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Identification of new members of the Escherichia coli K-12 MG1655 SlyA regulon --Manuscript Draft--

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Abstract:	SlyA is a member of the MarR family of bacterial transcriptional regulators. Previously, SlyA has been shown to directly regulate only two operons in Escherichia coli K-12 MG1655, fimB and hlyE (clyA). In both cases SlyA activates gene expression by antagonizing repression by the nucleoid associated protein H-NS. Here the transcript profiles of aerobic glucose-limited steady-state chemostat cultures of E. coli K-12 MG1655, slyA mutant and slyA over-expression strains are reported. The transcript profile of the slyA mutant was not significantly different to that of the parent; however, that of the slyA expression strain was significantly different from that of the vector control. Transcripts representing 27 operons were increased in abundance, whereas 3 were decreased. Of the 30 differentially regulated operons, 24 have been previously associated with sites of H-NS binding, suggesting that antagonism of H-NS repression is a common feature of SlyA-mediated transcription regulation. Direct binding of SlyA to DNA located upstream of a selection of these targets permitted the identification of new operons likely to be directly regulated by SlyA. Transcripts of four operons coding for cryptic adhesins exhibited enhanced expression and this was consistent with enhanced biofilm formation associated with the SlyA over-producing strain.

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Identification of new members of the *Escherichia coli* K-12 MG1655 SlyA regulon

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- 10
- 11 Key words: biofilm, gene expression, MarR family, transcription regulation
- 12
- 13 Abbreviations used: A₆₀₀; absorbance at 600 nm; Amp^R, ampicillin resistance; Btn, biotin;
- 14 Cm^R, chloramphenicol resistance; CRP, cyclic-AMP receptor protein; EMSA, electrophoretic
- mobility shift assay; Kan^R, kanamycin resistance; OD₆₀₀; optical density at 600 nm; Tet^R,
- 16 tetracycline resistance; T^s, temperature sensitive
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30 SlyA is a member of the MarR family of bacterial transcriptional regulators. Previously, SlyA 31 has been shown to directly regulate only two operons in Escherichia coli K-12 MG1655, fimB 32 and *hlyE* (*clyA*). In both cases SlyA activates gene expression by antagonizing repression by the nucleoid associated protein H-NS. Here the transcript profiles of aerobic glucose-33 limited steady-state chemostat cultures of E. coli K-12 MG1655, slyA mutant and slyA over-34 expression strains are reported. The transcript profile of the *slyA* mutant was not significantly 35 different to that of the parent; however, that of the *slyA* expression strain was significantly 36 different from that of the vector control. Transcripts representing 27 operons were increased 37 38 in abundance, whereas 3 were decreased. Of the 30 differentially regulated operons, 24 39 have been previously associated with sites of H-NS binding, suggesting that antagonism of H-NS repression is a common feature of SlyA-mediated transcription regulation. Direct 40 41 binding of SlyA to DNA located upstream of a selection of these targets permitted the 42 identification of new operons likely to be directly regulated by SlyA. Transcripts of four 43 operons coding for cryptic adhesins exhibited enhanced expression and this was consistent with enhanced biofilm formation associated with the SlyA over-producing strain. 44

45

46 **INTRODUCTION**

47 The MarR family of transcription regulators are widespread throughout the Bacterial and Archeal 48 kingdoms [1]. MarR family members are homodimeric and bind to palindromic DNA sequences 49 within regulated promoters using a characteristic winged-helix-turn-helix DNA-binding domain [2]. 50 These regulators repress gene expression by promoter occlusion (e.g. MarR; [3]), or activate gene 51 expression by stabilizing RNA polymerase-promoter DNA interactions (e.g. OhrR; [4]), or by 52 antagonizing the action of repressors (e.g. RovA; [5]). These activities of MarR proteins are inhibited 53 upon interaction with cognate signalling molecules, although for many members the natural ligand is 54 unknown [6].

The Salmonella enterica serovar Typhimurium LT2 SlyA protein is one of the best 55 56 characterized members of the MarR family. The S. enterica serovar Typhimurium slyA mutant is attenuated for virulence, is hypersensitive to oxidative stress and is impaired for survival in 57 58 macrophages [7, 8]. A consensus DNA binding site has been proposed, TTAGCAAGCTAA [9, 10], and proteomic and transcriptomic comparisons of parent and slyA mutant strains suggest that SlyA 59 can act as both a negative and positive regulator of gene expression, with significant overlap with 60 genes of the PhoPQ regulon involved in cell envelope function, virulence, resistance to anti-microbial 61 62 peptides and regulation of small RNAs [11-15]. Salmonella enterica serovar Typhimurium 14028s 63 SlyA has also been linked to the stringent response by binding ppGpp resulting in enhanced DNA-64 binding [16, 17]. The expression of many SlyA-regulated genes is subject to H-NS-mediated silencing and activation of these genes generally involves an element of antagonism of H-NS repression bySlyA; e.g. [11, 18-22].

The SlyA protein of *E. coli* MG1655 is 91% identical, 95% similar (over 142 amino acids) to the *S. enterica* serovar Typhimurium LT2 protein, but is much more poorly characterized. Only two genes, *hlyE* and *fimB* (as well as autoregulation of *slyA*) have been shown to be directly regulated by SlyA [19, 21, 23]. In some other *E. coli* strains, SlyA regulates capsule synthesis and lipid A palmitoylation in biofilms [18, 19, 22]. Here transcriptional profiling of parent, *slyA* mutant and *slyA* over-expression strains reveals the breadth of the *E. coli* MG1655 SlyA regulon, indicating roles in activating expression of cryptic fimbrial-like adhesins that contribute to enhanced biofilm formation.

74

75 **METHODS**

76 Bacterial strains, plasmids, oligonucleotides and growth conditions. The bacterial strains 77 plasmids and oligonucleotides that were used are listed in Table 1. Bacterial strains were routinely 78 cultured in Luria Bertani broth or on Luria Bertani agar plates [24]. Aerobic glucose-limited steady-79 state chemostat cultures of E. coli were established in Evans minimal medium [25] in Labfors 3 80 fermentation vessels (Infors-HT, Switzerland) with a 1 L working volume, 0.2 h⁻¹ dilution rate, 37°C, pH 6.9, 400 rpm stirring rate and sparging with 1 L min⁻¹ air. Evans minimal medium consists of: 10 81 mM NaH₂PO₄, 10 mM KCl, 1.25 mM MgCl₂, 20 mM NH₄Cl, 0.02 mM CaCl₂, 0.1 mM Na₂SeO₃, 1.5 82 83 mM monosodium nitrilotriacetate, 20 mM glucose and 100 ml trace element solution. The trace 84 element solution consisted of (g L⁻¹): ZnO (0.412), FeCl_{3.6}H₂O (5.4), MnCl_{2.4}H₂O (2.0), CuCl_{2.2}H₂O 85 (0.172), CoCl₂.6H₂O (0.476), H₃BO₃ (0.064), Na₂MoO₄.H₂O (0.004) in 0.3% v/v HCl. For generation of cell paste for purification of His-tagged SlyA, E. coli BL21 (\lambda DE3) transformed with pGS2469 86 87 was grown in auto-induction medium supplemented with ampicillin (100 mg L⁻¹) [26]. Resistance to 88 chloramphenicol was tested by inoculating Luria Bertani broth (2 ml) containing kanamycin (30 µg ml⁻¹) and either 0, 1, 2, 3 or 4 µg ml⁻¹ chloramphenicol with 10 µl of overnight starter cultures (E. coli 89 K-12 MG1655 pET28a or E. coli K-12 MG1655 pGS2468). Triplicate cultures were grown under 90 91 aerobic conditions for 6 h at 37°C before measuring OD_{600} as a measure of growth. The experiment 92 was carried out twice.

