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1	Real-time kinetic studies of Mycobacterium tuberculosis LexA-DNA interaction
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16 Transcriptional repressor, LexA, regulates the "SOS" response, an indispensable bacterial 17 DNA damage repair machinery. Compared to its *E.coli* ortholog, LexA from 18 Mycobacterium tuberculosis (Mtb) possesses a unique N-terminal extension of additional 24 19 amino acids in its DNA binding domain (DBD) and 18 amino acids insertion at its hinge 20 region that connects the DBD to the C-terminal dimerization/autoproteolysis domain. Despite 21 the importance of LexA in "SOS" regulation, Mtb LexA remains poorly characterized and the 22 functional importance of its additional amino acids remained elusive. In addition, the lack of 23 data on kinetic parameters of Mtb LexA-DNA interaction prompted us to perform kinetic 24 analyses of Mtb LexA and its deletion variants using Bio-layer Interferometry (BLI). Mtb 25 LexA is seen to bind to different "SOS" boxes, DNA sequences present in the operator 26 regions of damage-inducible genes, with comparable nanomolar affinity. Deletion of 18 27 amino acids from the linker region is found to affect DNA binding unlike the deletion of the 28 N-terminal stretch of extra 24 amino acids. The conserved RKG motif has been found to be 29 critical for DNA binding. Overall, this study provides insights into the kinetics of the 30 interaction between Mtb LexA and its target "SOS" boxes. The kinetic parameters obtained 31 for DNA binding of Mtb LexA would be instrumental to clearly understand the mechanism of 32 "SOS" regulation and activation in Mtb.

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40 The expression of DNA damage and stress response genes, which serve to preserve genome 41 integrity upon exposure to DNA damaging agents, is controlled by the "SOS" response 42 pathway. Activation of the "SOS" response helps the bacteria to develop resistance to 43 antibiotics, making it indispensable for survival and growth under adverse conditions (1-3). 44 This pathway is regulated by two key players, namely, RecA and LexA. LexA binds to a 45 consensus sequence of DNA known as the "SOS" box located in the operator region of 46 several genes and transcriptionally represses them under normal physiological conditions. 47 However, under stress conditions, LexA falls off from the operators leading to activation of these genes to facilitate DNA repair (4). The "SOS" regulons exhibit significant variations 48 49 across the bacterial kingdom, reflecting their overall complexity. For example, 50 while Bacillus subtilis harbors only 33 genes in its "SOS" regulon, E.coli contains over 45 51 genes (5). Most of the "SOS" regulons include genes that encode for error-prone DNA 52 polymerases, LexA, RecA, and proteins involved in the nucleotide excision repair pathway, 53 although exceptions are known to exist (6).

54 "SOS" activation occurs in the following sequence of events— (i) RecA interacts 55 with single-stranded DNA to form activated nucleoprotein filament complex, (ii) activated 56 RecA directly interacts with LexA leading to autoproteolytic cleavage of the latter, and 57 finally (iii) LexA falls off from the operator regions causing transcriptional de-repression of 58 the damage-inducible genes (7). In *E. coli*, the LexA repressor binds to consensus "SOS" box 59 sequence present in different operators, with variable affinity (7). Genes with lower operator-60 repressor affinity are activated early on in the "SOS" pathway when compared to genes 61 having tightly bound operators. For instance, genes such as lexA, uvrA, uvrB, uvrD, and 62 recA express early on, while those encoding for error-prone polymerases, (dnaE2, umuD) and

*sulA* express much later in the cascade (7). Altogether, the differential binding of LexA to
different operators results in a highly complex but well-coordinated process that protects the
cellular machinery during DNA damage.

66 LexA homologs are widespread across bacterial genomes and have remained 67 evolutionarily conserved (8). While most of the DNA damage-inducible genes in E.coli are 68 under the direct control of LexA/RecA, such is not in the case of 69 Mycobacterium tuberculosis (Mtb), wherein several DNA damage-inducible genes are 70 independent of LexA regulation (9). Among the 21 genes regulated by LexA/RecA in Mtb, 71 only a few have been characterized, limiting our present understanding of the molecular 72 mechanisms underlying the regulatory role of LexA. The functional characterization of 73 LexA, including estimation of kinetic parameters obtained from binding to different "SOS" 74 boxes, will provide new insights in mechanistically understanding mycobacterial "SOS" 75 regulation.

76 Well-studied LexA homologs from E.coli and B.subtilis are known to exhibit 77 nanomolar binding affinities towards "SOS" box DNA with an apparent K<sub>D</sub> of 0.8 nM and 78 2.3 nM respectively (10, 11). However, a lack of kinetic analyses for Mtb LexA-DNA 79 interaction greatly limits our understanding of mycobacterial "SOS" activation. Nearly 25 in-80 binding sites of LexA were identified in the Mtb genome by Davis and colleagues in 2012 81 (12). The study demonstrates differential expression of the damage-inducible genes upon 82 Mitomycin C-induced stress (12). Precise quantitation of such molecular interactions of Mtb 83 LexA with DNA could reveal the importance of structural integrity and the mechanism of 84 "SOS" induction. To address this, it is imperative to study the kinetics of the interaction of 85 LexA with its target binding sites and thus we carried out real-time Mtb LexA-DNA 86 interaction studies.

87 Although the C-terminal domain (CTD) crystal structure of Mtb LexA provides a 88 molecular framework for its architecture and assembly, the DNA binding domain (DBD) 89 remained inaccessible for structural studies (13). Mtb LexA has an N-terminal DBD and a C-90 terminal dimerization/catalytic domain separated by a linker region similar to its orthologs 91 from other bacteria. Notably, sequence comparison between Mtb and E.coli LexA revealed a 92 unique N-terminal extension of 24 amino acids, which came into light after the re-annotation 93 of Mtb LexA by Smollett et al., in 2009 (14), and a longer linker region with additional 18 94 amino acids in Mtb LexA. Further, the functional importance of these extra stretches of 95 amino acids in and around the DBD and linker region of Mtb LexA remains unclear. 96 Therefore, in this study, we investigated whether these unique regions play any role in DNA 97 binding. We have generated Mtb LexA mutants that lack 24 amino acids from its N-terminal 98 domain (NTD), 18 amino acids from its linker region, in addition to mutating the conserved 99 DNA binding residues "RKG" to "AAA". We have compared the DNA binding property of 100 these mutants with wild-type Mtb LexA qualitatively using EMSA and quantitatively by Bio-101 layer Interferometry (BLI) to obtain real-time binding kinetic parameters. The DNA binding 102 assessed using BLI revealed a comparable binding affinity of wild-type Mtb LexA and its N-103 terminal 24 amino acids truncation variant. Deletion of 18-amino acids from the linker 104 resulted in 16 times reduced binding, whereas RKG mutant showed no binding, suggesting 105 their importance in DNA binding. The kinetic parameters of Mtb LexA-DNA interaction 106 obtained from this study provide new functional insights crucial for an understanding of 107 mycobacterial "SOS" regulation.

#### **109** Materials and Methods

#### 110 Plasmids, strains, and reagents

Bacterial strains and plasmids used in the study are listed in Table S1 of Supplementary Material. Primers and oligonucleotides were purchased from Sigma Aldrich and are listed along with the constructs generated in the study in Table S2 of Supplementary Material. All the reagents, media, and chemicals were purchased from Sigma Aldrich, Hi-Media, and SRL, and enzymes were purchased from New England Biolabs.

#### 116 Construction and analysis of phylogenetic tree

Multiple sequence alignment of LexA proteins from diverse bacterial groups was created using Clustal X and the phylogenetic tree was constructed using RaxML (15). The evolutionary history of the taxa has been inferred from 1000 replicates of the bootstrap consensus tree. The percentage of replicates in which clustering of related taxa took place is mentioned next to the branches. 283 positions were found in the final dataset. iTOL was used to visualize the generated tree (16).

# 123 Over-expression and purification of proteins

The Mtb LexA and its Δ24aa variant were individually cloned between NdeI and BamHI
restriction sites, respectively in pET28a(+) bearing N-terminal 6x His tag. Mtb LexAΔ18aa
and Mtb LexA RKG/AAA mutants were generated by the overlap PCR method. The primers
used have been listed in Table S2 of Supplementary Material. All constructs generated
have been confirmed by sequencing.

