Homologs of the small RNA SgrS are broadly distributed in enteric bacteria but have diverged in size and sequence

Richard S. P. Horler and Carin K. Vanderpool*

Department of Microbiology, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA

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ABSTRACT

Sugar phosphate stress in *Escherichia coli* is sensed and managed by the transcriptional regulator SgrR and the small RNA (sRNA) SgrS. SgrS is a dual function RNA that performs base pairing-dependent regulation of mRNA targets and encodes a small protein, SgrT. Homologs of SgrR were analyzed for gene synteny and inter-homolog identity to identify those that are likely to be functionally analogous. These 22 SgrR homologs were used to manually locate adjacent sRNAs functionally analogous to SgrS. SgrS homologs shared little sequence identity with *E. coli* SgrS, but most shared several structural features. The most conserved feature of SgrS homologs was the base pairing region while the most variable feature was the sgrT-coding sequence. Analyses of predicted interactions between SgrS:ptsG mRNA pairs in different organisms revealed interesting differences in the patterns of base pairing interactions. RNA pairs with more interrupted regions of complementarity had a higher proportion of G:C base pairs than those with longer contiguous stretches of complementarity. The identification of this set of homologous sRNAs and their targets sets the stage for future studies to further elucidate the molecular requirements for regulation by SgrS.

INTRODUCTION

Small RNAs (sRNAs) are ubiquitous and have been identified in the genomes of organisms from all three domains of life. Numerous studies using a variety of global approaches (1–3) have identified more than 80 sRNAs in the genome of *Escherichia coli* K12. The most abundant class of sRNAs is comprised of those that post-transcriptionally regulate expression of target genes by a base pairing-dependent mechanism that requires the RNA chaperone Hfq (referred to as riboregulation). The most extensively studied member of this class is RyhB, which was first identified in *E. coli* K12 and has been shown to regulate numerous genes involved in cellular iron homeostasis (4–6). RyhB homologs have been identified in *Erwinia chrysanthemi* (7), *Shigella dysenteriae* (8), *Vibrio cholerae* (9,10) and *Salmonella* (11,12) species on the basis of nucleotide sequence identity with the *E. coli* RyhB. Functionally analogous sRNAs have been identified by other means in more distantly related species such as *Pseudomonas aeruginosa* (13) and *Bacillus subtilis* (14). These RyhB-like sRNAs are expressed under the same conditions (iron limitation) and regulate the same types of target genes as the RyhB sRNAs in *E. coli* and related organisms yet share little sequence similarity. How widely RyhB and other base pairing-dependent sRNAs occur in the bacterial world remains to be determined, and this is no small task given the divergence of primary sequences of sRNAs and the inherent difficulty of locating sRNAs in genomes of distantly related organisms. Nevertheless, elucidating the conserved structural and functional features of bacterial sRNAs is a crucial step toward understanding how they evolve.

In this study, the species distribution and conservation of the sRNA SgrS was investigated. SgrS is an Hfq-dependent sRNA that was first identified in *E. coli* (15,16). SgrS expression is induced by a metabolic stress referred to as glucose-phosphate stress, so named because stress is associated with intracellular accumulation of glucose-6-phosphate (G6P) or its analog α-methyl glucoside-6-phosphate (αMG6P) (15,17). SgrS is unique from other characterized Hfq-dependent sRNAs because it has two distinct functions. The first is a base pairing dependent riboregulation function that results in translational repression and degradation of mRNAs that encode glucose transporters of the phosphoenolpyruvate phosphotransferase system (PTS) (15,18). The second is an mRNA function whereby SgrS encodes the 43-amino-acid polypeptide SgrT. SgrT inhibits glucose transport by a mechanism independent of the riboregulation activity (19). Together, these two functions serve to prevent...
further uptake of G6P or α-MG6P and thus promote recovery from glucose-phosphate stress (20).

The goals of the present study were 2-fold: (i) to define the distribution of SgrS and SgrT homologs and (ii) to analyze the conservation of determinants known from other studies (15,18,19) to be important for function. The genetic linkage of sgrS to sgrR, the gene encoding its regulator, greatly facilitated identification of SgrS homologs. Since sRNAs are typically not conserved at the primary nucleotide sequence level except between very closely related organisms, most homologs are not identifiable by BlastN searches. However, we were able to utilize sgrR as a genomic ‘handle’ to identify SgrS homologs that we would not have found by BlastN searches. This approach was used successfully to identify homologs of the sRNA GcvB, which, like SgrS, is encoded divergently from the gene encoding its transcriptional regulator, GcvA (21,22). In another study (C. S. Wadler and C. K. Vanderpool, submitted for publication), the functions of a subset of the SgrS homologs identified here were tested in a heterologous host (E. coli) in order to confirm their function and gain insight into the requirements for riboregulation and SgrT in the glucose-phosphate stress response.