93

Biofilm assay. Biofilm assays were performed using 96-well plates essentially as described by Tagliabue *et al.* [27] using M9 minimal medium with 20% (w/v) glucose and 50 μ g ml⁻¹ kanamycin. Wells containing 200 μ l of medium were inoculated (1:10) from an overnight culture of *E. coli* K-12 MG1655 pET28a or *E. coli* K-12 MG1655 pGS2468 and then incubated for 16 h under aerobic conditions at 37°C. Growth of cultures was monitored by measuring OD₆₀₀. The planktonic cells were removed and the remaining biofilm was stained for 5 min with 200 μ l 1% (w/v) crystal violet solution. Excess stain was removed by three washes with deionized water before the plate was 101 air-dried. To quantify the extent of staining, 200 μ l ethanol:acetone (4:1) was added to each well, and 102 after incubating for 20 min the amount of biofilm was estimated by measuring A₆₀₀. Adhesion units 103 were calculated by dividing the A₆₀₀ values for crystal violet-stained adhered cells by the OD₆₀₀ values 104 for the corresponding planktonic cells.

105

106 Creation of *E. coli* K-12 MG1655 *slyA* mutant. A PCR-amplified DNA fragment containing 107 the kanamycin cassette from pKD4 flanked by 40 bp DNA homologous to regions surrounding the slyA gene was synthesized using oligonucleotide primers TC7 and TC8 (Table 1). The purified 108 (QiaQuick PCR cleanup, Qiagen) PCR product (5 µg) was used to transform E. coli JRG6072 by 109 electroporation (Hybaid Cell Shock unit; 1800 V, 1 mm path length). The E. coli JRG6072 competent 110 cells were prepared from aerobic Luria Bertani broth batch cultures supplemented with ampicillin 111 112 (100 mg L⁻¹) at 30°C that had been induced to express the λ red recombinase by addition of Larabinose (1 mM). Kanamycin resistant mutants were selected on Luria Bertani agar plates 113 containing kanamycin (30 mg L^{-1}) at 37°C. Mutation of the *slyA* gene by insertion of the kanamycin 114 115 resistance cassette was confirmed by colony PCR using oligonucleotides TC9 and TC10. The slyA 116 mutation was then transduced using bacteriophage P1 to E. coli MG1655 [24].

117

Transcriptional profiling. Transcriptomic analyses were carried out as described by Rolfe *et al.* 118 119 [28] using directly quenched samples from glucose-limited steady-state chemostat cultures (dilution rate 0.2 h⁻¹) for the three E. coli K-12 MG1655 strains; parent, slyA mutant (JRG6457) and slyA over-120 121 producer (JRG6636). RNA samples were labelled with Cy5 and the reference E. coli K-12 MG1655 122 genomic DNA was labelled with Cy3. In total, two independent biological replicates were performed that were hybridised in duplicate (technical replicates) giving four replicates. After hybridization and 123 124 image capture, data were extracted from the raw image files using Agilent Feature Extraction v11.5 125 software and analyzed using GeneSpring v7.3.1. Transcriptomic data have been deposited with 126 ArrayExpress (accession E-MTAB-5220).

127

Purification of SlyA and Western blotting. Cultures (500 ml auto-induction medium 128 129 supplemented with ampicillin in 2 L conical flasks) of E. coli BL21 (λ DE3) pGS2469 were grown at 37°C for 24 h with shaking (250 rpm). Bacteria were collected by centrifugation, the pellet was re-130 suspended in 15 ml of breakage buffer (20 mM Tris-HCl, 500 mM NaCl, 5% v/v glycerol, pH 7.5), 131 132 the bacteria were lysed by two passages through a French pressure cell (16,000 psi) and the extract clarified by centrifugation (27,000 g, 15 min, 4°C). The His-tagged SlyA protein was isolated from 133 134 the cell-free extract by affinity chromatography on a HiTrap chelating column (1 ml) attached to an 135 AKTA prime according to the standard manufacturer's protocol (GE Healthcare). The eluted SlyA was buffer exchanged into 20 mM Tris-HCl, pH 7.4 containing 200 mM NaCl by repeated dilution 136

137 and Vivaspin 6 concentration (Sartorius Stedim Biotech). The protein was judged to be >90% pure by 138 Coomassie blue-stained SDS-PAGE and protein concentration was estimated by the BioRad protein 139 reagent protocol [29]. SlyA protein was detected by Western blotting after separation of polypeptides 140 by SDS-PAGE and electrophoretic transfer (100 V for 1 h; transfer buffer: 5.8 g L⁻¹ Tris, 2.9 g L⁻¹ 141 glycine, 20% v/v methanol, 0.037% w/v SDS) to Hybond-C Extra nitrocellulose membranes (GE 142 Healthcare). The membranes were soaked in a blocking solution, which contained 5% w/v dried 143 skimmed milk in PBS (10 mM phosphate buffer, 137 mM NaCl, 2.7 mM KCl, pH 7.4) and 0.05% v/v Tween 20, for 16 h at 4°C. The blocking solution was then removed and the membranes washed in 144 PBS containing 0.05% v/v Tween 20 before exposure to a 1:1000 dilution of the SlyA antibody 145 (raised in rabbit and provided by Prof. Ian Blomfield, University of Kent) in blocking solution for 1 h 146 at room temperature. After four washes with PBS containing 0.05% v/v Tween 20, the membranes 147 148 were soaked in blocking solution containing anti-rabbit secondary antibody provided in the Pierce ECL Western Blotting kit and the presence of SlyA was visualized according to the manfacturer's 149 150 standard protocol (Thermo Scientific).

151

Electrophoretic mobility shift assays (EMSA). The LightShift Chemiluminescent EMSA kit 152 153 (Thermo Scientific) was used according the manufacturer's intstructions. Biotin-labelled DNA of 154 target promoter regions was amplified from genomic DNA using the appropriate oligonucleotide primer pairs (Table 1). The core binding assays (20 µl) contained: 2 µl 10x binding buffer (100 mM 155 Tris-HCl, pH 7.5, containing 500 mM KCl and 10 mM dithiothreitol and 1 µg poly (dI•dC). The 156 DNA concentration was ~1 nM and the concentration of SlyA ranged from 0 to 500 nM as indicated. 157 158 Mixtures were incubated at 25°C for 30 min before separation of SlyA-DNA complexes by native gel electrophoresis, followed by transfer to Hybond-N+ nylon membranes, UV-crosslinking for 60 s at 159 160 120 mJ cm⁻² and detection of labelled DNA using the Nucleic Acid Detection Module (Thermo 161 Scientific).

162

163 **RESULTS AND DISCUSSION**

164 Enhanced expression of *slyA* in *E. coli* K-12 MG1655 results in altered abundance of

165 transcripts from 30 operons

Previous work has shown that SlyA directly activates the expression of two genes in *E. coli* K-12 (*hlyE* and *fimB*) by antagonising H-NS repression [20, 21, 23]. However, in *S. enterica* serovar Typhimurium the influence of SlyA is much more extensive, with at least 31 regulated genes resulting in hypersensitivity to reactive oxygen species and attenuation in infection models [7, 8, 12]. The initial aim of this work was to apply transcript profiling to determine the extent of the *E. coli* K-12 MG1655 SlyA regulon by comparison of steady-state glucose-limited aerobic chemostat cultures of wild-type and *slyA* mutant strains. Comparison of transcript profiles of wild-type and *slyA* mutant

cultures grown at a dilution rate of 0.5 h⁻¹ (equivalent to a doubling time of 1.4 h) revealed no 173 significant (≥ 2 -fold; $p \leq 0.05$) changes in transcript abundance. Because SlyA translation might be 174 enhanced at low growth rates, due to its unusual UUG start codon [21], steady-state cultures at 175 176 dilution rates 0.2, 0.1 and 0.05 h⁻¹ were established (equivalent to doubling times of 3.5, 6.9 and 13.8 h, respectively). However, once again, when the transcript profiles and growth characteristics of the 177 178 wild-type and *slyA* mutant cultures were compared no significant differences were detected. These observations indicated that, under the conditions tested, deletion of the *slyA* gene had no significant 179 180 effect on gene expression in E. coli K-12 MG1655, even at low growth rates.