The recombinant WT and its variant proteins were over-expressed in *E.coli* BL21
(DE3) cells. Cultures were induced with 0.5 mM IPTG at O.D<sub>600</sub> 0.6. The cells were pelleted

131 after 4 hrs by centrifugation at 5000×g following which the pellet was resuspended in Lysis 132 Buffer composed of 50 mM Tris-Cl (pH 8.0), 150 mM NaCl and 10 mM imidazole (pH 8.0), 133 5% glycerol, 1mM Phenylmethylsulfonyl fluoride (PMSF). After lysing the cells by 134 sonication on ice (10" On, 30" Off cycles), the clarified lysate was centrifuged for 1h at 135 20,000g. The clarified supernatant was passed through a pre-equilibrated HisTrap HP column 136 at 5ml/min. Column was washed with 50 mM Tris-Cl, 500 mM NaCl, 30 mM imidazole (pH 137 8.0) and proteins were subsequently eluted under a gradient in 25 mM Tris-Cl (pH 8.0), 150 138 mM NaCl and 750 mM imidazole (pH 8.0). Pure fractions were pooled, diluted with a low 139 salt buffer (25 mM Tris-Cl (pH 8), 100 mM NaCl, 1 mM EDTA, 5% glycerol), and loaded 140 onto Q-Sepharose column for anion exchange chromatography. Proteins were eluted in 141 gradient by passing a high salt buffer (25 mM Tris-Cl (pH 8.0), 1 M NaCl, and 1 mM 142 EDTA). The pure fractions were concentrated using Gel Filtration Buffer (20 mM Tris-Cl 143 (pH 8.0), 100 mM NaCl and 5% glycerol) and separated using Superdex 75 10/300 GL for 144 gel filtration. The purified proteins were run on 12% SDS-PAGE for analyzing their purity 145 and concentrations were determined via spectrophotometric analysis.

# 146 Cross-linking reactions

147 The cross-linking reactions were performed by incubation of each of the proteins at 5  $\mu$ M 148 final concentration in presence of 0.01% v/v glutaraldehyde in 10 mM HEPES (pH 8.0), 50 149 mM NaCl for 30 mins on ice. The reactions were stopped with 25 mM of DTT. Samples were 150 separated on a 12% SDS-PAGE.

- 151 Circular dichroism (CD)
- 152 CD spectra were recorded from 195-280 nm using a Jasco J-815 spectropolarimeter. A 1 mm
  153 pathlength quartz cuvette was used. Resolution up to 0.2 nm was maintained with a scan rate

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of 100 nm/min. 25°C was maintained for all experiments. 5 μM of each protein in 10 mM
Tris-Cl 50 mM NaCl (pH 7.5) was taken for analysis. The data presented is an average of
three scans after correction for the buffer baseline. Recorded spectra were analyzed using
Origin 8.1 software.

# 158 Extrinsic Fluorescence

Extrinsic fluorescence spectra were obtained using a Jobin-Yvon Fluorometer FluoroMax3, at 25°C. 5  $\mu$ M of each of the proteins in 10 mM Tris-Cl (pH 7.5), 50 mM NaCl were incubated with 44 mer ds *dnaE2* "SOS" box containing DNA (sequence given in **Table 1**) at 1:2 ratio for 30 mins at 37°C. The samples were incubated with 40  $\mu$ M of ANS in dark for 10 mins. Samples were excited at 350 nm, and in the range of 400 to 600 nm, emission spectra were recorded. Measurements were corrected for fluorescence intensity of buffer, DNA, and ANS intrinsic fluorescence.

#### 166 Electrophoretic mobility shift assay

WT and its variants at 128 nM were incubated in presence of 3.5 nM of end-labeled (nonbiotinylated) 44 mer ds *dnaE2* "SOS" box containing DNA (sequence given in **Table 1**) in 10 mM HEPES (pH 7.5), 50 mM NaCl for 30 mins on ice. The unbound DNA and DNA-protein complexes were resolved on 8% native PAGE at 100V for 1 h in cold. Gels were dried and autoradiographed. The same procedure was followed for EMSA analysis in which increasing concentrations (0-128 nM) of WT Mtb LexA was incubated with <sup>32</sup>P end-labeled ds 44 mer ds *dnaE2* "SOS" box containing DNA.

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177 The ForteBio Octet RED 96 (Forte Bio, USA) platform was used to conduct interaction 178 studies between LexA and its variants with biotinylated ds 44mer of different "SOS" boxes 179 containing sequences (listed in Table 1). Streptavidin matrix-coated sensor chip (SA) was 180 equilibrated in 10 mM HEPES (pH 7.5), 50 mM NaCl followed by immobilization of 100 nM 181 of biotinylated ds DNA on it. Increasing concentrations of WT and mutant proteins were 182 passed onto the chip and change in response units (RU) was analyzed. The program 183 comprises one-minute stabilization of the baseline with the buffer followed by ten-minute loading of sensors with biotinylated DNA, a five-minute association enabling interaction 184 185 between the protein and DNA, a five-minute dissociation step finally followed by a five-186 second regeneration step (unless mentioned otherwise). A reference sensor dipped in the 187 buffer was used as a background control. All analyses were carried out at 25°C. A 1:1 188 binding model was applied to globally fit the binding isotherms and kinetic parameters such as kon, koff, and K<sub>D</sub> were obtained. The experiments were performed in triplicates. 189

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### 191 Results and Discussion

192

# 193 Mtb LexA features distinct characteristics from its counterparts

LexA is present in most bacterial species and phyla (17). Evolutionarily, the protein has retained its two distinct domains, the NTD, involved in DNA binding, and CTD, which is responsible for dimerization and autoproteolytic cleavage. Alpha helices involved in DNA binding and the residues critical for autoproteolysis have remained well conserved across different species, thereby preserving the overall functions of the protein. 199 From an evolutionary viewpoint, a comparison between LexA homologs from 200 selected representatives belonging to major classes of Gram-positive, Gram-negative, 201 Archaebacterial, and Actinobacterial phyla reveals the discrete clustering based on their 202 classification, evident from the phylogenetic tree constructed (Figure 1A). This tree has been 203 deduced by comparing sequences from 24 representative bacterial species (shown 204 in Figure S1 of Supplementary Material). LexA homologs from Actinobacteria closely 205 resemble homologs from Gram-positive Firmicutes which is consistent with their relatedness 206 at the species level. Interestingly, LexA from members of Actinobacteria that include the 207 pathogenic tuberculous mycobacteria such as Mtb, M.canetti, M.bovis, shows significant 208 similarity, suggesting a possible link between pathogenicity and sequence evolution of 209 LexA.

210 Subsequent comparison of the LexA sequences among some of the well-known 211 tuberculous and non-tuberculous mycobacteria revealed interesting results. Although the C-212 terminal regions remain almost identical, N-terminal regions exhibit sequence variations 213 especially at the terminal end. Pathogenic mycobacteria have relatively smaller genome sizes 214 (18) and are not expected to code for unwanted additional stretches of amino acids in their 215 proteins unless they prove advantageous for their survival. Interestingly, tuberculous 216 mycobacteria except for M.bovis were found to possess additional amino acids at the N-217 terminal end of LexA, unlike their E.coli counterpart, implying unexplored adaptive 218 functions. Another region of less conservation spans the linker region that connects the NTD 219 and CTD of the protein. While the latter half of the linker sequences (towards the CTD) 220 exhibits more conservation, the initial half varies among different mycobacterial species. 221 Although the linker is identical between Mtb and *M.bovis* harboring 25 amino acids, the 222 number of residues and sequence conservation vary among the other mycobacteria (Figure 223 **1B**). In *M.leprae* and *M.haemophilum*, the linker can extend up to 28 amino acids long. LexA

224 possessing additional stretches of amino acids triggers the curiosity to explore their 225 mechanistic roles in "SOS" induction that could help mycobacterial survival and evolution.

226 Presumably, these stretches of amino acids in Mtb LexA that remain uncharacterized 227 may confer additional functions unique to mycobacterial species when compared to its well-228 characterized homologs. LexA is a global repressor controlling the expression of DNA repair 229 genes. Hence, assessing whether these additional stretches of amino acids may influence 230 interactions with "SOS" boxes could provide new information related to "SOS" regulation. 231 Considering both the additional stretches of amino acids to lie in proximity to the DNA 232 binding regions of the protein, we deleted these stretches to explore their impact on DNA 233 binding. Subsequently, the stretch of 24 amino acids (residues 1-24) was deleted to generate 234 LexA $\Delta$ 24aa and LexA $\Delta$ 18aa was generated by deleting 18 amino acids spanning the hinge 235 region (residues 94-111) of the protein. This long hinge region separates the NTD from its 236 CTD in Mtb LexA. This is in sharp contrast to the much shorter hinge region of just 4 amino 237 acids (O70-E74) present in *E.coli* (Figure 1B). The functional relevance of such a long hinge 238 region in Mtb has not been explained. Possibly, the longer length of this inter-domain linker 239 in Mtb LexA can enhance its flexibility to attain suitable conformations for binding to DNA 240 (13). Next, we have attempted to shed light on this aspect in the present study by 241 characterizing and comparing the DNA binding ability of the variants with that of WT LexA. 242 RKG motif involved in DNA binding contains Arg 52, Lys 53, and Gly 54 in E.coli LexA 243 lying in the third alpha helix of the protein, (19) and its corresponding Arg 75, Lys 76 and 244 Gly 77 residues of Mtb LexA have remained conserved. We have mutated these residues to 245 assess whether they are essential for DNA binding in Mtb LexA.

