of cis-aconitate
189 at 240nm in a UV spectrophotometer (Thermo Scientific Biomat 3S, USA) as described [67].
190 One unit (U) of aconitase activity is defined as 1 μ mol cis-aconitate formed or converted per
191 minute. Reaction mixtures (1ml) for Acn contained 25mM Tris-HCl (pH 8.0), 100 mM NaCl,
192 and 50 μ g *Mtb* cell lysates. Reactions were initiated by adding 0.15 mM cis-aconitate and
193 monitored by following the disappearance of cis-aconitate at 240 nm after every 15 sec for total
194 30 min. Absorbance at 240 nm was plotted against time. Acn activity was calculated from linear
195 portion of the curve in initial 5 min when reaction follows 0th order of reaction. An extinction
196 coefficient of 3,500 M⁻¹cm⁻¹ was used to calculate the rates.

197

198 **Western blot**

199 Whole cells lysate (50 µg) was separated on 12% SDS-PAGE and then transferred onto a PVDF
200 membrane (GE Healthcare, Piscataway, NJ, USA). Membrane were blocked in 5% (w/v) nonfat
201 dry milk and incubated for 3 h at room temperature with primary antibody (Acn and Cbs 1:10000
202 dilution). After washing with 1XTBST, membranes were incubated in goat anti-rabbit IgG HRP-
203 conjugated secondary antibody (1:10000 dilution) for 1 h. The autoradiography signals were
204 visualized using ECL advance Western blotting detection kit (BioRad, USA).

205

206 **OCR and ECAR measurements**

207 The *Mtb* strains adhered to the bottom of a XF cell culture microplate (Agilent technologies,
208 USA), at 2X10⁶ bacilli per well by using Cell-Tak (a cell adhesive). OCR and ECAR were
209 measured using Agilent XF Extracellular Flux Analyser. Assays were carried out in unbuffered
210 7H9 media (pH 7.35) with glucose 2 mg/ml as carbon source. Basal OCR and ECAR were
211 measured for initial 21 min before the automatic addition of freshly prepared DETA-NO (0 mM,
212 0.25 mM, 0.5 mM and 1 mM) in 7H9 unbuffered media, through port A of cartridge plate. Three
213 measurements were taken after 1 h of incubation. CCCP (Sigma-Aldrich, India) was added at 10
214 µM concentration to achieve maximum rate of respiration. Raw data of OCR and ECAR was
215 CFU normalised for 2X10⁶ CFU/ well. Spare respiratory capacity, was calculated from % OCR
216 value, by subtracting third basal reading (normalized as 100%) from first point after CCCP
217 addition.

218

219 **CellRox Deep Red Staining and Flow Cytometry**

220 Logarithmically growing *Mtb* strains were treated with DETA-NO (0 mM, 0.25 mM, 0.5 mM
221 and 1 mM) and then incubated at 37 °C with shaking for 2 h. 200 µl cells were treated with
222 CellROX® Deep Red reagent (Thermo Fisher, USA) at a final concentration of 5 µM for 30
223 minutes at 37°C and analysed on BD FACSVerse flow cytometer with 640/665 nm excitation
224 and emission respectively. We collected 5000-10,000 events for each sample wherever possible.

225

226 **Animal experiments**

227 For the chronic model of infection, 5- to 6-week-old female BALB/c, C57BL/6 and iNOS^{-/-} mice
228 (n = 6 per group) were infected by aerosol with approximately 100 bacilli per mouse with the
229 *Mtb* strains using a Madison chamber aerosol generation. At indicated times post infection, mice
230 were euthanized, and the lungs were harvested for bacillary load, tissue histopathology analysis,
231 and pathological scoring as described [23]. The remaining tissue samples from each mouse were
232 homogenized and bacillary load was quantified by plating serial dilutions of tissue homogenates
233 onto Middlebrook 7H11-OADC agar plates supplemented with lyophilized BBL MGIT PANTA
234 antibiotic mixture ((polymyxin B, amphotericin B, nalidixic acid, trimethoprim, and azlocillin, as
235 supplied by BD; USA). Colonies were observed and counted after 4 weeks of incubation at
236 37°C.

237

238 **Construction of *Mtb* SufT knockdown strain**

239 For construction of *Mtb* SufT knock down strain (SufT-KD), we have followed CRISPR
240 interference (CRISPRi) technology was utilized as described previously [68].
241 Anhydrotetracycline (ATc, Cayman, USA) 200 ng/ml was added for the induction of *sufT*

242 specific-guide RNA (sgRNA) and dCas9 every 48 h. SufT-KD culture was divided equally when
243 A_{600} reached 0.1–0.2 and cultured in the presence or absence of ATc. The dCas9 was expressed
244 by pRH2502 from a TetR-regulated *uvtetO* promoter and sgRNA in pRH2521 under the control
245 of a TetR-regulated *smyc* promoter (P*myc1tetO*). To deplete *sufT*, gene specific sgRNAs were
246 designed for two regions between 98-127 bp and 171-191 bp of *sufT* and cloned in pRH2521.
247 Depletion of *sufT* was verified by qRT-PCR and based on the significant repression of *sufT*, we
248 chose sgRNA targeting *sufT* region between 98-127 bp for further study. For qRT-PCR analysis,
249 total RNA was extracted from SufT-KD and control strains after 24 h of ATc treatment. After
250 DNase treatment, total 600 ng of RNA was used for cDNA synthesis by using Random hexamer
251 oligonucleotide primer (iScript Select cDNA Synthesis Kit, BioRad, USA). Gene specific
252 primers (Table S2) and iQ SYBER Green Supermix (BioRad, USA) were used for RT-PCR
253 (StepOne Plus, Thermo, USA). Gene expression was normalized to *Mtb* 16S rRNA expression
254 level.

255

256 **NO exposure and recovery**

257 Logarithmically grown *Mtb* strains (OD_{600} of 0.8) were diluted to an OD_{600} of 0.1 and exposed to
258 six doses of 100 μ M DETA-NO, once every 6 h. Recovery from exposure to DETA-NO was
259 monitored by recording the OD_{600} after each addition of DETA-NO, as well as every 24 h after
260 addition. Two biological replicates were used for each strain. For single dose experiments,
261 different doses of DETA-NO (0.5 mM, 1.25 mM, 2.5 mM and 1mM) were added. After 4 and 24
262 h cells were harvested, washed with 1X PBS and plated on ADS-7H11 plates. Colonies were
263 counted after 3–4 weeks of incubation at 37°C.

264

265 **Cell line experiments**

266 RAW264.7 murine macrophage cell line was activated by treatment with IFN γ (100U/mL,
267 Invitrogen, USA) for 12 h before infection and LPS (100 ng/mL, Sigma-Aldrich, India) for 2 h
268 before infection. Activated RAW264.7 macrophages were infected with wt *Mtb*, *MtbΔsufR*, and
269 *sufR-Comp* strains at multiplicity of infection (MOI) 2 for 4 h, followed by washing thoroughly
270 to remove extracellular bacteria with warm DMEM medium and suspended in the same
271 containing 10% FBS. For CFU determination, macrophages were lysed using 0.06 % SDS-7H9
272 medium diluted in PBS/Tween and plated on OADC-7H11 at indicated time points. Colonies
273 were counted after 3–4 weeks of incubation at 37°C.

274

275 **Purification of SufR under anaerobic conditions**

276 The entire ORF of *Mtb* *sufR* (Rv1460) was PCR-amplified using gene-specific oligonucleotides
277 (pET28asufRF and pET28asufRR; Table S2), digested with *Nde*I-*Hind*III, and ligated into
278 similarly digested His-tag-based expression vector, pET28a (TAKARA BIO, Clontech
279 Laboratories, CA, USA) to generate pET28a:SufR. A N-terminal histidine-tagged SufR was
280 overexpressed in *E. coli* BL21 λDE3 by 0.6 mM IPTG [Isopropyl β- d-1-thiogalactopyranoside;
281 MP Biomedicals, USA (60 min, 30 °C)]. To facilitate Fe-S cluster formation, cultures were
282 incubated on ice for 18 min prior to induction and were supplemented with 300 μM ferric
283 ammonium citrate and 75 μM L-methionine (Amresco, USA) and purified as described [24].
284 Purification was performed under strict anaerobic conditions inside an anaerobic glove box
285 (PLas-Labs, Lansing, MI, USA) maintaining ≈2.0 ppm O₂ by volume, and buffers and solutions
286 were appropriately deoxygenated. The *sufR* gene on pET28a-SufR was mutated using
287 oligonucleotide-based site-directed mutagenesis approach to create individual cysteine to alanine

288 substitutions. After the PCR, *DpnI* was added into the reaction mixture to digest the wild- type
289 plasmid that was used as the template. The reaction mixture containing the mutated *sufR* gene
290 was used to transform *E. coli* BL21 λDE3. Sequences of primers used to create mutations are
291 shown in (Table S2). Resulting clones were verified by sequencing, and the mutant Cys variants
292 of the wt SufR were purified as described earlier. In order to generate apo-SufR, the holo-SufR
293 was incubated with EDTA (Ethylenediaminetetraacetic acid, Sigma-Aldrich, India) and
294 potassium ferricyanide in a molar ratio of protein: EDTA: ferricyanide in 1:50:20 at 25°C and
295 incubated for 20-30 min till the extensive loss of color. The solution was passed through PD10
296 desalting column and stored at -80 °C.

297

298 **UV-Vis, CD analysis, and gel filtration of SufR**

299 The UV-visible absorption spectroscopy was carried out in a Thermo scientific
300 spectrophotometer (Thermo scientific, USA) at 25°C. Absorption spectra of SufR WT
301 (native/holet) and mutants were recorded immediately as the elution fractions were collected
302 during the purification. In order to study the effect of air oxidation, on Fe-S cluster stability,
303 freshly purified holo-SufR was transferred to an anaerobic quartz cuvette, exposed to air by
304 opening the cap and mixing by pipetting for 2 min. The cuvette was then sealed and monitored
305 by UV-visible spectroscopy over time (Thermo scientific, USA). To study the effect of NO,
306 DTH and H₂O₂ on [4Fe-4S] cluster of SufR, the absorption spectra of freshly purified protein
307 were recorded at indicated concentration and different time intervals in anaerobic quartz cuvette.
308 CD measurements were conducted in a Jasco J-715 spectropolarimeter (Jasco, USA). Far-UV
309 spectra were measured from 190 to 250 nm range and near-UV spectra from 300 nm to 650 nm
310 range. Protein concentration used for the far-UV CD measurements was 10-20 μM and for the

311 near-UV measurements was 140-150 μ M. Cells of 1.0 cm path length were used for the
312 measurements of the far- and near-UV spectra, respectively. Three repeat scans were obtained
313 for each sample. The averaged baseline spectrum was subtracted from the averaged sample
314 spectrum. The protein was dissolved in 5 mM phosphate buffer pH 7.4 and 150 mM NaCl.
315 Results are expressed as molar ellipticity $[\theta]$ (deg cm² dmol⁻¹), calculated from the following
316 formula $[\theta]_{\lambda} = \theta / [c] \square l \square 10 \square n$, where θ is the measured ellipticity in degrees at wavelength λ , c is
317 the protein concentration in mg ml⁻¹, l is the light path length in centimeters and n is the number
318 of amino acids. CD Pro software was used to analyze the data.

319

320 To determine the molecular mass of apo- and holo-SufR protein, analytical size-exclusion
321 chromatography experiments were performed with Superdex 200 increase, 10/300 GL analytical
322 column (GE Healthcare Life Sciences, USA). The column was pre-equilibrated and eluted with
323 the running buffer (10 mM phosphate buffer, 10% glycerol and 100 mM NaCl at pH 7.4) at a
324 constant flow rate of 0.5 ml/min. Molecular mass of the proteins was determined by using gel
325 filtration molecular mass standard (Carbonic Anhydrase, Albumin, Alcohol Dehydrogenase, β -
326 Amylase, Apoferritin and Thyroglobulin). V_e/V_o was plotted as a function of $\log_{10}M_r$ of the
327 standard protein where V_e is the elution volume of the protein, V_o is the void volume of the
328 column and M_r is the molecular weight of the particular protein. Blue dextran was used to
329 determine the void volume (V_o). Running buffers were purged with nitrogen gas and were
330 degassed thoroughly to remove any dissolved atmospheric oxygen. The experiments were
331 conducted under anaerobic conditions.

332

333 **EPR spectroscopy of SufR**

334 For EPR spectroscopy, holo SufR was treated inside the anaerobic glove box with aliquots of
335 freshly prepared proline NONOate (Cayman Chemicals, Ann Arbor, MI, USA). Aliquots of
336 SufR were placed in an anaerobic cuvette and titrated by injection with aliquots of a 2.5 mM
337 stock solution of proline NONOate (Cayman chemicals, USA). Samples were then transferred to
338 EPR tubes and immediately frozen in liquid nitrogen. EPR characteristics of NO-treated SufR
339 were analyzed by subjecting the samples to continuous-wave spectrometers at liquid nitrogen
340 temperature as described previously [25]. NO-treated SufR was analyzed by continuous-wave
341 EPR on a perpendicular mode X-band EPR spectrometer operating at 100-kHz modulation
342 frequency and equipped with liquid nitrogen cryostat and a dual mode X-band cavity (JES200
343 ESR spectrometer, JEOL, USA). Field calibration was done by using a standard NMR G meter.
344 The background signal from the buffer was subtracted from the spectra.

345

346 **Chemical analysis of iron and sulfide**

347 The total iron content and acid-labile sulfide content of holo-SufR, was measured using a
348 previously described procedure [69]. For iron estimation, freshly purified holo-SufR (0.1 mL)
349 was heated at 95°C degree for 30 min after treatment with 22% HNO₃ (0.1 mL). Samples were
350 cooled to ambient temperature followed by addition of 0.6 mL ammonium acetate (7.5 % w/v),
351 0.1 mL freshly prepared ascorbic acid (12.5% w/v) and 0.1mL ferene (10 mM). The
352 concentration of iron present in the protein was determined by measuring the absorbance of the
353 product, iron- ferene complex at 593 nm, which was compared with a standard curve prepared
354 from dilutions of freshly prepared Fe(III) solution in the range of 0-200 μM. To measure acid-
355 labile sulphide content, freshly prepared Na₂S.9H₂O solution was used to prepared standard
356 solution in the range of 52-260 μM of S²⁻. Protein sample/standard (200 μL) was mixed with 0.6

357 mL of zinc acetate (1% w/v) followed by addition of 50 μ L of NaOH (12% w/v). After
358 incubation of 15 min, 0.15 mL of N, N-dimethyl-p-phenylenediamine dihydrochloride (0.1% w/v
359 dissolved in 5 M HCl) and 0.15 mL of freshly prepared 10 mM FeCl₃ (dissolved in 1 M HCl)
360 was added. The reaction mixture was further incubated for 30 min at room temperature and the
361 absorbance of the product, methylene blue, was measured at 670 nm. For iron and sulfide
362 estimation three independent preparations of holo-SufR were analyzed to ensure the
363 reproducibility and three dilution of each sample were considered.

