A highly conserved protein of unknown function in *Sinorhizobium meliloti* affects sRNA regulation similar to Hfq

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**ABSTRACT**

The SMc01113/YbeY protein, belonging to the UPF0054 family, is highly conserved in nearly every bacterium. However, the function of these proteins still remains elusive. Our results show that SMc01113/YbeY proteins share structural similarities with the MID domain of the Argonaute (AGO) proteins, and might similarly bind to a small-RNA (sRNA) seed, making a special interaction with the phosphate on the 5'-side of the seed, suggesting they may form a component of the bacterial sRNA pathway. Indeed, eliminating *SMc01113/YbeY* expression in *Sinorhizobium meliloti* produces symbiotic and physiological phenotypes strikingly similar to those of the *hfq* mutant. Hfq, an RNA chaperone, is central to bacterial sRNA-pathway. We evaluated the expression of 13 target genes in the *smc01113* and *hfq* mutants. Further, we predicted the sRNAs that may potentially target these genes, and evaluated the accumulation of nine sRNAs in WT and *smc01113* and *hfq* mutants. Similar to *hfq*, *smc01113* regulates the accumulation of sRNAs as well as the target mRNAs. AGOs are central components of the eukaryotic sRNA machinery and conceptual parallels between the prokaryotic and eukaryotic sRNA pathways have long been drawn. Our study provides the first line of evidence for such conceptual parallels. Furthermore, our investigation gives insights into the sRNA-mediated regulation of stress adaptation in *S. meliloti*.

**INTRODUCTION**

In a symbiotic event, rhizobial bacteria, e.g. *Sinorhizobium meliloti*, colonize the legume roots to fix biological nitrogen. This symbiosis is extremely important not only from agricultural, economical and environmental perspectives, but also the interaction of rhizobial bacteria with its legume host is a valuable model system to study chronic infection processes (1). The rhizobial bacteria alternate between a free living phase in the soil/rhizophere and a symbiotic phase within the host plant cells where they differentiate into nitrogen fixing bacteroids (2). Symbiosis comprises a complex series of events that begin with the infection process, which is initiated with the exchange of signals between the bacteria and the plant hosts (3,4). The signal components include the NOD factors, secreted proteins, polysaccharides, flavonoids, ligand-gated-ion channels, the calcium-calmodulin dependent protein kinases and calcium spiking (3,5). Invasion of plant roots by bacteria involves the development of an infection thread and cytoskeleton remodeling. During the invasion, bacteria encounter a plant innate immune response- a burst of reactive-oxygen-species (ROS) (4). Further along, the infection thread is targeted to the cortical tissues, where the rhizobia are endocytosed resulting in formation of symbiosome, followed by their differentiation into nitrogen-fixing bacteroids. Adaptation to the extracellular environment of the host cells requires lipopolysaccharide (LPS) and non-LPS factors such as those needed for coping with ion-stress (3,4). Bacteria enclosed within the symbiosome membrane encounter very low oxygen levels, where they can express enzymes of the nitrogenase complex and begin nitrogen fixation (4). Fixed carbon provided by plants in the form of...
dicarboxylic acids is metabolized to provide the energy necessary for nitrogen fixation.

Large-scale transcriptional, proteomic and metabolic rearrangements underlie the complex transition from the free-living state to the symbiotic state, as well as the changes when the free-living bacteria adapt to changing environmental conditions (2,6–9). For example, a ‘base-line proteomic output’ evaluated proteins from rhizobia grown in several conditions: an accumulation of a total of 2224 proteins, derived from 810 distinct genes and resulting in alteration of 53 metabolic pathways was determined (8). Analyses comparing transcription during microoxic versus oxic conditions and between free-living versus symbiotic states in S. melliloti revealed hundreds of genes, including several potential regulators, being differentially regulated (2). The microoxic and bacteroid transcriptomes partially overlapped, suggesting complex modes of gene regulation in different environments. Furthermore, the development of a ‘dual-genome symbiosis chip’ facilitated identification of changes in approximately 5000 transcripts during symbiosis and when S. melliloti is grown in different environments (9). Thus, complex transcriptional rearrangements occur in S. melliloti during its adaptation to environmental stresses and symbiosis. How these large scale transcriptional and proteomic responses are coordinated remains largely unknown. The magnitude of responses suggests that transcriptional regulation by certain proteins may not be the only mechanism of gene regulation.

RNA silencing has emerged as a fundamental regulatory process affecting many layers of endogenous gene expression; non-coding small-RNAs (sRNAs) appear to be important regulators of gene expression during stress responses in both eukaryotes and bacteria. It is only relatively recently that the pervasiveness, importance and roles of the sRNA-mediated regulation in bacteria have become apparent. They modulate transcription, translation, mRNA stability and genomic integrity via several mechanisms including changes in RNA conformation, protein binding, base pairing with target RNAs and interactions with DNA (10). Most known sRNAs regulate gene expression by base pairing with their target mRNAs that is facilitated by the RNA chaperone, Hfq; Hfq is implicated as a central player in the sRNA-mediated regulation during stress; non-coding small-RNAs (sRNAs) appear to be important regulators of gene expression during stress responses in both eukaryotes and bacteria. It is only relatively recently that the pervasiveness, importance and roles of the sRNA-mediated regulation in bacteria have become apparent. They modulate transcription, translation, mRNA stability and genomic integrity via several mechanisms including changes in RNA conformation, protein binding, base pairing with target RNAs and interactions with DNA (10). Most known sRNAs regulate gene expression by base pairing with their target mRNAs that is facilitated by the RNA chaperone, Hfq; Hfq is implicated as a central player in the sRNA-mediated regulations (11–14).

Although much is understood about the role of sRNAs in modulating gene expression in the model bacteria, Escherichia coli and Salmonella (10,15), comparatively less is known about the role of the sRNA-mediated interaction in the model symbiotic bacteria, S. melliloti. A few studies have recently appeared that use comparative genomics and prediction approaches to identify sRNAs in the S. melliloti genome (16–18). Although expression of few of the predicted sRNAs was verified in these studies, the physiological roles of most of these sRNAs, their possible targets and the pathways that may be under their control remain largely unknown. In our and related recent studies, mutation of hfq resulted in S. melliloti that are highly compromised in their symbiotic ability (19–22), suggesting that the sRNA-pathways form a central component of the gene regulation during symbiosis and stress adaptation. Characterization of the symbiotically deficient hfq mutant provided important clues in understanding the underlying molecular mechanisms of regulation during symbiosis and stress adaptation. Nevertheless, hardly any specifics for Hfq-dependent sRNA–mRNA interactions are known in S. melliloti.