181 Anti-SlyA serum was used to determine whether SlyA was detectable in E. coli K-12 MG1655 cells grown in glucose-limited chemostats at a dilution rate of 0.2 h⁻¹. In accordance with 182 the transcript profiling, SlyA was not detected (Fig. 1). This suggests that the expression of SlyA is 183 184 regulated and switched on under conditions other than those imposed here; for example, SlyA protein 185 has been detected by Western blotting extracts from E. coli batch cultures grown in minimal medium 186 with glycerol as the carbon and energy source [21]. To overcome any regulatory barrier to identifying 187 genes potentially controlled by SlyA, a plasmid (pGS2468) to express *slyA* under the control of its own promoter was constructed. Western blotting showed that SlyA protein was now readily 188 detectable in the transformed E. coli K-12 MG1655 cells grown in glucose-limited chemostats at a 189 190 dilution rate of 0.2 h⁻¹ (Fig. 1). The growth characteristics of the vector control and the *slyA* expression strains were essentially the same, with similar yields $(1.4 \pm 0.2 \text{ g cell dry weight per litre})$ 191 192 and no detectable glucose or over-metabolites in the culture supernatants. Therefore, the transcript profiling experiments were carried out with these strains grown in aerobic glucose-limited chemostats 193 194 at a dilution rate of 0.2 h^{-1} . The transcript profile of the SlyA over-production strain was significantly different from that of the vector control. Transcripts representing 27 operons were increased in 195 196 abundance and 3 were decreased (Table 2; Fig. 1c). The transcripts exhibiting decreased abundance 197 were: the sgc operon (sgcXBCQAER), which encodes a phosphotransferase system for the uptake of 198 an unknown sugar; *fecIR* the membrane-bound sensor (FecR) that receives signals from the outer 199 membrane ferric citrate uptake receptor (FecA) for transmission to FecI (σ^{19}), which activates 200 transcription of the *fecABCDE* operon encoding components of a cytoplasmic membrane bound ferric 201 citrate uptake system; and yecH, which encodes a predicted protein of unknown function (Table 2) 202 [30, 31].

Amongst the up-regulated transcripts were the previously identified SlyA-regulated gene *hlyE* and *slyA* itself (Table 2). The latter was not surprising as the *slyA* gene was present in multi-copy, but despite this the *slyA* transcript only increased ~3-fold in abundance, yet the SlyA protein level increased from being undetectable in the control to a level equivalent to ~1.5 μ M in the cytoplasm (based on the dry weight of *E. coli* being 3 x 10⁻¹³ g with an aqueous volume of 7 x 10⁻¹³ ml per cell; [32]). The relatively low level of induction of the *slyA* transcript when present in multi-copy but much greater induction of SlyA protein suggests that the *slyA* promoter is subject to auto-regulation, consistent with the reported SlyA binding at the *slyA* promoter [19]. It was also notable that the *ydhI*-K operon, which is divergently transcribed from *slyA* and not present on the *slyA* expression plasmid, also exhibited enhanced transcript abundance, suggesting that SlyA is capable of activating expression from divergent promoters; an assertion supported by the enhanced abundances of the divergently

transcribed *hlyE* and *C0299* (encodes a small RNA) genes in the presence of SlyA (Table 2).

215 Twenty-four of the 30 operons (~80%) that showed altered transcript abundance upon overproduction of SlyA have also been shown to be associated with H-NS binding sites (Table 2). Thus, it 216 217 appears that H-NS repressed genes are over-represented in the set of transcripts that increase in 218 abundance when SlyA is expressed, suggesting that SlyA acts by antagonizing H-NS repression at the corresponding promoters; a mechanism that is established for hlyE [20]. H-NS binds DNA by 219 220 recognizing the structure of A-T-rich minor grooves and silences the expression of horizontally 221 acquired A-T-rich genes (reviewed by [33]). H-NS is thus considered crucial in permitting the 222 acquisition of new genes whilst counteracting the potentially detrimental effects of inappropriate 223 expression of these genes. Counter-silencing by H-NS antagonists, such as SlyA, provides a route to 224 integrate expression of the genes into the regulatory circuits of E. coli under appropriate conditions. 225 Horizontally acquired genes are located within genomic islands, which are regions of bacterial 226 chromosomes containing that are often associated with drug resistance, metabolic adaptability, stress 227 tolerance and pathogenesis. Genomic islands can be recognized by their sequence composition and 228 increased transcript start point densities [32]. The analysis tools GIST (Genomic-island Identification 229 by Signals of Transcription) and IslandViewer have been used to map the genomic islands of E. coli 230 K-12 MG1655 [34]. Notably, 13 of the 30 differentially regulated operons overlapped predicted 231 genomic islands, suggesting a general role for SlyA in the counter-silencing of H-NS repressed 232 horizontally acquired genes under conditions when *slyA* is up-regulated (Table 2).

233 The H-NS-repressed casABC operon was up-regulated by SlyA (Table 2). This operon encodes proteins involved in maintaining and utilising the library of foreign genetic elements 234 235 interspersed between CRISPR sequences which act as the immune system memory of Bacteria and 236 Archaea [35]. CRISPR loci, in general, consist of closely spaced direct repeats separated by short spacer regions of variable sequence. Spacer regions mostly correspond to sections of foreign plasmid 237 238 or viral sequences which have been integrated. The CRISPR loci are found adjacent to the casABC 239 operon. The fact that the casABC operon was significantly up-regulated by SlyA suggests that this 240 regulator may contribute to viral resistance and immunity in E. coli K-12 MG1655.

Other transcripts that exhibited increased abundance in the presence of SlyA were associated with uptake and metabolism of phenylacetic acid (*paaA-K*), utilization of alkanesulfonates as alternative sulfur sources (*ssuEADCB*; divergently transcribed from the *elf* operon; see below), a cryptic galactosamine transport and catabolism system (*agaS-I*) and a 2-O- α -mannosyl-D-glycerate phosphotransferase and α -mannosidase (Table 2) [36-39]. Hence, it appears that SlyA plays a role in 246 regulating systems that expand the repertoire of substrates utilized by E. coli. Increased abundance of 247 the *mdtM* transcript suggests a role for SlyA in enhancing expression of this multidrug transporter that 248 confers resistance to ethidium bromide and chloramphenicol with mutants exhibiting attenuated 249 growth at alkaline pH [40]. However, simple growth inhibition studies suggested that *slyA* expression 250 led to increased sensitivity to chloramphenicol (growth yield after 6 h at 37°C in Luria Bertani broth was lowered to $\sim 50\%$ by 2 µg ml⁻¹ for the wild-type carrying the empty vector compared to 1 µg ml⁻¹ 251 for the wild-type carrying the slyA expression plasmid), rather than increased resistance, perhaps 252 253 reflecting the complexity of the phenotype of the *slyA* expression strain.

254 Several of the SlyA-regulated operons code for proteins involved in membrane function. In S. 255 enterica serovar Typhimurium the majority of genes affected by SlyA encode proteins associated with the bacterial cell envelope and are important for virulence and survival within murine macrophages. 256 257 Although it has been previously shown that the majority of genes regulated by SlyA in S. enterica 258 serovar Typhimurium are not present in E. coli K-12 [12, 15], a similar propensity for cell envelope 259 proteins being regulated by the E. coli SlyA was evident here. Thirteen (43%) of the 30 operons that 260 exhibited altered expression in SlyA-expressing bacteria were associated with cell-surface/membrane 261 functions (Table 2).

The *gspC-O* operon is cryptic membrane-associated, H-NS-repressed, transcription unit that was up-regulated by SlyA (Table 2). The *gspC-O* operon encodes a Type-II Secretion System (T2SS) for the export of endogenous proteins and formation of structural elements of the Gsp secreton, which is thought to facilitate the export of the endogenous endochitinase ChiA, a product of another H-NS silenced gene [41, 42].