364

365 **Electrophoretic mobility shift assay (EMSA)**

366 For EMSA, promoter fragment of *sufR* (170 bp upstream of ATG) and *blaC* (200 bp upstream of
367 ATG) were PCR amplified from the *Mtb* genome. The 5' end was labeled by [³²P]-ATP using
368 T4 polynucleotide kinase (MBI Fermentas, USA) as per the manufacturer's instructions. The
369 labeled oligonucleotides were passed through a 1 mL Tris-EDTA, pH 7.5 equilibrated Sephadex
370 G-50 column and elute was vacuum dried. Blunt ended duplex DNA was prepared by annealing
371 radiolabelled and complementary cold oligonucleotides in a 1:2 molar ratio in 50 μ L reaction
372 system containing TE and 1X saline-sodium citrate buffer (3 M sodium chloride and 0.3 M
373 sodium citrate). The reaction mixture was heated to 95 °C for 5 min and then cooled in a thermal
374 cycler. Annealed mixtures were resolved by 10% native PAGE in 1X Tris-borate EDTA (TBE).
375 Assembled DNA substrates were visualized on an X-ray film and substrates were subsequently
376 purified by excising the respective desired bands. DNA substrates were eluted by incubating
377 corresponding gel pieces in TE at 4°C for 6 h. Binding reactions were performed in binding
378 buffer (25mM Tris-HCl, 1mM DTT, 0.1 mg/ml BSA, 5 mM MgCl₂; pH 7.4) for 30 min at 4°C
379 and 6% polyacrylamide gel was used to resolve protein-DNA complexes in an anaerobic

380 chamber by electrophoresis at constant voltage (50 V). 0.5 nM ^{32}P -labeled DNA substrates were
381 incubated with indicated concentrations of various forms of SufR (apo, holo, H_2O_2 -, and NO-
382 treated). For competition with unlabeled DNA, fragments of *sufR* and *blaC* (~200–250 bp
383 upstream of translational start codon) were PCR amplified from the *Mtb* genome and used in
384 various amounts to outcompete binding of holo-SufR to ^{32}P -labelled DNA fragments. Gels were
385 exposed to auto radiographic film and visualized by a phosphor imager (Typhoon FLA-9000, GE
386 Healthcare Life Sciences, USA). All the above reactions and separations were performed under
387 anaerobic condition inside glove box (Plas-Labs, Lansing, MI, USA).

388

389 ***In vitro* transcription assays**

390 The DNA templates (180 bp) including the *sufR* promoter regions were PCR amplified using
391 primers PsufR F1/PsufR R1 (Table S2). The amplicons (50 nM) were pre-incubated with
392 different concentration of holo- and apo-SufR in the transcription buffer (50 mM Tris HCl, (pH
393 8.0), 10 mM magnesium acetate, 100 mM EDTA, 100 mM DTT, 50 mM KCl, 50 mg/ml BSA,
394 and 5% glycerol) for 30 min at room temperature. Single-round transcription reactions were
395 initiated with the addition of 100 nM *Mtb* RNAP- σ A holo enzyme, 100 μM NTPs, 1 μCi [α -
396 ^{32}P]-UTP, 50 μg ml $^{-1}$ heparin and incubated at 37°C for 20 min. The reactions were terminated
397 by addition of 2X formamide dye (95% formamide, 0.025% (w/v) bromophenol blue, 0.025%
398 (w/v) xylene cyanol, 5 mM EDTA and 0.025% SDS and 8 M urea) and heated at 95 °C for 5
399 min followed by snap chilling in ice for 2 min. The transcripts were resolved on an 8% TBE-
400 urea-PAGE gel. To study the effect of NO, transcription assays were performed as mention
401 above after treating the holo-SufR with proline NONOate for 5 min followed by purification. All

402 the treatments and the reactions were performed inside anaerobic glove box under anoxic
403 condition.

404

405 **DNase I Footprinting**

406 5 nM of ³²P-labeled DNA substrate was incubated with increasing concentration of purified
407 holo-SufR in binding buffer (25 mM Tris- HCl (pH 7.5), 1 mM DTT, 100 µg/ml BSA, 5 mM
408 MgCl₂, 5 mM CaCl₂), and samples were incubated for 30 min at 4°C inside glove box under
409 anaerobic condition. Reactions were initiated by the addition of DNase I to a final concentration
410 of 0.05 units and incubated for 2 min at room temperature. The reactions were terminated by the
411 addition of 150 µL of TE (pH 7.5) followed by incubation at 75°C for 15 mins to deactivate
412 DNase I enzyme. The sample was further subjected to vacuum evaporation and the pellet thus
413 obtained was re-suspended in loading dye (80% (v/v) formamide, 0.1% (v/v) BPB, and 0.1%
414 (v/v) xylene cyanol) and analyzed on a denaturing 15% polyacrylamide gel containing 7M urea.
415 The gel was dried, and the bands were visualized with a Typhoon FLA-9000 phosphor imager
416 (GE Healthcare Life Sciences, USA). Maxam and Gilbert A+G ladder was prepared as described
417 previously [70]. A custom python script was developed to search for the SufR binding site and
418 find its relative positions with respect to genes across the genome. A sequence of 15 nt length
419 was searched in the region 500 bp upstream of the start codon and considered as a potential SufR
420 binding site if it shares 80% identity with 'ACACT' or 'TGTGA' at the ends.

421

422 **Ethics**

423 Animal experimentation: This study was carried out in strict accordance with the guidelines
424 provided by the Committee for the Purpose of Control and Supervision on Experiments on
425 Animals (CPCSEA), Government of India. The protocol of animal experiment was approved by
426 animal ethical Committee on the Ethics of Animal Experiments, Indian Institute of science
427 (IISc), Bangalore, India (Approval number: CAF/Ethics/544/2017). All efforts were made to
428 minimize the suffering.

429

430 **Statistical Analysis**

431 All data were graphed and analyzed with Prism v8.0 (GraphPad) unless otherwise stated.
432 Statistical analyses were performed using Student's t-test (two-tailed). Where comparison of
433 multiple groups was made either one-way or two-way ANOVA with Bonferroni multiple
434 comparisons was performed. Differences with a p value of <0.05 were considered significant.
435 Statistical significance for RNA-seq was calculated using the QL F-test followed by Benjamini-
436 hochberg method of multiple testing correction.

437

438 **Miscellaneous**

439 The molar extinction coefficient per $[4\text{Fe}-4\text{S}]^{2+}$ cluster ($\epsilon_{413}=16200 \text{ M}^{-1} \text{ cm}^{-1}$) was determined
440 from *in vitro* reconstituted protein. Protein concentration of SufR throughout the study was
441 calculated using A_{280} reading. Anti-SufR polyclonal antibody was used to monitor the expression
442 and purification of wt SufR and mutant proteins by Western blot. Samples (5 mg of protein per
443 slot) were resolved by 12% reducing SDS-PAGE and transferred to nitrocellulose membrane.

444 Blots were probed with primary and secondary (horseradish peroxidase-conjugated anti-rabbit
445 IgG) antibodies and processed.

446

447 **Data availability**

448 The RNA-sequencing data presented in the manuscript are deposited into the NCBI Gene
449 Expression Omnibus (GEO) (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE154169>
450 with the accession number GSE154169 and reviewer token kbcdmqecrlotzgp.

451

452 **Results and Discussion**

453 ***Mtb* SufR contains a 4Fe-4S cluster**

454 To understand the role of *Mtb* SufR in sensing NO and regulating Fe-S cluster homeostasis, we
455 first examined the biochemical and biophysical characteristics of the SufR Fe-S cluster. We
456 anaerobically purified histidine-tagged SufR from *E. coli* cultured in growth conditions
457 optimized for the maximum incorporation of Fe-S clusters *in vivo* [24]. As isolated, native SufR
458 (holo-SufR) displayed a characteristic straw brown color, with an absorption maximum
459 indicative of a 4Fe-4S cluster at 413 nm (molar absorption coefficient [ϵ] at 413 = 16200 M⁻¹ cm⁻¹)
460 (Fig 1A and Fig. S1 A-C) [25]. Treatment of holo-SufR with a one-electron donor, sodium
461 dithionite (DTH), caused partial bleaching of brown color and loss of absorbance at 413 nm,
462 consistent with the presence of a redox-responsive 4Fe-4S cluster [25] (Fig. 1A). Gel filtration
463 performed under anaerobic conditions suggested a molecular mass of ~ 54 kDa for native SufR,
464 consistent with a dimer. Clusterless SufR (apo-SufR) also eluted as a dimer, indicating that the
465 Fe-S cluster does not bridge SufR monomers (Fig. 1B). Moreover, the identity of cysteine

466 residues that coordinates the 4Fe-4S cluster was experimentally validated. The alanine mutant of
467 two putative cysteine ligands (C₁₇₅A_C₁₇₉A) was sufficient to disrupt the 4Fe-4S cluster as
468 evident from the loss of absorbance at 413 nm (Fig. 1C).

469

470 A recent study using aerobically purified SufR, demonstrated that SufR is a monomer and
471 coordinates a 2Fe-2S cluster upon Fe-S cluster reconstitution *in vitro* [19]. In general, Fe-S
472 clusters are inherently unstable under aerobic conditions [26] and stabilize only under anaerobic
473 conditions [24]. Fe-S clusters can be assembled *in vitro*, but this process has poor yield [24].
474 Therefore, the technique of assembling the Fe-S cluster *in vivo* followed by anaerobic
475 purification is more likely to provide native configuration of SufR Fe-S cluster. We clarified this
476 by analyzing native SufR using circular dichroism (CD). The far-UV CD-spectrum showed two
477 minima at 208 and 224 nm and indicated 70% α -helical content in the secondary structure (Fig.
478 1D). The near-UV CD-spectrum displayed two characteristic maxima at 330 and 420 nm and
479 two minima near 350 nm and 550 nm, as described earlier for [4Fe-4S] cluster (Fig. 1E) [24, 27]
480 As expected, the near-UV CD-spectrum of apo-SufR did not show a [4Fe-4S] cluster (Fig. 1E).
481 Lastly, the estimation of iron and sulfide ions in the native SufR revealed the association of 3.23-
482 3.75 iron atoms per SufR monomer and similar amount of sulfide ions per SufR monomer (Fig.
483 S1D). In sum, we demonstrate that *Mtb* SufR contains a 4Fe-4S cluster.

484

485 **SufR responds to NO through its 4Fe-4S cluster**

486 We exposed native SufR to the fast-releasing NO donor; proline NONOate ($T_{1/2} \sim 1.8$ s, pH 7.4)
487 and subjected it to UV-visible spectroscopy. Treatment with proline NONOate gradually reduced

488 absorbance at 413 nm and formed a new chromophoric feature at ~350 nm over time with a clear
489 isosbestic point (Fig. 1F). These spectral features were consistent with a dinitrosyl-iron dithiol
490 complex (DNIC), wherein the sulfide ligands of the 4Fe-4S clusters were displaced by NO to
491 form [Fe-(NO)₂] [28]. The DNIC formation was confirmed by subjecting NO-treated SufR to
492 continuous-wave electron paramagnetic resonance (cw-EPR). A strong EPR signal centered at
493 $g=2.03$ was observed in a dose-dependent manner, consistent with the formation of the
494 monomeric DNIC complex on SufR (Fig. 1G) [25].

495

496 The *suf* operon showed induction under oxidizing stress (e.g., H₂O₂) [29]. Therefore, we
497 determined whether the Fe-S cluster of SufR is responsive to atmospheric O₂ and H₂O₂.
498 Exposure to atmospheric O₂ resulted in a slow decline in absorbance at 413 nm (Fig. 1H). A plot
499 of $\Delta A_{413\text{nm}}$ against time revealed the loss of ~75% of the 4Fe-4S cluster only at 20 hours (h) post-
500 O₂ exposure. However, exposure of native SufR to 1 mM H₂O₂ resulted in the rapid loss of
501 absorption at 413 nm (Fig. 1I). A plot of $\Delta A_{413\text{nm}}$ against time revealed the loss of ~75% of the
502 4Fe-4S cluster in 20 min. Higher concentrations of H₂O₂ (10 mM and 100 mM) resulted in the
503 loss of the cluster within 1-2 min (Fig. S2A-C). Taken together, these data demonstrate that the
504 4Fe-4S cluster of SufR is sensitive to NO and H₂O₂.

505

506 **SufR binds upstream of *suf* operon promoter and represses expression**

507 To investigate if SufR mediates Fe-S cluster biogenesis by regulating the *suf* operon expression,
508 we first assessed the DNA-binding properties of SufR. We carried out electrophoretic mobility
509 shift assays (EMSA) using a radioactively (³²P)-labeled 170 bp DNA fragment encompassing the

510 promoter region of *suf* operon [19]. We used four different forms of SufR; holo-SufR, apo-SufR,
511 NO-treated SufR, and H₂O₂-treated SufR. Holo-SufR bound to the *suf* promoter region, whereas
512 apo-SufR, NO-treated SufR, and H₂O₂-treated SufR showed no DNA binding (Fig. 2A-C). DNA
513 binding was outcompeted by 50-100-fold excess of unlabeled *suf* promoter DNA, but not by an
514 unrelated promoter fragment (*blaC*) (Fig. 2D). These findings suggest that *Mtb* SufR is an NO-
515 and H₂O₂-responsive, sequence-specific DNA-binding protein.

516

517 We next performed *in vitro* transcription using *Mtb* RNA polymerase as described [30]. A single
518 round of transcription of the *suf* promoter fragment (180 bp [-73 to + 107 bp]) produced a single
519 transcript of 107 nucleotides, consistent with the leaderless transcription of the *suf* operon [31].
520 Addition of holo-SufR repressed transcription from the *suf* promoter in a dose-dependent
521 manner, whereas apo-SufR did not (Fig. 2E-F). Importantly, the treatment of holo-SufR with
522 proline NONOate (NO) reversed the repression of the *suf* promoter (Fig. 2G). These results
523 indicate that holo-SufR binds to the *suf* promoter and represses expression, whereas NO-
524 damaged or clusterless apo-SufR lacks DNA-binding and transcription repression. The data
525 suggest that occupancy of Fe-S cluster on SufR serves as an indicator of Fe-S cluster-sufficient
526 or -deficient conditions. During low Fe-S cluster demand, holo-SufR represses the Suf system to
527 limit excessive assembly of Fe-S clusters. In contrast, loss or damage of the Fe-S cluster in SufR
528 (apo-SufR) signals heightened demand for Fe-S clusters that results in the de-repression of the
529 *suf* operon and mobilization of Fe-S cluster assembly. In support of this, we depleted an essential
530 gene of the *suf* operon (Rv1466; *sufT*) involved in Fe-S cluster maturation [32] using CRISPR
531 interference (CRISPRi-*sufT*). The depletion of SufT is expected to perturb Fe-S cluster
532 biogenesis and increase the pool of apo-SufR, which could result in the de-repression of the *suf*

533 operon in *Mtb*. Consistent with this, expression of the *suf* operon was induced in CRISPRi-*sufT*
534 as compared to wild-type *Mtb*, indicating that *Mtb* de-represses the *suf* operon in response to
535 abnormal Fe-S cluster biogenesis caused by SufT depletion (Fig. S3).

