1	Autoregulation of the S. mutans SloR metalloregulator is constitutive and driven by an
2	independent promoter.
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16	

17 Abstract

Streptococcus mutans, one of ~600 bacterial species in the human oral cavity, is among the most 18 acidogenic constituents of the plaque biofilm. Considered to be the primary causative agent of 19 dental caries, S. mutans harbors a 25kDa SloR metalloregulatory protein which controls metal ion 20 transport across the bacterial cell membrane to maintain essential metal ion homeostasis. The 21 expression of SloR derives, in part, from transcriptional readthrough of the *sloABC* operon which 22 encodes a Mn^{2+}/Fe^{2+} ABC transport system. Herein, we describe the details of the *sloABC* 23 promoter that drives this transcription, as well as a novel independent promoter in an intergenic 24 25 region (IGR) that contributes to downstream *sloR* expression. RT-PCR studies support *sloR* transcription that is independent of *sloABC* expression, and the results of 5' RACE revealed a *sloR* 26 transcription start site in the IGR from which the -10 and -35 promoter regions were predicted. 27 The results of gel mobility shift assays support direct SloR binding to the IGR, albeit with lower 28 affinity than SloR binding to the *sloABCR* promoter. Function of the *sloR* promoter was validated 29 in qRT-PCR experiments. Interestingly, *sloR* expression was not significantly impacted when 30 grown in the presence of high manganese, whereas expression of the *sloABC* operon was repressed 31 under these conditions. The results of *in vitro* transcription studies support SloR-mediated 32 33 transcriptional-activation of *sloR* and -repression of *sloABC*. Taken together, these findings implicate SloR as a bifunctional regulator that represses *sloABC* promoter activity and encourages 34 *sloR* transcription from an independent promoter. 35

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Importance: Tooth decay is a ubiquitous infectious disease that is especially pervasive in
 underserved communities worldwide. *S. mutans*-induced carious lesions cause functional,
 physical, and/or aesthetic impairment in the vast majority of adults, and in 60-90% of

schoolchildren in industrialized countries. Billions of dollars are spent annually on caries treatment, and productivity losses due to absenteeism from the workplace are significant. Research aimed at alleviating *S. mutans*-induced tooth decay is important because it can address the socioeconomic disparity that is associated with dental cavities and improve overall general health which is inextricably linked to oral health. Research focused on the *S. mutans* SloR metalloregulatory protein can guide the development of novel therapeutics and so alleviate the burden of dental cavities.

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49 Introduction

Dental caries are important indicators of oral health and overall general health in both children and adults. Despite significant public health efforts aimed at reducing caries incidence, approximately 60-90% of school-age children worldwide experience caries, with 91% of adult caries in the United States involving the permanent dentition (1, 2). Of particular concern are children of socioeconomically disadvantaged families who are twice as likely to experience rampant caries in comparison with their wealthier counterparts, and who often present with severe clinical outcomes later in life (3).

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Among the early colonizers of the tooth surface is Streptococcus mutans, considered to be the 58 primary causative agent of dental cavities in humans (4). Ongoing research aimed at alleviating or 59 eliminating caries has given rise to valuable insights of S. mutans virulence properties, including 60 genes that mediate its obligate biofilm lifestyle, its ability to tolerate acid and oxidative stress, and 61 maintain metal ion homeostasis (5-12). The introduction of sucrose into the Western diet marked 62 a turning point for S. mutans, which metabolizes carbohydrates for energy production and 63 generates a lactic acid byproduct that demineralizes tooth enamel and drives the process of tooth 64 decay (13, 14). Taken together, S. mutans' arsenal of virulence attributes makes it an especially 65 resolute dental pathogen, and in dysbiotic plaque a primary instigator of caries formation (15, 16). 66 67

Among the evolutionary responses that are paramount for *S. mutans* survival and pathogenesis in dental plaque is tight regulation of essential metal ion transport across the bacterial cell membrane. To this end, *S. mutans* is endowed with metal ion uptake machinery which enables the import of divalent cations such as Mn^{2+} and Fe^{2+} that are essential for cellular and subcellular functions, and ultimately for bacterial cell viability. Aberrant metal ion uptake, however, can result in overaccumulation of intracellular metal ions and bacterial cell death owing to Fenton chemistry and the elaboration of toxic oxygen radicals (17–20). To counteract these deleterious effects and achieve intracellular homeostasis, *S. mutans* has evolved transport mechanisms that function to maintain appropriate metal ion uptake, which is especially paramount during periods of feast and famine in the oral cavity when Mn^{2+} concentrations can vary greatly.

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S. mutans metal ion uptake is mediated, in large part, by the *sloABCR* operon which encodes a 79 SloABC Mn²⁺/Fe²⁺ transporter and, via transcriptional readthrough, a 25 kDa SloR 80 metalloregulatory protein. As a transcription factor, SloR, modulates metal ion transport upon 81 binding to DNA in response to manganese availability (7, 20, 21). For instance, between 82 mealtimes exogenous metal ions are not readily available because they are sequestered to host 83 proteins, such as lactoferrin in saliva. Hence, under fasting conditions, S. mutans upregulates the 84 sloABC gene products which includes ATP- binding and -hydrolyzing proteins as well as a 85 transmembrane SloC lipoprotein that scavenges metal ions for uptake (20, 21). In contrast, during 86 a mealtime free metal ions are plentiful in the oral cavity, and in response S. mutans downregulates 87 its metal ion importers so as to avoid over-accumulation of intracellular metal ions and their 88 associated cytotoxic effects. We believe SloR mediates such metalloregulation by binding directly 89 to Mn²⁺ which, in turn, promotes SloR dimerization and a subsequent conformational change at 90 91 the N-terminus of the protein to facilitate DNA binding. Specifically, in a previous report we describe SloR-DNA binding upstream of the *sloABC* locus to a promoter-proximal SloR 92 Recognition Element (SRE) that represses *sloABC* transcription, presumably via a mechanism that 93 94 involves promoter exclusion to RNA polymerase (RNAP) (22, 23). Hence, the sloABC-encoded

95 metal ion uptake machinery in S. mutans is subject to transcriptional repression by SloR under conditions of Mn²⁺ availability, and conversely to de-repression when Mn²⁺ becomes limiting. 96 The transcriptional regulatory properties of the SloR protein are not limited to the *sloABC* locus. 97 In fact, work in our laboratory suggests that the SloR protein, a member of the DtxR family of 98 metalloregulators, may be involved in regulating as many as 200 genes in the S. mutans genome, 99 either directly or indirectly (24). The genes that are subject to SloR control belong to a variety of 100 different functional categories beyond metal ion homeostasis, and include gene products that 101 mediate S. mutans oxidative stress and acid tolerance, biofilm formation, and genetic competence, 102 all of which contribute to S. mutans virulence (7, 8, 24, 25). In addition, accumulating evidence 103 in our laboratory supports SloR as more than just a repressor of S. mutans gene expression. While 104 the results of expression profiling studies support SloR-mediated-repression of certain genes 105 106 (called Class I genes), the binding of SloR to other gene loci can culminate in gene activation (called Class II genes) (24). The mechanism by which SloR encourages gene transcription is 107 unknown and is currently under investigation in our laboratory. 108

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Despite the central importance of SloR in promoting *S. mutans* survival and virulence gene expression, surprisingly little is known of the regulatory mechanism(s) that modulate SloR itself. To date, the regulation of SloR in *S. mutans* has been shown to be manganese-dependent and driven, in part, by the *sloABC* promoter via transcriptional read-through of a weak terminator that is located 3' to the *sloC* gene (7, 20, 22). Hence, SloR levels that derive from *sloABC* promoter activity likely vary with Mn^{2+} availability between and during meal times. We propose however, that some constitutive baseline level of SloR is likely necessary to facilitate scavenging of essential