A screen that we had conducted to identify oxidative-stress-compromised symbiotically deficient mutants led us to identify SMc01113 gene as essential for symbiosis (23,24). SMc01113 encodes a protein of unknown function that is strongly conserved in most bacteria (24); Escherichia coli ortholog is named YbeY. The SMc01113/ YbeY proteins belong to the UPF0054 family. While the crystal structures of the YbeY/UPF0054 family members from E. coli, Aquifex aeolicus and Thermotoga maritima have been determined and suggest that YbeY is a metallo-protein (25–27), the cellular and biochemical function of this protein remains elusive. It has been annotated as a heat shock protein, as a peptidase or even as a protease (25,28,29). Recently, in E. coli, YbeY has been reported to be involved in translational regulation during high-temperature adaptation, presumably by affecting ribosome activity (28,29).

In the current investigation, we explore the possible cellular function of the S. melliloti SMc01113 gene that appears to be central to symbiosis. Our sequence and structure analysis suggested SMc01113’s similarity to a domain of Argonaute (AGO) proteins, which are central to sRNA mediated interaction in eukaryotes (30,31). AGOs are multi-domain-proteins comprising of N, PAZ, MID and PIWI domains: the N- (or the amino-terminal) domain projects a basic surface face for putative RNA recognition; the PAZ domain contains a specific binding pocket that anchors the characteristic 2-nt 3’-overhang that results from digestion of RNAs by RNase III (a step in the processing of sRNAs); the PIWI domain shows extensive homology to RNase H, and these are often referred to as ‘slicers’; the MID domain resides between the PAZ and PIWI domains and binds the characteristic 5'-phosphates of smRNAs, thus anchoring them onto the Ago protein; the MID domain has also been implicated in protein–protein interactions (32,33). AGO proteins constitute the key components of the ‘RNA-induced-silencing-complex’ (RISC) that modulates sRNA-dependent gene expression, and are mostly found in eukaryotes. This suggested the possibility that SMc01113 may form a component of, or be involved in, the interactions mediated by sRNA-pathways. We used a comparative approach with hfq as a reference to investigate this hypothesis. Our results suggest a possible role of SMc01113 in interactions mediated by the sRNA-pathway.

**MATERIALS AND METHODS**

**Strains, media and culture conditions**

Strains used in this study are detailed in the Supplementary Table S2. *Sinorhizobium melliloti* strains were grown aerobically at 30°C in the complex LB medium containing 5 ml/l of 0.5M CaCl₂ and 25 ml/l of...
MgSO₄ (LB-MC) to an optical density at 600 (OD₆₀₀) of 1.5–2; they were then inoculated in the LB-MC or rhizobial minimal medium (RMM) with glucose at an OD₆₀₀ of 0.1 (19,20). For selection of S. meliloti, streptomycin (500 µg/ml) and gentamycin (50 µg/ml) and neomycin (200 µg/ml) were used. Bacto agar was used at 1.5% for solid medium.

### Plant nodulation assay and fitness estimates

Nodulation assays for plants were conducted as previously described (19). Briefly, the alfalfa (Medicago sativa cv. Iroquois) seeds were surface sterilized and germinated on 0.8% agar in water in dark for 3 days; seedlings were placed on the top of a petri dish with 25 ml Jensen’s agar, and inoculated with 1 ml appropriate strain of S. meliloti grown to saturation in LB-MC. Plates were wrapped in aluminium foil and incubated at 25°C for 4 weeks, after which fitness parameters such as plant height, number of pink/white nodules, number of green leaves and plant dry weight were recorded.

### Physiological characterization following challenge with stresses

Bacterial response to various stresses for WT, smc01113 and hfq mutants S. meliloti was evaluated by analysis of the growth curves in liquid LB-MC or RMM. The effect of mutating SMc01113 on swarming behavior was evaluated on swarming plates by measuring the diameter of bacterial spreading as previously described (19) and compared to the WT and hfq mutant. Similarly, the effect of heat shock (50°C for 35–40 min) was evaluated on the WT, smc01113 and hfq mutants as previously described (19,20). Ability to respond to various other environmental stresses was evaluated by growing the WT, smc01113 and hfq mutants with or without Paraquat (100 mM), MMS (0.025%), ethanol (2.5%), SDS (0.1%), NaCl (50 mM) or cefotaxime (50 µg/ml) for 48 h, where the cells were incubated with shaking at 30°C. Aliquots were collected at different time intervals, OD₆₀₀ was measured and residual growth was determined (19,20).

### Sequence and structure analysis

Similarity between the SMe01113, YbeY and AGO protein sequences was determined by aligning the sequences with CLUSTALW (http://www.ebi.ac.uk/Tools/clustalw2/index.html). Structural homology search was performed using YbeY (PDB ID: 1XM5; chain A) as the search probe on DALi server (34) and pair-wise comparisons were done using DALIlite (35). Structural similarities between the YbeY (1XM5) and the A. aeolicus AGO (2NUB) protein were also determined by super-imposing the YbeY structure (27) with the structures of complete AGO protein and MID/PWI domain (32) using the Crystallographic Object-Oriented Toolkit [COOT; (36,37)]. YbeY structure was analyzed by Hotpatch server (http://hotpatch.mbi.ucsd.edu/) (38) for the DNA/RNA binding properties and for finding functionally important patches on its surface. YbeY (PDB ID: 1XM5; chain A) and 4mer RNA (generated from PDB ID: 2BGG, chain R) were docked by using the ClusPro 2.0 docking server (http://cluspro.bu.edu) in order to model YbeY and RNA interactions. A single phosphate (PO₄) ion was manually docked in COOT (36,37). Electrostatic surface calculation and all figures were made by PYMOL. Sequences homologous to SMc01113/YbeY in other bacterial species were extracted after performing a non-redundant global BLAST search with the SMc01113. Homologous sequences were extracted from NCBI, multiple alignments were made and phylogenetic tree was constructed using CLUSTALW. Similarly, homologous sequences were searched in closely related species (S. medicae, Rhizobium sp. NGR234, R. leguminosarum and R. etli) using the ‘genome BLAST’ tool at the NCBI (http://www.ncbi.nlm.nih.gov/sutils/genom_table.cgi). The homologous sRNA sequences were extracted, multiple alignments were made and phylogenetic trees were constructed using CLUSTALW.