267 Amongst the transcripts with increased abundance in the SlyA over-producing strain were 268 four cryptic operons (elfADCG-ycbUVF, sfmHF, yehDCBA and yadN) encoding fimbrial-like 269 adhesins (Table 2). These four operons were amongst seven putative chaperone-usher fimbrial 270 systems shown to be poorly expressed under laboratory conditions by Korea et al. [43]. Nevertheless, 271 when these operons were individually expressed by placing them under the control of a constitutive 272 promoter six were shown to be functional and expression of the *elf* (ycb), yad and yeh operons 273 resulted in enhanced biofilm formation on abiotic surfaces, whereas sfm promoted binding to 274 eukaryotic cells [43]. Moreover, all four operons were repressed by H-NS. The increased 275 abundances of the *elf*, *sfm*, *yad* and *yeh* transcripts upon expression of SlyA is consistent with the cryptic status of these genes under normal laboratory conditions and suggests that these chaperone-276 277 usher fimbriae are functional under environmental conditions that enhance *slyA* expression such that 278 SlyA can operate as an H-NS antagonist (Table 2).

279

280 SlyA over-production is associated with enhanced biofilm formation

The observation that SlyA increased transcription of four cryptic fimbrial-like adhesins suggested that
 the SlyA over-producing strain should exhibit enhanced biofilm production. This was tested using

static cultures of *E. coli* K-12 MG1655 transformed with pET28a (control) or the *slyA* expression
plasmid pGS2468 in conditions that mirrored the transcript profiling experiment. The data showed a
4-fold increase in biofilm formation when *slyA* was over-expressed, consistent with the transcript
profiling data (Fig. 2).

287

288 Identification of new *E. coli* K-12 MG1655 operons that are directly regulated by SlyA

289 The changes in transcript profiles that were observed upon over-production of SlyA could result from 290 direct interaction of SlyA with the promoter regions of the corresponding genes or indirectly via SlyA-regulated factors. For example, one of the genes up-regulated upon SlyA over-production, *leuO*, 291 encodes a transcriptional regulator that, like SlyA, operates by antagonising H-NS regulation [44, 45]. 292 Fourteen (52%) of the 27 transcripts that were increased in abundance when SlyA was expressed in E. 293 294 coli K-12 MG1655 were associated with LeuO binding sites identified in the SELEX-chip study of 295 Shimada et al. [44]. This strong correlation could arise from; (1) the positive effect SlyA has on the 296 expression of *leuO* resulting in an increase in expression of the entire LeuO regulon, i.e. indirect 297 regulation by SlyA; or (2) SlyA and LeuO have overlapping regulons as a consequence of the fact they both operate by antagonising H-NS-mediated repression. To further investigate the extent of 298 299 direct SlyA-mediated regulation in E. coli K-12 MG1655 binding of SlyA to ten promoter regions 300 was examined by electrophoretic mobility shift assays (EMSA).

301 Amongst the transcripts differentially regulated by over-production of SlyA there were three 302 arranged as divergent operons (Fig. 3). Binding of SlyA at the hlyE-C0299 intergenic region was 303 shown previously (Fig. 3) [20]. Two other examples of SlyA-activated divergent operons (slyA-304 ydhIJK and ssuE-B-elfADCG-ycbUVF) were shown to bind SlyA in EMSA (Fig. 3). Furthermore, SlyA bound at the casA, fecIR, gspCDEF, leuO, mdtM and paaA-K promoters (Fig. 3). The K_{d(app)} 305 306 values for SlyA binding at these promoters were similar at ~50-100 nM. These experiments indicate 307 that these operons are likely to be directly regulated by SlyA. The sgcXBCQ-sgcAER genes are separated by a sRNA ryjB on the opposite DNA strand (Fig. 3). It is suggested that the sgcXBCOAER 308 is a single transcription unit, but there is no high quality evidence to support this suggestion [46]. 309 310 Therefore, both the region upstream of sgcX and the intergenic region between sgcQ and sgcA were used in EMSAs with the SlyA protein. No specific interaction was observed with the region upstream 311 312 of sgcA but interaction, albeit weaker than that observed for the promoter regions analyzed above, was 313 observed when the DNA upstream of sgcX was tested (Fig. 3). These observations suggest that 314 sgcXBCQAER is a single SlyA-repressed transcription unit.

The EMSA experiments indicate that SlyA binds Pssu, Pcas, Ppaa, Pelf, PleuO and Pgsp, all of which are promoter regions of genes or operons proposed to be part of the LeuO regulon (Table 2). This suggests that, perhaps because of the similarity in their mode of action, i.e. antagonizing H-NS repression, the SlyA and LeuO regulons substantially overlap such that upon activation by their respective signals a similar transcriptional response is elicited. 320 A consensus binding site (TTAGCAAGCTAA) for the Salmonella enterica serovar 321 Typhimurium LT2 SlyA protein was proposed based on footprinting and a limited SELEX analysis 322 [10]. This consensus was further analyzed by site-directed mutagenesis, which suggested the 323 consensus sequence TTAN₆TAA [9]. All the DNA fragments that bound *E. coli* SlyA in EMSAs (Fig. 324 3) possessed DNA sequences similar to the previously proposed consensus sequences (Table 3). Site-325 directed replacement amino acid residues of Salmonella enterica serovar Typhimurium LT2 SlyA 326 identified 16 locations that impaired DNA-binding [9], all these amino acids are conserved in the E. 327 coli SlyA protein, suggesting that these closely related proteins recognize similar DNA motifs.

328

329 Concluding remarks

SlyA proteins have been shown to play important roles in regulating gene expression in a wide range 330 331 of bacterial species. The most common mechanism for SlyA-mediated activation of gene expression 332 is through antagonism of H-NS repression. Here transcript profiling has revealed the breadth of the 333 SlyA regulon (directly and indirectly regulated genes) in E. coli K-12 MG1655 cultures grown under 334 precisely controlled conditions such that any potential effects associated with changes in growth rate/growth phase could not confound the interpretation of the data obtained. Enhanced transcript 335 336 abundance for several cryptic fimbrial operons in a SlyA over-producing strain and an over-337 representation of H-NS repressed genes were consistent with the current model of SlyA-mediated 338 gene activation. The SlyA protein was shown to bind at 9 intergenic regions controlling the 339 expression of 11 operons, thus expanding the number of known directly SlyA-regulated genes in E. 340 coli MG1655 from 2 to 13.

341

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346

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350

351 CONFLICTS OF INTEREST

- 352 The authors declare no conflicts of interest.
- 353
- 354 **REFERENCES**
- 355