Here we describe the identification of SgrR, SgrS and SgrT homologs in a variety of bacterial species. SgrR homologs are found in bacterial species belonging to a number of different phyla, however, those that we believe are most likely functionally analogous to E. coli SgrR are distributed only among the γ-Proteobacteria, primarily in enteric species. Likewise, our findings suggest that SgrS and SgrT homologs are mainly confined to enteric bacteria. Predictions for SgrS:target mRNA interactions suggest that the riboregulation function is relatively conserved, while comparison of SgrT homologs revealed that the mRNA function of SgrS is more variably distributed.

MATERIALS AND METHODS

Identification of SgrR homologs

The SgrR amino-acid sequence was retrieved from EchoBASE (23) and used as a query in a BlastP search against the non-redundant protein sequence databases at NCBI, UniProt database at EBI and the OMNIOME pep database at CMR. Additional BlastP searches using the same databases were performed using a subset of proteins obtained in the results of the primary search as queries. Results from all searches were curated to remove redundancy, which produced a list of 60 unique SgrR homologs. The Blast searches were completed in November 2008. The amino-acid sequences of the 60 SgrR homologs were aligned using ClustalW (Gonnet protein weight matrix) (The FASTA format amino-acid sequence alignment is available in Supplementary Figure S1). Phylogenetic and molecular evolutionary analyses were conducted using MEGA version 4 with the UPGMA, Neighbor Joining and Minimum Evolution algorithms (24). Homologs were divided into clades (Figure 1A and Supplementary Figure S2) based on the consensus among the trees. For analyses of gene synteny, chromosomal sequences were obtained from NCBI Entrez and EBI Genomes for each species containing a SgrR homolog. These GenBank files were analyzed using Artemis (Version 10) (25) from the Sanger Centre to examine gene context surrounding the sgrR homolog and obtain DNA sequences of the region between the nearest conserved upstream and downstream genes, tbaA and leuD, respectively. Where tbaA and/or leuD could not be found in close proximity to a sgrR homolog, Artemis (or WebACT) and colIBASE were used to identify their locations elsewhere on the chromosome. To further investigate the putative functions of genes in the sgrR neighborhood in different organisms, both colIBLAST (26) and EchoBASE (23) were used to find the corresponding E. coli homolog of the gene.

Identification of SgrS and SgrT homologs

The homologs of SgrR that appeared to be functionally analogous to E. coli K12 SgrR based on the above analyses were used as landmarks to look for a small RNA. The DNA sequence upstream of and on the opposite strand from the sgrR coding sequence through the next annotated gene was analyzed. All open reading frames within the retrieved sequences were compared against the SgrT amino-acid sequence of E. coli K12 using BL2Seq (27). In all genomes where a peptide with identity to SgrT was found, Artemis was used to define the start and stop codons for the open reading frame. Promoter elements and sgrS transcription start sites were identified primarily by alignment of sequences with the E. coli K12 sgrS gene and upstream sequences for which these elements have been experimentally determined (15,28). In addition, promoter predictions were generated using the Softberry BPROM network server and the Neural Network Promoter Prediction by the BDGP using prokaryotic settings. The sequences analyzed by these programs encompassed the region between sgrR and sgrT homologs. The predictions generated by the BPROM network server supported predictions derived from sequence alignments more frequently than predictions generated by the BDGP algorithm. Putative intrinsic terminators for sgrS were identified by examining sequences downstream of the sgrT stop codon manually and searching for an inverted repeat followed by a run of Ts. Mfold (29) and RNAalifold (30) were used to determine the secondary structure of this region to confirm the predicted stem loop structure.

Identification of base pairing determinants in SgrS homologs

Homologs of ptsG were identified in each genome containing an sgrR homolog by BlastP using the E. coli PtsG protein as the query. (The FASTA format alignment is given in Supplementary Figure S3.) Gene synteny around these homologs was analyzed using colIBASE for localized genome alignments and WebACT for genome wide alignments between pairs of genomes downloaded from GenBank.
ClustalW alignments were performed to localize base pairing determinants in SgrS homologs. These analyses identified a highly conserved region near the 3′ end corresponding to the region required for the base pairing function of *E. coli* K12 SgrS (15,18). To define interactions between this region of SgrS and *ptsG* mRNA, a Microsoft Excel Macro (created by R.S.P.H., available upon request) was used to produce an alignment that yielded the maximal complementarity between the base pairing region of SgrS and the 5′ untranslated region (UTR) of *ptsG* mRNA (defined for this purpose as a region from −100 through +30 with respect to the start codons). The macro determined maximal complementarity by optimizing both the total number of interactions and the maximum length of contiguous interactions over a 50-nt window.

**Alignments of SgrT and sgrR−sgrS intergenic region**

The amino-acid sequences of the SgrT homologs were aligned using ClustalW. DNA sequences of the intergenic region between the initiation codons of SgrR and SgrT were aligned using ClustalW. (This alignment is given in FASTA format in Supplementary Figure S4.)