- 117 Mn^{2+} and/or Fe²⁺ by the SloC metal ion importer regardless of exogenous metal ion availability,
- and that such fine-tuning could involve additional mechanisms of control.
- 119
- 120 In the present study, we set out to determine whether a 184 base pair intergenic region (IGR) that
- is located immediately downstream of the *sloC* coding region and upstream of the *sloR* gene,
- harbors a specific promoter that drives *sloR* expression independent of *sloABC* promoter activity.
- 123 We propose that together with a unique and as-yet-unidentified SRE in this IGR, a bifunctional
- role for SloR as both a repressor and activator of *S. mutans* gene expression will be supported (26).
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126 **Results**

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SloR homodimers bind to a 72bp SRE. In a previous report, we describe the thermodynamic 128 binding properties of the S. mutans SloR metalloregulator to its cognate SRE within the sloABC 129 promoter region. Herein, the results of fluorescence anisotropy studies conducted with 1mM 130 manganese and various SRE-containing DNA fragments revealed tight binding of the SloR protein 131 to a 72bp DNA fragment (Kd = 32nM). Combined with the results of EMSA and DNA 132 footprinting experiments which support the binding of at least two SloR dimers to this 72bp 133 134 sequence, we hypothesized that SloR may bind as a set of homodimers to each of three inverted hexameric repeats on this 72bp target DNA, each with the consensus sequence AATTAA or some 135 modification thereof (Figure 1). To test this hypothesis, we used the SloR protein, 1mM Mn^{2+} , 136 137 and the predicted 72bp SloR recognition element in negative staining and electron microscopy The resulting dataset comprised of 55 total images was classified into twoexperiments. 138 dimensional class averages using a PARTICLE software package (www.image-analysis.net/EM/). 139 The dominant pattern that was revealed by the class averages share a common shape with three 140 distinct ellipsoidal regions presumed to be SloR dimers (Figure 1 inset, arrowheads), tilted off of 141 the DNA axis by $\sim 32^{\circ}$. The SloR binding pattern occupies a total length of 239 Angstroms on the 142 DNA with each SloR dimer measuring ~90 Angstroms across, and the distance from the center of 143 one dimer to the center of the next measures 75 Angstroms (equal to 22bp). Taken together, the 144 145 binding pattern and low-resolution image of the SloR-SRE interaction are consistent with the binding of three SloR dimers to the 72bp target sequence. 146

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148 SloR binding to the 72bp SRE is cooperative. Equilibrium binding of SloR to fluoresceinated duplex DNA containing sequences from the *sloABC* promoter region (Table S1) was measured 149 using fluorescence anisotropy with SloR binding to fluorescently-labeled duplex DNA containing 150 either one, two, or three 22bp sequences identified previously on the 72bp SRE (labeled A, B and 151 C in Figure 1). Region B, which includes a pair of inverted repeats, was the only site that 152 demonstrated SloR-specific binding under conditions as high as 250mM NaCl (data not shown). 153 Sequences A and C, which deviate from the consensus 22bp sequence at two and three nucleotide 154 positions respectively, do not measurably bind SloR under these same assay conditions. Saturation 155 156 binding to these sequences was observed however, when the salt concentration was lowered to 50mM, albeit along with some non-specific interactions. With that said, when SloR was titrated 157 into solutions containing 46bp duplexes harboring two contiguous pairs of inverted repeats, either 158 159 *sloA* AB or *sloA* BC, specific cooperative binding was observed under high salt conditions, with Hill coefficients of 1.8 and 1.7, respectively. This result indicates that SloR is capable of binding 160 to the A and C sites with high affinity if the B site is already occupied, consistent with SloR 161 interactions at the B-site that recruit the additional dimers to the flanking A and C sites. In addition, 162 we measured 50% occupancy of the two sites on *sloA* AB and *sloA* BC at 26nM and 10nM SloR, 163 164 respectively (data not shown). This result corroborates the observed binding of SloR dimers to all three sites on the 72bp duplex described above in the negative staining experiments. 165

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167 Differential expression of the *S. mutans sloABC* and *sloR* genes suggests the presence of a 168 *sloR*-specific promoter. To determine whether expression of the *sloABC* and *sloR* genes is 169 coordinated under conditions of low versus high manganese, we performed qRT-PCR 170 experiments, the results of which reveal different transcription profiles (Table 2) despite the

derivation of these genes from a single polycistronic mRNA. Specifically, expression of the 171 *sloABC* operon was repressed 3-fold under conditions of high manganese availability (Student's 172 *t*-test, p<0.05), whereas transcription of the *sloR* gene was not significantly altered under these 173 174 same conditions (Student's *t*-test, p>0.05). That expression of the *sloABC* and *sloR* genes is different under conditions of high Mn²⁺ suggests an additional control mechanism for *sloR* 175 transcription that may involve an independent promoter. We predict this promoter is located 176 within the 184bp intergenic region (IGR) that separates the *sloC* gene of the *sloABC* operon from 177 the downstream *sloR* gene on the *S. mutans* chromosome. 178

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The results of 5' RACE reveal the location of a transcription start site in the intergenic region 180 between the S. mutans sloC and sloR genes. To investigate whether a promoter might exist on 181 the IGR that separates the *sloABC* and *sloR* genes, we performed 5' RACE experiments to identify 182 a putative +1 transcription start site. The results revealed that transcription of the 654bp sloR-183 specific transcript begins at an adenosine residue located 19bp upstream of the ATG translation 184 start codon. Mapping the cDNA sequence back to the S. mutans UA159 reference genome allowed 185 us to predict and annotate the -10 and -35 sites of the predicted *sloR* promoter (Figure 2). The 186 putative -10 site aligns precisely with the conserved prokaryotic consensus sequence (TATAAT) 187 whereas there is variation in the sequence between the predicted -35 site and its consensus 188 sequence in other prokaryotes. A putative extended -10 element which is characterized by a TGN 189 sequence and the presence of two poly T tracts in the spacer region may compensate for degeneracy 190 in the -35 promoter. 191

193 The *S. mutans sloR* gene is transcribed even in the absence of a functional *sloABC* promoter.

To investigate the impact of promoter/SRE variants on transcription of the S. mutans sloABCR 194 operon, we introduced transition mutations into the 72bp SRE at positions 11 and 11d, both of 195 which share overlap with the -35 and -10 promoter sites upstream of *sloABC*, respectively. 196 Notably, T-to-C mutations at these sites in the resulting GMS611 and GMS611d strain variants 197 culminated in significantly compromised *sloABC* transcription (Student's t-test, p<0.0001) in 198 qRT-PCR experiments, with Cq values approaching those of the no template and reverse 199 transcriptase-minus controls (data not shown). This is consistent with disruption of the *sloABC* 200 promoter in the GMS611 and GMS611d strain variants. While *sloABC* transcription in these 201 mutant variants was greatly reduced, transcription of *sloR* was diminished to a much lesser extent 202 (Student's t-test, p<0.05) that we propose is the result of continued transcription from an 203 independent, *sloR*-specific promoter (Table 2). 204

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206 The S. mutans sloR gene is transcribed from the sloABC promoter as well as from an independent promoter on the 184bp IGR. To determine whether *sloR* transcription is indeed 207 driven by an independent *sloR*-specific promoter, we performed reverse transcriptase PCR (RT-208 PCR) experiments with cDNAs that we generated from the S. mutans GMS611 and GMS611d 209 *sloABC* promoter variants and their UA159 wild-type progenitor. As noted above, expression of 210 the *sloABC* operon in the GMS611 and GMS611d is virtually nil, indicating successful disruption 211 of the *sloABC* promoter in these strains. In S. mutans, expression of the *sloABC* and *sloR* genes 212 derives from a 3.4Kb polycistronic mRNA owing to transcription off of the UA159 chromosome 213 that is driven by the upstream *sloABC* promoter and transcriptional readthrough of a weak 214 terminator at the 3' end of the *sloC* gene (21, 32, 33). Herein, we expect to generate a 2.7Kb 215