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All proteins were purified to ~98% purity (Figure 1C). Near and far UV spectra of Mtb LexA and its mutants were used to assess changes in their secondary structures due to mutations (Figure 2A). Interestingly, both far (195-250 nm) and near (250-280 nm) UV spectra from Circular Dichroism (CD) studies revealed the comparable secondary structures of WT Mtb LexA and its mutants.

254 All the variants predominantly exist as dimers in solution as analyzed from profiles of 255 gel filtration chromatography (Figure S2 of Supplementary Material). To confirm this 256 further, the purified proteins were subjected to cross-linking using the chemical cross-linker 257 glutaraldehyde. Upon cross-linking, the predominant form appeared to be dimeric in all cases 258 as evident from the top band running between 48 and 63kDa (Figure 2B). Structural analysis 259 of the C-terminal segment of Mtb LexA revealed residues 229-236 along with residues 139-260 153 from its NTD to be involved in dimer formation between two LexA monomers (13). The 261 mutants generated in this study have no overlapping sequences with the afore-mentioned 262 residues and therefore, all of them retained the ability to form dimers. Therefore, a 263 comparable profile of results in CD spectroscopy and protein dimerization experiments 264 revealed that the selected regions had minimal or no influence on the secondary structure of 265 Mtb LexA.

Next, we used fluorescence spectroscopy to assess the structural changes of Mtb LexA and its variants upon interaction with DNA. Hydrophobic extrinsic fluorescent dye 8anilino-1-naphthalenesulfonic acid (ANS) has been widely used for screening the alterations in the tertiary structure of proteins and to monitor their ligand-binding events such as proteinnucleic acid interactions (20). The ligand displaces the fluorescent dye upon binding to the protein, resulting in a quench in fluorescence. Changes in fluorescence intensity are a direct

272	readout of protein-DNA binding. The maximum quench in fluorescence intensity was noted
273	when WT LexA formed a complex with DNA (Figure 2C). A similar quench in fluorescence
274	intensity was observed for LexA $\Delta$ 24aa and the WT protein upon DNA binding (Figure 2F),
275	suggesting that the 24 amino acids extension is not crucial for DNA binding. However, we
276	cannot overlook the possibility that it may have a regulatory role in DNA binding which
277	could be dependent on the "SOS" box sequences (21). In striking contrast to LexA $\Delta$ 24aa,
278	deleting 18 amino acids from the linker region has significantly reduced the fluorescence
279	quenching to nearly half compared to that for the WT, suggesting that this linker region may
280	play a prominent role in DNA binding (Figure 2E). The deletion of 18-amino acids had weak
281	or no effect on the secondary structures; however, the observed reduction in nucleic acid
282	binding affinity may be influenced by the Van der Waals interactions offered by either
283	glutamic acid or aspartic acid residues (4 out of 18 amino acid residues) within the linker
284	region. Lastly, RKG/AAA mutant showed no significant fluorescence quench upon
285	interaction with DNA (Figure 2D), thereby establishing that RKG residues play a critical role
286	in DNA binding in Mtb LexA similar to the other orthologs. Further, we have carried out
287	electrophoretic mobility shift assays (EMSA) to see the DNA binding of WT LexA to the
288	dnaE2 "SOS" box. The shift in the <sup>32</sup> P labelled $dnaE2$ "SOS" box DNA suggests that Mtb
289	LexA can bind to "SOS" box with nM affinity (Figure 3A). We have also performed EMSA
290	with Mtb LexA variants. EMSA analysis revealed that LexA $\Delta$ 24aa and LexA $\Delta$ 18aa showed
291	mobility shift, whereas RKG/AAA mutant did not show any DNA binding (Figure 3B). This
292	further confirms the importance of the RKG motif in DNA binding. Based on these
293	observations, we further decided to quantitate the real-time kinetic parameters of LexA-DNA
294	interactions with WT LexA and its variants.

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We used BLI for determining the DNA binding affinity for Mtb LexA and its mutants. First, we characterized the interaction between WT Mtb LexA and its variants to the perfectly palindromic *dnaE2* "SOS" box ("SOS" box sequence given in **Table 1**). WT and its variant, LexA $\Delta$ 24aa, exhibit comparable DNA binding affinity (K<sub>D</sub> of 2.16±0.01 nM for WT and 4.94±0.03 nM for LexA $\Delta$ 24aa, respectively). Further, LexA $\Delta$ 24aa with and without Nterminal 6x His tag showed comparable DNA binding kinetics (**Figure S3**).

303 LexA18aa $\Delta$  variant shows significantly reduced affinity as seen from the sensograms 304 depicted in Figure 4. Deleting 18 amino acids from the linker connecting NTD and CTD 305 resulted in a more than fifteen times reduction in affinity when compared to the full-length 306 protein, with an obtained K<sub>D</sub> of 34.4±0.19 nM (Table 2). The rate of association (k<sub>on</sub>) was highest in the order of  $10^6$  for full-length Mtb LexA and decreased to the order of  $10^4$  in the 307 308 case of the  $\Delta 18aa$  variant. While the association rate constants varied, the dissociation rate 309 constants (koff) did not change significantly. We speculate that deleting the 18 amino acids 310 from the linker affected the conformation to bind DNA suitably, thereby reducing its 311 association rate (Figure 4B, Table 2). We did not observe any detectable binding for 312 RKG/AAA, even after doubling the time of interaction (from 300 to 600 seconds of 313 association) (Figure 4G), indicating the RKG motif in DBDs is critical for the DNA binding. 314 Altogether, Mtb LexA exhibits similarity to *E.coli* counterpart in terms of binding to its 315 cognate "SOS" box with nanomolar affinity. Additionally, the presence of the longer linker in 316 Mtb LexA is found to positively affect its DNA binding ability. The kinetic parameters 317 obtained by performing experiments at physiological pH intrigued us to investigate how they 318 would vary when subjected to acidic conditions. The rationale behind choosing a highly 319 acidic pH condition to monitor changes in DNA binding affinity of Mtb LexA arises from the

fact that Mtb is known to face a hostile environment of acidic pH inside host macrophages and is challenged to maintain internal pH homeostasis for survival (22). Although internal pH lower than 6 is noted to be lethal for mycobacteria, it has to endure external pH as low as 4 (22). We wanted to assess whether Mtb LexA and its variants could exhibit DNA binding even at an extreme pH such as pH 4.

325 We found that Mtb LexA exhibited maximal and optimal binding with the perfectly palindromic dnaE2 "SOS" box near physiological pH (pH 7.5). Surprisingly, Mtb LexA 326 327 retained the ability to bind DNA even at pH 4, although with reduced affinity (Figure 5). The 328 dissociation constants changed from  $2.16\pm0.01$ ,  $4.94\pm0.03$  and  $34.4\pm0.19$  nM (at pH 7.5) to 329  $75.99\pm0.27$ ,  $29.94\pm0.13$  and  $111\pm0.33$  nM (at pH 4) for WT, LexA $\Delta$ 24aa and LexA $\Delta$ 18aa, 330 respectively. Binding at acidic pH (pH 4) can thus be said to have reduced by nearly 35 times 331 for the wild-type protein, nearly 6 times for LexA $\Delta$ 24aa, and nearly 3.2 times for LexA $\Delta$ 18aa 332 when compared to physiological conditions (Table 2). From the kinetic parameters observed, we notice a significant reduction in association rate constants (kon) at low pH (pH 4) 333 334 compared to physiological conditions (pH 7.5) for wild-type protein and its  $\Delta 24aa$  variant 335 while the change is not so pronounced in the case of the  $\Delta 18aa$  variant. Dissociation rate 336 constants (k<sub>off</sub>) in the case of WT Mtb LexA remained comparable in both the pH conditions 337 tested but reduced by a power of 10 for the mutant proteins at low pH. Mutants seem to both 338 associate and dissociate faster at physiological pH as compared to low pH and as for the wild-339 type protein, there has been a drastic reduction only in its rate of association to bind DNA at 340 low pH conditions.

Our observations results corroborate with those made from *in-vitro* studies that have assessed the effect of variations in pH in regulating the "SOS" response for *E.coli* LexA (23-25). Relan and co-workers reported that *E.coli* LexA bound to its operator maximally near Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BCJ20210434/924128/bsr-2021-14194.pdf by guest on 16 April 2024

# 347 DNA binding kinetics of Mtb LexA with different "SOS" boxes

348 The differential gene expression profile following DNA damage led to the identification of 349 genes that fall under direct regulation of LexA in Mtb (21). However, DNA binding kinetics 350 of LexA to "SOS" boxes of these DNA damage-inducible genes remained uncharacterized. 351 We, therefore, determined the DNA binding affinity for LexA and its mutants to different 352 mycobacterial "SOS" boxes (the kinetic parameters determined for the interaction of mutants 353 to different "SOS" boxes are provided in Figure S4 and Table S3 of Supplementary 354 Material). The "SOS" boxes chosen have unique characteristics (Table 1). While the *dnaE2* 355 "SOS" box is a perfect palindrome throughout (as mentioned in the previous section), lexA 356 and recA "SOS" boxes have one mismatch towards their 3'ends (on the flank). rv3074 "SOS" 357 box is unique in displaying a perfect palindrome of sequences on either side repeat flanks but 358 showing mismatches in sequences between the flanks. All the genes whose "SOS" boxes have been chosen for the present study are highly induced following DNA damage in Mtb 359 360 (21).