**RESULTS AND DISCUSSION**

**Identification of homologs of SgrR**

To determine the extent of conservation of SgrR, SgrS and SgrT, a systematic search for homologs was undertaken. This search identified a total of 60 potential SgrR homologs. Alignments and phylogenetic analyses allowed us to divide these 60 members into seven clades that contained homologs from more than one genus (Figure 1A and B; Supplementary Figure S2); an additional clade contained two homologs found only in *Chromobacterium violaceum*. Representative proteins from each of the seven main clades show between 24% and 39% identity to SgrR of *E. coli* K12 (DR_B042 from *Deinococcus radiodurans* and STM3860 from *Salmonella enterica* serovar typhimurium, respectively; Figure 1B). Within the clade defined by *E. coli* SgrR (Clade 1, Figure 1A), identity with SgrR of *E. coli* K12 ranges from 57% to 98% (plu3677 from *Photorhabdus luminescens* and SSON_0075 from *Shigella sonnei*, respectively). Clade 2 members are homologous to an *E. coli* protein of unknown function, YbaE, and can be found in a variety of other enteric species. Proteins belonging to Clade 3 are more closely related to those of Clade 1 (SgrR) than those in other clades (Figure 1B and Supplementary Figure S2); these proteins, found in *Salmonella* species, *Chromobacterium violaceum*, *Yersinia enterocolitica* and *Aeromonas* species, may have originated from the duplication of an ancestral sgrR-like gene. Homologs belonging to Clades 4, 5 and 6 are narrowly distributed among gram negative enteric bacteria and Clade 7 encompasses proteins from gram positive organisms (Figure 1A).

The helix-turn-helix (HTH) motif weight and frequency tables of Dodd and Egan (31) revealed that homologs from Clades 5 and 7 were not predicted to contain a functional HTH motif characteristic of DNA-binding proteins (indicated in Figure 1A). Thus, these proteins may not be transcription factors like *E. coli* SgrR. Alternatively, they may be *bona fide* DNA-binding proteins that are too divergent from the proteins in the training set (31) to be accurately analyzed. Of the SgrR homologs that contain a conserved HTH motif, several are located on the opposite DNA strand divergent from genes that encode proteins with putative functions related to sugar transport or metabolism. By analogy with the *E. coli* sgrR-sgrS organization and pattern of regulation, we hypothesize that the genes encoded divergently from sgrR homologs are targets of transcriptional regulation by these homologs.

To narrow the list of homologs to those likely to be functionally analogous to *E. coli* SgrR, we focused on members of Clades 1 and 3 (Supplementary Figure S2) and further analyzed gene synteny and identity to SgrR of *E. coli* K12. We also examined genes divergently transcribed from each of these homologs and looked for characteristics of small RNAs. This analysis cut the list of proteins likely to be functional analogs of SgrR to 22. (Hereafter, when referring to ‘SgrR homologs’, we mean this subset likely to be functionally analogous to *E. coli* K12 SgrR.) The gene neighborhoods of putative SgrR homologs are depicted in Figure 2. Two genes that flank the sgrR region in all but three genomes and have no obvious link to the function of SgrRST are *tbpA*, which encodes the extracytoplasmic solute receptor of the thiamine ABC transporter, and *leuD*, encoding a leucine biosynthesis enzyme. Both of these genes were present near the location of all putative sgrR homologs, except those of *Aeromonas* species and *Photorhabdus luminescens*. In *Aeromonas* genomes, the *tbpA* and *leuD* genes are present in close proximity to one another but distant from the sgrR homolog. In *P. luminescens*, the *leuD* gene is encoded upstream of the sgrR homolog, but a genome rearrangement appears to have translocated *tbpA/thiPQ* and *mutH/ygdB* (Figure 2). The sgrR homologs of these organisms have the lowest levels of identity to *E. coli* K12 SgrR out of the set of 22 homologs. Although there are contradictory indicators for the sgrR homologs from *Aeromonas* species and *P. luminescens*, we retained them on the list of putative functional analogs because we found signatures of sgrS and/or sgrT adjacent to them.

Analysis of regulatory sequences adjacent to sgrR homologs supported our assignment of this subset of 22 proteins as likely functionally analogous to *E. coli* SgrR. The region upstream of the sgrR coding sequence is highly conserved (Supplementary Figure S5A and B). This is particularly true for the sequences that we have previously shown are important for transcriptional activation of sgrS (15,28). Furthermore, based on the conservation of the sequences that comprise the −10 element of the sgrS promoter in *E. coli* (15), we could assign a putative transcription start site for sgrS homologs from all organisms except *Y. enterocolitica* (Supplementary Figure S5A).