### Expression analysis by quantitative real-time PCR

The smc01113 and hfq mutants and the WT S. meliloti were grown aerobically at 30°C in the LB-MC medium to an OD₆₀₀ of 1.5–1.8; they were then inoculated in RMM with 0.5% glucose, and grown for 24 or 48 h, after which the cells were harvested by centrifugation. Total RNA was isolated by TRI reagent (T 9424; Sigma, St Louis, MO, USA) using manufacturer’s protocol. SYBR green assays were developed using Express One-step SYBRGreenER kit (#11780-200; Invitrogen; http://www.invitrogen.com) following the manufacturer’s protocol. All the quantitative real-time PCR (qPCR) assays were performed with 50 ng total RNA. Three to four biological replicates were used for each genotype; each biological replicate was used twice on the qPCR plate. The 2⁻ΔΔCT method (relative quantitation) was used for data analysis. Briefly, the Ct (threshold cycle) values of target genes were normalized to the endogenous reference gene (ΔCt = Ct_target−Ct_reference) and compared with those values obtained from the calibrator (∆∆Ct = ∆Ct_sample−∆Ct_calibrator). To simplify data interpretation, expression levels in control samples (WT, Rm1021) were fixed to 1 and relative expression levels were calculated with respect to this reference value in the smc01113 and hfq mutants. A list of primers used in this study is available in the Supplementary Table S3. All the gene-specific primers were designed with Primer3 software (http://frodo.wi.mit.edu/primer3/), and all the assays were run on the ‘AB7500 fast real-time PCR system’ (Applied Biosystems; http://www.appliedbiosystems.com/absite/us/en/home.html).

### Bioinformatic analysis of sRNAs-target interactions

In order to determine if the S. meliloti sRNAs targeted the genes differentially regulated in the smc01113 and hfq mutants, and informed by the fact that sRNAs regulate the gene expression by mostly base pairing with their target mRNA sequences, we used the concept of ‘seed pairing’, already used in some other bacteria (39–42). Complete coding sequences, obtained from the Rhizobase database (http://genome.kazusa.or.jp/rhizobase/), were
used for genes differentially regulated in the smc01113 and hfg mutants. Using custom-written perl scripts, ‘seed-pair analysis’ of sRNAs was conducted for the target genes: small ‘seed’ sequences, >7 nt, starting from the first as well as the second nucleotide at the 5'-end of the sRNA, with a floating window of 1 nt, were generated. Then, sequences from the 5'-ends of the target mRNA was compared for perfect complementarities with sRNA-seed-sequences from the 5'-ends. These seeds were mapped to the sequences of the target genes for ‘Watson–Crick’ base pairing, and a hit-map was generated (Supplementary Table S1). Most trans-acting sRNAs have the seeds >9 nt (10,13,43), therefore we used a length of 10 or more perfectly matching nucleotides as a cut-off. To test the statistical significance, similar analysis of predicting sRNA-complementary motifs was performed on a set of unrelated and randomized mRNA sequences [from plants; (44)]. Significantly low number of hits were obtained for seeds >10 nt (Fisher’s exact test, P ≤ 0.05).

In order to determine the conservation of the sRNA-target interaction sites for specific sRNAs and target mRNAs, we used sequences from three closely related species: S. meliloti, S. medicae and Rhizobium sp. NGR234. We extracted the longest ‘seed’ sequence from the target as well as sRNA. Sequences homologous to these ‘seed sequences’ were extracted from the aforementioned multiple alignments of the sRNAs and targets. Separate conservation profiles for sRNAs and targets were generated using the ‘WebLogo’ program (45,46). It is advantageous to use this methodology of determining the conservation as it gives frequencies of conservation of a given nucleotide at a particular position, consensus sequence and the total sequence conservation.

**Over-expression of sRNAs**

For over-expression of sra16 and sra35, genomic DNA corresponding to the appropriate sra coding sequence was PCR amplified from isolated S. meliloti genomic DNA and with engineered 5'-XhoI and 3'-KpnI restriction sites. These DNA fragments were then cloned into XhoI/KpnI-digested pMS03 (47) under the control of the constitutive Trp promoter. Transfer of pMS03-sra16 and pMS03-sra35 into S. meliloti (SRA16 and SRA35) was achieved by tri-parental mating followed by selection of S. meliloti harboring the plasmid on LB/MC containing 100 μg/ml spectinomycin and 500 μg/ml streptomycin. WT Rm1021 strains were also grown with streptomycin. Over-expression of sra16 and 35, and its effects on their predicted targets were evaluated by qPCR analyses after growing the cells to late exponential/early stationary phase. We evaluated the affect of pMS03 vector in the Rm1021 by similarly generating control Rm1021 strains with empty vector. No significant difference in growth or gene-expression was observed between the WT and the control WT strain harboring empty vector, which confirmed that presence of the vector had no effects on bacteria (data not shown). Similarly, over-expression of sra35 had no affect on growth of S. meliloti; sra16-over-expressing strain grew marginally better than WT or WT with empty vector controls (data not shown).

**Yeast two-hybrid analysis**

For cloning, XL1Blue E. coli were plated on LB media supplemented with 100 μg/ml of ampicillin and incubated at 37°C. Yeast were grown on synthetic complete medium containing 2% dextrose (SCD) and lacking the appropriate nutrient (48). Plates were supplemented with 2% agar. Yeast cultures and plates were incubated at 30°C.

Plasmids were constructed by PCR amplification of the targeted gene from E. coli genomic DNA (strain MC4100) using oligonucleotides containing homology to the gene of interest flanked by either EcoRI for the 5'-primer or BglII for the 3'-primer (Supplementary Table S3). The 5'-primer lacked homology to the start codon of the amplified ORF to insure correct fusion to the Gal4 binding domain (BD) or activation domain (AD). After amplification, the amplicon was digested with EcoRI and BglII, and ligated to EcoRI and BglII-digested pGBD-C2 and pGAD-CD plasmids (Supplementary Table S2). Correct clones were confirmed by DNA sequencing.

To check for protein interactions, the budding yeast strain PJ69-4A (49) (MATa trp1-901 leu2-3, 112 ura3-52 his3-200 gal4Δ gal80Δ lys2::GAL1-HIS3 GAL2-ADE2 met2::GAL7-lacZ) was transformed sequentially using a standard lithium acetate protocol (50). In the first round, PJ69-4A was transformed with plasmids pGBK-C2, pM122, pGAD-C2 and pM130 selecting for either Trp+ (pGBK-C2, pM122) or Leu+ transformants (pGAD-C2 and pM130) on SCD–Trp or SCD–Leu, respectively. Colonies from the first round were subsequently transformed with the second plasmid containing the putative interacting protein and selected for on SCD–Leu–Trp. Finally, multiple Leu+ Trp+ colonies from each transformation were pooled and resuspended in 100 μl of ddH2O. Ten microliter of the resuspended colonies were spotted on SCD–Leu–Trp to confirm growth and SCD–Leu–Trp–Ade to select for protein–protein interactions. Plates were incubated at 30°C for 5 days before interpretation of data.