Increasing concentrations of WT LexA and its variants (analytes) were allowed to interact with biotinylated "SOS" boxes till saturation in binding was achieved. WT LexA was found to bind to different "SOS" boxes with close affinities (**Figure 6, Table 3**).  $K_D$  values ranged from 0.98±0.01 nM for the *lexA* "SOS" box to 3.86±0.03 nM as noted for the *rv3074* "SOS" box. The association rate was relatively higher for *dnaE2* and *lexA* "SOS" boxes as compared to the other two "SOS" boxes. The perfectly palindromic nature of the dnaE2 "SOS" box facilitates faster association with LexA. *dnaE2* encodes an error-prone DNA Although the association rate constants  $(k_{on})$  varied for different "SOS" boxes, the corresponding dissociation rate constants  $(k_{off})$  are also seen to change proportionately; hence, the overall  $K_D$  is not widely altered for different "SOS" boxes tested here. Though, no drastic changes in binding affinity to different "SOS" boxes are noted here, studying these repressor-DNA binding events extensively in the cellular context can reveal additional modes of regulation or factors modulating the expression patterns of "SOS" responsive genes at the time of "SOS" activation.

### 379 Conclusion

Mtb LexA controls gene expression patterns of the crucial "SOS" response pathway that facilitates mycobacterial adaptation to stress (12). However, lack of thorough understanding at the molecular level, taking into account the unique regions of Mtb LexA that could potentially influence its interaction with DNA, prompted us to execute this study by analyzing the impact of such truncations/ mutations on Mtb LexA-DNA interaction. Together, we present our detailed analysis of Mtb LexA and the role of its additional stretches of amino acids in regulating the "SOS" response.

To begin with, the deletion and mutated variants displayed comparable secondary structure as that of wild-type Mtb LexA protein, inferred from the ellipticity measurements carried out using circular dichroism. Moreover, they retained the ability to form dimers as observed from size exclusion chromatography and crosslinking studies. Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BCJ20210434/924128/bsr-2021-14194.pdf by guest on 16 April 2024

391 Qualitative estimation and comparative analysis of protein-nucleic acid interaction of 392 the variants compared to the WT revealed that while the 24-amino acid extension at the N-393 terminal is not critical for Mtb LexA-DNA association, deletion of the 18 amino acids linker 394 connecting the NTD and CTD of the protein resulted in a marked reduction in DNA binding 395 compared to the full-length protein. The 18 amino acids present in the linker most likely 396 accounts for the conformational flexibility of Mtb LexA to suitably bind DNA. Moreover, 397 mutating the RKG motif in the DNA binding helix abolished LexA-DNA binding, 398 highlighting the significance of strong evolutionary conservation of this motif across different 399 organisms. 400 The quantitation of DNA binding in real-time has been carried out using BLI and

401 kinetic parameters of Mtb LexA-DNA interaction have been determined. The binding affinity of WT Mtb LexA (K\_D 2.16 $\pm$ 0.01 nM) and LexA $\Delta$ 24aa (K\_D 4.94 $\pm$ 0.03 nM) was within two-402 403 fold range, while a seventeen-fold reduction with LexA $\Delta$ 18aa (K<sub>D</sub> 34.4±0.19) and no 404 observable binding with RKG/AAA mutant was observed. Mtb LexA was found to bind 405 different "SOS" boxes under mycobacterial "SOS" regulation with comparable affinity. 406 Association to perfectly palindromic sequence was found to be stronger. However, since 407 association and dissociation rates changed proportionately for all the "SOS" boxes, the 408 overall affinities were found to fall in a close range. Although Mtb LexA binds to different 409 "SOS" boxes with comparable affinity *in-vitro*, in the cellular context, the time and spatial 410 regulation of "SOS" genes might be altered by other transcription factors, intracellular pH, 411 specific cations, and anions. In-vivo studies may uncover their actual regulation under DNA 412 damaging and normal conditions. DNA binding assays under both physiological and extreme 413 acidic pH conditions in-vitro revealed that mycobacterial LexA retains DNA binding even at 414 pH as low as 4, albeit with reduced affinity as compared to its optimum binding at 415 physiological pH. Taken together, our study provides a better understanding of the real-time Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BCJ20210434/924128/bsr-2021-1419-t.pdf by guest on 16 April 2024

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422 Figure 1. Phylogenetic analysis and domain architecture of Mtb LexA. (A) Phylogenetic 423 tree of LexA protein sequence of 24 species. The protein sequences are clustered into 4 424 groups: Actinobacteria, Proteobacteria, Euryarchaeota, and Firmicutes that include the 425 representatives from some of the major groups of bacteria. (B) Comparison of LexA 426 sequences among tuberculous, non-tuberculous mycobacterial species and model Gram-427 negative organism, E.coli. The first 24 amino acids extension, the N-terminal DNA binding 428 domain, linker region, the stretch of 18 amino acids insert in the linker and C-terminal 429 dimerization domain are shown as a bar representation below the sequence in magenta, light 430 blue, dark blue, yellow, and light green respectively. DNA binding residues chosen for 431 mutation are shown with a brown bar representation below. Sequence alignment was done 432 using Clustal Omega and ESPript was used to generate the figure. (C) Representation of 433 constructs generated for this study is shown and the gel picture shows purified Mtb LexA and 434 its variants resolved and visualized on 12% SDS PAGE.

435 Figure 2. Biochemical analysis and DNA binding property of Mtb LexA and its 436 variants. (A) The secondary structure of WT Mtb LexA, LexA $\Delta$ 24aa, LexA $\Delta$ 18aa, and 437 LexA RKG/AAA were compared using CD spectroscopy, monitored at wavelengths ranging 438 from 195-280 nm. (B) Gel image showing glutaraldehyde cross-linking of Mtb LexA and its 439 variants. Dimeric states of proteins are boxed. (C-F) Changes in extrinsic fluorescence 440 spectra of the proteins, (C) WT Mtb LexA, (D) LexA RKG/AAA, (E) LexA $\Delta$ 18aa, and (F) 441 LexA $\Delta$ 24aa, as seen when incubated at 1:2 ratios with non-biotinylated *dnaE*2 "SOS" box 442 (sequence given in Table 1) indicates the conformational changes of the proteins upon DNA 443 binding. Fluorescence intensity is shown in arbitrary units.

**Figure 3. DNA binding properties of Mtb LexA and its variants compared by EMSA analysis.** (A) EMSA analysis of WT Mtb LexA binding to <sup>32</sup>P end-labeled ds 44 mer ds *dnaE2* "SOS" box containing DNA at 0, 1, 2, 4, 8, 16, 32, 64, 128 nM is shown. (B) 128 nM of WT, LexA $\Delta$ 24aa, LexA $\Delta$ 18aa, and LexA RKG/AAA were incubated with 3.5 nM of <sup>32</sup>P end-labeled ds 44 mer ds *dnaE2* "SOS" box containing DNA (sequence given in **Table 1**) and EMSA was carried out according to standardized conditions mentioned in Materials and Methods.

and EMSA was carried out according to standardized conditions mentioned in Materials and Methods. **Figure 4. Kinetic analysis of DNA binding of Mtb LexA and its variants to** *dnaE2* **"SOS" box at physiological pH.** Representative BLI sensograms showing the real-time concentration-dependent (each concentration depicted by a different color indicated in the box below) binding of WT Mtb LexA and its variants, LexA $\Delta$ 18aa, and LexA $\Delta$ 24aa to biotinylated 44 mer *dnaE2* "SOS" box at physiological pH (A, B, C), the kinetic parameters obtained from which are tabulated in **Table 2**. The corresponding response versus concentration curves has been plotted in (D, E, F) from the results of three independent experiments. (G) Representative BLI sensogram showing absence of binding of LexA RKG/AAA to biotinylated 44 mer *dnaE2* "SOS" box at physiological pH.

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Figure 5. Kinetic analysis of DNA binding of Mtb LexA and its variants to dnaE2"SOS" box at acidic pH. Representative BLI sensograms showing the real-time concentration-dependent (each concentration depicted by a different color indicated in the box below) binding of WT Mtb LexA and its variants, LexA $\Delta$ 18aa, and LexA $\Delta$ 24aa to biotinylated 44 mer dnaE2 "SOS" box at pH 4 (A, B, C), the kinetic parameters obtained from which are tabulated in Table 2. The corresponding response versus concentration curves has been plotted in (D, E, F) from the results of three independent experiments.