**Identification of homologs of SgrS and SgrT**

To find SgrS homologs, the region between sgrR and *leuD* was examined for features indicative of a small RNA.
These features included: (i) a promoter that could not be assigned to any other known gene, (ii) a small open reading frame encoding a protein with identity to SgrT, (iii) a region with complementarity to E. coli ptsG mRNA or with the species’ cognate ptsG mRNA and (iv) an inverted repeat followed by a run of T residues (putative intrinsic terminator). We identified putative SgrS homologs that met at least three of the four criteria in all of the genomes we investigated. All SgrS homologs also contained SgrR homologs (Figure 2, Table 1). In several cases, not all features were present, for example several putative SgrS homologs did not encode SgrT. Alignments of all SgrS homologs showed significant levels of identity between SgrS sequences only from organisms most closely related to E. coli (Supplementary Figure S6A). Although sequence identity across the whole length of sgrS homologs was minimal, a short stretch (13 nt) near the 3′ end was nearly invariant. This region corresponds to the sequences of E. coli K12 SgrS that constitute the determinants for base pairing with ptsG mRNA. The consensus sequence for this region is 5′-CUAGAUUUGUG-3′ where the central ‘u’ is the only position that varies among all SgrS homologs except those from Aeromonas and Photorhabdus species. The 22 SgrS homologs are found exclusively in species of γ-Proteobacteria, and with the exception of Aeromonas species, all are members of the Enterobacteriaceae. This distribution is similar to that of several other Hfq-dependent sRNAs (32–34), but is narrower than that of the GcvB sRNA (21) which is found in a broader range of γ-Proteobacteria.

The structure of a riboregulator plays a major role in its function, particularly for regions that interact with target
mRNAs. Although there are several bacterial riboregulators whose structures have been mapped, for those that have, base pairing sequences have often been localized to single-stranded regions. These single-stranded regions are sometimes localized to loop regions of stem-loop (SL) structures and are important for making initial interactions with loop sequences of their targets through formation of so-called “kissing complexes” (35). Loop sequences of sRNAs can also interact with non-loop single-stranded regions of their mRNA targets (33). Alternatively, single-stranded regions of sRNAs important for interactions with targets may be located adjacent to SL structures (21). Structural rearrangements promoted by the RNA chaperone Hfq may subsequently allow more extensive sRNA–mRNA interactions beyond those made by initial contacts (36).

Structural predictions for SgrS homologs were made by the RNA folding algorithms MFOLD (29), which calculates secondary structure based only on thermodynamic properties and RNAalifold (30), which predicts a consensus structure based on a set of aligned RNA sequences (Supplementary Figure S6A) and uses information from covariation to compute the secondary structure and energy score. The predictions of the two folding programs were highly similar, and revealed several conserved structural features that are shown schematically in Figure 3B. The 5' end of the hairpin at the 3' end. Most of the homologs are predicted to have an additional SL preceding the terminator. The base pairing sequences of most homologs (Table 1) are localized to a single-stranded region adjacent to this hairpin; in a number of cases some of the predicted base pairing residues extend into one side of the stem of this hairpin (Figure 3B). Only in the case of the Entero bacter SgrR homolog is the base pairing region predicted to be fully contained within a hairpin structure. Several SgrS homologs, including those of E. coli K12 and Shigella and Citrobacter species, have an additional small hairpin at the 3' end of the sgrT-coding sequence; the significance of this structure, if any, is not known. The 5'
Table 1. Characteristics of SgrS homologs and base pairing to ptsG