536

537 Lastly, we performed DNase I footprinting to identify the binding site of holo-SufR on the
538 promoter region of the *suf* operon (Fig. 3A). A clear region of protection from DNase I digestion
539 was evident with increasing molar ratios of SufR:DNA (Fig. 3A-B). The protected region
540 contains a perfect inverted repeat (ACACT-N₅-TGTGA) separated by 5 bp (Fig. 3A-B). This
541 inverted repeat forms a part of a larger inverted repeat (ATTTTGTACACACT-N₅-TGTGAAAAT).
542 Consistent with the footprinting data, mutations in the inverted repeat completely abolished
543 binding, thus confirming that the SufR binds to the palindrome ACACT-N₅-TGTGA (Fig. 3 C-
544 E). Altogether, using multiple techniques, we confirmed that SufR functions as a NO-sensitive
545 DNA-binding transcription factor in *Mtb*.

546

547 **NO regulates expression of the *suf* operon and Fe-S cluster pathways in *Mtb***

548 Having shown that SufR responds to NO via its 4Fe-4S cluster and directly regulates the
549 transcription of *suf* operon promoter, we next asked if SufR coordinates *Mtb*'s adaptation under
550 NO stress. To examine this possibility, we first investigated the role of SufR in regulating
551 transcriptome of *Mtb* in response to NO. All genes of the *suf* operon are essential except *sufR*.
552 [16]. Therefore, we utilized *sufR*-deficient (*MtbΔsufR*) strain of *Mtb* [23]. We generated the *sufR*-
553 complemented strain (*sufR-Comp*) by integrating *sufR* (Rv1460) along with its native promoter
554 (~500 bp upstream of *sufR*) in the genome of *MtbΔsufR*. Using qRT-PCR, we confirmed the
555 restoration of *sufR* expression to wild type (wt) *Mtb* levels in *sufR-Comp* (Fig S4.). Since

556 exposure to 0.5 mM of NO donor diethylenetriamine-nitric oxide (DETA-NO) arrested the
557 growth of wt *Mtb*, *MtbΔsufR*, and *sufR-Comp* without affecting viability (Fig. S5), we performed
558 RNA-sequencing (RNA-seq) on *Mtb* strains treated with 0.5 mM of DETA-NO for 4 hours.

559
560 Our RNA-seq data recapitulated the previously published NO-responsive transcriptome of *Mtb*,
561 with the induction of the DosR, IdeR, Zur, and CsoR regulons, and reduced expression of RNA
562 and protein biosynthesis pathways (fold change ≥ 2 ; FDR ≤ 0.05 ; Fig. S6 and Table S1A) [13].
563 These pathways were similarly affected in *MtbΔsufR* and *sufR-Comp* in response to NO (Fig. S6
564 and Table S1B-C). Since Fe-S cluster proteins are the most susceptible targets of NO [33], we
565 found that the expression of ~50% of genes encoding Fe-S cluster proteins was altered in wt
566 *Mtb*, *MtbΔsufR*, and *sufR-Comp* under NO stress (Fig. 4 and Table S1D). The strong reactivity of
567 NO towards heme iron of primary cytochrome oxidase is thought to arrest respiration [2].
568 However, we found that NO does not affect the expression of genes encoding primary terminal
569 oxidase (cytochrome bc1-aa3 complex; *qcrABC*) in *Mtb* (Fig. 4). Nonetheless, downregulation of
570 Fe-S cluster-containing NADH dehydrogenase complex I (*nuo* operon) and other genes encoding
571 Fe-S cluster proteins involved in central metabolism (*acn*, *prsA*, and *udgA*) are consistent with
572 the inhibition of primary respiration by NO. Moreover, the expression of genes associated with
573 alternate form of respiration (e.g., NADH dehydrogenase type II [*ndh*], cytochrome BD oxidase
574 [*cydABCD*], nitrate reductase [*narH*], sulfite reductase [*sirA*], hydrogenase [*hycP*]) [34], carbon
575 catabolism (*korAB*, *frdB*, *icl1*, and *pckA*), and branched chain amino acid (BCAA) biosynthesis
576 [35] were induced by NO. (Fig. 4).

577

578 The association between NO, respiration, and metabolism is further indicated by the upregulation
579 of Fe-S cluster enzymes associated with the biosynthesis of respiratory cofactors such as
580 isoprenoid (*lytB2* and *gcpE*), molybdopterin (*moaA2*), thiamin (*thiC*), quinolinate (*nadA-B*), and
581 sulfur metabolites (*cysH*) [36-39] (Fig. 4; Table S1D). Also, NO induced the expression of
582 transcription factors containing redox-sensitive Fe-S clusters (e.g., *whiB1*, *whiB2*, and *whiB3*)
583 [25, 40, 41] (Fig. 4). We found that NO induced comparable expression changes in *Mtb* and
584 *MtbΔsufR* (Fig. 4), indicating that SufR is not a global regulator of *Mtb*'s transcriptome under
585 NO stress.

586 Upregulation of the Fe-S cluster transcriptome indicates an increased demand for Fe-S cluster
587 biogenesis. In agreement, expression of the *suf* operon (*sufRBDCSUT*; Rv1461-Rv1466) was
588 stimulated 15- to 25-fold by NO in *Mtb* (Fig. 4). Also, NO induces the cysteine biosynthetic
589 machinery that supplies sulfur for Fe-S cluster biogenesis (Fig. 4). However, genes coordinating
590 biosynthesis of Fe-binding heme (*hemZ*) and encoding Fe-storage proteins (bacterioferretin;
591 *bfrA-B*) were downregulated (Fig. 4). These observations suggest that *Mtb* prioritizes the
592 assembly of Fe-S clusters under NO stress. In contrast to wt *Mtb*, the *suf* operon remains basally
593 expressed in NO-treated *MtbΔsufR* (Fig. 4). A direct comparison of the expression data
594 confirmed a uniformly reduced expression of the *suf* operon in *MtbΔsufR* as compared to wt *Mtb*
595 under standard growing conditions and NO stress (Fig. 5A-D). Since SufR is a putative repressor
596 of the *suf* operon [19], the operon's diminished expression in *MtbΔsufR* was unexpected. One
597 possibility is that by replacing 345 bp fragment internal to the *sufR* (+308 to +653 bp) with the
598 hygromycin resistance cassette (*lox-hyg'gfp-lox*)[23], we might have interrupted the NO-
599 inducibility of the downstream *suf* genes (*sufBDCSUT*) in *MtbΔsufR* (Fig. 5A). Consistent with
600 this, the partial transcript of *sufR* originated from the undeleted region (+1 to +307 bp) that is

601 present upstream to the deleted fragment (+308 to +653 bp) retained NO-inducibility in
602 *MtbΔsufR* (Fig. 5C-D). The basal expression of the *sufBDCSUT* genes in *MtbΔsufR* is likely due
603 to alternative transcription start site (TSS2) present upstream of *sufB* (Fig. 5A). As expected,
604 *sufR-Comp* expressing a native copy of *sufR* restored the induction of *sufR* in response to NO,
605 whereas rest of the operon remained basally expressed (Fig. 5B-D and Fig. S7). While the *suf*
606 operon was not induced, the expression of pathways requiring Fe-S clusters remained induced in
607 NO-treated *MtbΔsufR* and *sufR-Comp*. Thus, we anticipate that the heightened demand for Fe-S
608 clusters under NO stress is unlikely to be satisfied in *MtbΔsufR* or in *sufR-Comp*. The restoration
609 of NO-inducibility of *sufR* but not of the *suf* operon in *sufR-Comp* provides an opportunity to
610 assess the contribution of SufR other than regulating Fe-S cluster homeostasis under NO stress.
611 Altogether, the transcriptomic data indicate that SufR is mainly required to adjust the NO-
612 responsive expression of the *suf* operon in *Mtb*.

613

614 **NO irreversibly damages Fe-S clusters of aconitase in *MtbΔsufR***

615 Altered expression of the Fe-S pathways involved in metabolism and respiration by NO indicates
616 that NO might modulate these processes in *Mtb*. To investigate this idea, we evaluated a 4Fe-4S-
617 containing enzyme aconitase (Acn) activity, which functions as a critical gatekeeper of the TCA
618 cycle, and shows sensitivity to NO due to a solvent-exposed Fe atom [42]. A gradual decrease in
619 Acn activity over time was observed in *Mtb* exposed to 0.5 mM of DETA-NO, indicating Fe-S
620 cluster damage. At 12 h post-exposure, DETA-NO triggered a 40% reduction in Acn activity
621 without decreasing its abundance (Fig. 6A). Importantly, re-culturing DETA-NO-treated *Mtb* in a

622 DETA-NO-free medium significantly restored Acn activity, indicating efficient mobilization of
623 Fe-S cluster regeneration machinery (Fig. 6A).

624

625 We next examined if defective induction of the *suf* operon impairs regeneration of NO-damaged
626 Fe-S clusters in *MtbΔsufR*. The Acn activity in *MtbΔsufR* was similar to wt *Mtb* under aerobic
627 growing conditions (Fig. 6B). Moreover, like wt *Mtb*, Acn activity decreased in *MtbΔsufR* under
628 NO stress over time (Fig. 6B). However, in contrast to wt *Mtb*, reactivation of Acn upon removal
629 of NO stress was absent in *MtbΔsufR* (Fig. 6B). Importantly, *sufR-Comp* that maintains *sufR*
630 expression but lacks NO-inducibility of the *suf* operon also failed to reinstate Acn activity (Fig.
631 S8A). Data suggest that NO-mediated induction of the *suf* operon rather than SufR alone is
632 critical for the repair of NO-damaged Fe-S clusters in *MtbΔsufR*. A previous study indicated that
633 Acn activity is dependent upon a stand-alone cysteine desulfurase (IscS) in *Mtb* under standard
634 culture conditions [43]. These findings, along with our data, suggest that *Mtb* prefers IscS under
635 aerobic conditions and Suf system under NO stress for biogenesis of Fe-S clusters. Similar roles
636 were assigned for Isc and Suf systems in *E. coli* [44].

637

638 **NO depletes spare respiratory capacity and perturbs redox homeostasis of *MtbΔsufR***

639 Fe-S cluster-containing enzymes are crucial for maintaining carbon catabolism, oxidative
640 phosphorylation (OXPHOS), and redox balance [36, 45]. Therefore, we exploited Seahorse XF
641 Flux technology to analyze the influence of NO on oxygen consumption rate (OCR) and
642 extracellular acidification rate (ECAR), which are measurable readouts of OXPHOS and
643 glycolysis, respectively [46]. To quantify the basal and maximum rates of OCR and ECAR, we

644 cultured *Mtb* and *MtbΔsufR* in 7H9-glucose in an XF microchamber, then exposed it to DETA-
645 NO, and finally to the uncoupler carbonyl cyanide m-chlorophenyl hydrazine (CCCP). Addition
646 of CCCP stimulates respiration to the maximal capacity manageable by *Mtb*. The difference
647 between basal and CCCP-induced OCR provides an estimate of the spare respiratory capacity
648 (SRC) available for sustaining stress-mediated bioenergetics (e.g., nitrosative and oxidative
649 conditions) [47]. Under normal growing conditions, *Mtb* displayed a basal OCR of 20±0.14
650 pmoles/min, which increased to 85±11.7 pmoles/min in response to uncoupling stress by CCCP
651 (Fig. 6C). This indicates that *Mtb* normally functions at a submaximal OXPHOS (~25%)
652 capacity. Under similar conditions, *MtbΔsufR* utilizes ~ 35% of its maximal respiratory capacity,
653 which is more than wt *Mtb* (Fig. 6D). Similar to the uncoupler CCCP, NO also depolarizes the
654 cytoplasmic membrane to arrest respiration and growth [48, 49]. Consistent with this, and as
655 seen with CCCP, pretreatment with DETA-NO also increased basal OCR of wt *Mtb* and
656 *MtbΔsufR* (Fig. 6C, D). However, while 1 mM of DETA-NO was required to increase basal OCR
657 of wt *Mtb* significantly, 0.25 mM was sufficient for *MtbΔsufR* (Fig. 6C, D). Data suggest that NO
658 stimulated basal OCR, possibly by collapsing proton motive force (PMF), and that *MtbΔsufR* is
659 more sensitive to membrane depolarization by NO.

660

661 We also found that DETA-NO pretreatment progressively reduced the ability of bacteria to
662 increase OCR in response to CCCP (Fig. 6C, D). As a result, DETA-NO significantly decreased
663 SRC in a dose-dependent manner in both *Mtb* and *MtbΔsufR* (Fig. 6E). However, SRC of
664 *MtbΔsufR* was significantly lower than wt *Mtb* under normal culture conditions and upon
665 exposure to 0.25 mM DETA-NO (Fig. 6E). These results indicate that *Mtb* mobilizes its reserved
666 respiratory capacity to sustain bioenergetics in response to NO. Data also suggest that the

667 inherently reduced SRC of *MtbΔsufR* due to diminished Fe-S cluster biogenesis increased its
668 vulnerability towards bioenergetic exhaustion by NO.

669
670 Measurement of basal ECAR with and without CCCP treatment indicated that wt *Mtb* and
671 *MtbΔsufR* operate at a suboptimal glycolytic capacity of 20% and 15%, respectively (Fig. S8B-
672 C). Similar to OCR, DETA-NO pretreatment progressively reduced the ability of *Mtb* to increase
673 ECAR in response to CCCP (Fig. S8B). However, unlike wt *Mtb*, *MtbΔsufR* significantly
674 increased basal ECAR and retains CCCP-induced ECAR in response to 0.25 mM and 0.5 mM of
675 DETA-NO (Fig. S8C). This suggests an increased reliance of *MtbΔsufR* on glycolysis to handle
676 the bioenergetic needs under NO stress. Similar to *MtbΔsufR*, the NO-induced changes in OCR,
677 SRC, and ECAR were recapitulated in *sufR-Comp* (Fig. S8D-F), indicating that the increased
678 expression of the *suf* operon rather than *sufR* alone is critical for *Mtb*'s response to NO.