**RESULTS**

Smc01113 shows sequence and structural similarities to the AGO proteins

Smc01113 homologs are found in nearly every bacterium including symbiotic and pathogenic bacteria [Supplementary Figure S1; (24,27)]. Smc01113 and its E. coli homolog, YbeY, show amino acid similarity of >70% (Figure 1A). Expression of the E. coli homolog complements the symbiotic defects of the smc01113 mutant of S. meliloti (24,51). Sequence and structural similarities of the YbeY to AGO protein were readily apparent when we performed homology studies with the YbeY. Sequence alignment of Smc01113/YbeY showed similarities with AGO protein sequences from...
Neurospora crassa, A. Aeolicus and other species. In particular, SMc01113/YbeY showed sequence similarity with the MID domain of the AGO protein (Figure 1). The MID domain contains the binding site for the 5' end of the eukaryotic sRNAs, by binding to the PO4 group of a nucleic acid. The structure of *E. coli* YbeY (27) has been solved, as have the structures of AGO proteins from *A. aeolicus* and the *Thermus thermophilus* (32,33). Recently, the crystal structure of a eukaryotic MID-domain, recognizing the 5'-terminal phosphate of a guide RNA, has also been solved, and it highly resembles the previous structures of MID domains from other species (52).

![Figure 1](https://academic.oup.com/nar/article-abstract/39/11/4691/1145450)

*Figure 1. Sinorhizobium meliloti* ORF, SMc01113/ybeY, shares similarities with Ago protein. (A) Alignment of the protein sequences of the *S. meliloti* and *E. coli* homologs, SMc01113 and YbeY. (B) Alignment of protein sequences of the YbeY and Ago-Mid domain. Structural alignment of Ago-Mid domain (red) and YbeY (green) from *N. crassa* (C) and *A. aeolicus* AGO (2NUB) proteins (D). Manually docked PO4 shows potential interactions with the Arg59, K61 residues (E) in the positively charged cavity (F). (G) The lowest energy docked 4mer RNA (Cluspro 2.0 web server) onto YbeY surface fits nicely into the protein cavity, with negatively charged RNA backbone aligned towards the positively charged protein surface.
A structural homology search with YbeY (PDB ID: 1XM5) by using the DALI server (34) against the full PDB database and PDB90 (34) resulted in multiple hits corresponding to a wide variety of domains with RNA-binding, enzymatic and regulatory functions. Among all the DALI hits, the MID domain of N. crassa AGO (PDB code: 2XDY), Pyrococcus furiosus AGO (PDB code: 1Z26) and the MID domain from human AGO (PDB ID: 3LUJ) show statistically significant Z-scores of 3.6, 3.2 and 2.9, respectively. Further, a pair-wise structure alignment using DAiLiLite (35) yielded a significant Z-score of 4.7 for YbeY (PDB ID: 1XM5) and the N. crassa AGO MID domain (PDB ID: 2XDY). Indeed, there are 83 equivalent C-alpha positions between YbeY and the N. crassa AGO MID domain with a root mean square deviation (RMSD) of 3.4 Å (Figure 1C). Further, we tested the structural similarities of YbeY with the MID-domain of the A. aeolicus AGO protein; structural superimposition showed RMSD of 2.6 Å for 72 aligned residues indicating structural similarity (Figure 1D).

Encouraged by these results, we further explored the potential of YbeY to bind RNA. YbeY and a 4mer RNA (52) were docked by using the fully automated, web-based program ClusPro version 2.0 (53,54) with default parameters (Figure 1E–G). The ClusPro docking server yielded 69 top-scoring solutions based on the balanced, electrostatic-favored, hydrophobic-favored and van der Waals plus electrostatic-favored coefficients (53,54). The 4mer RNA fits nicely into the protein cavity (Figure 1G) and the negatively charged RNA backbone is aligned towards the positively charged protein surface of the cleft. Most strikingly, in the backbone is aligned towards the positively charged cavity (Figure 1G) and the negatively charged RNA cients (53,54). The 4mer RNA fits nicely into the protein and van der Waals plus electrostatic-favored coefficients (53,54). The 4mer RNA fits nicely into the protein cavity (Figure 1G) and the negatively charged RNA backbone is aligned towards the positively charged protein surface of the cleft. Most strikingly, in the top-scoring model (Figure 1E–G) the free PO₄ group of RNA interacts with the Arg59, Lys61 (Figure 1E–F). The docking results suggest a probable RNA binding site in the YbeY protein. We further subjected the YbeY structure to analysis of surface binding properties by Hotpatch webserver (38). HotPatch finds unusual patches on the surface of proteins, and statistically computes how unusual they are (patch rareness), and how likely each patch is to be of functional importance [functional confidence (FC)]. The statistical analysis is done by comparing protein’s surface against the surfaces of a large set of proteins whose functional sites are known. We analyzed the YbeY structure to find surface patches with putative DNA/RNA binding properties. DNA/RNA-binding refers to proteins that interact with oligonucleotides either catalytically or non-catalytically. Interestingly, Hotpatch identified three patches with DNA/RNA binding properties and they fall in the same protein cavity where we have docked the RNA 4mer (Supplementary Figure SIC and D). A fourth functionally important patch, corresponding to a His-triad (H114, H118, H124) and possibly coordinating a metal ion, was identified (Supplementary Figure SID). This may provide a hydrolytic function to YbeY. These observations raised the intriguing possibility that SMc01113/YbeY may be involved in the process of sRNA-mediated interactions, as are the AGO proteins.

The S. meliloti smc01113 mutant shows symbiotic and physiological phenotypes highly similar to those of hfq mutant

Parallels between the prokaryotic and eukaryotic sRNA machinery have been noted (13), with the RISC complex serving as the protein scaffold for pairing of the sRNA with its target in eukaryotes, while the Hfq protein has been proposed to play a conceptually analogous role in bacteria. Members of biological pathways deregulated when the RNA chaperone, Hfq, is mutated, are most likely targets of sRNA-machinery (15). We reasoned that if SMc01113 is involved in sRNA-mediated interactions, physiological and molecular loss-of-function phenotypes of smc01113/ybeY should be comparable to those of other established components of sRNA-pathway such as hfq. To begin with exploring the hypothesis that SMc01113 might participate in the processes related to sRNA-mediated interactions, we compared the smc01113 mutant phenotypes to that of an hfq mutant. Loss of Hfq (19) as well as SMc01113 (24) makes S. meliloti similarly symbiotically deficient, resulting in high fitness costs for both the interacting partners (Supplementary Figure S2). Swarming is an important multicellular phenomenon characterized by the coordinated and rapid movement of bacteria across semisolid surfaces, which may help the bacteria to adapt to extra- and intra- cellular niches (55). Both, hfq and smc01113 mutants were equally defective in their ability to swarm (Figure 2A, Supplementary Figure S3).