<table>
<thead>
<tr>
<th>SgrS ortholog species</th>
<th>SgrS length</th>
<th>SgrT (identity / similarity to E. coli K12)</th>
<th>Base pairing with cognate ptsG mRNA (RBS underlined, AUG highlighted)</th>
<th>Terminator</th>
<th>Promoter/ Base pairing scores</th>
</tr>
</thead>
<tbody>
<tr>
<td>Escherichia coli K12</td>
<td>227 nt</td>
<td>43 aa (100%/100%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.86 BP: ΔG: -24.2</td>
</tr>
<tr>
<td>Escherichia coli O157:H7</td>
<td>226 nt</td>
<td>N/A</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.86 BP: ΔG: -24.2</td>
</tr>
<tr>
<td>Escherichia coli CFT073</td>
<td>228 nt</td>
<td>43 aa (84%/93%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.86 BP: ΔG: -24.2</td>
</tr>
<tr>
<td>Shigella ( flexneri, boydii and sonnei)</td>
<td>227 nt</td>
<td>43 aa (100%/100%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.86 BP: ΔG: -24.2</td>
</tr>
<tr>
<td>Salmonella (typhimurium and enterica)</td>
<td>239 nt</td>
<td>40 aa (53%/72%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.95 BP: ΔG: -22.2</td>
</tr>
<tr>
<td>Klebsiella pneumonia</td>
<td>410 nt</td>
<td>50 aa (36%/60%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGATTATACGTGCTGTTTTTTTTT</td>
<td>Prom: 0.77 BP: ΔG: -20.1</td>
</tr>
<tr>
<td>Enterobacter sp. 638</td>
<td>246 nt</td>
<td>50 aa (44%/62%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAAATGTTTTTGTGCTGTTTTTT</td>
<td>Prom: 0.78 BP: ΔG: -26.0</td>
</tr>
<tr>
<td>Citrobacter koseri</td>
<td>236 nt</td>
<td>40 aa (58%/70%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGTTATTCTGCTGTTTTTTTTT</td>
<td>Prom: 0.93 BP: ΔG: -24.5</td>
</tr>
<tr>
<td>Citrobacter rodentium</td>
<td>242 nt</td>
<td>40 aa (58%/79%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGTTATTCTGCTGTTTTTTTTT</td>
<td>Prom: 0.87 BP: ΔG: -20.6</td>
</tr>
<tr>
<td>Erwinia carotovora</td>
<td>260 nt</td>
<td>54 aa (44%/63%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGCAAGTTATTCTGCTGTTTTTTTTT</td>
<td>Prom: 0.63 BP: ΔG: -22.6</td>
</tr>
<tr>
<td>Serratia proteamaculans</td>
<td>249 nt</td>
<td>57 aa (32%/51%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGTAACCCTGCTGTTTTTTTTTTTT</td>
<td>Prom: 0.75 BP: ΔG: -27.3</td>
</tr>
<tr>
<td>Serratia marcescens</td>
<td>267 nt</td>
<td>57 aa (32%/53%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGTGCGGCCACCCATGTTTTTTTTTT</td>
<td>Prom: 0.90 BP: ΔG: -25.5</td>
</tr>
<tr>
<td>Yersinia (pestis and pseudotuberculosis)</td>
<td>139 nt</td>
<td>N/A</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGTGGGCACCCATGTTTTTTTTTTT</td>
<td>Prom: 0.65 BP: ΔG: -28.1</td>
</tr>
<tr>
<td>Yersinia enterocolitica</td>
<td>84 nt</td>
<td>N/A</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GCCAGTGGTTATACGTGCTGCTGCTGTTTT</td>
<td>Prom: 0.62 BP: ΔG: -24.0</td>
</tr>
<tr>
<td>Aeromonas hydrophila</td>
<td>330 nt</td>
<td>58 aa (19%/41%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GATTTGAGCCTCTACCCATCATAAA</td>
<td>Prom: BP: ΔG: -16.0</td>
</tr>
<tr>
<td>Aeromonas salmonicida</td>
<td>325 nt</td>
<td>58 aa (21%/45%)</td>
<td>pteG 5' sgrT 3' AAAAGAACAGACACGACGCAGCGACGCCCAUUUAAAACAAAAAAAAAAAAAA</td>
<td>GATTTGAGCCTCTACCCATCATAAA</td>
<td>Prom: BP: ΔG: -16.0</td>
</tr>
</tbody>
</table>

Twenty-two homologs of SgrS were identified; 17 were unique as indicated by the rows in the table that contain more than one strain name. The SgrS homolog of E. coli K12 is denoted as SgrT and the cognate ptsG mRNA was predicted with base pairing interactions experimentally demonstrated for E. coli K12 (18) and automated alignment of the two RNA sequences (this study). For ptsG mRNA, the ribosome binding site is underlined and initiation codon is highlighted; for SgrS: sequences in blue represent the conserved base pair region. The predicted terminator sequence is shown with the inverted repeat of the SL underlined; the distance from the end of the predicted base pair region is indicated. The promoter scores were determined by Neural Network Promoter Prediction algorithm at BDGP. A score closer to 1.00 indicates a stronger promoter; this algorithm did not predict a promoter at the correct position relative to the probable +1 of SgrS in Aeromonas (based on alignment with E. coli K12 SgrS). The base pairing (BP) score was determined using UNAfold at the DINAMelt Server (http://dinamelt.bioinfo.rpi.edu/hybrid2.php) to predict AG for each pairing interaction.

The additional length for SgrS homologs is due to the presence of the sgrT coding sequence (CDS) (ranging from 120 to 171 bases) at the 5' end of most homologs. Homologs from Yersinia species, Photobacterium phosphoreum and Klebsiella pneumoniae deviate from this size range for reasons that will be discussed below. SgrT homologs range in size from 40 to 57 amino acids.
There are a number of conserved leucines and a repeating motif of a polar residue followed by an aromatic residue, but minimal conservation of other residues in SgrT (Figure 4 and Supplementary Figure S4). Despite the relatively low level of primary amino-acid sequence identity with SgrT of E. coli K12 (Table 1), SgrT homologs from S. typhimurium, K. pneumoniae and E. carotovora complement an E. coli sgrST mutant (C. S. Wadler and C. K. Vanderpool, submitted for publication), indicating conservation of SgrT function.