679
680 Lastly, we asked whether NO perturbed redox homeostasis in *Mtb*. We used a genetic biosensor
681 (Mrx1-roGFP2) to measure the redox potential of a physiologically relevant antioxidant,
682 mycothiol (MSH), as a proxy for the cytoplasmic redox potential (E_{MSH}) of *Mtb*[50]. Ratiometric
683 measurements of emission at 510 nm after excitation at 405 and 488 nm can easily quantify any
684 changes in redox physiology [50]. In response to an oxidant or a reductant, the biosensor ratio
685 showed a rapid increase or decrease, respectively [50]. *Mtb* and *MtbΔsufR* expressing Mrx1-
686 roGFP2 were treated with 0.25 mM, 0.5 mM, 1 mM DETA-NO, and the biosensor ratio was
687 measured. Exposure of *Mtb* to NO did not increase the Mrx1-roGFP2 ratio, indicating that wt
688 *Mtb* robustly maintains cytoplasmic E_{MSH} in response to NO (Fig. 7A). In contrast, NO induces a

689 slightly higher oxidative shift in E_{MSH} of $Mtb\Delta sufR$ than wt Mtb in a dose-dependent manner
690 (Fig. 7A). Using a ROS-sensitive fluorescent dye Cell ROX, we confirmed that NO induces
691 oxidative stress in $Mtb\Delta sufR$ but not in wt Mtb (Fig. 7B). Altogether, our data indicate that
692 the *suf* operon's NO-mediated induction is required to regenerate Fe-S clusters, maintain
693 respiratory reserves, and buffer redox imbalance.

694

695 **SufR is required to recover from NO-induced growth arrest and persistence *in vivo***

696 Next, we investigated the biological consequence of compromised Fe-S homeostasis and
697 bioenergetics by assessing the phenotype of $Mtb\Delta sufR$ under NO stress. First, we investigated the
698 survival phenotype of $Mtb\Delta sufR$ under NO stress *in vitro*. A single dose of various concentrations
699 of DETA-NO (0.5 mM, 1.25 mM, and 2.5 mM) did not influence the survival of $Mtb\Delta sufR$ (Fig.
700 8A and Fig.S9). Repeated exposure to low doses of NO is known to arrest Mtb 's growth for an
701 extended duration [51]. Administration of 0.1 mM of DETA-NO every 6 h for 36 h induces an
702 extended period of growth arrest followed by recovery of wt Mtb at day 7 post-treatment (Fig.
703 8B). In contrast, $Mtb\Delta sufR$ resumed growth only at day 16 post-treatment with NO (Fig. 8B).
704 The *sufR-Comp* strain showed a recovery defect largely similar to $Mtb\Delta sufR$, reinforcing that the
705 NO inducibility of the entire *suf* operon rather than only *sufR* is necessary for the timely
706 resumption of growth. We confirmed this using CRISPRi-*sufT* strain, which expresses reduced
707 levels of another Fe-S cluster assembly factor- SufT. Similar to $Mtb\Delta sufR$, diminished levels of
708 *sufT* delayed recovery of Mtb from NO-induced growth inhibition (Fig. 8C).

709

710 Immunologically activated macrophages are known to induce nitrosative stress in *Mtb* [52]. In
711 line with this, *MtbΔsufR* showed 10- and 15-fold reduced survival in immune-activated RAW
712 264.7 macrophages as compared to wt *Mtb* at day 2 and 4 post-infection, respectively (Fig. 8D).
713 These observations prompted us to investigate the NO-dependent phenotype of *MtbΔsufR* *in*
714 *vivo*. Previous studies have reported the requirement of SufR for persistence of *Mtb* in mice [22,
715 53]. However, it remains to be addressed if SufR coordinates pathogen's persistence in response
716 to NO. wt*Mtb* and *MtbΔsufR* showed comparable growth in the lungs of BALB/c mice during the
717 acute phase of infection (0-2 weeks) (Fig. 8E). However, *MtbΔsufR* was cleared progressively
718 from the lungs, with a more than 1.5-log decline in bacterial burden by 8 weeks (Fig. 8E). The
719 histopathological changes observed in animals' lungs at 8 weeks post-infection were
720 proportionate to the bacterial burden (Fig. S10). The magnitude of pulmonary damage was
721 highest in case of wt *Mtb* (2.75±0.5), intermediate in *sufR*-Comp (2.0±0.0), and lowest in
722 *MtbΔsufR* (0.25±0.50) (Fig. S10B). Lastly, to clarify NO's role in the persistence defect
723 of *MtbΔsufR*, we infected a highly susceptible mouse strain lacking inducible nitric oxide (iNOS⁻
724 ^{+/−}) [5]. We found that the persistence defect of *MtbΔsufR* was abolished in iNOS^{−/−} mouse,
725 indicating the requirement of SufR for the persistence of *Mtb* in response to NO (Fig. 8F-G).

726

727 Surprisingly, the persistence defect of *MtbΔsufR* was somewhat rescued in animals and
728 macrophages infected with *sufR*-Comp (Fig. 8D-E). Since *suf* operon's expression is also
729 responsive to Fe-limitation encountered *in vivo* (41), it is possible that SufR induces the
730 expression of *suf* operon under Fe-limitation from a promoter that is distinct from NO-responsive
731 promoter in *sufR*-Comp. Consistent with this, overexpression of SufR led to its binding inside the
732 ORF of Rv1461 (*sufB*) in *Mtb* [54]. Moreover, several other transcription factors (*e.g.*, *Rv0081*,

733 *Rv0023, Rv1189, Rv3765c, Rv3849, Rv0260c*, and *glnR*) bind and alter the expression of *suf*
734 operon [54, 55]. Some of these transcription factors, along with SufR, could regulate *suf*
735 operon's expression to restore persistence of *sufR-Comp* in animals and macrophages.
736 Alternatively, other enzymes involved in Fe-S cluster coordination in *Mtb* (e.g., IscS) (42)
737 partially counterbalance the repressed Suf system's effect in *sufR-Comp* *in vivo*. Consistent with
738 this, our unpublished data suggest that IscS and Suf systems compensate for the loss of each
739 other in mediating *Mtb*'s survival inside macrophages. Altogether, data show that SufR enables
740 NO-dependent persistence of *Mtb* during infection.

741

742 **Conclusions**

743 Fe-S cluster production is tightly regulated to promote Fe-S formation when the necessity for the
744 clusters is heightened (e.g., ROI/RNI/iron-limitation) and to limit unnecessary production when
745 the demand is low (e.g., hypoxia) (43). Deregulation of Fe-S cluster biogenesis can lead to toxic
746 accumulation of iron and polysulfides inside cells (43). Therefore, the calibrated expression of
747 Fe-S cluster biogenesis is important. Here we show that *Mtb* SufR is required to regulate Fe-S
748 cluster biogenesis in *Mtb* under NO stress. Our findings provide mechanistic insights into how
749 *Mtb* exploits Fe-S cluster regulation and biogenesis under NO stress to favor the pathogen's
750 persistence.

751

752 We found that NO more severely inhibits spare respiratory capacity of *MtbΔsufR* as compared to
753 wt *Mtb*. Since spare respiratory capacity depends mainly on the recruitment of previously
754 inactive respiratory complexes (55), the SUF system's sustained activation can provide a reserve

755 of Fe-S cluster-containing respiratory complexes to maintain electron transfer in response to NO.
756 Induction of the *suf* operon in response to conditions that damage Fe-S clusters (*e.g.*, H₂O₂, NO,
757 iron-starvation, antibiotics, phagosomal pH, and sputum) [22, 23, 29, 30, 56-59] indicates that
758 *Mtb* relies on Suf-dependent Fe-S cluster coordination to maintain persistence. The capacity of
759 SufR to sense and respond to a range of cues, such as NO, H₂O₂, and iron limitation, possibly
760 empowers *Mtb* to transduce different redox signals into transcriptional responses crucial for
761 persistence *in vivo*.

762

763 One limitation of our study is the lack of complementation in various *in vitro* assays. Since the
764 polar effects of *sufR* disruption interfered with NO-inducibility of the downstream *suf* genes, the
765 phenotypic changes exhibited by *MtbΔsufR* were mainly due to basal expression of the *suf*
766 operon. Consistent with this, the restoration of expression of *sufR* alone did not rescue the
767 phenotype of *MtbΔsufR* *in vitro*. Surprisingly, while both *MtbΔsufR* and *sufR-Comp* showed
768 defective recovery from NO-mediated growth arrest, the mutant strain recovered earlier than the
769 complemented strain. In this context, we noticed that NO exposure upregulated the DOS
770 dormancy regulon more in *MtbΔsufR* as compared to wt *Mtb* but below 2-fold (FDR< 0.05)
771 cutoff (Fig. S11). Notably, the expression of the DOS regulon was restored to wt *Mtb* levels in
772 *sufR-Comp* under NO stress (Fig. S11). Therefore, marginally better recovery of *MtbΔsufR* than
773 *sufR-comp* could be a consequence of elevated DOS regulon in the mutant. Agreeing to this, an
774 *Mtb* strain lacking DOS dormancy regulator (*MtbΔdosR*) showed defective recovery from NO-
775 mediated growth cessation [51]. Interestingly, a previous study reported overexpression of the
776 *suf* operon in *MtbΔdosR* under hypoxia [15], signifying a regulatory loop between SufR and
777 DosR in *Mtb*. In addition to SufR and DosR/S/T system, the Fe-S cluster containing regulators

778 such as WhiB3 and WhiB1 also respond to NO [25, 41, 60]. Further, using bacterial-one-hybrid
779 system, another study reported binding of WhiB3 to the promoter region of *sufR* [55], suggesting
780 that further experiments are needed to fully understand the mechanism underlying the regulation
781 of the *suf* operon in *Mtb*.

782

783 Previous studies on the *sufR* mutant did not clarify the polar effects on the downstream *suf* genes.
784 Pandey *et al.*, reported a marginal induction of *sufD*, *sufC*, and *sufT* and a basal expression of
785 *sufB*, *sufS*, and *sufU* in the *sufR*-deleted strain ($\Delta SufR_{TB}$) [22]. $\Delta SufR_{TB}$ grew similar to wt *Mtb*
786 under standard growth conditions but showed survival defect under redox stress and inside
787 macrophages [22]. The mutant also displayed persistence defect in mice [22]. In contrast to our
788 findings, Pandey *et al.*, reported a significantly better survival of the *sufR*-complemented strain
789 ($\Delta SufR_{TB}::pJEBsufRTB$) than wt *Mtb* under diverse *in vitro* stress conditions and macrophages,
790 and full rescue of the persistence defect in mice [22]. Another study generated three identical
791 truncated mutants of *sufR* ($\Delta Rv1460$ stop_1.19, $\Delta Rv1460$ stop_5.19, and $\Delta Rv1460$ stop_5.20) by
792 introducing a premature stop codon at position 122 [19]. Surprisingly, $\Delta Rv1460$ stop_1.19 and
793 $\Delta Rv1460$ stop_5.19 grew slowly than wt *Mtb* under standard growing conditions, whereas growth
794 of $\Delta Rv1460$ stop_5.20 was comparable to wt *Mtb*. Intriguingly, the activity of Fe-S cluster
795 enzymes was not affected in $\Delta Rv1460$ stop_5.20 but diminished in the reported *sufR*-
796 complemented strain [19]. None of these studies examined the expression of full *suf* operon both
797 in the *sufR* mutant and the complemented strain under normal and/or NO stress conditions to rule
798 out polar effects. We believe that the reported discrepancies in the *sufR* complementation could
799 be due to the use of non-native promoters (*e.g.*, mycobacterium optimum promoter [MOP] [22]
800 and *hsp60* [19]) to restore SufR expression in previous studies rather than the NO-responsive

801 native *sufR* promoter used in this study. Altogether, future work is required to explore the
802 breadth of SufR-mediated gene regulation and the role of additional regulators in coordinating
803 the expression of the *suf* operon.

804

805 Lastly, the *suf* operon was uniformly induced in clinical isolates of *Mtb* belonging to five
806 globally circulating lineages during survival inside macrophages [61]. These results indicate that
807 regulation of Fe-S cluster biogenesis is a part of core processes that remain conserved in diverse
808 *Mtb* lineages evolved under selection pressure inside the human host. Altogether, we propose a
809 new model of mycobacterial persistence in which SufR senses NO through its Fe-S cluster to
810 coordinate Fe-S cluster biogenesis and regulate metabolism, respiration, and redox balance (Fig.
811 9).

812

813 ASSOCIATED CONTENT

814 Supporting Information

815 Supporting text files containing supplementary figures and supplementary tables information

816 Figure S1-S11

817 RNA Seq dataset-Table S1A-D (.Xlsx)

818 List of oligonucleotides used in this study-Table S2 (.Xlsx)

819

820 Author Contributions

821 KA, AT, and AS participated in the design of the study. KA, AT, KS, NM, AJ, RKJ, RSR, and
822 SNC, carried out the experiments. AR, VN, BG, GN, and ASN contributed to reagents and
823 analyzed the data. KA and AS conceived the study, supervised the project, analyzed the data and
824 drafted the manuscript. All authors read and approved the final manuscript.

825

826 **Funding Sources**

827 The *Mtb* work was supported by the following Wellcome Trust/DBT India Alliance Grants,
828 IA/S/16/2/502700 (AS), IA/E/16/1/503017 (KA), and in part by DBT grants
829 BT/PR13522/COE/34/27/2015, BT/PR29098/Med/29/1324/2018, and
830 BT/HRD/NBA/39/07/2018-19 (A.S.), DBT-IISc Partnership Program grant 22-0905-0006-05-
831 987 436, and the Infosys Foundation. AS and KA are senior- and early-career fellows of
832 Wellcome Trust/DBT India Alliance.

833

834 **Notes**

835 The funders had no role in study design, data collection and analysis, decision to publish, or
836 preparation of the manuscript. We confirm that no competing financial interests exist.