We evaluated the susceptibility of the hfq and smc01113 mutants to several stresses (Supplementary Figure S3) and observed that they displayed strikingly similar phenotypes (Figure 2). Escherichia coli YbeY has been implicated with adaptation to heat-shock (29); analogously, the loss of hfq or smc01113 makes S. meliloti similarly susceptible to heat shock (Figure 2). Further, the smc01113 and hfq mutants displayed similar sensitivities to agents like SDS and ethanol that may change the surface potential of the membranes and affect cellular osmoregulation (Figure 2). Resistance to oxidative stress, resulting in generation of reactive-oxygen-species is critical to S. meliloti (56). We evaluated the sensitivity of smc01113 and hfq mutants to oxidative stress (Paraquat): both were similarly sensitive (Figure 2). We have previously reported that both the smc01113 mutant (24) and the hfq mutant (20) are sensitive to hydrogen peroxide. The smc01113 and hfq mutants were similarly sensitive to a DNA damaging agent, methyl methanesulfonate (MMS) (Figure 2). Additionally, we tested the sensitivity of the two mutants to the β-lactam antibiotic Cefotaxime, which affects bacterial cell wall synthesis, and to elevated salt concentration. Both the mutants were similarly susceptible to Cefotaxime treatment, but not to NaCl (Figure 2). Taken together, these observations indicate it is plausible that Hfq and SMc01113 play roles that are physiologically related.

SMc01113 and Hfq regulate similar molecular targets

To test this hypothesis of functionally related roles for these two proteins, we assayed mRNA accumulation patterns in the smc01113 mutant for the genes that are
Figure 2. Loss of hfq or SMc01113 makes S. meliloti similarly sensitive to environmental stresses. (A) smc01113 mutant swarms as less efficiently as the hfq mutants than the wild-type strain. Strains were plated on the swarming plates (0.3% agar) and incubated at 30°C. Results of swarming for the three strains were evaluated by measuring colony progression every day for 3 days. (B and C) show sensitivity of the smc01113 and the hfq mutants towards variety of stresses. Strains were grown with or without stress agents as described in the text and OD600 were measured, and percent residual growth (B) after 24 and 48 h (for paraquat, MMS, ethanol and NaCl) or (C) 48 h (for heat shock, SDS and cefotaxime) was calculated.
differentially regulated in the hfq mutant and observed striking overlaps (Figures 3 and 4). We recently identified proteins significantly affected in their expression in the S. meliloti hfq mutant (20). Of the 55 differentially accumulated proteins in the hfq proteome, 30 are annotated genes. Furthermore, we found that the activity of KatB and SodC was reduced in the hfq mutant compared to a wild-type (WT) strain (20). In order to understand how loss of SMc01113 affected the transcriptional accumulation of the targets that are under Hfq control, we performed qPCR analyses for 13 of the genes (namely, sodC, katB, agpA, atpD, ppiA, ehuB, cysK1, livK, dppA1, dppA2, aapJ, glnA and thiR), which are associated with diverse processes such as transport and oxidative/superoxide resistance. Most of these genes have been confirmed to show differential accumulation in hfq mutants in independent studies, in S. meliloti, E. coli, Vibrio cholerae and Salmonella (20–22) at RNA or protein levels in independent studies compared to WT Rm1021 strain, accumulation of eight of these genes was similarly down-regulated (Figure 3) and four similarly up-regulated (Figure 4) in both mutants, whereas one gene showed a differential accumulation pattern between the hfq and smc01113 mutants.

Compared to the WT Rm1021 strain, the genes involved in protection from oxidative/superoxide stresses such as katB (catalase B), sodC (superoxide dismutase C), cysK1 (Cysteine synthase A) were down-regulated in both

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**Figure 3.** Loss of hfq or SMc01113 similarly down-regulates target transcript accumulation. WT and the smc01113 and the hfq mutants were grown in the RMM till exponential (ppiA, katB and cysK1) or stationary phase (sodC, agpA, livK, ehuB and atpD). RNA was extracted and qPCR was performed. Transcript accumulation in the WT strain was fixed to 1 and relative transcript abundance to WT in the two mutants was evaluated. ‘Single’ and ‘double asterisk’ shows significant difference at $P \leq 0.05$ and $P \leq 0.01$, respectively.
the smc01113 and hfq mutants (Figure 3). Similarly, the expression levels of agpA [α-galactoside-binding protein; periplasmic binding protein gene required for the utilization of sugars (57)], atpD (F0F1 ATP synthase beta subunit), ppiA [peptidyl-prolyl cis–trans isomerase; involved in proper folding of newly synthesized proteins, resistance to antibiotics and in cell cycle (58,59)] and ehuB [Ectonine binding protein; belonging to ATP-binding cassette transporters, a central to osmoprotection under hyper-osmotic stress in S. meliloti and provides substrate specificity (60,61)] were down-regulated in both the mutants, when compared to WT Rm1021 (Figure 3).

Somewhat surprisingly, the expression of livK (which was up-regulated in the hfq proteome) was transcriptionally down-regulated in both the smc01113 and hfq mutants (Figure 3).

Transporters involved in peptide [dppA1 and dppA2; (62)], amino acids and solute transport [aapJ; (63)] were up-regulated in both of the smc01113 and hfq mutants when compared to their transcript levels in WT Rm1021 (Figure 4). Similarly, glnA (glutamine synthetase 1, a key enzyme in nitrogen assimilation) was up-regulated in both the mutants (Figure 4). However, thtR (a sulfotransferase) mRNA showed a difference in accumulation pattern between the hfq and smc01113 mutants. No difference in mRNA levels was observed between the WT and hfq mutant, whereas smc01113 mutant showed enhanced thtR mRNA levels, compared to WT and hfq mutant (Figure 4). Taken together, these observations suggest that bacterial SMc01113/YbeY proteins play regulatory roles that are highly related, but not completely identical to, those played by the RNA chaperone, Hfq.

Figure 4. Up-regulated target transcript accumulation in the S. meliloti mutated for hfq or SMc01113. WT and the smc01113 and the hfq mutants were grown in the RMM till stationary phase, RNA was extracted and qPCR was performed. Transcript accumulation in the WT strain was fixed to 1 and relative transcript abundance to WT in the two mutants was evaluated.
**In silico** identification of possible sRNAs and their binding sites in the target genes differentially regulated in *hfq* and *smc01113* mutants

Above-mentioned results prompted us to search for the sRNAs that may regulate the altered expression of the above said targets in *hfq* and *smc01113* mutants. Our hypothesis suggested that we would observe similar patterns of accumulation of sRNAs between the *hfq* and *smc01113* mutants. We, therefore, undertook a computational approach to identify possible sRNAs that might target the 13 genes tested above.

sRNA-mediated regulation of gene expression falls into two broad categories (10). First is by the *cis*-encoded anti-sense sRNAs, which have extensive complementarity (>75 bp) with their targets, and second, by the *trans*-encoded sRNAs, which have more limited complementarity. The *trans*-encoded sRNAs have 10–25 bp complementary ‘seed sequences’ to their target mRNAs in discontinuous patches, where only a core of nucleotides seems to be critical for regulation. These sRNAs regulate translation by targeting regions in the 5’ UTRs, inside and upstream of the ribosome-binding sites (RBS)/Shine–Dalgarno sequences, occluding or sequestering RBS, as well as binding within the coding sequences of the targets largely leading to their degradation (13,43,64,65).