216 polycistronic mRNA by RT-PCR given the positioning of the P1 and P3 primers that span the *sloABC* operon and the downstream IGR (Figure 3a). In fact, the results of RT-PCR indicate the 217 presence of a 2.7kb cDNA product in S. mutans UA159, and the absence of this product in 218 219 GMS611 and GMS611d (Figure 3b). In addition to this polycistronic mRNA however, we noted the presence of a 250bp cDNA product in all three S. mutans strains with the P2/P3 primer pair. 220 The presence of this amplicon in GMS611 and GMS611d indicates the production of a sloR-221 specific transcript even in the absence of a functional *sloABC* promoter and supports *sloR* 222 transcription from an independent promoter. Importantly, PCR products deriving from genomic 223 224 DNA when used as the amplification template confirm the specificity of the respective primer pairs (Figure 3b). 225

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SloR binds directly to the intergenic region between the S. mutans sloC and sloR genes. To 227 determine whether the impact of SloR binding on *sloR* transcription is direct, we performed EMSA 228 experiments, the results of which support direct SloR binding to the intergenic region between the 229 S. mutans sloC and sloR genes. Specifically, we observed protein-IGR binding when SloR was 230 provided at concentrations as low as 400nM, but not at concentrations of 200nM or less (Figure 231 4a). This contrasts with the SloR binding we observed at the *sloABC* promoter region which 232 occurred with as little as 60nM SloR protein. These findings support SloR binding to the IGR 233 with lower affinity than that of SloR binding to the *sloABC* promoter region. In addition, SloR-234 IGR binding was abrogated upon the addition of 1.5mM EDTA, consistent with the metal ion-235 dependence of the interaction. 236

We also used EMSA to determine the region on the IGR to which SloR binds. To this end, we generated a series of IGR fragments with serial deletions at their 5' or 3' ends (Figure 4a).

Interestingly, robust band shifts were generated with DNA fragments harboring promoter-distal hexameric repeat sequences that are located at least 62bp upstream of the +1 transcription start site, but not with DNA fragments less than 62bp from the +1 start site that lack these repeat sequences (Figure 4b).

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Expression of the S. mutans sloR gene is subject to positive autoregulation. Direct binding of 244 SloR to promoter-proximal sequences at the *sloABC* locus and to promoter-distal sequences at the 245 *sloR* locus, coupled with their negative and positive effects on global gene expression respectively. 246 247 led us to hypothesize a role for SloR as a bifunctional regulator in S. mutans. To validate such a dual role for SloR, we performed *in vitro* transcription (IVT) experiments, the results of which 248 demonstrate unequivocally that *sloABC* transcription is repressed by SloR while transcription of 249 250 the *sloR* gene is facilitated by SloR (Figure 5). These findings, quantified with ImageJ software and in combination with the results of binding studies, indicate that SloR can either repress or 251 encourage gene expression via direct binding to DNA. To our knowledge, this is the first 252 experimental evidence to demonstrate bifunctional regulation of gene expression by the SloR 253 metalloregulator in S. mutans. 254

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257 Discussion

258 Until recently, work in our laboratory was focused primarily on understanding the mechanism(s) 259 of SloR binding at the *sloABC* locus, which is subject to transcriptional repression by SloR. 260 Herein we present evidence that supports an updated model for SloR binding at this locus, consistent with the direct binding of SloR homodimers to binding sites that span a 72bp region of 261 262 DNA which includes the -10 and -35 sloABC promoter. In previous work (22), we described a pattern for SloR binding at the *sloABC* promoter site that involved two SloR dimers binding to two 263 264 sets of inverted repeats, each 6bp in length and separated by 8bp. Subsequent analysis of the 72bp 265 region upstream of the *sloABC* operon suggested that, in fact, SloR binds to three distinct but abutting 22bp sites in that region, referred to herein as A, B, and C. Each binding site is comprised 266 267 of two inverted hexameric repeats with an AATTAA consensus separated by 6bp, thereby defining putative 6-6-6 motif for SloR binding (Figure 1). This 268 a sequence pattern (xxAATTAAxxxxxTTAATTxx, where "x" is a non-conserved nucleotide) is similar to that of 269 the SloR homolog in S. gordonii, called ScaR, which binds to two adjacent 22bp sites upstream of 270 the scaABC operon (34) and for which the binding pattern was confirmed by negative staining and 271 electron microscopy (unpublished observations). We therefore expanded what we previously 272 273 thought to be a 42bp SRE in the *sloABC* promoter region to include at least 30 additional base pairs. 274

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While SloR binds to the central 22bp SRE (region B) with strong affinity when provided as a template in isolation, we measured cooperative interactions between SloR homodimers when bound to adjacent SRE sequences (regions A and C). These cooperative interactions are strong enough to support high affinity binding between SloR homodimers, presumably because of

interactions involving the initial binding of SloR homodimers to the B site. Such cooperativity
has likewise been observed for the *S. gordonii* ScaR protein (32), suggesting that this property may
be a common feature among interactions involving other streptococcal manganese-dependent
regulators and their promoter/operator sequences.

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285 In the present study, we extend our SloR binding observations to include the details of protein 286 binding to the IGR that immediately precedes the S. mutans sloR gene. Based on accumulating evidence presented herein, we propose that the location of the SloR-DNA binding element relative 287 288 to the promoter sequences that modulate downstream *sloABC* and *sloR* gene transcription contributes to SloR's ability to differentially down-regulate sloABC promoter activity and up-289 290 regulate *sloR* promoter activity. Interestingly, an *in silico* analysis of the 184bp IGR that precedes the S. mutans sloR gene failed to reveal a recognizable SRE like the one we describe above for the 291 *sloABC* locus, consistent with differential regulation by SloR at these two loci. 292

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The transcription start site for the S. mutans sloR gene occurs within the 184bp IGR that separates 294 *sloR* from the *sloABC* operon immediately upstream, as determined in 5'RACE experiments. 295 296 From the +1 transcription start site, -10 and -35 promoter regions were predicted and a 19bp 5' untranslated region (UTR) was defined. Notably, the hexameric -10 region shares 100% sequence 297 identity with the canonical prokaryotic consensus sequence for a -10 promoter (TATAAT) (35, 298 299 36). An *in silico* analysis of the *sloR* promoter revealed a putative extended -10 element in the IGR which is absent from the *sloABC* promoter region. Reports in the literature describe such a 300 TGN motif immediately upstream of the -10 sequence as an element that could facilitate 301 downstream transcription by stabilizing the open complex during initiation, and by shortening the 302

distance between the -10 and -35 sites (34, 37). The contact that RNA polymerase makes with the nonamer that defines the extended -10 site could compensate for the suboptimal contact that the polymerase makes with the degenerate -35 sequence (37). The TGN motif that we noted on the *sloR*-containing IGR is similarly located 14-16 nucleotides upstream of the transcription start site. Previous studies have also noted that short poly(T) tracts are characteristic to the spacer region of *E. coli* TG promoters (38). We similarly noted the presence of two poly(T) tracts centered at positions -18 and -29 in the spacer region of *sloR*'s TG promoter.