837

838 **ACKNOWLEDGMENT**

839 We are thankful to Awadhesh Pandit and Next Generation Genomics Facility (NGGF) at
840 National Centre for Biological Sciences, Bangalore, for conducting the RNA-sequencing. We
841 thank BSL3 facilities at CIDR, IISc Bangalore. We thank A. Varalakshmi (Sophisticated

842 Analytical Instrument Facility, IIT Chennai, India) for excellent technical help in EPR
843 experiments.

844

845 **ABBREVIATIONS**

846 *Mtb*, *Mycobacterium tuberculosis*; NO, nitric oxide; iNOS, inducible nitric oxide synthase;
847 DNIC, dinitrosyl-iron dithiol complex; DTH, sodium dithionite; CD, circular dichroism; EMSA,
848 electrophoretic mobility shift assay; DETA-NO, diethylenetriamine-nitric oxide; OCR, Oxygen
849 Consumption Rate; SRC, spare respiratory capacity; CCCP, carbonyl cyanide m-chlorophenyl
850 hydrazine; ECAR, extracellular acidification rate

851

852 **REFERENCES**

- 853 1. Mishra, B.B., et al., *Nitric oxide controls the immunopathology of tuberculosis by*
854 *inhibiting NLRP3 inflammasome-dependent processing of IL-1beta*. *Nat Immunol*, 2013.
855 **14**(1): p. 52-60.
- 856 2. Voskuil, M.I., et al., *Inhibition of respiration by nitric oxide induces a Mycobacterium*
857 *tuberculosis dormancy program*. *J Exp Med*, 2003. **198**(5): p. 705-13.
- 858 3. Nathan, C. and M.U. Shiloh, *Reactive oxygen and nitrogen intermediates in the*
859 *relationship between mammalian hosts and microbial pathogens*. *Proc Natl Acad Sci U S*
860 *A*, 2000. **97**(16): p. 8841-8.
- 861 4. MacMicking, J., Q.W. Xie, and C. Nathan, *Nitric oxide and macrophage function*. *Annu*
862 *Rev Immunol*, 1997. **15**: p. 323-50.
- 863 5. MacMicking, J.D., et al., *Identification of nitric oxide synthase as a protective locus*
864 *against tuberculosis*. *Proc Natl Acad Sci U S A*, 1997. **94**(10): p. 5243-8.
- 865 6. Nicholson, S., et al., *Inducible nitric oxide synthase in pulmonary alveolar macrophages*
866 *from patients with tuberculosis*. *J Exp Med*, 1996. **183**(5): p. 2293-302.
- 867 7. Mattila, J.T., et al., *Microenvironments in tuberculous granulomas are delineated by*
868 *distinct populations of macrophage subsets and expression of nitric oxide synthase and*
869 *arginase isoforms*. *J Immunol*, 2013. **191**(2): p. 773-84.
- 870 8. Wang, C.H., et al., *Increased exhaled nitric oxide in active pulmonary tuberculosis due to*
871 *inducible NO synthase upregulation in alveolar macrophages*. *Eur Respir J*, 1998. **11**(4):
872 p. 809-15.
- 873 9. Schon, T., et al., *Arginine as an adjuvant to chemotherapy improves clinical outcome in*
874 *active tuberculosis*. *Eur Respir J*, 2003. **21**(3): p. 483-8.

875 10. Schon, T., et al., *Effects of a food supplement rich in arginine in patients with smear*
876 *positive pulmonary tuberculosis--a randomised trial.* *Tuberculosis (Edinb)*, 2011. **91**(5):
877 p. 370-7.

878 11. Kumar, A., et al., *Mycobacterium tuberculosis DosS is a redox sensor and DosT is a*
879 *hypoxia sensor.* *Proc Natl Acad Sci U S A*, 2007. **104**(28): p. 11568-73.

880 12. Sardiwal, S., et al., *A GAF domain in the hypoxia/NO-inducible Mycobacterium*
881 *tuberculosis DosS protein binds haem.* *J Mol Biol*, 2005. **353**(5): p. 929-36.

882 13. Cortes, T., et al., *Delayed effects of transcriptional responses in Mycobacterium*
883 *tuberculosis exposed to nitric oxide suggest other mechanisms involved in survival.* *Sci*
884 *Rep*, 2017. **7**(1): p. 8208.

885 14. Shiloh, M.U., P. Manzanillo, and J.S. Cox, *Mycobacterium tuberculosis senses host-*
886 *derived carbon monoxide during macrophage infection.* *Cell Host Microbe*, 2008. **3**(5):
887 p. 323-30.

888 15. Park, H.D., et al., *Rv3133c/dosR is a transcription factor that mediates the hypoxic*
889 *response of Mycobacterium tuberculosis.* *Mol Microbiol*, 2003. **48**(3): p. 833-43.

890 16. Huet, G., M. Daffe, and I. Saves, *Identification of the Mycobacterium tuberculosis SUF*
891 *machinery as the exclusive mycobacterial system of [Fe-S] cluster assembly: evidence for*
892 *its implication in the pathogen's survival.* *J Bacteriol*, 2005. **187**(17): p. 6137-46.

893 17. Sassetti, C.M., D.H. Boyd, and E.J. Rubin, *Genes required for mycobacterial growth*
894 *defined by high density mutagenesis.* *Mol Microbiol*, 2003. **48**(1): p. 77-84.

895 18. Hickok, J.R., et al., *Dinitrosyliron complexes are the most abundant nitric oxide-derived*
896 *cellular adduct: biological parameters of assembly and disappearance.* *Free Radic Biol*
897 *Med*, 2011. **51**(8): p. 1558-66.

898 19. Willemse, D., et al., *Rv1460, a SufR homologue, is a repressor of the suf operon in*
899 *Mycobacterium tuberculosis.* *PLoS One*, 2018. **13**(7): p. e0200145.

900 20. Coldren, C.D., H.W. Hellinga, and J.P. Caradonna, *The rational design and construction*
901 *of a cuboidal iron-sulfur protein.* *Proc Natl Acad Sci U S A*, 1997. **94**(13): p. 6635-40.

902 21. Nanda, V., et al., *Structural principles for computational and de novo design of 4Fe-4S*
903 *metalloproteins.* *Biochim Biophys Acta*, 2016. **1857**(5): p. 531-538.

904 22. Pandey, M., et al., *Iron homeostasis in Mycobacterium tuberculosis is essential for*
905 *persistence.* *Sci Rep*, 2018. **8**(1): p. 17359.

906 23. Mishra, R., et al., *Targeting redox heterogeneity to counteract drug tolerance in*
907 *replicating Mycobacterium tuberculosis.* *Sci Transl Med*, 2019. **11**(518).

908 24. Crack, J.C., et al., *Characterization of [4Fe-4S]-containing and cluster-free forms of*
909 *Streptomyces WhiD.* *Biochemistry*, 2009. **48**(51): p. 12252-64.

910 25. Singh, A., et al., *Mycobacterium tuberculosis WhiB3 responds to O₂ and nitric oxide via*
911 *its [4Fe-4S] cluster and is essential for nutrient starvation survival.* *Proc Natl Acad Sci*
912 *U S A*, 2007. **104**(28): p. 11562-7.

913 26. Imlay, J.A., *Iron-sulphur clusters and the problem with oxygen.* *Mol Microbiol*, 2006.
914 **59**(4): p. 1073-82.

915 27. Freibert, S.A., et al., *Biochemical Reconstitution and Spectroscopic Analysis of Iron-*
916 *Sulfur Proteins.* *Methods Enzymol*, 2018. **599**: p. 197-226.

917 28. Cruz-Ramos, H., et al., *NO sensing by FNR: regulation of the Escherichia coli NO-*
918 *detoxifying flavohaemoglobin, Hmp.* *EMBO J*, 2002. **21**(13): p. 3235-44.

919 29. Voskuil, M.I., et al., *The response of mycobacterium tuberculosis to reactive oxygen and*
920 *nitrogen species.* *Front Microbiol*, 2011. **2**: p. 105.

921 30. Mishra, S., et al., *Efficacy of beta-lactam/beta-lactamase inhibitor combination is linked*
922 *to WhiB4-mediated changes in redox physiology of Mycobacterium tuberculosis*. Elife, 2017. **6**.

924 31. Cortes, T., et al., *Genome-wide mapping of transcriptional start sites defines an extensive*
925 *leaderless transcriptome in Mycobacterium tuberculosis*. Cell Rep, 2013. **5**(4): p. 1121-31.

927 32. Tamuhla, T., et al., *SufT is required for growth of Mycobacterium smegmatis under iron*
928 *limiting conditions*. Microbiology, 2020. **166**(3): p. 296-305.

929 33. Crack, J.C., et al., *Iron-sulfur clusters as biological sensors: the chemistry of reactions*
930 *with molecular oxygen and nitric oxide*. Acc Chem Res, 2014. **47**(10): p. 3196-205.

931 34. Beites, T., et al., *Plasticity of the Mycobacterium tuberculosis respiratory chain and its*
932 *impact on tuberculosis drug development*. Nat Commun, 2019. **10**(1): p. 4970.

933 35. Baughn, A.D., et al., *An anaerobic-type alpha-ketoglutarate ferredoxin oxidoreductase*
934 *completes the oxidative tricarboxylic acid cycle of Mycobacterium tuberculosis*. PLoS Pathog, 2009. **5**(11): p. e1000662.

936 36. Palde, P.B., et al., *First-in-Class Inhibitors of Sulfur Metabolism with Bactericidal*
937 *Activity against Non-Replicating M. tuberculosis*. ACS Chem Biol, 2016. **11**(1): p. 172-84.

939 37. Boshoff, H.I., et al., *Biosynthesis and recycling of nicotinamide cofactors in*
940 *mycobacterium tuberculosis. An essential role for NAD in nonreplicating bacilli*. J Biol Chem, 2008. **283**(28): p. 19329-41.

942 38. Buchko, G.W., et al., *Solution-state NMR structure and biophysical characterization of*
943 *zinc-substituted rubredoxin B (Rv3250c) from Mycobacterium tuberculosis*. Acta Crystallogr Sect F Struct Biol Cryst Commun, 2011. **67**(Pt 9): p. 1148-53.

945 39. Brown, A.C., et al., *The nonmevalonate pathway of isoprenoid biosynthesis in*
946 *Mycobacterium tuberculosis is essential and transcriptionally regulated by Dxs*. J Bacteriol, 2010. **192**(9): p. 2424-33.

948 40. Ren, B., et al., *Nitric oxide-induced bacteriostasis and modification of iron-sulphur*
949 *proteins in Escherichia coli*. Mol Microbiol, 2008. **70**(4): p. 953-64.

950 41. Kudhair, B.K., et al., *Structure of a Wbl protein and implications for NO sensing by M.*
951 *tuberculosis*. Nat Commun, 2017. **8**(1): p. 2280.

952 42. Tortora, V., et al., *Mitochondrial aconitase reaction with nitric oxide, S-*
953 *nitrosoglutathione, and peroxynitrite: mechanisms and relative contributions to*
954 *aconitase inactivation*. Free Radic Biol Med, 2007. **42**(7): p. 1075-88.

955 43. Rybniker, J., et al., *The cysteine desulfurase IscS of Mycobacterium tuberculosis is*
956 *involved in iron-sulfur cluster biogenesis and oxidative stress defence*. Biochem J, 2014. **459**(3): p. 467-78.

958 44. Mettert, E.L. and P.J. Kiley, *How Is Fe-S Cluster Formation Regulated?* Annu Rev
959 Microbiol, 2015. **69**: p. 505-26.

960 45. Lill, R., *Function and biogenesis of iron-sulphur proteins*. Nature, 2009. **460**(7257): p.
961 831-8.

962 46. Zhang, J. and Q. Zhang, *Using Seahorse Machine to Measure OCR and ECAR in Cancer*
963 *Cells*. Methods Mol Biol, 2019. **1928**: p. 353-363.

964 47. Dranka, B.P., B.G. Hill, and V.M. Darley-Usmar, *Mitochondrial reserve capacity in*
965 *endothelial cells: The impact of nitric oxide and reactive oxygen species*. Free Radic Biol
966 Med, 2010. **48**(7): p. 905-14.

967 48. Jones-Carson, J., et al., *Nitric oxide disrupts bacterial cytokinesis by poisoning purine*
968 *metabolism*. *Sci Adv*, 2020. **6**(9): p. eaaz0260.

969 49. Jones-Carson, J., et al., *Nitric oxide from IFN γ -primed macrophages modulates the*
970 *antimicrobial activity of beta-lactams against the intracellular pathogens *Burkholderia**
971 *pseudomallei and Nontyphoidal *Salmonella**. *PLoS Negl Trop Dis*, 2014. **8**(8): p. e3079.

972 50. Bhaskar, A., et al., *Reengineering redox sensitive GFP to measure mycothiol redox*
973 *potential of *Mycobacterium tuberculosis* during infection*. *PLoS Pathog*, 2014. **10**(1): p.
974 e1003902.

975 51. Leistikow, R.L., et al., *The *Mycobacterium tuberculosis* DosR regulon assists in*
976 *metabolic homeostasis and enables rapid recovery from nonrespiring dormancy*. *J*
977 *Bacteriol*, 2010. **192**(6): p. 1662-70.

978 52. MacMicking, J.D., G.A. Taylor, and J.D. McKinney, *Immune control of tuberculosis by*
979 *IFN-gamma-inducible LRG-47*. *Science*, 2003. **302**(5645): p. 654-9.

980 53. Sassetti, C.M. and E.J. Rubin, *Genetic requirements for mycobacterial survival during*
981 *infection*. *Proc Natl Acad Sci U S A*, 2003. **100**(22): p. 12989-94.

982 54. Minch, K.J., et al., *The DNA-binding network of *Mycobacterium tuberculosis**. *Nat*
983 *Commun*, 2015. **6**: p. 5829.

984 55. Guo, M., et al., *Dissecting transcription regulatory pathways through a new bacterial*
985 *one-hybrid reporter system*. *Genome Res*, 2009. **19**(7): p. 1301-8.

986 56. Tyagi, P., et al., **Mycobacterium tuberculosis* has diminished capacity to counteract redox*
987 *stress induced by elevated levels of endogenous superoxide*. *Free Radic Biol Med*, 2015.
988 **84**: p. 344-54.

989 57. Kurthkoti, K., et al., *The Capacity of *Mycobacterium tuberculosis* To Survive Iron*
990 *Starvation Might Enable It To Persist in Iron-Deprived Microenvironments of Human*
991 *Granulomas*. *mBio*, 2017. **8**(4).

992 58. Van den Bossche, A., et al., *Transcriptional profiling of a laboratory and clinical*
993 **Mycobacterium tuberculosis* strain suggests respiratory poisoning upon exposure to*
994 *delamanid*. *Tuberculosis (Edinb)*, 2019. **117**: p. 18-23.

995 59. Kumar, M., et al., *Identification of *Mycobacterium tuberculosis* genes preferentially*
996 *expressed during human infection*. *Microb Pathog*, 2011. **50**(1): p. 31-8.

997 60. Mehta, M. and A. Singh, **Mycobacterium tuberculosis* WhiB3 maintains redox*
998 *homeostasis and survival in response to reactive oxygen and nitrogen species*. *Free Radic*
999 *Biol Med*, 2019. **131**: p. 50-58.