Figure 6. DNA binding kinetics of Mtb LexA to various "SOS" boxes. Representative 469 470 BLI sensograms showing the real-time concentration-dependent (each concentration depicted 471 by a different color indicated in the box below) binding of WT Mtb LexA to 44 mer 472 biotinylated "SOS" boxes of (A) lexA, (B) rv3074, and (C) recA, the kinetic parameters 473 obtained from which are tabulated in Table 3. (D) Comparison in binding by plotting 474 association constants (Ka) of WT Mtb LexA to different "SOS" boxes from 3 independent 475 experiments is shown. Association constants (Ka) are inverse of equilibrium dissociation 476 constants or K<sub>D</sub> values obtained from kinetic analyses.

477 Associated Content. Supplementary Material. The Supplementary Material file is478 available with this manuscript. It includes the following:

## 479 Supplementary Table Captions:

#### 480 Table S1: Bacterial strains and plasmids.

481 Table S2: Oligonucleotide primers used in this study.

482 Table S3: Assessment of DNA binding properties of Mtb LexA variants to various

483 "SOS" boxes: Kinetic parameters obtained from binding studies performed using BLI.

# 484 Supplementary Figure Captions:

Figure S1. Multiple sequence analysis of LexA protein. Comparison between sequences of Mtb LexA with LexA proteins from representatives from Actinobacteria, Proteobacteria, Euryarchaeota, and Firmicutes are shown. The first 24 amino acids extension, the N-terminal DBD, linker region, and CTD are represented as colored bars below the sequences in magenta, light blue, dark blue, and light green respectively. Conserved DNA binding residues Downloaded from http://port.silverchair.com/bioscirep/article-pdf/doi/10.1042/BCJ20210434/924128/bsr-2021-14194.pdf by guest on 16 April 2024

490 are shown with a brown bar representation below. Sequence alignment was done using491 Clustal Omega and ESPript was used to generate the figure.

Figure S2. Analysis of the oligomeric state of Mtb LexA and its variants. Gel Filtration
analysis of (A) WT Mtb LexA and its variants, (B) RKG/AAA, (C) LexAΔ18aa and (D)
LexAΔ24aa. 500 µl was injected and run in Superdex 75 10/300 column pre-equilibrated
with 20 mM Tris-Cl, pH 7.5, 100 mM NaCl, and 5% glycerol. (E) Approximate molecular
weight of Mtb LexA and its variants plotted by comparing with standard markers.

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**Figure S3.** Assessment of DNA binding property of LexA $\Delta 24aa$  (without N-terminal 6x His tag). Representative BLI sensogram showing the real-time concentration-dependent (each concentration depicted by a different color indicated in the box below) binding of LexA $\Delta 24aa$  variant to biotinylated 44 mer lexA "SOS" box at physiological pH, the kinetic parameters obtained from which are tabulated and shown in the inset (A). The corresponding response versus concentration curve has been plotted in (B) from the results of three independent experiments.

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Figure S4. Assessment of DNA binding properties of variants of Mtb LexA to various
"SOS" boxes. BLI sensograms showing the real-time concentration-dependent (each
concentration depicted by a different color indicated in the box below) binding of
LexAΔ24aa to 44 mer biotinylated "SOS" boxes of (A) *lexA*, (B) *rv3074* and (C) *recA* and of
LexAΔ18aa to 44 mer biotinylated "SOS" boxes of (D) *lexA*, (E) *rv3074* and (F) *recA*,
respectively, the kinetic parameters obtained from which are tabulated in Table S3..

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530 prepared constructs used in the study. DP performed bioinformatic analyses. CC wrote the 531 original draft. CC, SM, and MS edited the manuscript. MS supervised the study.

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# 533 Data availability statement

All supporting data and sequence information are included within the main article and itssupplementary material.

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# 537 **Conflicts of interest**

538 The authors declare no conflict of interest.

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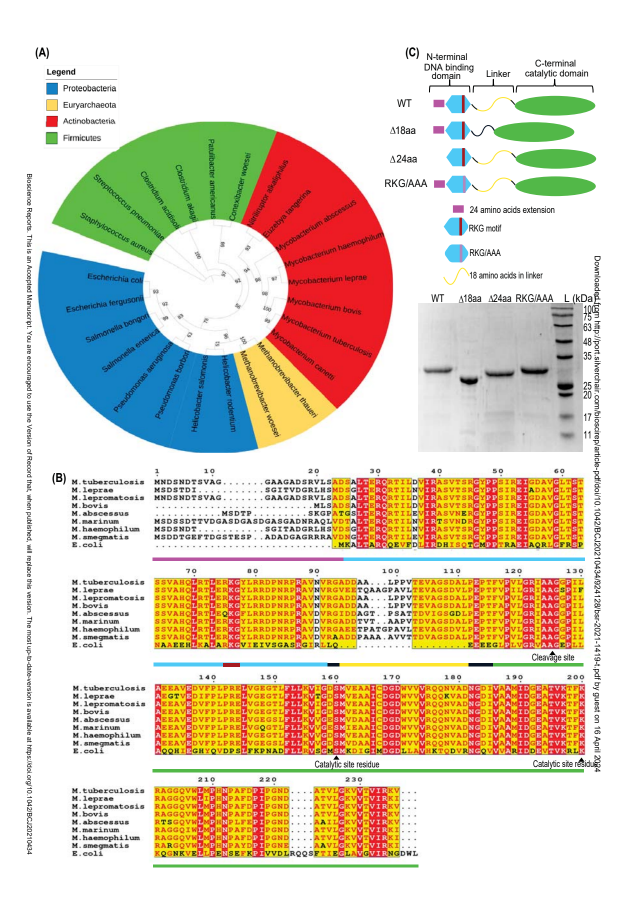
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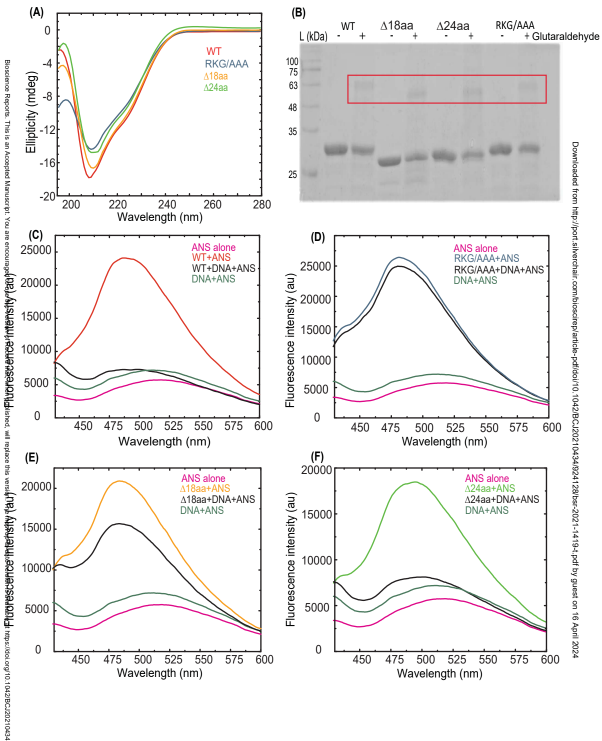
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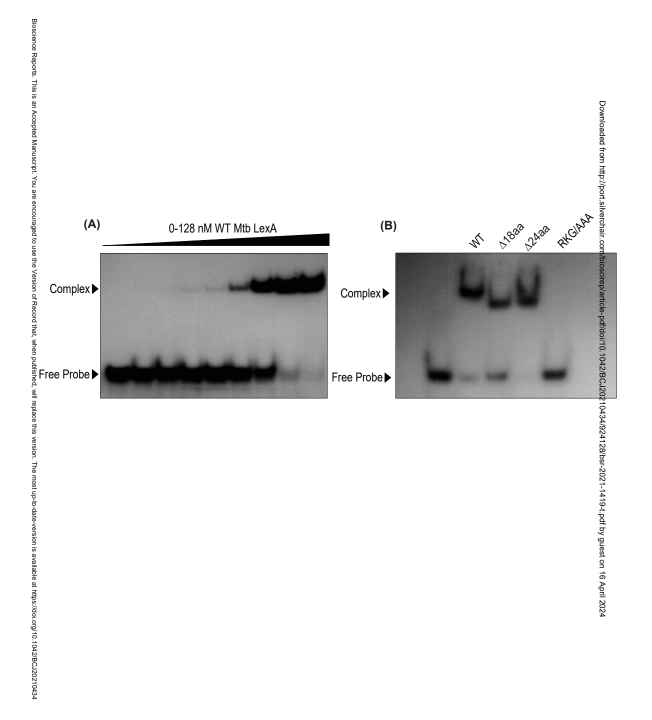
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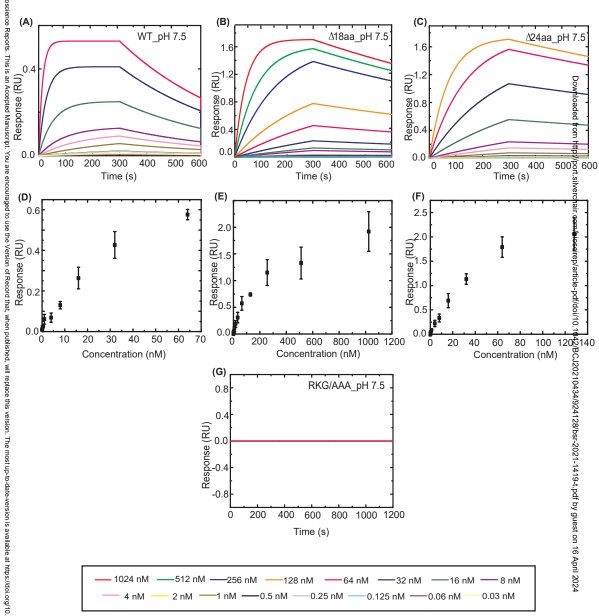
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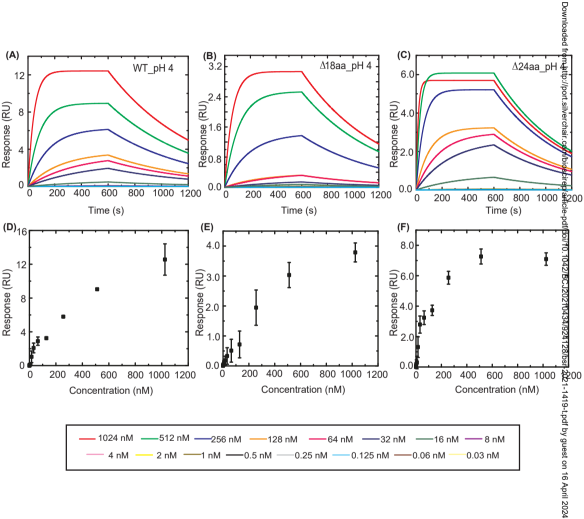