The *trans*-encoded sRNAs are functionally analogous to eukaryotic microRNAs (miRNAs), often require Hfq for their action (11,13), may pair targets in the coding region (43) and thus could be the preferred mode of regulation of the differentially accumulated targets identified in *hfq* and *smc01113* mutants. The principle of seed pairing has been extensively and effectively used for determining sRNA–mRNA interaction in eukaryotes (39,41,44,66,67) as well as in some prokaryotes (40,42). In general, the longer the seed sequence (and thus the greater the perfect complementation), the higher the chances of an effective sRNA-target interaction.

Informed by these considerations, we performed a bioinformatic analysis to identify *S. meliloti* sRNAs (17) that could potentially bind to the genes differentially regulated in the *hfq* and *smc01113* mutants. Our analysis revealed that these genes have a large number of potential sRNA binding sites (Supplementary Table S1). Genes with multiple sRNA-binding sites of >10 nt, for specific sRNAs were identified; e.g. *dppA1* had three different complementary sites for the sRNA, *sra16* (Figure 6B). It was strikingly apparent that a single sRNA has the potential to target more than one gene, and that a single gene can be targeted by several sRNAs (Supplementary Table S1; Figure 6A). For instance, *sodC* had a 13-nt sRNA complementary site for one sRNA, 12-nt seeds for five sRNAs, and 11-nt seeds for eight sRNAs (Supplementary Table S1). The potential for single sRNA to target multiple mRNAs is also striking. For example, our analysis indicated that *sra35* potentially targets three genes, *sra03* may target six genes and *sra16* may target 10 genes (Supplementary Table S1, Figure 6A).

Similar observations of ‘one-to-many’ sRNA–mRNA interactions, common in eukaryotes [e.g. (39)], have been previously reported in some other bacterial species, and would have profound effects on cellular physiology and stress adaptation (43,65,68–70). Using a combination of such criteria of longest seed length and potential to target several genes, we selected 9 sRNAs for their expression analysis in WT, a *smc01113* mutant and an *hfq* mutant.

**Expression of sRNAs is similarly affected in *S. meliloti smc01113* and *hfq* mutants**

Using qPCR analyses, we next investigated the relationship between changes in sRNA accumulation in *smc01113* and *hfq* mutants, and the changes in the various potential mRNA targets. Of the nine sRNAs tested, the accumulation of eight sRNAs (*sras 03, 11, 16, 45, 47, 51, 63 and 65*) was similarly reduced in both the *hfq* and *smc01113* mutants compared to the WT Rm1021 (Figure 5). In contrast, *sra35* was strongly up-regulated in both *hfq* and *smc01113* mutants relative to the WT Rm1021 strain (Figure 5). The pattern of the variation of sRNAs relative to particular mRNA target indicated both, inverse correlations (up-regulation of target corresponding to down-regulation of the sRNAs and vice versa) as well as direct correlations (change of target and the targeting sRNAs in the same direction; Figure 6A).

sRNA-mediated-interactions in bacteria results in both the inhibition and the stimulation of translation (10,13,43,64,65,69). Inverse correlations of expression of sRNAs and their potential targets (red/green regions in Figure 6A) suggest inhibitory action of these sRNAs on their targets; conversely similarly-regulated sRNA-target pairs (yellow regions, Figure 6A) indicate a stimulatory action of sRNAs in these interactions. Thus, these patterns suggest the existence of complex interactions between the sRNAs and their targets that are necessary for modulating and fine tuning the expression of genes to optimum levels.

To gain further insight into the biological relevance of the putative sRNA-target interactions described above (Figure 6A, Supplementary Table S1), we performed phylogenetic comparisons of the sRNAs and sRNA-target-motifs in mRNAs found in our *in silico* analysis in multiple rhizobium species. Homologous sequences to several of these differentially regulated sRNAs were identified in three or more related *Rhizobium* species (Supplementary Figures S4 and S5), suggesting their wide organismic relevance in symbiosis. Some of these sRNA sequences have been reported in other *α*-proteobacterial species (17). Conservation in mRNA-targets with complementarity to the seed (2–7 nt) of the miRNAs in closely related species has been used as one of the main criteria of identifying miRNA-target interactions in eukaryotes (41,71–73). On the other hand, sRNAs/targets in viruses are not well conserved (74). When we performed an analysis of conservation of the complementary sites in the mRNA as well as in sRNAs in the three closely related rhizobial bacteria, we found that the motifs of sRNAs as well as their complementary sequences in the targets were fairly well conserved (Figure 6C). SNPs in both sRNAs as well as mRNAs in other rhizobial species were seen (Figure 6C). Single-base substitutions from T to G/C (and vice versa) were seen in few cases.
Such mismatches in these sRNA–mRNA motifs may be analogous to eukaryotic miRNA target sites with seeds having G:U base pairs or single nucleotide bulges. These results suggest a large degree of conservation of these sites among closely related rhizobia and are thus interesting targets for future study of sRNA–mRNA interaction in rhizobia species.

Over-expression of sRNAs

In order to evaluate our predictions of sRNA-targets, we over-expressed sra16 and sra35 (Figure 7A and B) and evaluated the effects on the targets we initially predicted (Figure 6). Sra16 was down-regulated whereas sra35 was up-regulated in hfq- and smc01113- mutants, compared to WT Rm1021 (Figure 5). We tested the accumulation of 4 of the 10 predicted targets of sra16 and all the three predicted targets of sra35 (Figure 6). Putative target sites for sra16 and sra35 did not overlap with the ribosome binding sites. Three sra16-targets (glnA, dppA2 and katB) were down-regulated in the sra16-over-producing strain (SRA16; Figure 7C, E and G) compared to WT, whereas the fourth (dppA1) did not show a difference between the two strains (data not shown). Over-expression of sra35 (SRA35; Figure 7) resulted in up-regulation of glnA (Figure 7D), whereas two other targets (ppiA and atpD) showed a trend of down-regulation (Figure 7F).
and H). These results show that our predictions were identifying biologically relevant sRNA–mRNA interactions and that individual sRNAs have multiple targets. Sras16 and 35 differently regulated the accumulation of a common target, *glnA*: whereas *glnA* was down-regulated in bacteria over-expressing sra16, over-expression of sra35 up-regulated *glnA* expression, providing evidence that multiple sRNAs may regulate a given target differently, and that this regulation of expression of genes by sRNAs is complex. Thus, we were able to largely confirm the results of our predictions of sRNA-targets as well as target-sRNA relationships.