**Descriptions of individual SgrS and SgrT homologs**

While most SgrS homologs share major functional elements such as the presence of the sgrT CDS sequence, base pairing region and terminator, there are a number
of notable differences, particularly for homologs from organisms more distantly related to *E. coli*. Below and in Table 1 we have summarized the characteristics of individual homologs that may have relevance for their physiological functions. We begin the descriptions with *E. coli* SgrS and proceed to homologs from organisms that are increasingly more distantly related to *E. coli* (37).

Among complete and partially sequenced *Escherichia* genomes, we identified approximately 20 SgrS homologs. These homologs range in identity with genomes, we identified approximately 20 SgrS homologs.

![Figure 4. ClustalW alignment of SgrT homologs. The 14 unique SgrT sequences identified are aligned. (Of the 22 SgrS homologs, five do not encode SgrT and three others encode SgrT homologs identical to one of those shown.) The conserved leucines (l), and repeating polar (p) and aromatic (a) residue motifs are indicated above the alignment. The ClustalW consensus in red indicates the conserved residues when SgrT sequences from *Aeromonas* sp. are excluded; the consensus in black includes alignment of *Aeromonas* SgrTs. Abbreviations are as follows: EcK12, *Escherichia coli* K12; EcCFT, *Escherichia coli* CFT073; Shidy, *Shigella dysenteriae*; Shift, *Shigella flexneri*; Salty, *Salmonella typhimurium*; Citko, *Citrobacter koseri*; Citro, *Citrobacter rodentium*; Klebp, *Klebsiella pneumonia*; Enter, *Enterobacter sp.* 638; Serma, *Serratia marcesens*; Serpr, *Serratia proteamaculans*; Erwca, *Erwinia carotovora*; Aeros, *Aeromonas hydrophila*; Aeros, *Aeromonas salmonicida*.

| Sequence | Consensus | S. enterica serovar typhimurium LT2 and *S. enterica* serovar choleraesuis (referred to hereafter as *S. typhimurium* and *S. enterica*, respectively) contain identical SgrS and SgrT homologs. The *Salmonella* SgrT homolog is slightly shorter (40 aa) than *E. coli* SgrT (43 aa). The predicted secondary structure, shown schematically in Figure 3B, shows that the *Salmonella* homologs lack the SL structure that occurs directly upstream of sgrT in the *E. coli* K12 SgrS homolog. This suggests that the ribosome-binding site of *Salmonella* SgrS would not be occluded, a prediction supported by experimental data showing higher levels of SgrT production from *S. typhimurium* SgrS compared with *E. coli* SgrS (C. S. Wadler and C. K. Vanderpool, submitted for publication). The SL terminator for *Salmonella* SgrS is positioned 22-nt downstream of the base pairing region, rather than the 10 nt that is more common among the homologs; the significance of this is unknown. The additional nucleotides are predicted to participate in formation of the SL preceding the terminator (Figure 3B).

The *K. pneumoniae* SgrS homolog contains a longer SgrT (50 aa) than *E. coli*; the well-conserved base pairing region is located in the usual position 3′ of the sgrT CDS. However, the *K. pneumoniae* homolog does not contain a terminator SL in the typical position. Termination at the nearest predicted factor-independent terminator downstream would result in an sRNA of 410 nt. When the *K. pneumoniae* homolog is expressed in *E. coli*, an RNA of ~400 nt is produced (C. S. Wadler and C. K. Vanderpool, submitted for publication), consistent with the idea that transcription of this sgrS homolog terminates at the distal structure, which is 174-nt downstream of the base pairing region. There are no additional ORFs that would encode peptides longer than 15 aa located within this sequence, and the physiological role of this additional sequence is unknown.
The Enterobacter SgrS homolog also encodes a longer SgrT peptide (50 aa) compared with E. coli SgrT, but otherwise contains the same functional elements as E. coli SgrS. The base pairing region is positioned at the usual location 3' of sgrT; however, one predicted structural difference is the localization of base pairing elements to an SL structure. This differs from most homologs where the base pairing nucleotides are mostly unpaired (Figure 3B). In some cases, Hfq binding to sRNAs has been shown to remodel their secondary structure (36,38). Perhaps the Enterobacter SgrS undergoes more extensive Hfq-induced conformational changes than other homologs in order to interact with its cognate ptsG mRNA.

The two homologs from the Citrobacter genus both encode a 40 aa SgrT and have similar base pairing sequences complementary to their cognate ptsG mRNAs (Table 1). The position of the terminator relative to the base pairing region differs between these two homologs: the Citrobacter rodentium SgrS has a 19-nt spacer between these elements in contrast to the 6-nt spacer in Citrobacter koseri SgrS.