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In contrast to the -10 promoter region, the predicted -35 site (TATCCA) shares only 50% sequence 311 identity with the typical prokaryotic promoter sequence (TTGACA) (35). This is not surprising 312 given frequent reports of sequence variation in and around the -35 promoter region within and 313 314 across prokaryotic species (39). Since promoter strength is, in part, determined by conservation of the -10 and -35 promoter sequences (36) one might expect RNAP to have only moderate binding 315 affinity for the relatively divergent *sloR* promoter (TATAAT and TATCCA) as compared to 316 RNAP binding at the more highly conserved *sloABC* promoter (TATATT and TTGACT) (22), 317 and accordingly, weaker transcription from the former as compared to the latter. The results of 318 DNA binding and expression profiling experiments reported herein support these predictions and 319 suggest a role for divergent *sloABC* and *sloR* promoter sequences in fine-tuning metal ion transport 320 and minimizing the toxic effects of metal ion hyper-accumulation. Additional layers of gene 321 322 control involving SloR likely evolved at these loci given the importance of maintaining metal ion homeostasis under conditions as transient as those in the oral cavity. 323

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325 The absence of a recognizable SloR binding motif in the IGR that precedes *sloR* is consistent with differential gene regulation and SloR binding at this locus versus that at the *sloABC* locus. That 326 is, three adjacent palindromes on the 72bp SRE appear to be absent from the IGR, although a pair 327 of consensus palindromes with the sequence AATTAA appear to be uniquely located 44-50bp and 328 94-100bp distal to the *sloR* promoter, respectively. Interestingly, reports in the literature describe 329 330 AT-rich sequences, including palindromes like those in the *sloABC* and *sloR* promoters, that can engender intrinsic curvatures in the DNA (24, 38). To assess inherent DNA curvature in the 331 sloABC and sloR promoter regions, we applied a BEND algorithm (40) to the 72bp sloABC SRE 332 333 and the 184bp IGR, the results of which revealed high fidelity alignment of AT-rich palindromes with predicted sites for SloR binding (data not shown). Hence, DNA curvature that localizes to 334 the paired palindromic repeats at the *sloABC* and *sloR* loci supports a SloR-DNA interaction that 335 is not strictly defined by nucleotide sequence specificity, but by DNA conformation as well. 336 EMSA studies are currently underway to determine what impact, if any, these AT-rich palindromes 337 may have on SloR-DNA binding, and whether the DNA curvature they can instigate contributes 338 to SloR's function as a repressor versus an enhancer of gene transcription. 339

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Differential expression of the *sloABC* and *sloR* genes was especially pronounced in *in vitro* transcription (IVT) assays where we used the 5' end of the *sloA* or *sloR* coding regions and up to 200bp of upstream DNA sequence as the DNA template. Pixel counting of the resulting mRNA transcripts on autoradiograms, performed with Image J, revealed considerably more robust *sloR* transcription in the presence of exogenous SloR than in its absence (Figure 5). This contrasts with the *in vitro* transcription we observed for the upstream *sloABC* operon which, as expected, was repressed by the presence of SloR. Taken together with the EMSA results that support direct SloR 348 binding at these loci, these data demonstrably support a role for SloR as a bifunctional regulator of S. mutans gene transcription. While the mechanism for sloABC repression likely derives from 349 SloR binding to an SRE that shares overlap with the *sloABC* promoter, thereby excluding RNAP 350 351 from promoter access, *sloR* transcription is the likely result of de-repression with SloR binding to promoter-distal sites. In future work, we will consider Mn^{2+} status as a potential contributor to 352 differential SloR binding and gene transcription outcomes. In fact, binding of the MntR 353 metalloregulator to different sequences upstream of the mntABCD locus in Bacillus subtilis is 354 known to be Mn^{2+} concentration-dependent (41). 355

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In summary, the results of the present study support SloR-mediated transcriptional events at the 357 *sloR* locus that are different from those at the *sloABC* locus and lend further support to a role for 358 359 SloR as a bifunctional regulator of gene transcription. It's tempting to suggest that the *sloR*specific promoter on the 184bp IGR evolved to ensure at least some level of constitutive SloR 360 production. Accordingly, when free metal ions are introduced into the oral cavity during a 361 mealtime, S. mutans is poised and ready to modulate the controlled uptake of the exogenous Mn^{2+} 362 and Fe²⁺ it needs for survival. Hence, while the sudden introduction of metal ions into the mouth 363 could prove damaging to some constituents of the oral microbiota, S. mutans can exploit these 364 conditions with a metal ion-dependent SloR regulator that can repress the cytotoxic effects of 365 excessive metal ion import, while maintaining baseline levels of SloR from an ancillary promoter. 366 367 Such fine-tuned gene regulation can impact cell function beyond the scope of metal ion homeostasis, and influence processes like adherence, acid production and the oxidative stress 368 response that more directly contribute to S. mutans-induced disease (7, 8, 24, 25). In conclusion, 369 370 we propose that S. mutans coordinates the regulated expression of its metal ion transport machinery

- 371 with that of its virulence attributes. An improved understanding of *sloR* autoregulation is
- 372 significant because it can elucidate the mechanisms that fine-tune the regulated expression of metal
- ion homeostasis and virulence in an important oral pathogen. Moreover, from these investigations
- we may better elucidate the details of SloR-mediated gene regulation that can benefit the design
- of an anti-SloR therapeutic aimed at alleviating and/or preventing caries.

376 Materials & Methods

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Bacterial strains, plasmids, and primers. The bacterial strains used in this study are listed in
Table 1. Working stocks of each bacterial strain were prepared from overnight cultures and stored
in 20% or 50% sterile glycerol at -20°C or -80°C, respectively.

The primers used in this study are shown in Table 1. All primers were designed using the Primer Blast tool from the NCBI website. A RefSeq record was used as the input with forward and reverse primer locations specified by the user. Lack of secondary structure was confirmed using the Oligo Evaluator tool (Sigma) and primers were checked for specificity against the *S. mutans* UA159 genome with the NCBI Basic Local Alignment Search Tool (BLAST).

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Negative-staining and EM analysis. A SloR-Mn²⁺-72bp SRE complex was prepared *in vitro* 387 assuming a 3:1 stoichiometry of dimers to DNA. The reaction mixtures were then systematically 388 diluted to 25, 50 and 100nM for negative-stain EM grid preparation using 1% uranyl acetate. After 389 screening for the optimal staining quality and particle concentration, single-particle data were 390 collected on the 50 nM specimen grid using an FEI T12 electron microscope at 120KeV and 391 68,000x nominal magnification, producing 108 micrographs at 3.15 Å/pixel in the image with 392 varying defocus between 0.9 and 1.8 µm. Then, 8,500 particles of the SloR-DNA complex were 393 selected for 2D classification using the PARTICLE (www.sbgrid.org/software/titles/particle) 394 395 program.

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397 **Construction of** *S. mutans* **promoter variants.** To generate specific mutations in the promoters 398 that drive *sloABC* transcription, we adopted a markerless mutagenesis strategy similar to that 399 described by Xie *et al.* (27). This involved constructing promoter variants with an IFDC2 cassette

400 inserted within the *sloABC* promoter region for subsequent CSP-transformation into S. mutans UA159 to generate the erythromycin-resistant and p-4-chlorophenylalanine-sensitive GMS602 401 strain. The double-crossover event was confirmed by polymerase chain reaction (PCR) and Sanger 402 sequencing. Derivatives of GMS602 were generated by overlap extension PCR (OE-PCR) with 403 the reverse primers harboring a single point mutation in the predicted 72bp SRE that precedes the 404 sloABC genes (Table 1). S. mutans GMS611 was generated with degenerate primers that 405 introduced a point mutation into the SRE at nucleotide position 11 that we predict shares overlap 406 with the *sloABC*-specific -35 promoter region (22). Likewise, GMS611d was generated with a 407 408 different degenerate primer set that introduced a point mutation at position 11d into the SRE, which we predict shares overlap with the *sloABC*-specific -10 promoter region. Thymine to cytosine 409 transition mutations were generated in both S. mutans strains and validated by Sanger sequencing 410 411 (22).

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Chromosomal DNA Isolation. S. mutans grown overnight at 37°C and 5% CO₂ in 14ml Todd-413 Hewitt Yeast Extract (THYE) broth were pelleted by centrifugation at 7000rpm for 5 minutes in a 414 Sorvall RCB centrifuge after which the cells were resuspended in 1ml Tris-EDTA buffer (10mM 415 Tris, 1mM EDTA) and chemically disrupted according to established protocols (8, 22). Genomic 416 DNA was purified in subsequent rounds of phenol-chloroform extraction, ethanol precipitated, and 417 resuspended overnight in nuclease-free water (Ambion) at 4°C with gentle agitation (28). Nucleic 418 419 acid yield and purity were assessed with a NanoDrop Lite spectrophotometer (Thermo Fisher 420 Scientific) and the samples were stored at -20° C.