1000 61. Homolka, S., et al., *Functional genetic diversity among *Mycobacterium tuberculosis**
1001 *complex clinical isolates: delineation of conserved core and lineage-specific*
1002 *transcriptomes during intracellular survival*. *PLoS Pathog*, 2010. **6**(7): p. e1000988.

1003 62. Gerrick, E.R., et al., *Small RNA profiling in *Mycobacterium tuberculosis* identifies MrsI*
1004 *as necessary for an anticipatory iron sparing response*. *Proc Natl Acad Sci U S A*, 2018.
1005 **115**(25): p. 6464-6469.

1006 63. Li, H. and R. Durbin, *Fast and accurate short read alignment with Burrows-Wheeler*
1007 *transform*. *Bioinformatics*, 2009. **25**(14): p. 1754-60.

1008 64. Li, H., et al., *The Sequence Alignment/Map format and SAMtools*. *Bioinformatics*, 2009.
1009 **25**(16): p. 2078-9.

1010 65. Quinlan, A.R. and I.M. Hall, *BEDTools: a flexible suite of utilities for comparing*
1011 *genomic features*. *Bioinformatics*, 2010. **26**(6): p. 841-2.

1012 66. Robinson, M.D., D.J. McCarthy, and G.K. Smyth, *edgeR: a Bioconductor package for*
1013 *differential expression analysis of digital gene expression data.* Bioinformatics, 2010.
1014 **26**(1): p. 139-40.

1015 67. Tian, J., et al., *Variant tricarboxylic acid cycle in Mycobacterium tuberculosis:*
1016 *identification of alpha-ketoglutarate decarboxylase.* Proc Natl Acad Sci U S A, 2005.
1017 **102**(30): p. 10670-5.

1018 68. Singh, A.K., et al., *Investigating essential gene function in Mycobacterium tuberculosis*
1019 *using an efficient CRISPR interference system.* Nucleic Acids Res, 2016. **44**(18): p. e143.

1020 69. Crack, J.C., et al., *Techniques for the production, isolation, and analysis of iron-sulfur*
1021 *proteins.* Methods Mol Biol, 2014. **1122**: p. 33-48.

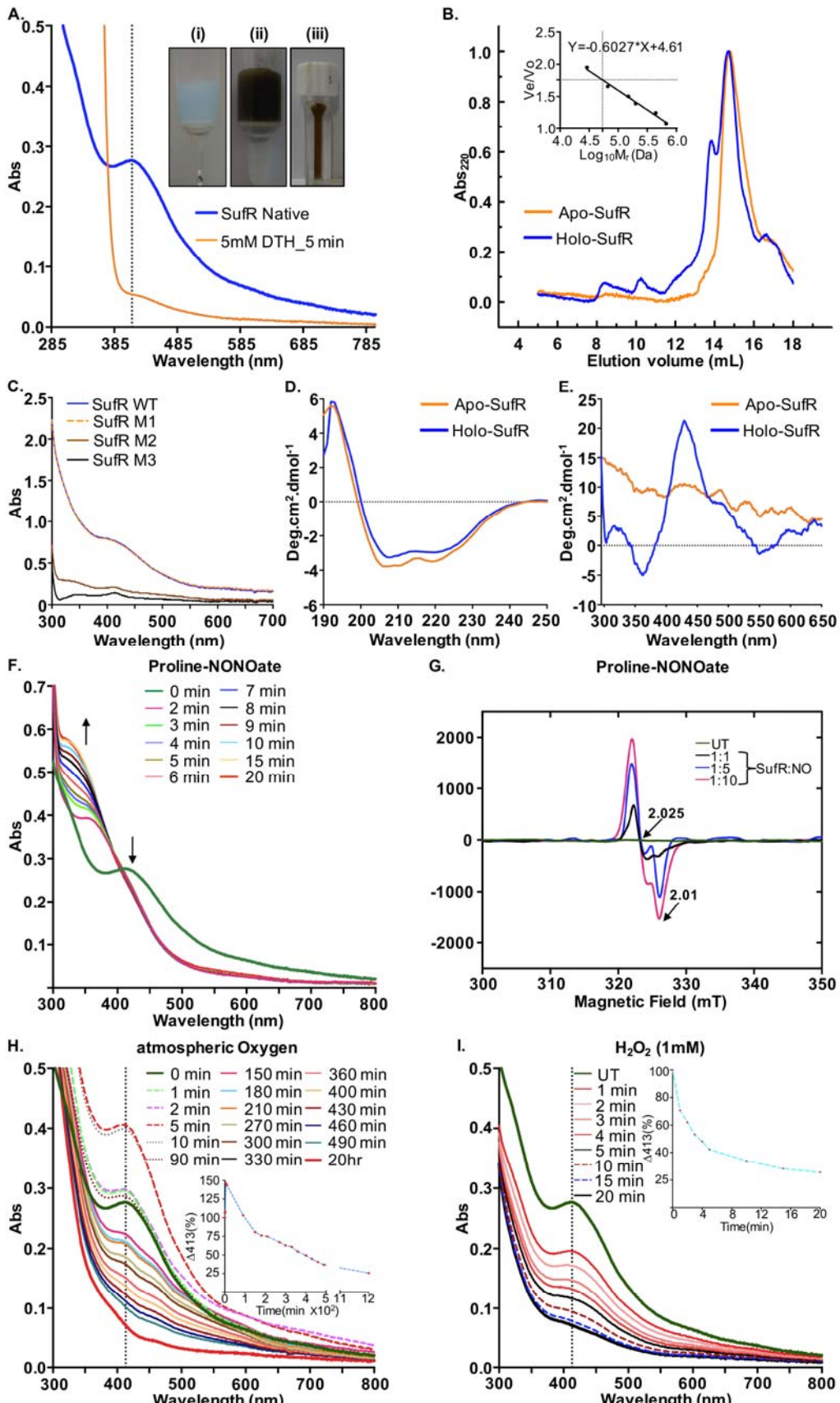
1022 70. Maxam, A.M. and W. Gilbert, *Sequencing end-labeled DNA with base-specific chemical*
1023 *cleavages.* Methods Enzymol, 1980. **65**(1): p. 499-560.

1024

1025

1026

1027



1029 **Figure 1: *Mtb* SufR coordinates 4Fe-4S cluster that responds to NO and H₂O₂. (A)**

1030 Anaerobically-purified SufR was subjected to UV-visible spectroscopy. A clear peak at 413 nm
1031 indicates presence of a 4Fe-4S cluster. The Fe-S cluster was reduced by addition of 5 mM DTH.
1032 (*Inset*) (i) unbound Ni²⁺-NTA, (ii) brown colored SufR bound to Ni²⁺-NTA beads, and (iii)
1033 anaerobically purified holo-SufR. (B) Superdex 200 HR gel filtration chromatography of SufR.
1034 (**Inset**: standard curve of the protein standards versus their elution parameter Ve/Vo where, Ve is
1035 the elution volume and Vo is void volume). (C) UV-visible spectroscopy of cysteine mutants of
1036 SufR (M1-C₁₇₅A, M2-C₁₇₅A_C₁₇₉A, and M3- C₁₇₅A_C₁₇₉A_C₁₉₂A). (D) A far-UV and (E) Near-
1037 UV CD-spectrum of holo- and apo-SufR. (F) UV-visible spectra were acquired before and after
1038 addition of proline NONOate (2.5 mM). A time dependent increase in absorbance at 350 nm
1039 indicates monomeric DNIC formation. (G) Holo-SufR exposed to various concentration of
1040 proline NONOate (SufR:NO). EPR data were acquired using a continuous-wave EPR
1041 spectrometer at a microwave frequency of 9.667 GHz and microwave power of 2 mW at liquid
1042 nitrogen temperature. The peak at 2.025 g is consistent with the formation of monomeric DNIC.
1043 (H) UV-visible spectra of holo- SufR exposed to either atmospheric O₂ for 2 min or (I) H₂O₂ (1
1044 mM). (*Inset*) Rate of the 4Fe-4S cluster loss was determined by calculating the percent loss of
1045 absorbance at 413 nm upon exposure to air and H₂O₂.

1046

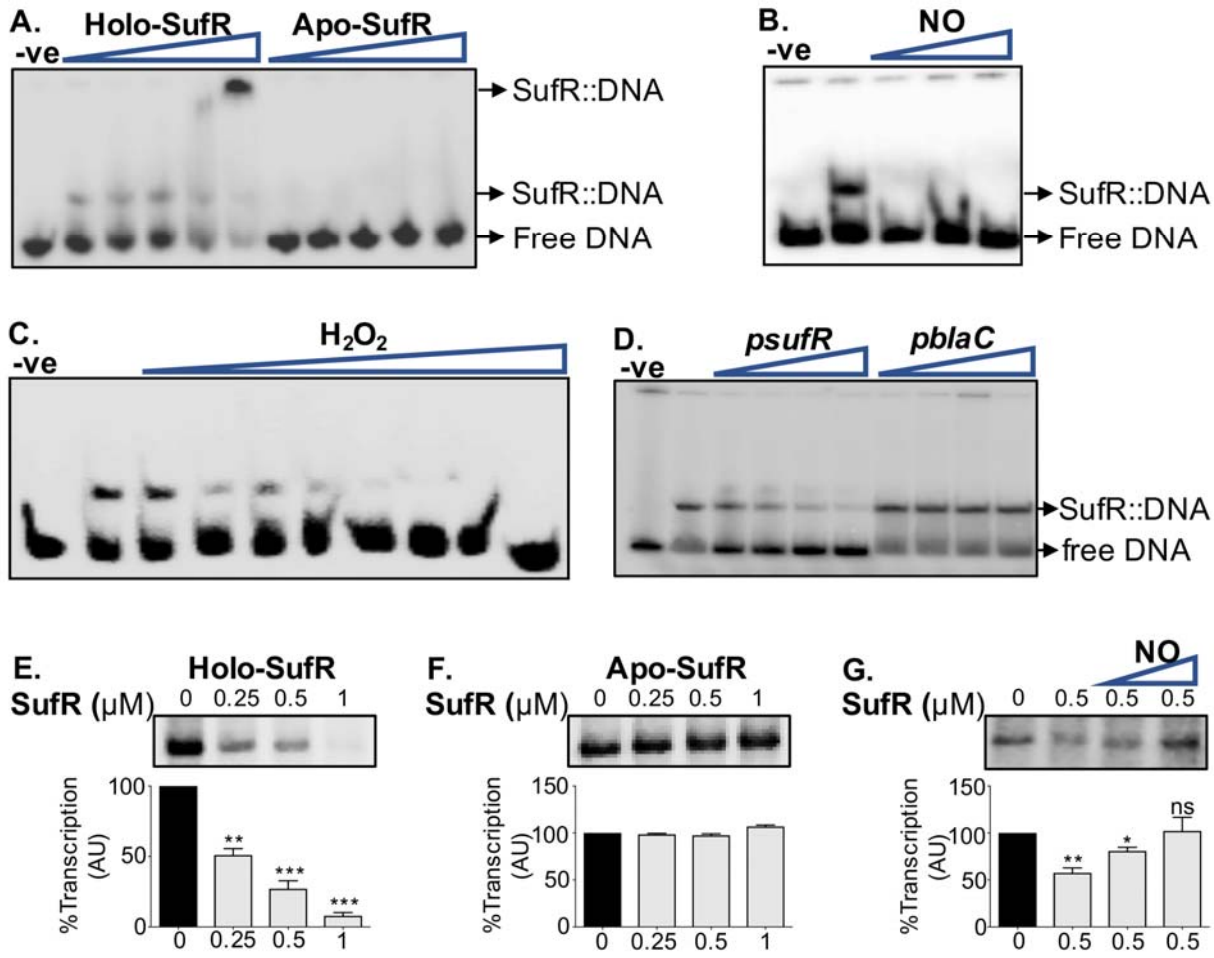
1047

1048

1049

1050

1051



1052

1053 **Figure 2: DNA binding activity of SufR.** (A) Binding of SufR to 32 P-labelled *suf* promoter (0.5 nM). The concentrations of SufR were 0.125 μ M, 0.25 μ M, 0.5 μ M, 1 μ M, and 2 μ M. Holo-1054 SufR (0.5 μ M) was pretreated with (B) proline NONOate (NO; 2.5-, 5-, and 10-fold molar 1055 excess) or (C) H_2O_2 (25, 50, 75, 100, 250, 500, 1000 and 5000 nM) and were incubated with 32 P-1056 labelled *suf* promoter. (D) Holo-SufR (0.5 μ M) was incubated with 32 P-labelled *suf* promoter and 1057 DNA binding was competed out using 12.5-, 25-, 50-, and 100-fold molar excess of unlabelled 1058 competitor promoter of *suf* operon and *blaC*. (E) Effect of SufR on transcription from *suf* 1059 promoter *in vitro*. Increasing concentrations of holo-SufR inhibit transcription. (F) Apo-SufR 1060

1061 was completely ineffective in inhibiting transcription. **(G)** Holo-SufR (0.50 μ M) pre-treated with
1062 proline NONOate (NO; 2.5- and 5-fold molar excess) stimulated transcription from *suf* promoter.
1063 The graph below indicates densitometry analysis of the *suf* transcript (n=2). One-way analysis of
1064 variance (ANOVA) with Bonferroni's post hoc was used to determine statistical significance.
1065 'ns' non-significant '*' p<0.05 '**' p<0.01 '***' p<0.001.