The SgrS homolog found in Erwinia carotovora encodes the second longest SgrT homolog at 54 aa. Structural analyses indicate a short SL structure upstream of the sgrT CDS similar to the one identified in E. coli K12 SgrS (Figure 3B). However, unlike the GC-rich stem in E. coli, the putative stem in E. carotovora contains two A:U base pairs that should weaken the interaction. In vivo complementation studies of an E. coli sgrST mutant with the E. carotovora sgrST strongly suggest that E. carotovora’s sgrT is well-translated and functional (C. S. Walder and C. K. Vanderpool, submitted for publication). These data argue against any significant translational inhibition mediated by this structure. The base pairing region and terminator of the E. carotovora sgrS are located at the typical 3' position.

SgrS homologs found in Yersinia species range in size from 84 to 139 nt owing to the lack of an sgrT. For Y. enterocolitica, this appears to be caused by a 5' truncation of sgrS resulting from insertion of two genes, ybiY and ybiW, in the sgrR-sgrS intergenic region (Figure 2). The ybiYW genes are uncharacterized, but they are predicted to encode a pyruvate formate lyase and its activating enzyme (23). Notably, the Y. enterocolitica sgrS homolog is the only one that is not divergently transcribed from sgrR. Whether ybiYW expression is controlled by SgrR is not known, though the conservation of regulatory sequences adjacent to sgrR and upstream of ybiY (Supplementary Figure S5A) suggests that this might be the case. Examination of the region immediately upstream of the Y. enterocolitica sgrS homolog (downstream of ybiW) did not reveal any additional putative SgrR regulatory sequences, suggesting that if Y. enterocolitica sgrS is expressed, it may be operonic with ybiYW. For Y. pestis and Y. pseudotuberculosis the cause of the 5' sgrS truncation is not obvious; there are no apparent insertion sequences or other signatures of mobile elements in this region. However, it has been reported that Yersinia genomes are extremely plastic and extensive rearrangements are common (39). Despite extensive differences at their 5' ends, the base pairing and terminator elements of Yersinia sgrS homologs are conserved and located at typical distances from one another. Another interesting characteristic of Yersinia SgrS homologs is their extensive pattern of predicted complementarity with their cognate ptsG mRNAs. The predicted interactions for Y. pestis and Y. pseudotuberculosis SgrS: ptsG mRNA pairs represent the longest contiguous stretches of complementarity for any homologous pair. In another study, it was shown that the Y. pestis sgrS homolog can complement an E. coli sgrST mutant, indicating that this truncated SgrS is still a fully functional riboregulator (C. S. Walder and C. K. Vanderpool, submitted for publication).

The Serratia proteamaculans and S. marcescens SgrS homologs encode the longest SgrT proteins at 57 aa. While the sRNA is slightly longer to compensate for the longer CDS, the base pairing region and terminator occur at typical positions relative to one another. The predicted base pairing interactions with the cognate ptsG mRNA are striking, as they are comprised of a single long stretch of complementarity similar to that predicted for Yersinia species (Table 1).

The SgrS homologs of Aeromonas species and P. luminescens are the most divergent from E. coli SgrS. The sgrT CDS is not present in the P. luminescens SgrS while the SgrT encoded by A. hydrophila and A. salmonicida shows very low identity with E. coli SgrT (Figure 4, Supplementary Figure S4). The genome of P. luminescens does not contain a ptsG homolog; however, it does contain genes encoding another PTS capable of transporting glucose, manXYZ. Results from our laboratory have shown that E. coli SgrS also regulates manXYZ mRNA (J. B. Rice and C. K. Vanderpool, unpublished results), so perhaps this regulation is conserved in P. luminescens. However, due to the poor similarity between P. luminescens and Aeromonas SgrS and E. coli SgrS, we cannot claim with certainty that these homologs are functionally analogous to E. coli SgrS.

**Base pairing of SgrS homologs to mRNA targets**

One of the great challenges in bacterial small RNA research has been to define rules that govern RNA:RNA interactions that result in positive or negative regulation of mRNA targets. Since we have identified a set of homologous sRNAs, we analyzed interactions between these sRNAs and their cognate targets in order to gain some insight into the requirements for regulation by SgrS. In order to conduct these analyses, we first identified ptsG homologs in the genomes where we had already identified SgrR, SgrS and SgrT homologs. For each of these genomes, with the sole exception of P. luminescens, a ptsG homolog with strong identity (Supplementary Figure S3) and gene synteny to E. coli ptsG was identified.

We previously predicted (15) that for E. coli, the SgrS:ptsG mRNA base pairing interactions occurred through short, interrupted segments of complementarity that we will refer to as a 5-8-4 pattern (referring the number of contiguous base pairs in each segment). The 5-8-4 pattern of complementarity sequesters most of the ribosome-binding site (RBS) of ptsG mRNA resulting in translation inhibition of ptsG (40). The base pairing
predictions were supported by genetic evidence obtained by another group (18). These results suggested that the central eight contiguous base pair segment was particularly important for regulation. However, one caveat to these experiments is that they only measured the effect on steady state levels of *ptsG* mRNA at 20 min after SgrS expression was induced. Thus, subtle contributions of the flanking regions of complementarity to the initial kinetics of regulation would not have been elucidated. At this time there have been no structural probing experiments to validate or refute the base pairing predictions.