421

RNA Isolation. RNA was isolated from *S. mutans* strains according to established protocols (8). Cells were grown to mid-logarithmic phase ($OD_{600nm} = 0.4-0.6$) before pelleting by centrifugation as described above and resuspending in RNAProtect (Qiagen). Total intact RNA was purified following cell lysis and DNase I treatment with a Qiagen RNeasy kit after which nucleic acid yield and purity were assessed with a NanoDrop Lite spectrophotometer (Thermo Fisher Scientific).

427 RNA samples were analyzed for integrity by agarose gel electrophoresis and stored at -80°C.

428

RT-PCR. Reverse-transcriptase PCR was performed with cDNAs that were reverse-transcribed 429 430 from RNA using a First Strand cDNA Synthesis kit according to the recommendations of the supplier (Thermo Fisher Scientific) or else with chromosomal DNA isolated from S. mutans 431 UA159, GMS611, or GMS611d according to a modification of Idone et al. (29). Q5 Hi-Fidelity 432 Polymerase was used for PCR in accordance with the recommendations of the supplier (New 433 England Biolabs). Each 50 µL PCR reaction consisted of 10 µL 5X Q5 Reaction buffer, 1 µL 434 10mM dNTPs, 2.5 µL sloA.RT_PCR.F or sloR.RT PCR.F, 2.5 µL sloA.RT PCR.R or 435 sloR.RT PCR.R [Table 1]), 200 ng of genomic DNA or 1 µL of cDNA product, nuclease-free 436 water up to 49.5 µL, and 0.5 µL of Q5 Hi-Fi Polymerase added to the reaction just prior to 437 amplification. PCR conditions were as follows: initial denaturation at 98°C for 30 s, followed by 438 25 cycles of denaturation at 98°C for 10 s, annealing at 67°C for 30 s, and extension at 72°C, 439 concluding with a final extension at 72°C for 80 s. An aliquot of each sample was visualized on 440 441 0.8% agarose gels as described above.

442

5' RACE. To reveal the *sloR* transcription start site and predict the -10 and -35 regions of the *sloR*promoter we performed 5' Rapid Amplification of cDNA Ends (5' RACE) with an Invitrogen 5'

445 RACE Kit in accordance with the manufacturer's protocol unless otherwise specified. RNA isolated from S. mutans GMS611 and GMS611d was reverse-transcribed into cDNA using a 446 sloR.[R].GSP1.A reverse primer and 200 U of Superscript II Reverse Transcriptase. After 447 S.N.A.P. column purification, the cDNA was poly(-dC) tailed using terminal deoxynucleotidyl 448 transferase as described in the manufacturer's protocol. The cDNA products were PCR amplified 449 using the kit-supplied Abridged Anchor Primer (AAP), a sloR.[R].GSP2.B reverse primer, and 450 Platinum Hi-Fi Taq polymerase, in a BioRad PCR machine programmed at 94°C for 2 minutes, 451 followed by 35 cycles of 94°C for 15 s, 58°C for 30 s, and 68°C for 1 minute, and a final hold at 452 4°C. An aliquot of the resulting PCR product was analyzed by agarose gel electrophoresis and the 453 remainder was purified using a MinElute PCR Purification kit in accordance with the 454 recommendations of the supplier (Qiagen). The purified amplicons were quantified using a 455 NanoDrop Lite spectrophotometer (Thermo Fisher Scientific) and sequenced (Eurofins) using the 456 reverse primers *sloR*.GSP2.A or *sloR*.nested.GSP. The 5' sequence of the mRNA transcript was 457 aligned with the S. mutans UA159 reference genome (RefSeq accession number NC 004350.2) 458 from NCBI to identify the nucleotide that immediately follows the poly(-dC)-tail as the 459 transcription start site (+1 site), predict the -10 and -35 promoter sequences in the 184bp intergenic 460 region, and reveal the 5' untranslated region of the *sloR* transcript (30). 461

462

EMSA. Electrophoretic mobility shift assays (EMSAs) were performed according to established protocols to determine whether SloR binding to the intergenic region upstream of the *sloR* gene is direct, and to narrow down the region of SloR binding at this locus (8, 22, 23). Primer design for the DNA binding template spanned the 184bp intergenic region (IGR) between *sloC* and *sloR* and included serial deletions thereof. PCR amplification was performed with Q5 polymerase

468 according to the manufacturer's protocol (New England Biolabs) using the following thermal 469 cycling conditions: initial denaturation at 98°C for 30 s, 35 cycles at 98°C for 10 s, annealing at 470 60°C for 30 s, and extension at 72°C for 30 s, with a final extension at 72°C for 2 minutes. The 471 resulting amplicons were PCR purified as described earlier, confirmed by agarose gel 472 electrophoresis, and quantified by NanoDrop spectrophotometry.

473

The resulting amplicons were end-labeled with $[\gamma^{-32}P]$ -dATP (Perkin-Elmer) in the presence of T₄ 474 polynucleotide kinase (New England BioLabs) after which they were centrifuged through a TE 475 Select-D G-25 spin column (Roche Applied Science) to remove the unincorporated ³²P-dATP. 476 Binding reactions were prepared as described previously (23) in a 16- μ l total volume containing 1 477 µl of end-labeled amplicon, purified native SloR protein at concentrations ranging from 60 nM to 478 479 400nM, and 3.2 µl of 5× binding buffer (42 mM NaH₂PO₄, 58 mM Na₂HPO₄, 250 mM NaCl, 25 mM MgCl₂, 50 µg/ml bovine serum albumin, 1 mg sonicated salmon sperm DNA, 50% [vol/vol] 480 glycerol, and 37.5 μ M MnCl₂). EDTA was added to a separate reaction mixture at a final 481 concentration of 1.5mM to validate SloR binding that is metal ion-dependent. An end-labeled *sloA* 482 promoter-containing amplicon was used as a positive control for SloR binding (7, 22, 23). Samples 483 were loaded onto 12% nondenaturing polyacrylamide gels (3 ml 20× bis-Tris borate [pH 7.4], 74 484 µl 100 nM MnCl₂, 1.5 ml 100% glycerol, 24 ml 30% acrylamide [37.5:1 acrylamide-bis], 31 ml 485 Millipore H₂O, 300 µl 15% ammonium persulfate [APS], 90 µl TEMED [N,N,N',N'-486 487 tetramethylethylenediamine]) and resolved at 300 volts for 1.5 hours. Gels were exposed to Kodak BioMax film for 24-72 hours at -80°C in the presence of an intensifying screen prior to 488 autoradiography. 489

490

491 Fluorescence anisotropy. Equilibrium binding of SloR to regions within the *sloABC* promoter/operator region was probed by titrating SloR onto fluoresceinated DNA and monitoring 492 binding by fluorescence anisotropy. Duplex DNA fragments containing relevant sequences were 493 prepared from oligonucleotides obtained from Integrated DNA Technologies (Coralville, IA). A 494 5' fluoresceinated oligonucleotide was annealed with a 10% excess of its unlabeled 495 complementary strand in 25mM HEPES, pH7.9, 50mM NaCl by heating to 90°C and cooling 496 slowly to room temperature (Table S1). Titrations were performed in either high or low salt 497 buffers; the high salt buffer contained 25mM HEPES, pH 8.0, 250mM NaCl, 10% glycerol and 498 1mM MnCl₂ whereas the low salt buffer contained 25mM HEPES, pH7.9, 50mM NaCl, and 1mM 499 MnCl₂. SloR was titrated into 1ml of 1nM fluoresceinated duplex DNA in buffer and anisotropy 500 measurements were made at 25°C using a Beacon 2000 fluorescence polarization instrument. Data 501 502 were fit to one of several equations related to simple 1:1 binding stoichiometry, where K_d was greater than 10nM (equation1), less than 10nM (equation 2) or to the Hill equation (equation 3). 503

504

505
$$r = \Delta r \times \frac{P}{K_d + P} + r_{min}$$
 (equation 1)

 $r = \Delta r \times \frac{K_d + D + P - \sqrt{(K_d + D + P)^2 - 4DP}}{2D} + r_{min}$ (equation 2)

507
$$r = \Delta r \times \frac{P^n}{K_h^n + P^n} + r_{min}$$
 (equation 3)

In these equations, r is anisotropy, Δr is the total change in anisotropy at saturation, K_d is the dissociation constant, K_h is the concentration of SloR dimers giving 50% maximal binding under cooperative conditions, n is the Hill coefficient, and r_{min} is the anisotropy obtained before addition of SloR. P is the concentration of SloR dimers and D is the concentration of duplex DNA. In addition, where non-specific binding prevented signal saturation, a term to model the slow linear

increase in anisotropy was added to equations 1 or 2, with the form $K_{ns}P$ where K_{ns} is the nonspecific binding constant. All curve fitting was performed in R software.