- Finn RD, Bateman A, Clements J, Coggill P, Eberhardt RY *et al.* Pfam: the protein families
 database. *Nucl Acids Res* 2014;42:D222-D230.
- 2. Perera IC, Grove A. Molecular mechanisms of ligand-mediated attenuation of DNA-binding by
 MarR family transcription regulators. *J Mol Cell Biol* 2010;2:243-254.
- 360 3. Martin RG, Rosen JL. Binding of the purified multiple antibiotic-resistance repressor protein
 361 (MarR) to *mar* operator sequences. *Proc Natl Acad Scis U S A* 1995;92:5456-5460.
- 4. Oh SY, Shin JH, Roe JH. Dual role of OhrR as a repressor and an activator in response to organic
 hydroperoxides in *Streptomyces coelicolor*. *J Bacteriol* 2007;189:6284-6292.
- 364 5. Cathelyn JS, Ellison DW, Hinchliffe SJ, Wren BW, Miller V. The RovA regulons of *Yersinia* 365 *enterolytica* and *Yersinia pestis* are distinct: evidence that many RovA-regulated genes were
 366 acquired more recently than the core genome. *Mol Microbiol* 2007;66:189-205.
- 367 6. Wilkinson, S. P. & Grove, A. (2006). Ligand-responsive transcriptional regulation by members of
 368 the MarR family of winged helix proteins. *Curr Issues Mol Biol* 8, 51-62.
- **7. Buchmeier N, Bossie S, Chen, CY, Fang FC, Guiney DG** *et al.* SlyA, a transcriptional regulator
 of *Salmonella typhimurium*, is required for resistance to oxidative stress and is expressed in the
 intracellular environment of macrophages. *Infect Immun* 1997;65:3725-3730.
- 8. Libby SJ, Goebel W, Ludwig A, Buchmeier N, Bowe F *et al.* A cytolysin encoded by *Salmonella*is required for survival within macrophages. *Proc Natl Acad Sci U S A* 1994; 91:489-493.
- **9. Haider F, Lithgow JK, Stapleton MR, Norte VA, Roberts RE** *et al.* DNA recognition by the *Salmonella enterica* serovar Typhimurium transcription factor SlyA. *Intl Microbiol* 2008;11:245250.
- **10. Stapleton MR, Norte VA, Read RC, Green J.** Interaction of the *Salmonella typhimurium*transcription and virulence factor SlyA with target DNA and identification of members of the SlyA
 regulon. *J Biol Chem* 2002;277:17630-17637.
- 11. Colgan AM, Kröger C, Diard M, Hardt WD, Puente J L *et al.* The impact of 18 ancestral and
 horizontally-acquired regulatory proteins upon the transcriptome and sRNA landscape of
 Salmonella enterica serovar Typhimurium. *PLoS Genetics* 2016;12;e1006258.
- 12. Navarre WW, Halsey TA, Walthers D, Frye J, McClelland M et al. Co-regulation of
 Salmonella enterica genes required for virulence and resistance to antimicrobial peptides by SlyA
 and PhoP/PhoQ. Mol Microbiol 2005;56:492-508.
- **13. Norte VA, Stapleton MR, Green J.** PhoP-responsive expression of the *Salmonella enterica* serovar Typhimurium *slyA* gene. *J Bacteriol* 2003;185:3508-3514.
- 14. Perez JC, Latifi T, Groisman EA. Overcoming H-NS-mediated transcriptional silencing of
 horizontally acquired genes by the PhoP and SlyA proteins in *Salmonella enterica*. *J Biol Chem* 2008;283:0773-10783.

- 15. Spory A, Bosserhoff A, von Rhein C, Goebel W, Ludwig A. Differential regulation of multiple
 proteins of *Escherichia coli* and *Salmonella enterica* serovar Typhimurium by the transcriptional
 regulator SlyA. *J Bacteriol* 2002;184:3549-3559.
- **16. Dalebroux ZD, Swanson MS.** ppGpp: magic beyond RNA polymerase. *Nat Rev Microbiol* 2012;10:203-212.
- 396 17. Zhao G, Weatherspoon N, Kong W, Curtiss R, Shi YX. A dual-signal regulatory circuit
 397 activates transcription of a set of divergent operons in *Salmonella typhimurium*. *Proc Natl Acad*398 *Sci U S A* 2008;105:20924-20929.
- 18. Chalabaev S, Chauhan A, Novikov A, Iyer P, Szczesny M et al. Biofilms formed by gramnegative bacteria undergo increased lipid A palmitoylation, enhancing *in vivo* survival. *MBio*2014;5:pii:e01116-14.
- 402 19. Corbett D, Bennett HJ, Askar H, Green J, Roberts IS. SlyA and H-NS regulate transcription of
 403 the *Escherichia coli* K5 capsule gene cluster, and expression of *slyA* in *Escherichia coli* is
 404 temperature-dependent, positively autoregulated, and independent of H-NS. *J Biol Chem*405 2007;282:33326-33335.
- 20. Lithgow JK, Haider F, Roberts IS, Green J. Alternate SlyA and H-NS nucleoprotein complexes
 control *hlyE* expression in *Escherichia coli* K-12. *Mol Microbiol* 2007;66:685-698.
- 408 21. McVicker G, Sun L, Sohanpal BK, Gashi K, Williamson RA *et al.* SlyA protein activates *fimB*409 gene expression and type 1 fimbriation in *Escherichia coli* K-12. *J Biol Chem* 2011;286:32026410 32035.
- 22. Xue P, Corbett D, Goldrick M, Naylor C, Roberts IS. Regulation of expression of the region 3
 promoter of the *Escherichia coli* K5 capsule gene cluster involves H-NS, SlyA and a large 5'
 untranslated region. *J Bacteriol* 2009;191:838-1846.
- 414 23. Wyborn NR, Stapleton MR, Norte VA, Roberts RE., Grafton J et al. Regulation of
 415 Escherichia coli hemolysin E expression by H-NS and Salmonella SlyA. J Bacteriol
 416 2004;186:1620-1628.
- 417 24. Sambrook J, Russell DW. Molecular cloning a laboratory manual, 3rd ed., Cold Spring. Harbor
 418 NY: Cold Spring Harbor Laboratory; 2001.
- 419 25. Evans CGT, Herbert D, Tempest DW. The continuous cultivation of micro-organisms. *Meth* 420 *Microbiol* 1970;2;278-327.
- 421 26. Studier FW. Protein production by auto-induction in high-density shaking cultures. *Prot Express* 422 *Purific* 2005;41:207-234.
- **27. Tagliabue L, Antoniani D, Maciag A, Bocci P, Raffaelli N** *et al.* The diguanylate cyclase YddV
 controls production of the exopolysaccharide poly-N-acetylglucosamine (PNAG) through
 regulation of the PNAG biosynthetic *pgaABCD* operon. *Microbiology* 2010;156:2901-2911.

- 28. Rolfe MD, Ter Beek A, Graham AI, Trotter EW, Shahzad Asif HM *et al.* Transcript profiling
 and inference of *Escherichia coli* K-12 ArcA activity across the range of physiologically relevant
 oxygen concentrations. *J Biol Chem* 2011;286:10147-10154.
- 429 29. Bradford MM. Rapid and sensitive method for quantification of microgram quantities of protein
 430 utilizing principle of protein-dye binding. *Anal Biochem* 1976;72:248-254.
- **30. Braun V, Mahren S, Sauter A.** Gene regulation by transmembrane signalling. *Biometals*2006;19:103-113.
- 433 **31. Reizer J, Reizer A, Saier MH.** Is the ribulose monophosphate pathway widely distributed in
 434 bacteria? *Microbiology* 1997;143:2519-2520.
- 32. Sundararaj S, Guo A, Habibi-Nazhad B, Rouani M, Stothard P *et al.* The CyberCell database
 (CCDB): a comprehensive, self-updating, relational database to coordinate and facilitate *in silico*modeling of *Escherichia coli*. *Nucl Acids Res* 2004;32:D293-D295.
- 438 33. Navarre WW. The impact of gene silencing on horizontal gene transfer and bacterial evolution.
 439 Adv Microb Physiol 2016;69:157-186.
- 440 34. Huang Q, Cheng, X, Cheung MK, Kiselev SS, Ozoline ON *et al.* High-density transcriptional
 441 initiation signals underline genomic islands in bacteria. *PLoS One* 2012;7:e33759.
- 442 35. Horvath P, Barrangou R. CRISPR/Cas, the immune system of Bacteria and Archaea. *Science*443 2010;327:167-170.
- 36. Brinkkötter A, Klöβ, H, Alpert CA, Lengeler JW. Pathways for the utilization of N-acetylgalactosamine and galactosamine in *Escherichia coli*. *Mol Microbiol* 2000;37:125-135.
- 446 37. Eichhorn E, van der Ploeg JR, Leisinger T. Deletion analysis of the *Escherichia coli* taurine
 447 and alkanesulfonate transport systems. *J Bacteriol* 2000;182:2687-2695.
- 38. Ismail W, Mohamed ME, Wanner BL, Datsenko KA, Eisenreich W et al. Functional
 genomics by NMR spectroscopy phenylacetate catabolism in *Escherichia coli. Eur J Biochem*2003;270:3047-3054.
- 39. Sampaio MM, Chevance F, Dippel R, Eppler T, Schlegel A *et al.* Phosphotransferase-mediated
 transport of the osmolyte 2-O-alpha-mannosyl-D-glycerate in *Escherichia coli* occurs by the
- product of the *mngA* (*hrsA*) gene and is regulated by the *mngR* (*farR*) gene product acting as
 repressor. *J Biol Chem* 2004;279:5537-5548.
- 40. Holdsworth SR, Law CJ. (2013). Multidrug resistance protein MdtM adds to the repertoire of
 antiporters involved in alkaline pH homeostasis in *Escherichia coli*. *BMC Microbiol* 13, 113.
- 457 **41. Francetic O, Belin D, Badaut C, Pugsley AP.** Expression of the endogenous type II secretion
 458 pathway in *Escherichia coli* leads to chitinase secretion. *EMBO J* 2000;19:6697-6703.
- 459 **42. Francetic O, Pugsley AP.** The cryptic general secretory pathway (*gsp*) operon of *Escherichia coli*
- 460 K-12 encodes functional proteins. *J Bacteriol* 1996;178:3544-3549.