The relative position of interacting sequences is conserved among all homologous pairs, i.e. 3’ of the *sgrT* CDS for SgrS and at least partially encompassing the RBS of *ptsG*. This strongly suggests that the regulatory outcome of translation inhibition and destabilization of *ptsG* mRNA is also conserved. The 5-8-4 pattern is conserved for SgrS:*ptsG* mRNA partners in *E. coli* and *Shigella* species, and is very similar (4-9-4) for *Salmonella* species (Table 1). Interestingly, the more divergent SgrS:*ptsG* mRNA pairs are predicted to have more extensive interactions. The *Erwinia*, *Yersinia* and *Serratia* homologs have a longer contiguous segment of complementarity that encompasses between 15 and 21 base pairs. For the most divergent SgrS homologs in *Aeromonas* species, the predicted interactions are somewhat shorter and more interrupted, but are still predicted to occlude the RBS.

There is currently little data to shed light on how the base composition of the base pairing region affects regulation by small RNAs. The mutations with the strongest negative effects on regulation by SgrS alter G:C base pairs in the central 8-bp region of SgrS:*ptsG* mRNA interactions in *E. coli* (18). As G:C base pairs form three hydrogen bonds versus the two hydrogen bonds contributed by A:U or G:U pairs, it would not be surprising if G:C base pairs make the most substantial contribution to the riboregulation. The *E. coli*, *Shigella* and *Salmonella* interaction regions are all composed of ~50% G:C base pairs. Interestingly, for homologous pairs with more extensive complementarity, e.g. *Erwinia*, *Yersinia* and *Serratia*, G:C base pairs comprise only ~35% of the interactions. Perhaps a more A:U/G:U-rich base pairing region requires a greater number of consecutive base pairs to achieve a rapid interaction *in vivo*.

To provide another qualitative way to compare these predicted sRNA:mRNA complexes, we utilized the DINAMelt server (41) to predict melting profiles for the short segments of SgrS and *ptsG* mRNA shown in Table 1. The ΔG values reported for each pair (Table 1) are meant only as a means to compare the relative stabilities of these short regions of complementarity for each homologous sRNA:mRNA pair. The ΔG values ranged from −16 to −28, with the majority falling between −22 and −26. The native *E. coli* K12 SgrS:*ptsG* mRNA pair had a ΔG value of −24.2. A single point mutation in SgrS that alters one G:C base pair in the central segment of complementarity reduced the ΔG value to −19; an additional mutation in sgrS that alters a second G:C base pair in this region further reduces the ΔG value to −10. We have shown that the *E. coli* SgrS single point mutant has a partial defect in regulating *ptsG* mRNA while the double point mutant is strongly defective for *ptsG* regulation [18] and C. S. Wadler and C. K. Vanderpool, submitted for publication]. Given this information, it is interesting to note that *K. pneumoniae* and *C. rodentium* SgrS:*ptsG* mRNA pairs have a ΔG value of −20 while the *A. hydrophila* ΔG value is only −16, suggesting that SgrS may not regulate *ptsG* in *A. hydrophila*. It will be interesting in the future to examine regulation by these SgrS homologs and determine how efficiently they regulate their cognate *ptsG* mRNAs.

**CONCLUSIONS**

We identified homologs of the transcription factor SgrR and the sRNA SgrS in a variety of enteric bacterial species. Most homologs appear to possess both of the functions that we have previously described for *E. coli* SgrS (15,19): the riboregulation function and ability to produce the SgrT protein. The bioinformatic analyses described in this study suggest that the riboregulatory function of SgrS is the more conserved of the two functions of SgrS, since in a few homologs the *sgrT* is missing or rendered non-functional by mutation of the start codon. Homologs of SgrR, SgrS and SgrT are more narrowly distributed than PtsG homologs, which are found in a much wider range of organisms. Not surprisingly, conserved sequences in the 5’ UTR of *ptsG* are found only in the same genomes where a SgrS homolog is found (Supplementary Figure S3 and data not shown). These observations suggest that either glucose-phosphate stress is not a problem for all organisms that use PtsG homologs to take up glucose, or that other mechanisms have evolved to deal with the stress.

To further test the roles of riboregulation and SgrT in the glucose-phosphate stress response, several of the homologs identified in this study were expressed in a heterologous (*E. coli*) *sgrST* mutant host. The experimental results of that study (C. S. Wadler and C. K. Vanderpool, submitted for publication) support the hypotheses generated by the identification and comparison of these homologs. Together, these studies reveal that despite low levels of nucleotide sequence similarity, SgrS and SgrT homologs are functionally interchangeable.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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