515

516 **Preparation of S.** *mutans* **RpoD.** An *E. coli* DH5α strain harboring plasmid pIB611 (kind gift of Indranil Biswas, University of Kansas) was streaked for isolation on L-agar plates supplemented 517 with ampicillin (100 µg/mL) and incubated overnight at 37°C. Resident on pIB611 is the S. mutans 518 rpoD gene cloned directly downstream of an inducible operon on vector pET-23d(+) and upstream 519 of a C-terminal His-tag (31). Importantly, the 6x His-tag was shown in *in vitro* transcription 520 521 experiments to have no physiological impact on RpoD functionality (31). Plasmid pIB611 was purified with a Qiagen miniprep kit according to the recommendations of the supplier and mapped 522 by restriction digestion (New England Biolabs). 523

524

Next, pIB611 was used to transform E. coli BL21 DE3 cells in accordance with the manufacturer's 525 protocol (Invitrogen). Successful transformants were selected after overnight growth on L-agar 526 plates supplemented with 100ug/mL ampicillin and used to inoculate a 25 mL starter culture for 527 yet another overnight incubation. This culture was then used to inoculate 500 mL of pre-warmed 528 L-ampicillin (100ug/ml) broth in a 2.8 L Fernbach flask which was incubated at 37°C with 529 continuous shaking at 200rpm. When the cells reached mid-log phase ($OD_{600}=0.4$), isopropyl β -530 D-1-thiogalactopyranoside (IPTG) was added at a final concentration of 0.5mM to induce 531 532 expression of the T7 RNA polymerase in the BL21 DE3 cells. Protein induction proceeded with continuous shaking for an additional 3.5 hours after which the cells were centrifuged as previously 533 described and stored as dry pellets at -80°C. 534

536 Cell pellets were resuspended in His-Binding Buffer (0.5M NaCl, 20mM Tris-HCl, 5mM imidazole) with Halt EDTA-free protease inhibitor at 1x concentration (Thermo Fisher Scientific). 537 The cell suspension was sonicated (Ultrasonic Power Corporation, model number: 2000U) at 60% 538 power for six 30-second cycles using a 0.5 second pulse, with samples maintained on ice between 539 runs. Cells were pelleted by centrifugation at 10,000 x g for 30 minutes at 4°C, after which aliquots 540 of the supernatant and 2X Laemmli buffer (4% SDS, 20% glycerol, 10% 2-mercaptoethanol, 541 0.004% bromophenol blue and 0.125 M Tris HCl, pH 6.8) were mixed in equal proportions and 542 resolved on 10% Bis-Tris polyacrylamide gels in 3-(N-morpholino) propanesulfonic acid (MOPS) 543 buffer. Proteins were visualized with Sypro Ruby Gel Stain (Thermo Fisher Scientific) according 544 to the manufacturer's instructions. Polyacrylamide gels were fixed for 45 minutes in fixative 545 solution (50% methanol, 7% acetic acid) with gentle shaking on an orbital shaker. The fixative 546 547 was subsequently decanted and replaced with 80 mL of Sypro Ruby Gel Stain. Gels were covered and left to stain on an orbital shaker overnight at room temperature. The gel stain was decanted 548 and wash solution (10% methanol, 7% acetic acid) added before UV visualization. 549

550

The remaining cell lysate was further purified by Ni²⁺-Nitrilotriacetic acid (Ni-NTA) column 551 chromatography (Thermo Fisher Scientific) at 4°C according to the manufacturer's instructions. 552 The columns were placed on a rotating platform for 30 minutes at 4°C to encourage RpoD binding 553 to the Ni-NTA resin before centrifugation to remove unbound protein from the column. After 554 555 elution, protein yield was determined using both NanoDrop Lite spectrophotometry and a bicinchoninic acid (BCA) protein determination assay (Thermo Fisher Scientific). RpoD purity 556 was assessed on SDS-PAGE gels following Sypro Ruby staining. Select fractions containing 557 558 RpoD were dialyzed using G2 Slide-A-Lyzer Cassettes with a 10 kDa molecular weight cut off (Thermo Fisher Scientific) in dialysis buffer (25% glycerol in 1x PBS). RpoD was assayed for
concentration as described above and stored at -20°C.

561

In vitro transcription. In vitro transcription was performed according to an adaptation of Kajfasz 562 et al (9). First, genomic DNA spanning approximately 100-200bp of the *sloA* or *sloR* coding 563 region and about 150-200bp of upstream sequence was amplified by PCR with Q5 polymerase 564 (New England Biolabs) and either PM.IVT.sloA.F and PM.IVT.sloA.R or PM.IVT.sloR.F and 565 PM.IVT.sloR.R (Table S1) according to the following thermal cycling conditions: initial 566 denaturation at 98°C for 30 s, 35 cycles at 98°C for 10 s, annealing (60°C for *sloA*, 67°C for *sloR*) 567 for 30 s, and extension at 72°C for 30 s, with a final extension at 72°C for 2 minutes. The resulting 568 amplicons were PCR purified as described earlier and confirmed by agarose gel electrophoresis. 569 570 DNA concentration was determined spectrophotometrically on a NanoDrop Lite spectrophotometer. Next, in 1.5 mL microfuge tubes, 10 nM DNA template (sloA or sloR), 1U E. 571 coli RNA Polymerase core enzyme (New England Biolabs), 20U of SUPERase RNase Inhibitor 572 (Thermo Fisher Scientific), 25 nM S. mutans RpoD extract (based on BCA assay), +/- 75nM 573 purified SloR were mixed in Reaction Buffer (10 Mm Tris-HCl [pH 8.0], 50 mM NaCl, 5 mM 574 MgCl₂, 50 µg/mL bovine serum albumin) to yield a final volume of 17.8 µL and incubated at 37°C 575 for 10 minutes. To generate an mRNA transcript, 2.2 µL of Nucleotide Mixture (200 µM ATP, 576 200 μ M GTP, 200 μ M CTP, 10 μ M UTP) and 5 μ Ci α^{32} P-UTP (Perkin Elmer) were then added 577 578 and the reaction was incubated at 37°C for 10 minutes. 10 µL of Stop Solution (1M ammonium 579 acetate, 0.1 mg/mL yeast tRNA (Ambion), 0.03M EDTA) was added to terminate transcription, after which 90 µL of ice-cold 99% EtOH was added for ethanol precipitation overnight at 4°C. 580 581 The following day samples were pelleted at 16,200 x g for 30 minutes, followed by three rounds

582 of washing with 70% EtOH, with additional centrifugation between washes. A final wash with 99% EtOH was performed after which the samples were lyophilized in a vacuum centrifuge 583 (Eppendorf) for 10 minutes. The radiolabeled cell pellet was resuspended in 5 uL of formamide 584 dye (0.3% xylene cyanol, 0.3% bromophenol blue, 12mM EDTA, dissolved in formamide) and 585 the samples were heated to 70°C in a water bath for 2 minutes before placing on ice for gel loading. 586 Samples were resolved on a Novex 10% TBE-7M Urea Gel in 1X TBE Buffer (10.8 g Tris, 5.5g 587 boric acid, 0.01M EDTA [pH 8.0]) and mRNA transcripts were visualized via autoradiography 588 using BIOMAX XAR Film (Thermo Fisher Scientific) exposed for up to 4 hours at -80°C in the 589 590 presence of an intensifying screen. Film was developed according to standard protocols and ImageJ software was used to quantify the band intensities between samples. 591