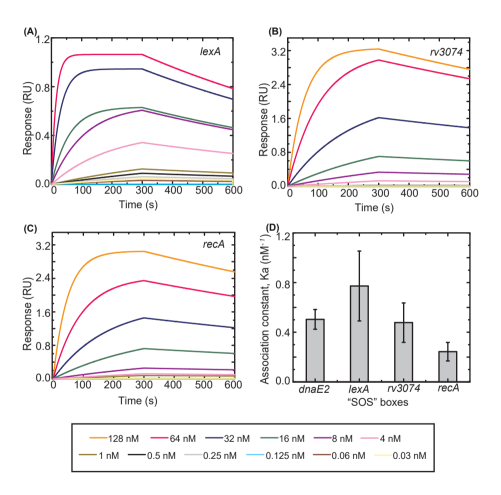




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Duplex	Oligonucleotides used for DNA binding analysis and kinetics*
dnaE2_44mer	FP- 5' Btn <sup>#</sup> -ACAACTGCGCTGTATCGAAC <u>AATT</u> GTTCGATATACTGTGGAATG 3'
	RP-3' TGTTGACGCGACATAGCTTG <u>TTAA</u> CAAGCTATATGACACCTTAC 5'
	(Perfect palindrome)
lexA_44mer	FP- 5' Btn-CCGGAACACGCCTGTCGAACACATGTTTGATTCTTGGTGCGAAT 3'
	RP-3' GGCCTTGTGCGGACAGCTTG <u>TGTA</u> CAAACTAAGAACCACGCTTA 5'
	(Imperfect palindrome)
recA_44mer	FP- 5' Btn-GTGTCACACTTGAATCGAACAGGTGTTCGGCTACTGTGGTGATC 3'
	RP-3' CACAGTGTGAACTTAGCTTG <u>TCCA</u> CAAGCCGATGACACCACTAG 5'
	(Imperfect palindrome)
rv3074_44mer	FP- 5' Btn-GCAGGCTGCTATTCTCGAACACATGTTCGAGACATTGACCGCGA 3'
	RP-3' CGTCCGACGATAAGAGCTTGTGTACAAGCTCTGTAACTGGCGCT 5' (Perfect palindrome on the repeat flanks, not in the sequences <u>in between</u> )

\*"SOS" box sequences are highlighted in **bold** <sup>#</sup>Btn stands for biotinylation

# Table 1: "SOS" boxes chosen for binding studies

k <sub>on</sub>	$k_{off} \left( 1/s \right)$	K <sub>D</sub> (nM)	R <sup>2</sup>
$(M^{-1}s^{-1}) \ge 10^5$	$(s^{-1}) \ge 10^{-3}$		
10.55±0.06	2.28±0.007	2.16±0.01	0.99
$0.22 \pm 0.0007$	$0.76 \pm 0.004$	34.4±0.19	0.99
1.06±0.003	0.52±0.003	4.94±0.03	0.99
0.20±0.0006	1.52±0.002	75.99±0.27	0.99
0.15±0.0004	1.64±0.002	111±0.33	0.99
0.61±0.003	$1.84 \pm 0.003$	29.94±0.13	0.98
	$(M^{-1}s^{-1}) \ge 10^{5}$ 10.55±0.06 0.22±0.0007 1.06±0.003 0.20±0.0006 0.15±0.0004	$(M^{-1}s^{-1}) \ge 10^5$ $(s^{-1}) \ge 10^{-3}$ $10.55 \pm 0.06$ $2.28 \pm 0.007$ $0.22 \pm 0.0007$ $0.76 \pm 0.004$ $1.06 \pm 0.003$ $0.52 \pm 0.003$ $0.20 \pm 0.0006$ $1.52 \pm 0.002$ $0.15 \pm 0.0004$ $1.64 \pm 0.002$	$(M^{-1}s^{-1}) \ge 10^5$ $(s^{-1}) \ge 10^{-3}$ $(10.55\pm0.06)$ $2.28\pm0.007$ $2.16\pm0.01$ $0.22\pm0.0007$ $0.76\pm0.004$ $34.4\pm0.19$ $1.06\pm0.003$ $0.52\pm0.003$ $4.94\pm0.03$ $0.20\pm0.0006$ $1.52\pm0.002$ $75.99\pm0.27$ $0.15\pm0.0004$ $1.64\pm0.002$ $111\pm0.33$

Table 2: Kinetic parameters obtained from binding studies performed using BLI.

"SOS" boxes	kon	k <sub>off</sub>	K <sub>D</sub> (nM)	$\mathbf{R}^2$
	$(M^{-1}s^{-1}) \ge 10^5$	$(s^{-1}) \ge 10^{-3}$		
dnaE2	10.55±0.06	2.28±0.007	2.16±0.01	0.99
lexA	10.27±0.04	$1.01 \pm 0.004$	0.98±0.01	0.99
rv3074	1.39±0.004	0.53±0.004	3.86±0.03	0.99
recA	1.61±0.005	0.59±0.004	3.63±0.02	0.99

 Table 3: Kinetic parameters obtained from binding studies of WT Mtb LexA with

 various "SOS" boxes performed using BLI.

Streptococcus\_pneumoniae Staphylococcus\_aureus Clostridium\_acidisoli Nitriliruptor alkaliphilu Patulibacter\_americanus Mycobacterium\_bovis Mycobacterium\_tuberculosi Mycobacterium\_canetii Mycobacterium\_leprae Mycobacterium\_haemophilum Mycobacterium\_abscessus Euzebya\_tangerina Conexibacter\_woesei Escherichia\_coli scherichia\_fergusonii almonella\_bongori Salmonella enterica a ⊉seudomonas\_aeruginosa ⊉seudomonas\_borbori Helicobacter\_salomonis Methanobrevibacter\_thauer