- 461 43. Korea CG, Badouraly R, Prevost MC, Ghigo JM, Beloin C. *Escherichia coli* K-12 possesses
 462 multiple cryptic but functional chaperone-usher fimbriae with distinct surface specificities. *Env*463 *Microbiol* 2010;12:1957-1977.
- 464 44. Shimada T, Bridier A, Briandet R, Ishihama A. Novel roles of LeuO in transcription regulation
 465 of *E. coli* genome: antagonistic interplay with the universal silencer H-NS. *Mol Microbiol*466 2011;82:378-397.
- 467 45. Stratmann T, Madhusudan S, Schnetz K. Regulation of the *yjjQ-bglJ* operon, encoding LuxR468 type transcription factors, and the divergent *yjjP* gene by H-NS and LeuO. *J Bacteriol*469 2008;190:926-935.
- 470 46. Keseler IM, Mackie A, Peralta-Gil M, Santos-Zavaleta A, Gama-Castro S *et al.* EcoCyc:
 471 fusing model organism databases with systems biology. *Nucl Acids Res* 2013;41:D605-D612.
- 472 47. Blattner FR, Plunkett G, Bloch CA, Perna NT, Burland V *et al.* The complete genome
 473 sequence of *Escherichia coli* K-12. *Science* 1997; 277:1453-1462.
- 474 48. Datsenko KA, Wanner BL One-step inactivation of chromosomal genes in *Escherichia coli* K475 12 using PCR products. *Proc Natl Acad Sci U S A* 2000; 97:6640-6645.
- 476 49. Grainger DC, Hurd D, Goldberg MD, Busby SJW. Association of nucleoid proteins with
 477 coding and non-coding segments of the *Escherichia coli* genome. *Nucl Acids Res* 2006;34:4642–
 478 4652.
- 479 50. Kahramanoglou C, Seshasayee ASN, Prieto AI, Ibberson D, Schmidt S *et al.* Direct and
 480 indirect effects of H-NS and Fis on global gene expression control in *Escherichia coli*. *Nucl Acids*481 *Res* 2011; 39:2073–2091.
- 51. Oshima T, Ishikawa S, Kurokawa K, Aiba H, Ogasawara N. *Escherichia coli* histone-like
 protein H-NS preferentially binds to horizontally acquired DNA in association with RNA
 polymerase. *DNA Res* 2006;13:141–153.

Table 1 Bacterial strains, plasmids and oligonucleotides

Strain or plasmid	Relevant characteristics ^a	Reference or source
Bacterial strain		
E. coli BL21	E. coli BL21 lysogen for inducible (IPTG) expression of the T7 RNA	Novagen
(λDE3)	polymerase	6
E. coli JRG6457	E. coli MG1655 slyA	This work
<i>E. coli</i> JRG6636	<i>E. coli</i> MG1655 pGS2468	This work
<i>E. coli</i> JRG6072	<i>E. coli</i> MG1655 pKD46	This work
<i>E. coli</i> MG1655	Genome sequenced parental strain	[47]
Plasmid	Schone sequenced parental strain	[+/]
pET28a	Multi-copy plasmid; Kan ^R	Novagen
pGS2468	pET28a derivative for expression of <i>slyA</i> from the <i>slyA</i> promoter; Kan ^R	This work
pGS2469	pLATE-51 derivative for over-production of SlyA; Amp ^R	This work
pKD4	Source of kanamycin resistance cassette; Amp ^R , Kan ^R	[48]
pKD46	Plasmid for inducible (L-arabinose) expression of the λ red recombinase;	[48]
pite io	Amp ^R , T ^s	[10]
pLATE-51	Expression vector for production of His-tagged proteins; Amp ^R	Thermo Scientific
Oligonucleotide	Expression vector for production of fils-tagged proteins, Amp	memo selenune
TC7	TAAAGCCGCATAATATCTTAGCAAGCTAATTATAAGGAGATTA	This work
10/	CACGTCTTGAGCGATT; creation of <i>slyA</i> mutant	THIS WOLK
TC8	TTGCGTGTGGTCAGGTTACTGACCACACGCCCCTTCATTCA	This work
100		THIS WOLK
TC9	ATGAATATCCTCCTTAG; creation of <i>slyA</i> mutant	This work
TC10	CTGACGGTAACCAAATGCAG; PCR of <i>slyA</i> locus	This work
TC49	TTTGCGTGTGGTCAGGTTAC; PCR of <i>slyA</i> locus [Btn]ACTCTCTCCTTATAACCAATTG; forward primer for PCR of	This work
1049		This work
TC50	biotin (Btn)–labelled 355 bp intergenic region between <i>ssuE</i> and <i>elfA</i>	
TC50	CGTTATCATCCTGATCTCTT; reverse primer for use with TC49	This work
TC51	[Btn]TGGTGAATATTATTGATCAATTAAT; forward primer for PCR	This work
TC 52	of biotin (Btn)–labelled 344 bp intergenic region between <i>leuO</i> and <i>leuL</i>	7 71 · 1
TC52	ACTTAACTCCACTGTCACACTTAA; reverse primer for use with	This work
T	TC51	T I 1
TC53	[Btn]TTGTTCTCCTTCATATGCTC; forward primer for PCR of biotin	This work
	(Btn)–labelled 414 bp intergenic region between <i>casA</i> and <i>cas3</i>	
TC54	CTTCGGGAATGATTGTTATC; reverse primer for use with TC53	This work
TC55	[Btn]TGTTGCTAATAGTTAAATCGC; forward primer for PCR of	This work
	biotin (Btn)-labelled 257 bp intergenic region between paaA and paaZ	
TC56	GTCATCACCTTTACGATTCC; reverse primer for use with TC55	This work
TC57	[Btn]AACAAACAACTCCTTGTCCG; forward primer for PCR of	This work
	biotin (Btn)–labelled 400 bp region upstream of <i>mdtM</i>	
TC58	CCCCGAGGCGCTTTCCAGGC; reverse primer for use with TC57	This work
TC59	[Btn]AGAACTTCCTGTTTTAATTATTG; forward primer for PCR of	This work
	biotin (Btn)-labelled 179 bp intergenic region between gspA and gspC	
TC60	GATGTATGTTCTAATAAAATAGATTG; reverse primer for use with	This work
	TC59	
TC61	[Btn]CCGTCGTTGACTCCATGC; forward primer for PCR of biotin	This work
	(Btn)–labelled 130 bp intergenic region between $sgcA$ and $sgcQ$	
TC62	GATGGGGATAAGCAGAGC; reverse primer for use with TC61	This work
TC63	[Btn]GCGGAGTGCATCAAAAGT; forward primer for PCR of biotin	This work
	(Btn)–labelled 291 bp intergenic region between <i>fec1</i> and <i>insA-7</i>	
TC64	GCAAGCACCTTAAAATCAC; reverse primer for use with TC63	This work
TC65	[Btn]TTTCATCTCCTTATAATTAGCTT; forward primer for PCR of	This work
	biotin (Btn)–labelled 200 bp intergenic region between <i>slyA</i> and <i>ydhI</i>	
TC66	AAAGTAGATTCCTTTACGACC; reverse primer for use with TC65	This work
TC70	[Btn]AGCTATCTCCGTAGACCGT; forward primer for PCR of biotin	This work
	(Btn)–labelled 400 bp region upstream of $sgcX$	
TC71	GATTATCTATACTCCCTCTGAATC; reverse primer for use with	This work
10/1	TC70	into work