592

593 Semi-quantitative real-time PCR (qRT-PCR). Total intact RNA was isolated from midlogarithmic phase S. mutans cultures of the wild-type UA159 strain grown in a semi-defined 594 medium (SDM) (22) supplemented with either 5uM (low) or 125uM (high) MnSO₄. The resulting 595 RNAs were analyzed for integrity on 0.8% agarose gels before reverse-transcribing 100ng of each 596 RNA sample into cDNA as described above. The cDNAs were used as templates for qRT-PCR 597 which was performed in accordance with established protocols (22) in a CXR thermal cycler 598 (BioRad). Specifically, *sloABC* and *sloR* transcription was assessed in three independent qRT-599 PCR experiments, each performed in triplicate and normalized against the expression of gyrA 600 601 (8,22).

602

To assess the impact of transition mutations in the SRE that precedes the *sloABC* operon on downstream *sloR* transcription, *S. mutans* GMS611 and GMS611d were grown as described above

- except without supplemental Mn^{2+} . Total intact RNA was isolated and reverse transcribed as
- described, and the results of qRT-PCR were normalized against hk11, the expression of which
- 607 does not change under the experimental test conditions.

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726	

728 Tables & Figures

729

730 Table 1. Bacterial strains used in this study.

732 Table 2. Fold change in expression for *S. mutans sloABCR*.

*Student's t-test, p<0.05; **Student's t-test, p<0.001; #Student's t-test, p<0.05

Figure 1. SloR-SRE binding model in the region of the *sloABC* promoter. Shown are each of 734 three inverted hexameric repeats (in blue, purple, and red) that span 72bp in the *sloABC* promoter 735 region, and to which SloR presumably binds as three homodimers. Affinity binding studies support 736 737 preferential binding of SloR homodimers to the hexameric repeat in region B (designated by the black bar), followed by cooperative binding of SloR dimers to regions A and C. Inset: Negative 738 staining of the SloR-SRE interaction in vitro supports homodimeric binding of SloR to a 72bp 739 SloR recognition element (SRE). Shown are three SloR homodimers bound to a DNA filament 740 containing the 72bp SloR binding element. The arrowheads denote the center of mass for each 741 742 SloR homodimer.

Figure 2. Shown is the intergenic region between the *sloR* and *sloC* genes which harbors a

recognizable promoter. The nucleotide sequence of this region was aligned with the S. mutans

- 746 UA159 genome from the NCBI GenBank Database (RefSeq accession number NC_004350.2).
- 747 The +1 transcription start site (designated by the arrow) marks the transcription start site of the
- *sloR* gene as defined by 5'RACE, and defines a 19bp 5' untranslated region (UTR). Also shown
- are the predicted -35 and -10 promoter regions, the predicted ribosome binding site (RBS), and the
- start codon (SC) of the 654bp *sloR* gene. A putative extended -10 element is denoted by the dashed
- 751 line.

752 Figure 3. The S. mutans sloR gene is transcribed even in the absence of a functional sloABC

promoter. a.) Map of the *sloABCR* operon and location of primer annealing sites. Primer P1 753 (sloA.RT PCR.F) anneals within the sloA coding sequence, and primers P2 and P3 754 (sloR.RT_PCR.F and sloR.RT_PCR.R, respectively) anneal within the sloR coding sequence. b.) 755 Products of reverse transcriptase PCR resolved on a 0.8% agarose gel. Amplification of cDNA 756 with the P1/P3 primer pair generated a 2745bp amplicon in UA159 but not in GMS611 or 757 GMS611d, consistent with disruption of the *sloABC* promoter in the mutant strains. In contrast, 758 cDNA amplification with the P2/P3 primer pair gave rise to a 250bp amplicon even in the *sloABC* 759 promoter mutants GMS611 and GMS611d, indicating the presence of a *sloR*-specific promoter in 760 the 184bp intergenic region that separates the *sloABC* operon from the downstream *sloR* gene 761 762 (gDNA = genomic DNA; cDNA = copy DNA).

763 Figure 4. SloR binds directly to the intergenic region (IGR) between the S. mutans sloC and sloR genes. a.) Shown are the results of EMSA which support direct SloR binding to 204bp and 764 155bp fragments of the *sloC-sloR* intergenic region at protein concentrations as low as 400nM. 765 766 SloR binding to a 95bp IGR derivative was relatively compromised however, and completely absent when a 62bp deletion derivative was used as the binding template. A 205bp amplicon that 767 includes the *sloABC* promoter was used as a positive control for SloR binding. EDTA was added 768 to select reaction mixtures in an attempt to abrogate metal ion-dependent binding. 12% non-769 denaturing polyacrylamide gels were run at 300 volts for 1.5 hours. Film exposure in the presence 770 771 of an intensifying screen proceeded for 48 hours at -80°C before development. b.) SloR binding to serial deletion fragments of the S. mutans IGR. The arrowheads facing inward represent 772 AATTAA hexameric repeats to which SloR putatively binds. The vertical red bars denote the 773 774 positioning of the -10 and -35 promoter sequences of the *sloR*-specific promoter. Whether or not SloR binds to the IGR fragment is shown with a (+) or (-) designation. 775

Figure 5. The results of *sloABCR* transcription experiments performed *in vitro* (IVT) support

SloR as a bifunctional regulator. Transcription of the *sloABC* and *sloR* genes was quantified using ImageJ software. SloR (75nM) was added to select reaction mixtures containing RNAse inhibitor and either *sloA* or *sloR* template DNA. Pixel counting was performed with ImageJ software and does not include unincorporated α^{32} P-UTP. Higher pixel counts indicate greater band intensity. Shown are the results of a single representative experiment (from a total of 3 independent experiments) which support repression of *sloA* and activation of *sloR* transcription by the SloR metalloregulator.

1 Tables

3 Table 1. Bacterial strains used in this study.

Strain	Description / Relevant Characteristics	Source	
E. coli	$hA2$ [lon] ompT gal(λ) (DE3) [dcm] Δ hsdS	Thermo Fisher	
BL21 (DE3)		Scientific	
S. mutans	Wild-type S. mutans,	ATCC 700610	
UA159	Serotype <i>c</i>		
S. mutans	UA159 derived, contains IFDC2 cassette in	Spatafora <i>et al.</i> (2015	
GMS602	the <i>sloA</i> promoter region, Em ^r , 4-Cl-Phe ^s	(22)	
S. mutans	Contains a markerless $T \rightarrow C$ mutation	This study	
GMS611	within the -35 region of the <i>sloABC</i>		
	promoter		
S. mutans	Contains a markerless $T \rightarrow C$ mutation	This study	
GMS611d	within the -10 region of the <i>sloABC</i>		
	promoter		

10 Table 2. Fold change in expression for *S. mutans sloABCR*.

Strain/Condition					
Locus	UA159	UA159	UA159	GMS611	GMS611d
	$(125 \ \mu M \ Mn^{2+})$	$(5 \ \mu M \ Mn^{2+})$	1 2 7 0 1 0 4 4	0.1.1 0.01.1.1.	0.07.0.0044
sloABC	$0.33 \pm 0.03*$	$1.02 \pm 0.09*$	1.35 ± 0.10 **	0.14 ± 0.01 **	0.05 ± 0.00 **
sloR	0.75 ± 0.09	1.02 ± 0.19	$1.35 \pm 0.14^{\#}$	$0.96 \pm 0.03^{\#}$	$0.81 \pm 0.10^{\#}$

* Student's t-test, p<0.05; **Student's t-test, p<0.0001; [#] Student's t-test, p<0.05