	1	10	20	30	40	50	60
		TKROSEIYDY	IKHVVOIKGYP	. PSVREIGE	A <mark>VGLAS</mark> S <mark>STV</mark> .		RDP
		TK <mark>RO</mark> SE <mark>I</mark> YNY	IKQV <mark>V</mark> QTKGYP	. PSVREIGE			RDP
		RRD <mark>VQ</mark> AE <mark>I</mark> YDF	IQSE <mark>V</mark> INK <mark>G</mark> YP	. PSVREICA			<b>RDP</b>
us	MTDPPVTDIAAARAANTGE	<mark>ITERQ</mark> RA <mark>I</mark> LEV				HAQ <mark>l</mark> et <mark>l</mark> ea <mark>kg</mark> y <mark>l</mark> rf	
			IGRY <mark>T</mark> TDF <mark>G</mark> YP				
	MLSADSA	LTE <mark>RO</mark> RT <mark>I</mark> LDV				AHQ <mark>l</mark> rt <mark>l</mark> er <mark>kg</mark> y <mark>l</mark> rf	
is	MNDSNDTSVAGGAAGADSRVLSADSA	LTE <mark>RO</mark> RT <mark>I</mark> LDV				ahq <mark>l</mark> rt <mark>l</mark> er <mark>kg</mark> y <mark>l</mark> rf	
	MNDSNDTSVAGGAAGADSRVLSAGSA	LTERORTILDV		. PSIREIGDA		AHQ <mark>L</mark> RT <mark>LERKG</mark> YLRF	
	MSDSTDISGITVDGRLHSMDSG	LTERORT I LNV				AHQ <mark>L</mark> RT <mark>LERKGYL</mark> RF AHOLRTLERKGYLRF	
m	MSDSNDTSGITADGRLHSVDSG	LTERORTILNV		. PSIREIGDA		AHQ <mark>LRTLERKGYL</mark> RF AHOLRTLEOKGFLRF	
			IOSTVAERGYP		AVGLISISSV.		
		TKROOEIFEF					
		TAROQEVFDL				EEHLKALARKGVÆEI	
		TVROOEVFDL				EEHLKALARKGVEI	
		TAROOEVFDL				EEHLKALARKGVEEI	
		TAROOEVFDL				EEHLKALARKGV	
		TPROAEILSF				EEH <mark>L</mark> KALARKGAZEM	1TP
		TS <mark>RQ</mark> AE	IKHCLDDNGYP			EEH <mark>l</mark> ka <mark>l</mark> ar <mark>kg</mark> a <mark>j</mark> em	1TP
		MLKEK	LKQLRNANNL.	TQEELAL	K <mark>CGV</mark> SLQ <mark>S</mark> IKRY	eseqksnitldt <mark>l</mark> ek <mark>lanal</mark> n <b>æ</b> di	HF
	MDLRERIKAARNNLGITQDE						
i						FSEEL <mark>N</mark> EE <mark>I</mark> CK <mark>YA</mark> L <mark>D</mark> SM	
ri	MAKKT	<mark>LAFI</mark> II <mark>I</mark> FIG	<mark>FS</mark> . A <mark>l</mark> FMI <mark>N</mark> . <mark>S</mark>	H <mark>DT</mark> VD <mark>VY</mark> LDO	G <mark>ENV</mark> S <mark>VE</mark> TKDFO	GNGNLDLL <mark>N</mark> QE <mark>I</mark> CD <mark>YV</mark> V <b>M</b> VM	IDD
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	70	80	90	100	110	120 1300	

Streptococcus\_pneumoniae Staphylococcus aureus Clostridium\_acidisoli Clostridium akaqii Nitriliruptor\_alkaliphilus Patulibacter\_americanus Mycobacterium\_bovis Mycobacterium\_tuberculosis ∰ycobacterium\_canetii Mycobacterium\_leprae Mycobacterium\_haemophilum Zycobacterium\_abscessus Euzebya\_tangerina Eonexibacter\_woesei zscherichia\_coli Escherichia\_fergusonii Salmonella\_bongori Salmonella\_enterica seudomonas\_aeruginosa Pseudomonas\_borbori Helicobacter\_salomonis Helicobacter\_rodentium Methanobrevibacter\_woesei Methanobrevibacter\_thaueri

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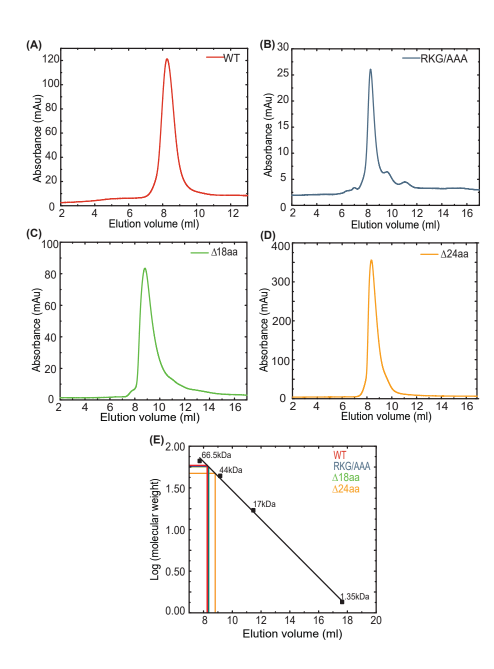
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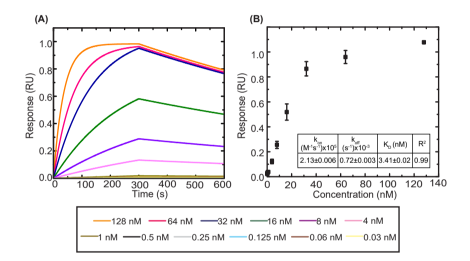
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	TKPRAIEIVSDO.NN	DMTREETIYVPV	IGKVTA.	<b>GVPITAVENIE</b> EN	YFPLPEHLTSTH	NSDVFILN	ICESVEEAGI
	TKPRAIEIVSDOTND		IGKVTA.	GVPITAVENIEES	YFP <mark>LP</mark> EHLTSTH	NSDIFILN	VGDSM EEAGI
	TKP <mark>RAIEL</mark> LRDT				SFT <mark>LP</mark> IOYTK.S	NKDLFMLKI	SGE <mark>SM</mark> ŽEAGI
	SKP <mark>RAIEL</mark> LKDS				YFT <mark>LP</mark> IQYTK.S	NKDLFMLKI	SGESM DAGI
ıs	TKP <mark>RALEM</mark> GRDPDTDL	AVRPS <mark>S</mark> GRN <mark>VPI</mark>	VGEIAA	GGPILAEERVESV	VYA <mark>LP</mark> KELVG		RGESMPEAGV
	SKP <mark>RAIEL</mark> LGRGVEQAV	/EGVRNAVG <mark>E</mark> SRR <mark>LPI</mark>	LGO <mark>V</mark> AA	<b>GPPILA</b> EEH <mark>VE</mark> EH	IVP <mark>VP</mark> DMAGG	AEGEYLLRI	RGESMHGAGI
;	NRP <mark>RAV</mark> NVRGADDAALPPVTI	EVAGSDALP <mark>E</mark> PTF <mark>APV</mark>	LGRIAA	<b>GGPILAEEAVE</b> DV	VFP <mark>LP</mark> RELVG		IGDSMVEAAI
is :	NRP <mark>RAV</mark> NVRGADDAALPPVTI	EVAGSDALP <mark>E</mark> PTF <mark>VPV</mark>	LGRIAA	<b>GGPILAEEAVE</b> DV	VFP <mark>LP</mark> RELVG	EGTL <mark>FLLK</mark>	IGDSMUEAAI
;	NRP <mark>R</mark> A <mark>V</mark> N <mark>V</mark> RGADDAALPPVTI	EVAGSDALP <mark>E</mark> PTF <mark>VPV</mark>	LGRIAA	<b>GGPILAEEAVE</b> DV	VFP <mark>LP</mark> RELVG	EGTL <mark>FLLK</mark>	IGDSMWEAAI
1	NRP <mark>R</mark> A <mark>V</mark> N <mark>V</mark> RGVEETQAAGPAVLTI	EVAGSDVLP <mark>E</mark> PTF <mark>VPI</mark>	LGRIAA	GSPIFAEGT <mark>VE</mark> DI	IFP <mark>LP</mark> RELVG	EGTLELK	TCDSMOEAT
n I	NRP <mark>R</mark> A <mark>V</mark> D <mark>V</mark> RGAEETPATGPAVLTI	EVAGSDVLP <mark>E</mark> PTF <mark>VPV</mark>	LGRIAA	GGPILAEEAVEDV	VFP <mark>LP</mark> RELVG	EGTL <mark>FLLK</mark>	VGDSNVEAAI
1	NRP <mark>R</mark> A <mark>V</mark> D <mark>V</mark> RGIDDAGTPSATTI	)VIGSGDLP <mark>E</mark> PTF <mark>VPV</mark>	LGRIAA	GGPILAEEAVEDV	VFP <mark>LP</mark> RELVG	EGSL <mark>FLLK</mark>	VGESMODAAI
	SRP <mark>R</mark> A <mark>I</mark> E <mark>V</mark> LAEFDED	APDTS <mark>P</mark> TRT <mark>VPV</mark>	VGEIA <mark>A</mark>	GGPILADQQIDE	HLV <mark>LP</mark> ESFVG	SGTI <mark>FALT\</mark>	RGESM TDAGV
	SKP <mark>R</mark> A <mark>I</mark> ELLDKAVDG	IKSIV <mark>K</mark> PAG <mark>LPI</mark>	VGQ <mark>V</mark> Q <mark>A</mark>	GQPVLAEEE IEDY	IETPAVAGG	AEGE <mark>YLLR</mark>	RGESM RDAGI
	GAS <mark>R</mark> G <mark>I</mark> R <mark>L</mark> L	QE <mark>E</mark> EEG <mark>LPI</mark>	VGR <mark>V</mark> A <mark>A</mark>	GEPLLAQQHIEGH	HYQ <mark>VD</mark> PSLFK	PNAD <mark>FLLR</mark>	/S <mark>GMSMKDI</mark> GI
	GAS <mark>R</mark> G <mark>I</mark> R <mark>L</mark> L	QE <mark>E</mark> EEG <mark>LPI</mark>	<mark>VG</mark> R <mark>V</mark> A <mark>A</mark>	GEPLLAQQHIEGH	HYQ <mark>VD</mark> PSLFK	PNAD <mark>FLLR</mark>	/S <mark>GMSMKDI</mark> GI
	GAS <mark>R</mark> G <mark>I</mark> R <mark>L</mark> L	QE <mark>E</mark> EEG <mark>LPI</mark>	VGR <mark>V</mark> A <mark>A</mark>	GEPLLAQQHIEGH	HYQ <mark>VD</mark> PSLFK		/S <mark>GMSMKDIGI</mark>
	GAS <mark>R</mark> G <mark>I</mark> R <mark>L</mark> L	QE <mark>E</mark> EDG <mark>LPI</mark>	<mark>VG</mark> R <mark>V</mark> A <mark>A</mark>	GEPLLAPQHIEGH	HYQ <mark>VD</mark> PSLFK	PSAD <mark>FLLR</mark>	/S <mark>GMSMKDI</mark> GI
	GAS <mark>R</mark> G <mark>I</mark> R <mark>I</mark> PGFEPHA				SCR <mark>IN</mark> PAFFN		/RGMSMEDIGI
	GAS <mark>R</mark> G <mark>I</mark> R <mark>I</mark> PGYEP				SCR <mark>IN</mark> PEFFH		/RGMSMKDIGI
	FTS <mark>T</mark> H <mark>R</mark> E <mark>V</mark> E						
	KSV <mark>S</mark> Q <mark>L</mark> S <mark>P</mark> NQKKSVPQSI						
	ISS <mark>NV.</mark> . <mark>.</mark>						
ri '	TTT <mark>N</mark> I <mark>.</mark>		IENICL	.K <mark>YGLD</mark> DPT <mark>VN</mark> II	DSS <mark>IG</mark> PDQI	P <mark>VIVY\</mark>	DGTSNUPT.L
			<b>▲</b>	<b>▲</b>			
			Cleavage site	residues		Cat	alytic site reștdue