^aAmp^R, ampicillin resistant; Kan^R, kanamycin resistant; T^s, temperature sensitive replication

Operonª	Fold- change ^b	Gene function ^c	H-NS regulon ^d	LeuO regulon ^e	Overlap with genomic island ^f
ybeT	4.1	conserved outer membrane protein	К	√	
trkG	3.8	Rac prophage potassium transporter subunit	K,O		IV
ssuEADCB	3.6	aliphatic sulfonate transport and metabolism	G,K,O	✓	
yehDCBA	3.6	chaperone-usher fimbrial operon (cryptic)	K,O	✓	GIST
mngAB	3.4	2-O- α -mannosyl-D-glycerate PTS and α -mannosidase		✓	
casABC	3.3	CRISPR associated genes	К	\checkmark	GIST, IV
yghS	3.1	predicted protein with nucleoside triphosphate hydrolase domain	K,O	\checkmark	
slyA	3.0	DNA-binding transcriptional activator	0		
yfbN	2.8	predicted protein	K,O	\checkmark	IV
paaA-K	2.8	phenylacetic acid degradation			
ybeU-hscD	2.8	predicted tRNA ligase and chaperone	K,O	\checkmark	
elfADCG-	2.7	predicted fimbrial-like adhesin	G,K,O	\checkmark	
ycbUVF		protein (cryptic)			
ygeG	2.7	predicted chaperone	G,K,O		GIST
crfC-yjcZ	2.6	clamp-binding sister replication fork co-localization protein and predicted protein	K,O		
sfmHF	2.6	protein predicted fimbrial-like adhesin protein (cryptic)	0		IV
agaS-kbaY-	2.5	predicted galactosamine-transport and		\checkmark	
aga B CDI		metabolism (cryptic)			
ydhYV-T	2.5	predicted oxidoreductase	G,K,O		GIST
yiiE	2.5	predicted transcriptional regulator	K,O		
mdtM	2.5	multidrug efflux system protein			
leuO	2.5	DNA-binding transcriptional activator	G,K,O	\checkmark	GIST
<i>C0299</i>	2.4	sRNA C0299	0		
ycj MN -V	2.4	predicted sugar transporter and metabolism	К,О	~	
yadN	2.4	predicted fimbrial-like adhesin protein (cryptic)	G,K,O	~	GIST
gspCDEF	2.4	type II secretion system (cryptic)	K,O	\checkmark	
ydhIJK	2.2	predicted proteins	0		
yfdM	2.1	CPS-53 (KpLE1) prophage predicted methyltransferase	0		GIST, IV

489 Table 2 Transcripts exhibiting altered abundance upon over-expression of *slyA* in *E. coli* MG1655

hlyE	2.0	hemolysin E (cryptic)	K,O	
yecH	0.5	predicted protein	0	GIST
sgcX B CQAER	0.5	predicted sugar transport and		IV
		metabolism		
fecI R	0.4	transcription regulation of ferric		IV
		citrate transport		

490

^aThe fold-change data shown are for the first gene in the operon except where indicated by bold typeface; note 491 that all genes in the operons followed the same pattern of regulation.

492 ^bFold-change (≥2-fold, *p*≤0.05) is the product of dividing the transcript abundance for the *slyA* over-expression

493 cultures by that for the control cultures.

494 °Gene functions as assigned in Ecocyc.org [46].

495 ^dGenes associated with H-NS binding were identified from Grainger et al. [49] (G), Kahramanoglou et al. [50] (K)

496 and Oshima et al. [51] (O).

497 eGenes located up- or down-stream of a LeuO binding site identified by Shimada et al. [44].

498 fGenes the overlap with genomic islands in *E. coli* K-12 MG1655 identified by GIST and/or IslandViewer (IV) [34].

Promoter region	Possible SlyA binding sites	Location of site relative to start codon
PcasA	TTATTG <u>A</u> ATTAA	100 bp upstream of <i>casA</i>
PssuE/elfA	TCA <u>G</u> GATGATAA	8 bp upstream of <i>elfA</i>
PgspC	TTATATTAGTAA	79 bp upstream of gspA
PpaaA	TTAAATC <u>GC</u> GAA	239 bp upstream of <i>paaA</i>
	TTA TA <u>AA</u> AA TA G	136 bp upstream of <i>paaA</i>
	TTACTT <u>A</u> A <u>C</u> TAT	81 bp upstream of <i>paaA</i>
PsgcX	TTATGCT <u>G</u> GGAA	336 bp upstream of <i>sgcX</i>
	TTTCA <u>A</u> CCATAA	188 bp upstream of <i>sgcX</i>
PfecI	TTA <u>G</u> A <u>AA</u> AACAA	109 bp upstream of <i>fecI</i>
PslyA	TTA <u>GCAAGC</u> TAA	22 bp upstream of <i>slyA</i>
	TTA <u>G</u> ATTAATAA	161 bp upstream of <i>slyA</i>
PleuO	TTAATGCATTAA	305 bp upstream of <i>leuO</i>
	TTAAAT <u>A</u> TATAA	297 bp upstream of <i>leuO</i>
PmdtM	TATA <u>CA</u> CCTTAA	249 bp upstream of <i>mdtM</i>

499 **Table 3** Candidate SlyA binding sites within the DNA fragments used for EMSA analyses

500

501 Sequences shown are those with the greatest similarity to the previously proposed consensus for the 502 *Salmonella enterica* serovar Typhimurium LT2 SlyA protein (Haider *et al.* [9]; **TTA**N₆**TAA**). Where

503 more than one possible site was present those with the greatest similarity to the consensus sequence

504 TTA<u>GCAAGC</u>TAA proposed by Stapleton *et al.* [10] are shown. Locations of sites are given as the

505 number of base pairs from the start codon of the specified gene to the centre of the proposed binding

506 site.

- 508 Figure legends
- 509

FIG. 1. Changes in the transcript profile of *E. coli* K-12 MG1655 over-producing SlyA. (a) 510 Transformation of *E. coli* K-12 MG1655 with a multi-copy plasmid expressing *slyA* under the 511 control of its own promoter results in detectable SIyA protein in lysed cell suspensions from 512 aerobic steady-state glucose-limited chemostat cultures. The upper panel shows the 513 Coomassie blue-stained SDS-polyacrylamide gel and the lower panel shows the relevant 514 region of a Western blot prepared with the same samples and loadings developed with SlyA 515 antiserum. The gels were loaded as follows: Lane M, SDS-PAGE markers (sizes, kDa, are 516 indicated); lanes 1 and 2, extracts from independent cultures of E. coli K-12 MG1655 517 transformed with the vector pET28a (SlyA^{WT}); lanes 3 and 4, extracts from independent 518 cultures of *E. coli* K-12 MG1655 transformed with the expression plasmid pGS2468 (SlyA⁺); 519 520 lane 5, purified (His)₆-SlyA (~10 ng protein loaded). (b) Western blot corresponding to the gel shown in (a). The locations of SlyA and purified (His)₆-SlyA are indicated. (c) Graphical 521 522 representation of the changes in transcript abundance occurring upon over-production of SlyA in E. coli K-12 MG1655. Comparison of the fold-changes in transcript abundance of 523 aerobic steady-state glucose-limited chemostat cultures of E. coli K-12 MG1655 transformed 524 with either the pET28a (SlyA^{WT}) or pGS2468 (SlyA⁺). Each line represents a gene that 525 exhibits a ≥ 2 -fold change in transcript abundance ($p \leq 0.05$) from two biological and two 526 527 technical replicates i.e. four measurements.