1 Figures

2

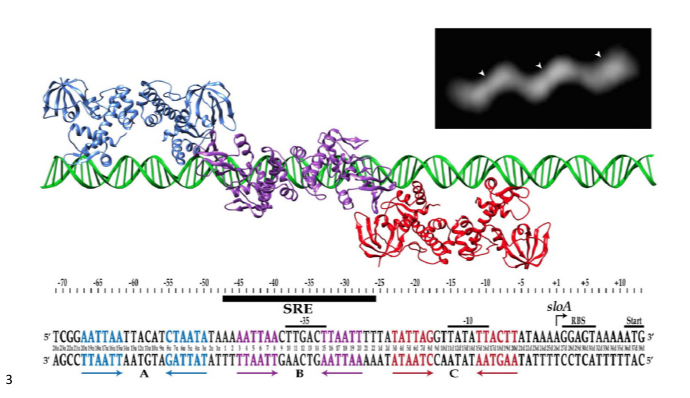


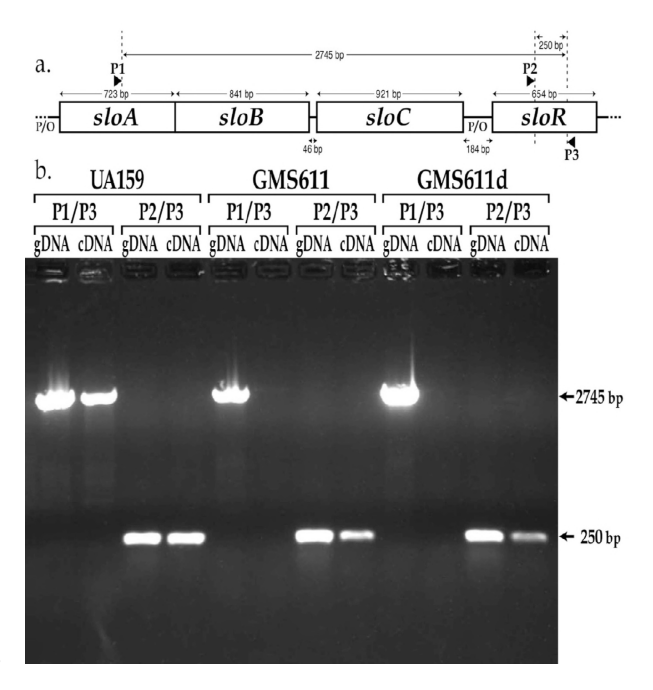
Figure 1. SloR-SRE binding model in the region of the *sloABC* promoter. Shown are each of 4 three inverted hexameric repeats (in blue, purple, and red) that span 72bp in the *sloABC* promoter 5 6 region, and to which SloR presumably binds as three homodimers. Affinity binding studies support preferential binding of SloR homodimers to the hexameric repeat in region B (designated by the 7 black bar), followed by cooperative binding of SloR dimers to regions A and C. Inset: Negative 8 9 staining of the SloR-SRE interaction in vitro supports homodimeric binding of SloR to a 72bp SloR recognition element (SRE). Shown are three SloR homodimers bound to a DNA filament 10 containing the 72bp SloR binding element. The arrowheads denote the center of mass for each 11 SloR homodimer. 12

<u>-35</u> <u>-10</u> +1 <u>RBS</u> <u>SC</u> gataattaaaaacttttatccacttttactgacattttgatataatgattttgacaaagaagaggataaactatgacac ctaataaagaagattaccttaaaattatttatgaactcagtgaacgtgatgagaagatcagcaacaagcaaattgct gagaaaatgtctgtatctgctcc

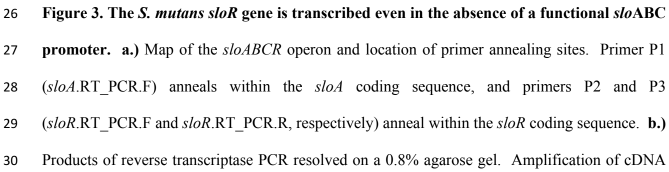
14

15	Figure 2. Shown is the intergenic region between the <i>sloR</i> and <i>sloC</i> genes which harbors a
16	recognizable promoter. The nucleotide sequence of this region was aligned with the S. mutans
17	UA159 genome from the NCBI GenBank Database (RefSeq accession number NC_004350.2).
18	The +1 transcription start site (designated by the arrow) marks the transcription start site of the
19	sloR gene as defined by 5'RACE, and defines a 19bp 5' untranslated region (UTR). Also shown
20	are the predicted -35 and -10 promoter regions, the predicted ribosome binding site (RBS), and the
21	start codon (SC) of the 654bp <i>sloR</i> gene. A putative extended -10 element is denoted by the dashed
22	line.

23







- 31 with the P1/P3 primer pair generated a 2745bp amplicon in UA159 but not in GMS611 or
- 32 GMS611d, consistent with disruption of the *sloABC* promoter in the mutant strains. In contrast,
- cDNA amplification with the P2/P3 primer pair gave rise to a 250bp amplicon even in the *sloABC*
- promoter mutants GMS611 and GMS611d, indicating the presence of a *sloR*-specific promoter in
- the 184bp intergenic region that separates the *sloABC* operon from the downstream *sloR* gene
- 36 (gDNA = genomic DNA; cDNA = copy DNA).

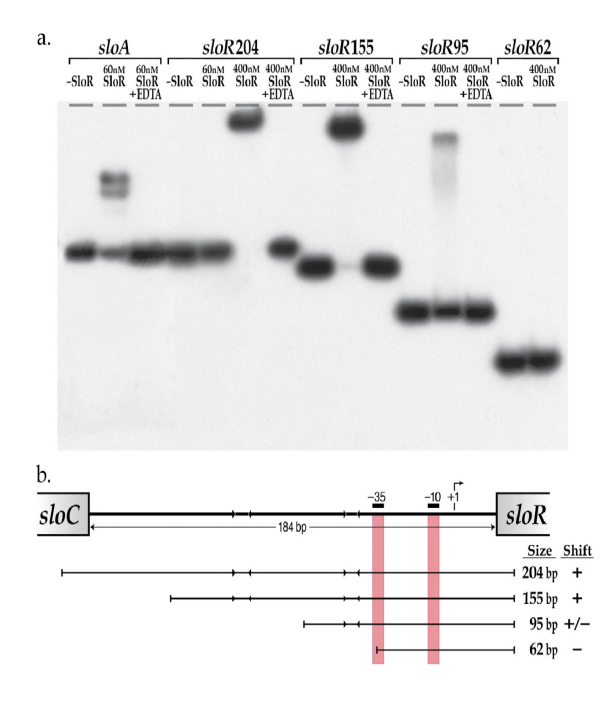


Figure 4. SloR binds directly to the intergenic region (IGR) between the *S. mutans sloC* and *sloR* genes. a.) Shown are the results of EMSA which support direct SloR binding to 204bp and 155bp fragments of the *sloC-sloR* intergenic region at protein concentrations as low as 400nM.
SloR binding to a 95bp IGR derivative was relatively compromised however, and completely absent when a 62bp deletion derivative was used as the binding template. A 205bp amplicon that

includes the *sloABC* promoter was used as a positive control for SloR binding. EDTA was added 44 to select reaction mixtures in an attempt to abrogate metal ion-dependent binding. 12% non-45 denaturing polyacrylamide gels were run at 300 volts for 1.5 hours. Film exposure in the presence 46 47 of an intensifying screen proceeded for 48 hours at -80°C before development. b.) SloR binding to serial deletion fragments of the S. mutans IGR. The arrowheads facing inward represent 48 AATTAA hexameric repeats to which SloR putatively binds. The vertical red bars denote the 49 positioning of the -10 and -35 promoter sequences of the *sloR*-specific promoter. Whether or not 50 SloR binds to the IGR fragment is shown with a (+) or (-) designation. 51