**Potential interactions of SMc01113/YbeY with other components of sRNA machinery/RNA-degradation**

Since our data suggested that the function of SMc01113/YbeY is related to Hfq, we next asked whether SMc01113/YbeY physically interacts with the Hfq protein or with

**Figure 6.** sRNA-target relation in *S. meliloti* and conservation of motifs in sRNAs and their possible targets in other related rhizobium species. (A) Correlation between the direction of accumulation of the sRNAs and their target in the *S. meliloti* strains mutated for *hfq* and SMc01113, as derived from Figures 3–5. Red and green shows inverse correlation, and yellow shows that the sRNAs and their predicted targets show accumulation in the similar direction. (B) Schematic representation of the three sra16 binding sites in the dppA1 gene. Sra16 was down-regulated whereas dppA1 was up-regulated in the *hfq* and *smc01113* mutants. (C) Conservation of the target-sRNA motif sequences in the three related rhizobium species. Weblogos were generated as described in the text.
other members of the sRNA machinery and RNA-degradation. The sRNA-machinery and the components of RNA-degradosome are well characterized in *E. coli* (75). RNase E (*rne*) has been implicated as an integral component of the sRNA-machinery and degrades targets of sRNAs (10,13,76,77). RNase III (*rnc*), whose eukaryotic analogs are the DICER/dicer-like (DCL) proteins, participates in sRNA-mediated interaction, along with the RNase E, by performing initial cleavage (78,79). The PIWI domain of the AGO proteins has the RNase H- (*rnhB*) type of catalytic domain (32). The two other RNases comprising important component of the RNA-degradosome during stress adaptation are RNase R and PNPase; both of these have overlapping functions (80). Informed by the fact that sRNAs can determine which proteins bind to Hfq (81), we asked if we could detect an interaction of YbeY with Hfq and other RNases in the absence of bacterial sRNAs. We therefore conducted a directed yeast two-hybrid screen to determine if YbeY interacts physically with any of these components. However no significant interaction with any of these was observed regardless of whether YbeY was fused with either the GAL4 activation or binding domains (Figure 8). This suggests that either (i) there is no physical interaction between YbeY/SMc01113 with Hfq and other RNases, or (ii) the interaction of YbeY/SMc01113 with these components is highly transient and/or may require additional facilitator proteins. Transient interaction with YbeY, mediated by other proteins would be consistent with the fact that there is a large difference in the number of Hfq protein molecules per cell [approximately 30–60000; (82)] relative to YbeY/SMc01113 (approximately 1000; Walker, G.C., unpublished data) molecules in a cell. The involvement of other co-factors in modulating protein–protein interactions during sRNA regulation is conceivable. For example, it has recently became apparent that the RraA regulatory protein modulates the helicase and RNA-binding activities of the RNA-degradosome (83). Furthermore, the association of the AGO-siRNA complex with AGO/binding proteins e.g. Giw1p for correct localization has been known (84). We also generated a *S. meliloti* double mutant of *hfq* and *smc01113*, which showed highly reduced growth and enhanced doubling time over both the single mutants (Figure 8). As discussed more fully below, these

![Figure 7](https://example.com/image7.png)

**Figure 7.** Over-expression of sRNAs affect accumulation of their predicted targets. (A and B) show the levels of over-expression of sra16 and 35, respectively. (C, E and G) show down-regulation of sra16 targets. (D, F and H) show the changes in putative sra35 targets. Sras 16 and 35 differently regulate the accumulation of common target, *glnA* (C and D). sRNA-targeting regions were predicted in the coding-region.

![Figure 8](https://example.com/image8.png)

**Figure 8.** (A) Directed yeast two hybrid of YbeY. No interaction was detected. (B) Growth of the WT, *smc01113* and *hfq* single, and *smc01113 hfq* double mutant in LB-MC. Insert shows doubling time of WT and each of the three mutants in LB-MC.
observations are consistent with SMc01113/YbeY and Hfq playing functionally related roles in sRNA regulation but with YbeY playing certain Hfq-independent roles (81) as well.

DISCUSSION

Taken together, our results suggest that an important, previously unrecognized function for the SMc01113/YbeY proteins, members of the UPF0054 family which is present in most bacterial species, is to participate in sRNA regulation. The SMc01113/YbeY proteins show intriguing structural similarities to the MID domain of the AGO proteins, which has a capacity to capture the 5'-phosphate of sRNA seed regions. Loss of SMc01113 results into perturbed sRNA accumulation, deregulated mRNA levels and consequently, physiological and symbiotic defects. Furthermore, there are striking similarities in the symbiotic, physiological and molecular profiles of the smc01113 and hfg mutants. Taken together, our results suggest that SMc01113 homologs may form a component of the sRNA-pathway and facilitate effective outcomes of the sRNA–mRNA interaction: in particular, we suggest that SMc01113/YbeY proteins could help in recognizing the prokaryotic sRNAs during their Hfq mediated interaction with the target mRNAs. In addition, this study also gives insights into the fundamentals of sRNA-mediated regulatory basis of stress adaptation in S. meliloti by identifying multiple sRNA-target interactions. Although sRNAs have been identified in a few genomic studies of S. meliloti (16–18), little is known about their functions, targets and regulation. Understanding regulatory mechanisms in S. meliloti are of utmost importance from agricultural, economical, environmental perspectives as well as for their implications for other symbiotic and pathogenic interactions.

An estimated 82 sRNAs were originally predicted in S. meliloti (16–18), whereas the most recent genome-wide survey puts the number at 1125 [which included trans-encoded sRNAs, antisense RNAs, putative mRNA leader and sense transcripts, which may be degradation products of mRNAs; (85)]. Genome-wide searches in E. coli estimate that the number of sRNAs is ~2% of the number of protein coding genes (77,86). In bacteria, a major class of these sRNAs, as in eukaryotes, act by base pairing with the target mRNAs. In response to environmental stress, the expression of several sRNAs may be stabilized by the RNA chaperone Hfq, which facilitates the trans-annaling of the sRNAs to their target mRNA in an anti-sense manner (70,87). This is followed by the recognition of the sRNA-target complex by a multi-membrane protein complex, resulting in mRNA degradation (11,75,88) or mRNA stabilization, facilitated ribosome assembly and enhanced translation (10,13,64,69).

However, these processes are still poorly understood, as little is known regarding the assembly of the sRNA–mRNA complex, except that Hfq is involved (11,13,81).

Conceptually, striking parallels exist between the prokaryotic and the eukaryotic sRNA-pathways in terms of synthesis of sRNAs, the presentation of sRNA and mRNAs onto the protein scaffold and the eventual outcomes of sRNA–mRNA interactions (13). Both, the sRNAs and the mRNAs bind to the Hfq protein, but at different sites (87). However, mechanistically, little is known about how the prokaryotic sRNAs are recognized and loaded on to the Hfq protein scaffold during their interaction with the target mRNAs. This is better understood in eukaryotes, where the si/miRNAs are loaded into AGOs, and then guided and assembled on to the targets by the RISC. The recognition of target is governed by the ‘seed sequence’ of the guide RNA strand associated with the MID domain of the AGO proteins that generates ‘an enhanced affinity anchor site promoting fidelity in target recognition’ (89). This stabilizes and guides the assembly of the active RISC (89). The structure of the MID domain associated with the dsRNA and ssDNA reveals that the MID-domain recognizes the 5'-phosphate specifically and anchors sRNAs on to the AGO/RISC (32,52). The 5'-end of the guide strand (of sRNA) could bind to the AGO in a stacked helical confirmation, both in the absence and presence of the partner mRNA strand, facilitating easy access for recognition of the mRNA target (32,52).