1066

1067

1068

1069

1070

1071

1072

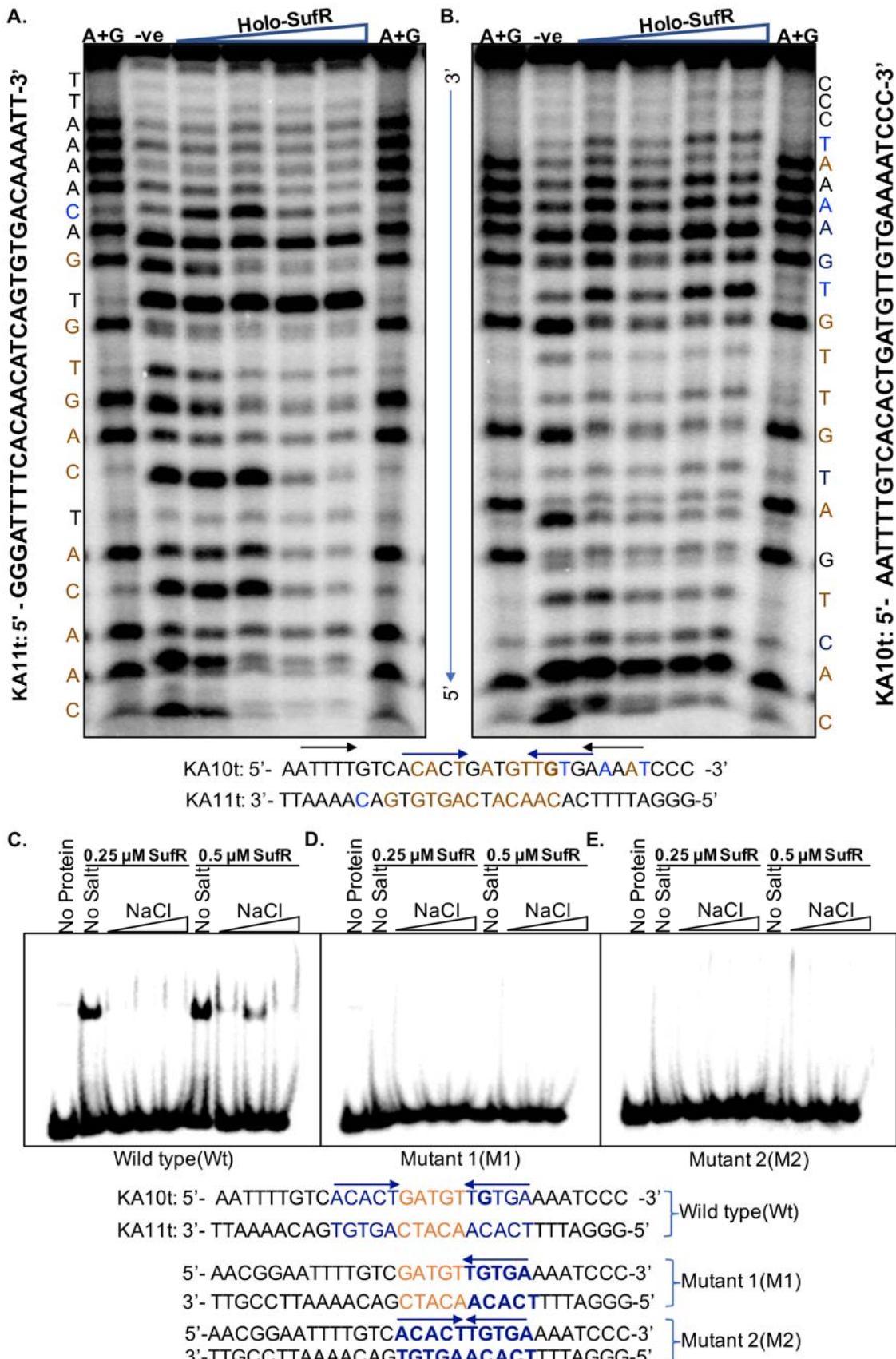
1073

1074

1075

1076

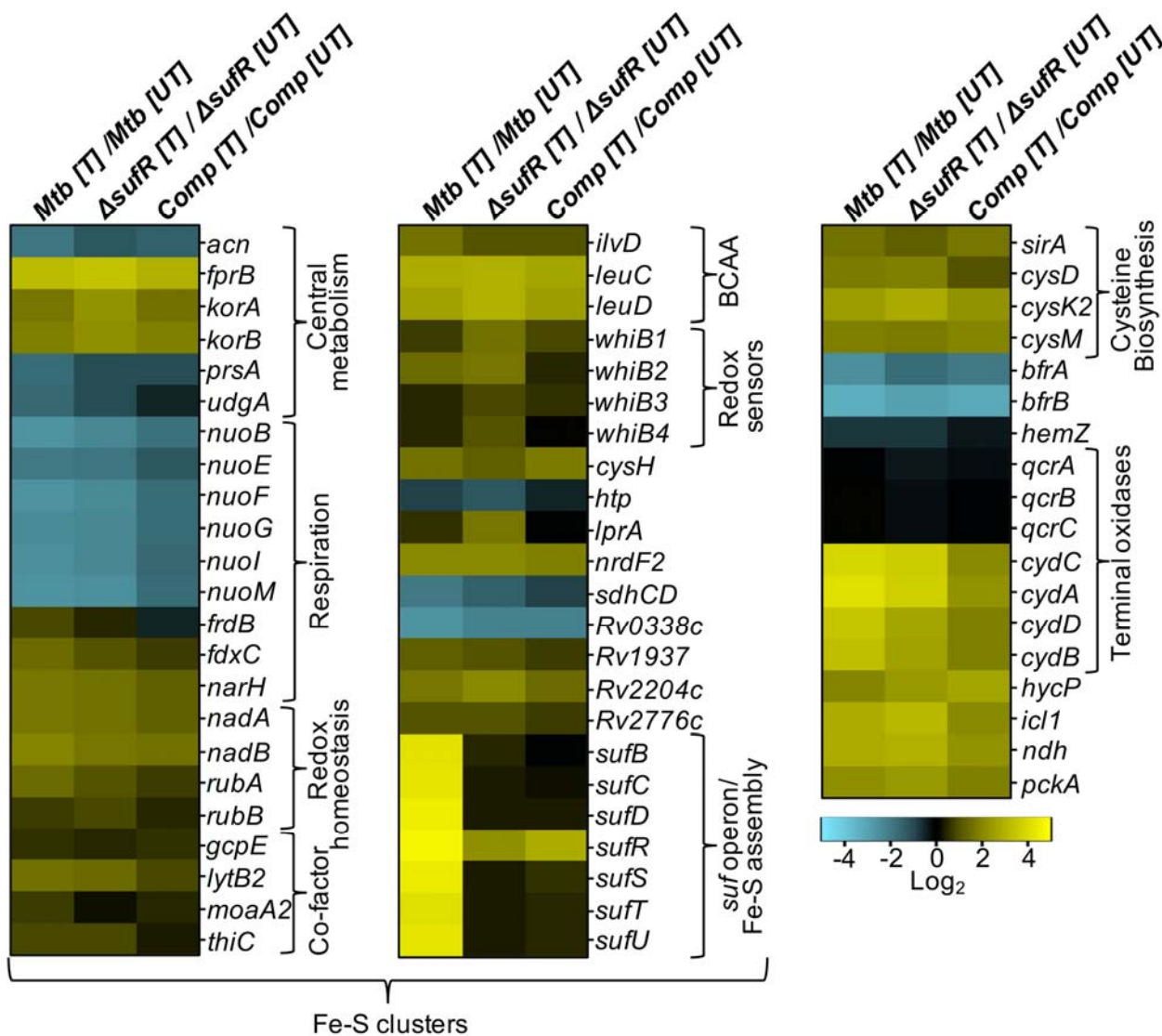
1077



1079 **Figure 3: DNase I footprint of holo-SufR bound to the *suf* promoter.** Sequence upstream of
1080 the *suf* operon (KA10t/KA11t) were used for DNase I footprinting. Bold letter G represents
1081 transcription start site (TSS) for the *suf* operon. **(A)** ^{32}P -KA10t was annealed with cold KA11t.
1082 **(B)** ^{32}P -KA11t was annealed with cold KA10t to examine footprint in the reverse strand. 5
1083 nM ^{32}P -labeled dsDNA was incubated either in absence or presence of holo-SufR (125 nM, 250
1084 nM, 500 nM, and 1 μM) and treated with 0.05 U of DNase I for 2 min. A+G is Maxam and
1085 Gilbert sequencing ladder. DNase I protected/unprotected/hypersensitive nucleotides are marked
1086 with brown/black/blue and two inverted repeats are marked by arrows (blue and black).

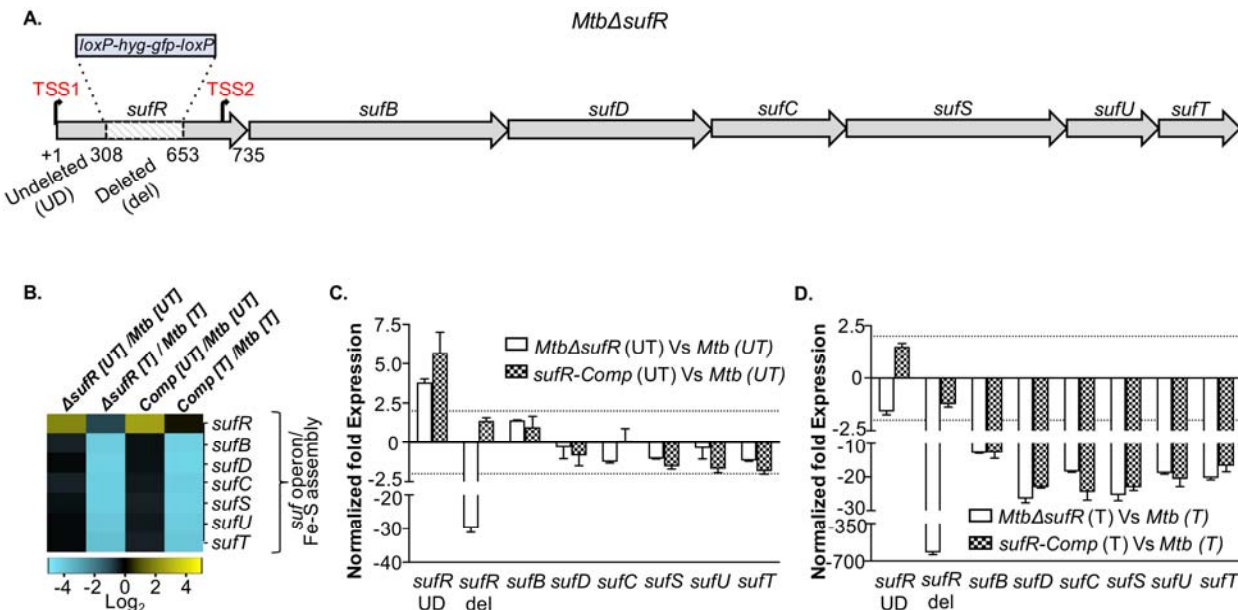
1087 **Mutations in the inverted repeat completely abolished binding of holo-SufR. (C-E)** EMSA
1088 was performed with wild type (KA10t and KA11t) and oligonucleotides with mutations in the
1089 inverted repeats (KA10tm1, KA11tm1, KA10tm2 and KA11tm2) at increasing concentrations of
1090 NaCl at two different concentrations of holo-SufR (0.25 μM and 0.5 μM).

1091



1093 **Figure 4: SufR mainly regulates Suf system involved in Fe-S cluster biogenesis in response**
1094 **to NO.** Total RNA from three biological replicates of untreated (UT) and DETA-NO treated (T)
1095 wt *Mtb*, *MtbΔsufR*, and *sufR-Comp* was isolated and subjected to RNA-seq. Heat maps indicate
1096 log₂ fold change of differentially expressed genes (DEGs) belonging to various functional
1097 categories (obtained from Mycobrowser, EPFL, Lausanne). Genes were considered differentially
1098 expressed on the basis of the false discovery rate (FDR) of ≤ 0.05 and absolute fold change of
1099 ≥ 1.5 . BCCA: branched-chain amino acid.

1100

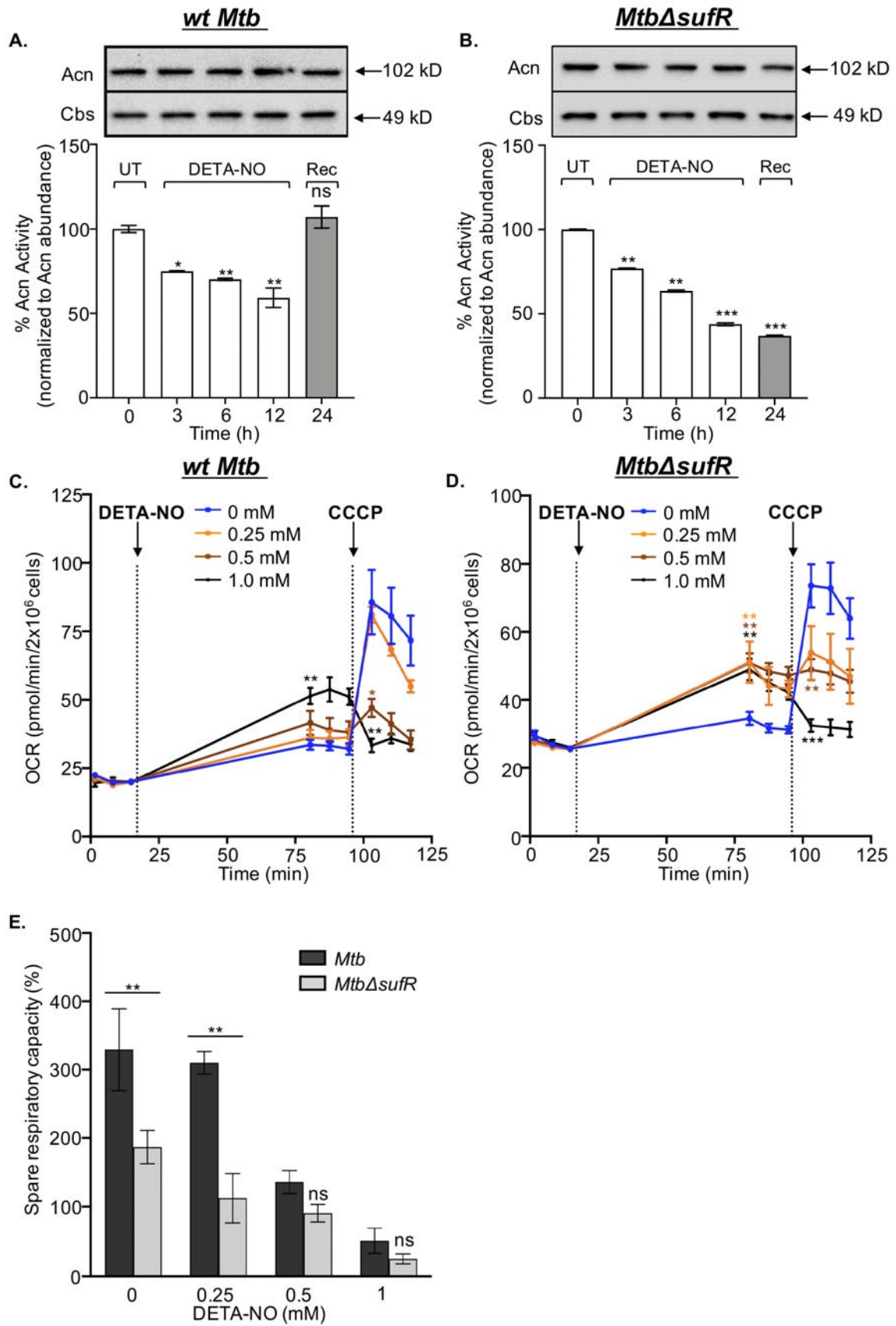


1101

1102 **Figure 5: Deregulation of the *suf* operon in *MtbΔsufR* and *sufR-comp*.** **(A)** Genomic
1103 organization of the *suf* operon upon disruption of *sufR*. The internal fragment of *sufR* ORF from
1104 +308 bp to +653 bp was deleted (del) and replaced by hygromycin resistance cassette (*loxP-hyg*^r
1105 *gfp-loxP*). The *sufR* sequence from +1 to +307 remained undeleted (UD) and resulted in the
1106 expression of an aberrant *sufR* transcript. The two transcription start sites TSS1 (coincides with
1107 the start codon ‘*gtg*’ of *sufR*) and TSS2 (upstream) of *sufB* were marked as per the published
1108 work [19]. **(B)** Exponentially grown cultures of wt *Mtb*, *MtbΔsufR*, and *sufR-comp* were either
1109 left untreated (UT) or treated (T) with 500 μ M DETA-NO for 4h. Total RNA was isolated and
1110 subjected to RNA-seq and qRT-PCR analysis. Heat map depicting expression of the *suf* genes
1111 (\log_2 fold-change ≥ 1 ; FDR ≤ 0.05). Note that *sufR* expression detected in *MtbΔsufR* samples
1112 originated from the reads mapping onto the undeleted (UD) region of *sufR*. **(C, D)** qRT-PCR
1113 data showing the expression of the *suf* genes where *MtbΔsufR* and *sufR-comp* were compared