528

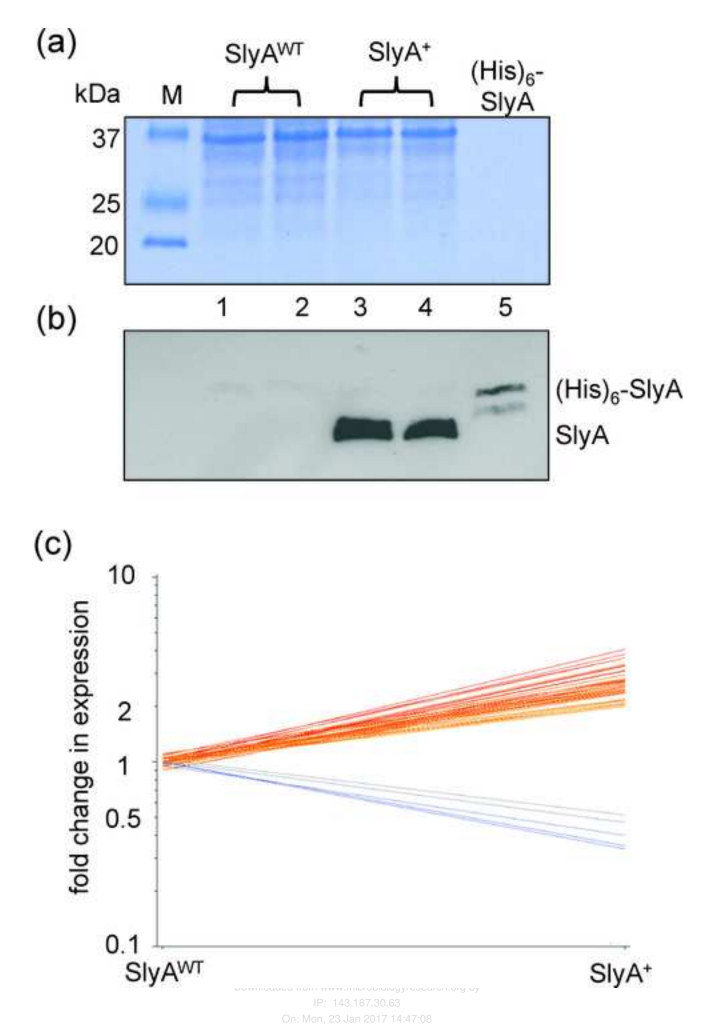
FIG. 2. Biofilm formation by *E. coli* K-12 MG1655 is enhanced by elevated *slyA* expression. Wells containing M9 minimal medium with 20% w/v glucose as a carbon source were seeded with 1:10 inocula of overnight cultures and incubated at 37°C for 16 h. The OD₆₀₀ of the planktonic bacteria was measured before a biofilm assay was carried out. Values shown are the mean and standard deviation (n = 12) *** denotes $p \le 0.00001$ in a Student's *t*-test.

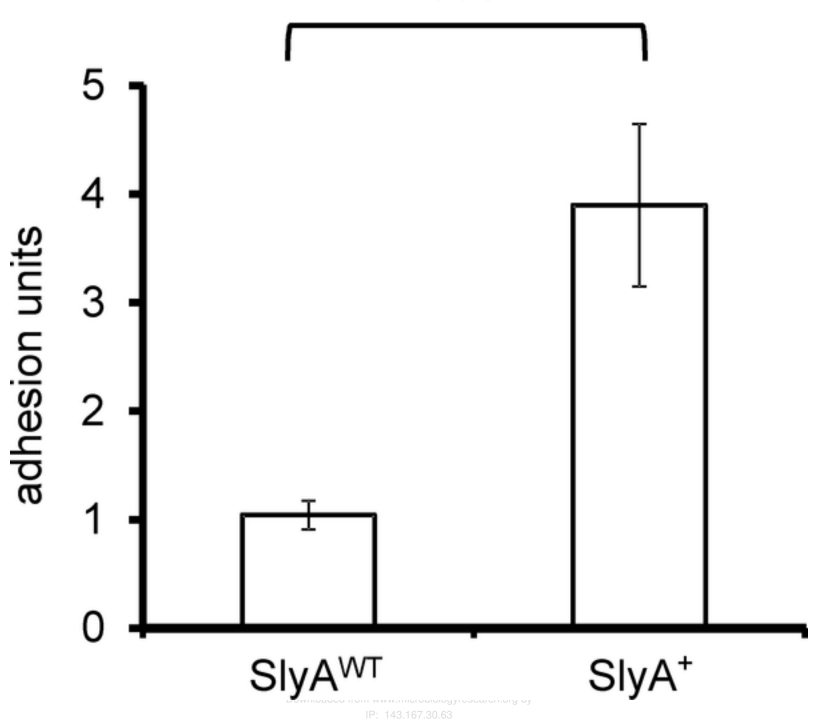
534

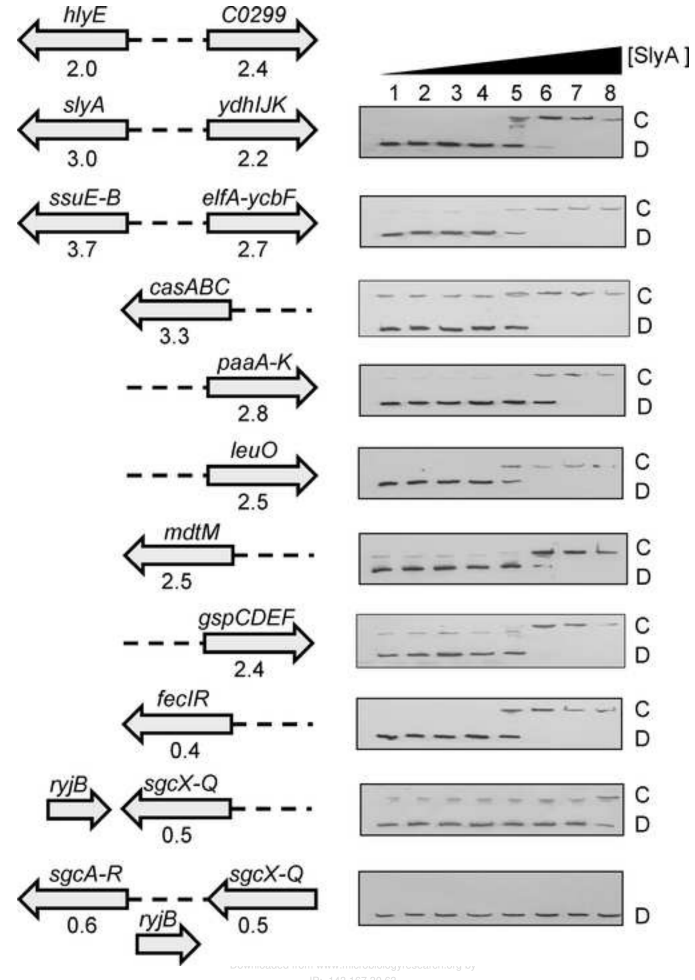
535 FIG. 3. Electrophoretic mobility shift assays show specific binding of SlyA to intergenic 536 regions of selected operons. The dashed lines in the diagrams on the left indicate the DNA 537 regions used in the EMSA shown on the right. The arrows indicate the polarity of the genes (names above the arrows). The numbers below the arrows representing genes are the fold-538 changes in transcript abundance observed upon over-production of SlyA (Table 2). SlyA-539 binding to the hlyE-C0299 intergenic region has been reported previously [20]. For the 540 EMSAs, biotin labelled intergenic DNA was prepared as described in the Methods. Labelled 541 DNA was incubated with increasing concentrations of purified SlyA protein and protein-DNA 542 complexes were separated by electrophoresis on native polyacrylamide gels. Lanes 1-8: 0, 1, 543 5, 10, 50, 100, 200, 500 nM SlyA. The locations of the free DNA (D) and the SlyA-DNA 544

545 complexes (C) are indicated. Note that binding at the *sgcX* upstream region was only 546 evident at the highest SlyA concentratrion tested and the complex (C) was located close to a 547 contaminating DNA species.









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