Moreover, the MID domain has been implicated in protein–protein interactions: interactors such as Tas3 in S. pombe form a so-called ‘Ago-hook’ that binds the MID domain (90). Whether the binding of the protein interactors and sRNAs occur simultaneously, or, is mutually exclusive remains unclear. Different AGO family members have diverse functions in a given organism (91). The MID domains are central to the functional distinctions that define the various AGO protein family members (92). Proteins that perform analogous functions in the rhizobial or other bacterial sRNA-pathways have remained unknown. Our structural analysis studies suggest that SMc01113/YbeY and their bacterial homologs might perform analogous functions of anchoring the seed sRNAs, which can help the sRNA access the mRNA. SMc01113/YbeY proteins have >50% sequence similarity to MID domains, and furthermore have similar structural folds to those of both the prokaryotic and eukaryotic MID domains [Figure 1; (52)]. Moreover, our structural analysis suggests that they similarly have a capacity to bind to the sRNA seeds, and regulate accumulation of sRNAs and target mRNAs (Figures 1, 3–5). Our analyses suggest that, similarly to the corresponding residues of the MID domain, R59 and K61 may undergo key interactions with a phosphate at 5'-end of an RNA sequence. The structure of YbeY suggests that it could accommodate an internal phosphate at the 5'-end of the seed sequence instead of requiring that the 5'-phosphate be at the 5'-terminus as well as a 5'-phosphate at the terminus of a sRNA. This suggests that SMc01113/YbeY homologs have a function that is analogous to the functions of AGO MID domain, facilitating the recognition of the sRNA, thereby guiding it to the target mRNA/Hfq and the assembly of active complex.

YbeY differs from the AGO MID domain in one interesting aspect: a metal coordinated ion, postulated to be zinc (27), is present on the other side of the cleft opposite R59 and K61 (Supplementary Figure S1D), along with a
highly conserved His-triad (H114, H118 and H124; Supplementary Figure S1). These could act catalytically in a hydrolytic reaction. In contrast to speculations of YbeY being a peptidase or a protease (28), our docking of the RNA and analysis of YbeY structure suggests that it is plausible that a phosphodiester bond could be located close enough to the presumed coordinated water for hydrolysis to occur. Thus, it is possible that YbeY could contribute catalytically, like an RNase, to RNA cleavage events after binding to a guide RNA. We hypothesize a model where a sRNA substrate would bind to the positively charged surface of YbeY with help of a phosphate that plausibly interacts with K59, L61, L63 and other residues on the positively charged surface (Figure 1, Supplementary Figure S1) followed by a metal-coordinated hydrolysis of phosphodiester bond by the His-triad (H114, H118 and H124; Supplementary Figure S1). Such metal-coordinated hydrolysis of phosphodiester bond by the His-triad (Supplementary Figure S1) followed by a metal-coordinated hydrolysis of phosphodiester bond by the His-triad (Supplementary Figure S1). Such metal-dependent RNA-binding and RNase activity is established for nucleases for instance in RNA-dependent RNA-binding and RNase activity is established.

Our proposal that SMc01113/YbeY proteins participate in the sRNA-pathway strengthens the parallels between the AGO/RISC functions and the bacterial sRNA–mRNA assembly process that have been previously noted (13,81). Furthermore, as in the eukaryotes, our computational analysis suggests a ‘one-to-many’ sRNA–target relationship in S. mellioti. Thus, this study gives the first line of evidence that supports more than a conceptual similarity (13) between the prokaryotic and eukaryotic RNAi machinery. The involvement of other analogous facilitator proteins (currently unknown), however, cannot be ruled out. The next step will be to work out the biochemical properties of the SMc01113/YbeY protein so that we can better understand the mechanism of such action.

For most of the phenotypes we tested, along with the underlying molecular targets and sRNAs, the smc01113 and hfq mutants displayed close similarity. Nevertheless, certain observations indicate that SMc01113 and Hfq also act independently. The difference in thtR levels in smc01113 and hfq mutants is consistent with this possibility, as is the reduced growth of smc01113 hfq double mutant compared to the single mutants in a rich LB-MC medium. Recent work from Rasouly et al. (28) and from our lab in E. coli rRNAs (51) suggests an additional function for YbeY as the deletion of ybeY from E. coli results in defects in ribosome assembly and activity as well as in attenuation of processing of ends of rRNAs (51). Interestingly, mutating the R59 residue, which our docking studies suggest may participate in coordinating the PO4 in a sRNA seed (Figure IE–G) on a multicopy plasmid did not affect rRNA processing but did affect the growth rate. The possibility that a member of sRNA-pathway could potentially be involved in maturation of rRNA-ends is not at all surprising. For example, RNase E, central to sRNA pathway (11,13,77,81), is also necessary for maturation of ends of rRNA (96,97). In another example, PNPase is indispensable for 3'-end maturation of 23S rRNA, as a component for mRNA degradation is sRNAs’ context, and is also a key regulator of sRNAs, (81,98). Even Hfq interacts with at least 24 ribosomal proteins (81), and also with RNase E and PNPase (81); in E. coli, YbeY shows genetic interactions with RNase E and PNPase (51). This overlap in function and sharing of components between sRNA-pathway and rRNA maturation and ribosome assembly may form a more general strategy of efficient post-transcriptional gene regulation by coupling transcription and translation (81). This warrants further investigation.

Our observations are also important because they verify the differential expression of genes in S. mellioti hfq proteome at the transcriptional level. Along with that, we have also identified several sRNAs that are Hfq-dependent. Mapping of these differentially accumulated sRNAs/mRNAs, the conservation of sRNA–mRNA complementary motifs in rhizobial species closely related to S. mellioti (Figure 6), and change in accumulation of predicted targets in bacteria over-expressing sRNAs (Figure 7) indicate functions for these sRNAs, a possible mode of regulation of these genes and a mechanistic relationship between the sRNAs and the targets. We have here further generated additional support for biological relevance of some S. mellioti sRNA–mRNA relationships (Figure 7) and identified other such candidate relationships for further investigation.

Our results also offer insights into the regulatory basis of stress adaptation in S. mellioti. Pathways such as transport system, oxidative stress resistance and nitrogen metabolism are coordinated and controlled by the sRNA-machinery. Along with identifying a new component, this study places the sRNA-pathway squarely in the centre of stress adaptation and regulation of plasticity of S. mellioti as the bacteria undergoes ecosystem shifts during its course of survival.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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