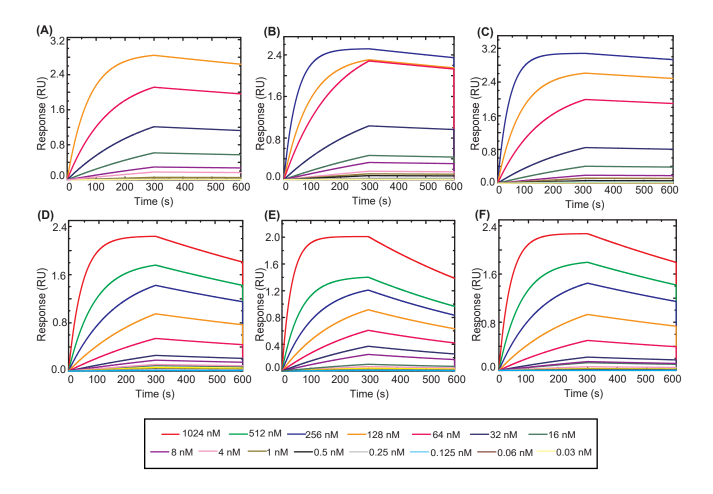
Š Streptococcus\_pneumoniae "staphylococcus\_aureus Clostridium\_acidisoli Clostridium\_akagii Witriliruptor\_alkaliphilus Patulibacter\_americanus Mycobacterium bovis Mycobacterium\_tuberculosis Mycobacterium\_canetii Mycobacterium\_leprae Mycobacterium\_haemophilum Mycobacterium\_abscessus auzebya\_tangerina Conexibacter\_woesei Escherichia\_coli zscherichia fergusonii Salmonella\_bongori Salmonella\_enterica eudomonas\_aeruginosa Seudomonas\_borbori Helicobacter\_salomonis ∄elicobacter\_rodentium Methanobrevibacter\_woesei Methanobrevibacter\_thaueri

140	150	160	170	180	190	200
L <mark>DGD</mark> K <mark>VIVR</mark> S	QTI <mark>AENG</mark> DII	VAMTEDDEAT		R <mark>Y</mark> R <mark>I</mark> Q <mark>P</mark> ENSAM	s <mark>pi</mark> yldnvt <mark>v</mark>	IGK <mark>VI</mark> G <mark>LYR</mark> EL
L <mark>DGD</mark> K <mark>VIVR</mark> S						IGK <mark>VI</mark> GLYREM
						IGELVGIYRKY
						<mark>VG</mark> Q <mark>LV</mark> G <mark>LYR</mark> RY
					E <mark>PI</mark> PVT . ADNDAH <mark>I</mark>	
						<mark>LG</mark> K <mark>VV</mark> GVLRKVG
						<mark>lg</mark> k <mark>vv</mark> tvirkv
						<mark>lg</mark> k <mark>vv</mark> tvirkv
						LGK <mark>VV</mark> TVIRKV
						LGK <mark>VV</mark> TVIRKI
						<mark>lg</mark> k <mark>vv</mark> tvirki
						LGK <mark>VV</mark> TVIRKV
						MGKVVAVLRKL
					E <mark>PI</mark> RTKEVQ <mark>I</mark>	
						EGLAVGVIRNGDWL EGLAVGVIRNGDWL
						EGLAVGVIRNGDWL
						EGLAVGVIRNGEWL
						EGLSVGAIRR
						EGLSVGAIRR
						LGR <mark>VI</mark> GVIRK
						IGRVVGYFKKTSL
						ITTWVDLSDIYGVVEDY
						ITTWVDISDINGVVIDY
Z <mark>POZ</mark> I <mark>VIII I</mark> I						

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Name	Characteristics	Source
<b>Bacterial Strains</b>		
<i>E.coli</i> DH5α	F– $\varphi$ 80lacZΔ M15 Δ ( <i>lacZYA-argF</i> ) U169 recA1 endA1 hsdR17 (rK–mK+) phoA supE44 λ- thi–1 gyrA96 relA1	Laboratory stock
E.coli BL21(DE3)	F- ompT hsdSB (rB-, mB-) gal dcm (DE3)	Laboratory stock
Plasmids		
pET28a(+)	Expression vector for low copy number (pBR322 ori), strong phage promoter (T7), IPTG induction (lac operon), and Kanamycin selection (Kan <sup>r</sup> )	Novagen
pET22b(+)	Expression vector (pBR322 ori), strong phage promoter (T7), IPTG induction (lac operon), and Ampicillin selection (Amp <sup>r</sup> )	Novagen

# Table S1: Bacterial strains and plasmids

Construct/Mutant	Forward Primer (FP) / Reverse Primer (RP) Sequences with restriction
	sites and base changes for mutation highlighted in bold
WT Mtb LexA	FP- 5' ATGCCG CATATG ATGAACGACAGCAACGAC 3'
	RP- 5' ATGCCG GGATCC TCAGACCTTGCGGATCAC 3'
Mtb LexA∆18aa	FP- 5' GTCAATGTGCGCGGTCCGGAACCCACCTTT 3'
	RP- 5' AAAGGTGGGTTCCGGACCGCGCACATTGAC 3'
Mtb LexA∆24aa	FP- 5' ATAT CATATG TCGGCGCTGACCGAGCGGCAA 3'
	RP-5' ATGCCG GGATCC TCAGACCTTGCGGATCAC 3'
	RP-5' CCC AAGCTT GACCTTGCGGATCACCGT 3'
Mtb LexA RKG/AAA	FP- 5' CTGCGCACCCTGGAGGCGGCGGCGTACCTACGCCGTGAC 3
	RP- 5' TCACGGCGTAGGTACGCCGCCGCCTCCAGGGTGCGCAG 3'

			0	e	
Protein	"SOS" boxes	k <sub>on</sub>	k <sub>off</sub>	$K_{D}\left( nM ight)$	$\mathbf{R}^2$
	DUXES	$(M^{-1}s^{-1}) \ge 10^5$	$(s^{-1}) \ge 10^{-3}$		
Mtb LexA∆18aa	dnaE2	0.22±0.0007	0.76±0.004	34.40±0.19	0.99
	lexA	0.21±0.0008	0.71±0.004	34.53±0.25	0.99
	rv3074	0.28±0.0011	1.24±0.005	43.50±0.24	0.99
	recA	0.24±0.0009	0.79±0.004	32.55±0.22	0.99
Mtb LexA∆24aa	dnaE2	1.06±0.003	0.52±0.003	4.94±0.03	0.99
	lexA	1.05±0.002	0.25±0.002	2.36±0.02	0.99
	rv3074	0.85±0.002	0.24±0.003	2.75±0.04	0.99
	recA	0.99±0.002	0.17±0.002	1.67±0.02	0.99

Table S3: Assessment of DNA binding properties of Mtb LexA variants to various "SOS"boxes: Kinetic parameters obtained from binding studies performed using BLI.

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