1114 with untreated and DETA-NO-treated wt *Mtb*. Results are expressed as mean \pm standard
1115 deviation (Mean \pm SD)

1116



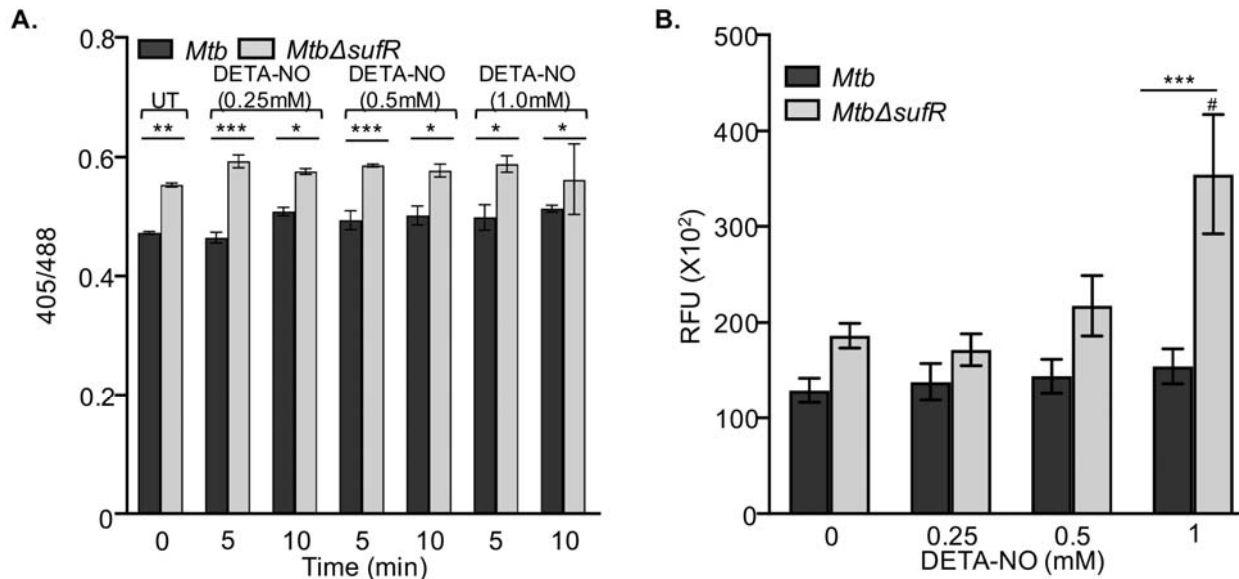
1118 **Figure 6: SufR coordinates Fe-S homeostasis, respiratory reserves, and redox balance in**
1119 **response to NO.** Aconitase (Acn) activity in response to 0.5 mM DETA-NO. **(A)** *wt Mtb* and
1120 **(B)** *MtbΔsufR*. At 12 h post-treatment, strains were cultured in a DETA-NO free 7H9 broth for
1121 24 h and recovery (Rec) of Acn activity was quantified. Intracellular levels of Acn remained
1122 unaltered. A non-Fe-S cluster protein cystathione- β -synthase (Cbs) of *Mtb* was used as the
1123 loading control. A change in oxygen consumption rate (OCR) of **(C)** *wt Mtb* and **(D)** *MtbΔsufR*
1124 was quantified upon addition of indicated concentrations of DETA-NO. The uncoupler CCCP
1125 was used to determine the spare respiratory capacity (SRC). The first and second vertical lines
1126 indicate point of addition of DETA-NO and CCCP, respectively. **(E)** SRC of *wt Mtb* and
1127 *MtbΔsufR* with increasing concentrations of DETA-NO. Percentage SRC was calculated by
1128 subtracting basal OCR (before adding DETA-NO) from CCCP-induced OCR considering basal
1129 OCR as 100%. Statistical significance for the OCR was obtained by comparing OCR upon
1130 treatment with different doses of DETA-NO with untreated (two-tailed, unpaired Student's t-
1131 test.). Comparisons whose P value is <0.05 were indicated with different symbols. Symbols: (*):
1132 comparison to 0.25mM; (*): comparison to 0.5mM; and (*): comparison to 1.0 mM */*/
1133 $p<0.05$; **/**/** $p<0.01$; ***/***/*** $p<0.001$. For SRC, one-way analysis of variance
1134 (ANOVA) with Bonferroni's post hoc was used to determine statistical significance. 'ns' non-
1135 significant '#' $p<0.05$ '##' $p<0.01$.

1136

1137

1138

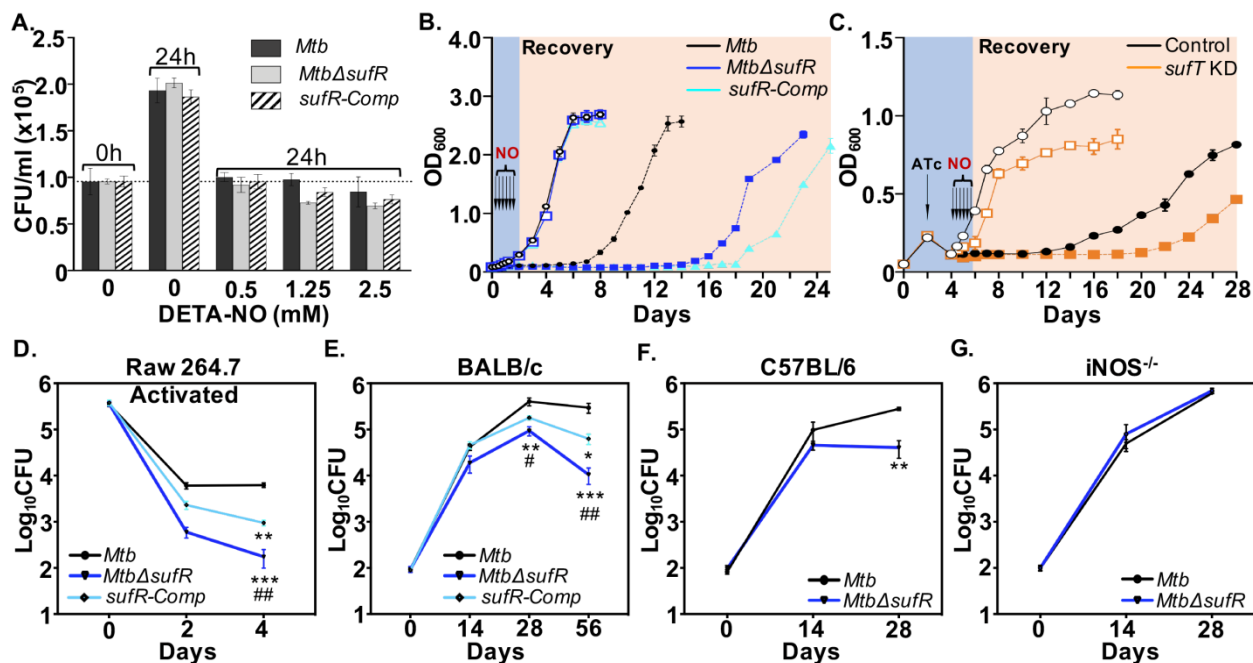
1139



1140

1141 **Figure 7: DETA NO induces oxidative stress in *MtbΔsufR*.** (A) wt *Mtb* and *MtbΔsufR*
1142 expressing Mrx1-roGFP2 were exposed to the indicated concentrations of DETA-NO and the
1143 biosensor ratio (405/488 nm) was measured using flow cytometry. Data shown are the result of
1144 three independent experiments performed in triplicate (mean \pm SD). One-way analysis of
1145 variance (ANOVA) with Bonferroni's post hoc test was employed to determine statistical
1146 significance. (B) wt *Mtb* and *MtbΔsufR* treated with the indicated concentrations of DETA-NO
1147 for 2 h and stained with CellRox Deep Red reagent to measure endogenous ROS. Data shown
1148 are the result of three independent experiments performed in triplicate (mean \pm SD). Two-way
1149 analysis of variance (ANOVA) with Bonferroni's post hoc test was employed to determine
1150 statistical significance between different doses of DETA-NO. Symbols: (*): comparison to *Mtb*,
1151 (#): comparison to untreated *MtbΔsufR*. * p < 0.05, ** p < 0.01, *** p < 0.001.

1152



1153

1154

1155 **Figure 8: Mtb Δ sufR displays defect in recovering from NO-induced growth arrest *in vitro***
 1156 **and NO-dependent persistence in mice. (A)** *Mtb* strains were exposed to DETA-NO for 24 h
 1157 and survival was monitored by enumerating CFUs. **(B)** *Mtb* strains were either left untreated
 1158 (UT; empty symbols) or treated (T; filled symbols) with repeated doses of 0.1 mM of DETA-NO
 1159 for 36 h at an interval of 6 h and growth was monitored over time. **(C)** *Mtb* strains expressing
 1160 CRISPRi vector without (Control) or with *sufT*-specific guide RNA (sgRNA, *sufT* KD) were left
 1161 untreated (UT) or treated (T) with 0.1 mM of DETA-NO at an interval of 6 h for 36 h and growth
 1162 was monitored over time. To deplete SufT, 200 ng/ml of anhydrotetracycline (ATc) was added
 1163 for the induction of *sufT* specific–guide RNA and dCas9 every 48 h. **(D)** RAW264.7
 1164 macrophages activated with IFN γ (100U/mL.) and LPS (100ng/ML.) were infected with *Mtb*
 1165 strains at a multiplicity of infection (moi) of 10 and survival was monitored by enumerating
 1166 CFUs. Data are the results of two independent experiments performed in triplicates. Inbred (E)

1167 BALB/c mice (n=6), (F) C57BL/6 (n=5), and iNOS knockout (iNOS^{-/-}, n=5) were given an
1168 aerosol challenge with *Mtb* strains and assessed for survival in lungs at indicated time points.
1169 Results are expressed as Mean±SD. Two-way analysis of variance (ANOVA) with Bonferroni's
1170 post hoc test was employed to determine statistical significance for the intramacrophage and
1171 pulmonic load between different strains. p value is <0.05 were indicated with symbols. Symbols:
1172 (*): comparison to wt *Mtb*; (#): comparison to *sufR-Comp*. */# p<0.05; **/## p<0.01; ***
1173 p<0.001. In fig B and C open and close symbol are untreated and DETA-NO treated respectively.

1174

1175

1176

1177

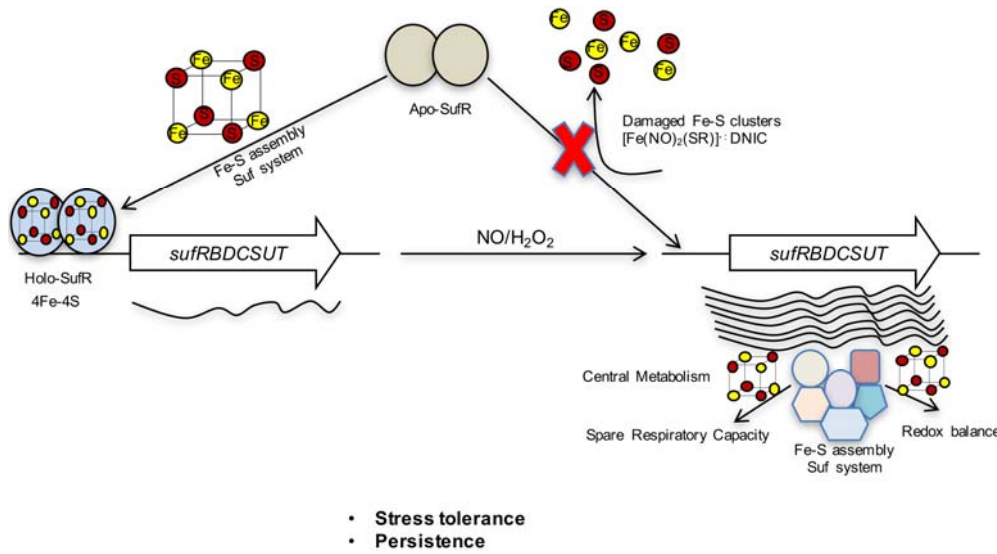
1178

1179

1180

1181

1182



1183

1184 **Figure 9: A model for SufR-mediated regulation of Fe-S cluster biogenesis in *Mtb*.** SufR
1185 appears to coordinate the expression of the *suf* operon in a negative feedback loop. Under Fe-
1186 sufficient conditions, the Fe-S cluster bound form of SufR (holo-SufR) binds to the promoter of
1187 the *suf* operon and represses Fe-S cluster biogenesis. Exposure to NO or H₂O₂ damages Fe-S
1188 clusters to generate DNIC or apo-forms of SufR, respectively, which abrogates SufR DNA-
1189 binding resulting in de-repression of the *suf* operon. The elevated expression of the SUF Fe-S
1190 assembly system regenerates Fe-S clusters of metabolic enzymes, respiratory complexes, and
1191 redox sensors to maintain Fe-S cluster homeostasis, spare respiratory capacity, and redox
1192 balance. The NO-sensing properties of SufR via its 4Fe-4S cluster and persistence defect
1193 exhibited by *MtbΔsufR* in mice in an iNOS-dependent manner suggest that SufR integrates host
1194 environmental signals to Fe-S cluster homeostasis in *Mtb*.

1195