

## A survey of sRNA families in $\alpha$ -proteobacteria

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We have performed a computational comparative analysis of six small non-coding RNA (sRNA) families in  $\alpha$ -proteobacteria. Members of these families were first identified in the intergenic regions of the nitrogen-fixing endosymbiont *S. meliloti* by a combined bioinformatics screen followed by experimental verification. Consensus secondary structures inferred from covariance models for each sRNA family evidenced in some cases conserved motifs putatively relevant to the function of *trans*-encoded base-pairing sRNAs i.e., Hfq-binding signatures and exposed anti Shine-Dalgarno sequences. Two particular family models, namely *ar15* and *ar35*, shared own sub-structural modules with the Rfam model *suhB* (RF00519) and the uncharacterized sRNA family *ar35b*, respectively. A third sRNA family, termed *ar45*, has homology to the *cis*-acting regulatory element *speF* (RF00518). However, new experimental data further confirmed that the *S. meliloti ar45* representative is an Hfq-binding sRNA processed from or expressed independently of *speF*, thus refining the Rfam *speF* model annotation. All the six families have members in phylogenetically related plant-interacting bacteria and animal pathogens of the order of the Rhizobiales, some occurring with high levels of paralogy in individual genomes. In silico and experimental evidences predict differential regulation of paralogous sRNAs in *S. meliloti* 1021. The distribution patterns of these sRNA families suggest major contributions of vertical inheritance and extensive ancestral duplication events to the evolution of sRNAs in plant-interacting bacteria.

Post-genomic research has rendered bacterial small non-coding RNAs (sRNAs) as major players in the post-transcriptional regulation of gene expression underlying a wide range of important cellular processes, e.g., general responses to abiotic stimuli, cell division, quorum sensing or virulence.<sup>1</sup> However, very little is known about the role of riboregulation in the control of symbiotic and pathogenic plant-microbe interactions.

The  $\alpha$ -subdivision of the proteobacteria includes Gram-negative microorganisms with diverse life styles, frequently involving long-term mutualistic or pathogenic interactions with higher eukaryotes.<sup>2</sup> *Sinorhizobium meliloti* is an environmentally and agronomically relevant  $\alpha$ -proteobacterium belonging to the order of the Rhizobiales. It is well recognized as a genetically tractable model microorganism for the investigation and exploitation of the nitrogen-fixing endosymbiosis with legume plants. The outcome of these interactions is the formation in the cognate legume (i.e., *Medicago* species for *S. meliloti*) of the so-called root nodules which finally host invading bacteria in their differentiated nitrogen-fixing competent form of bacteroids.<sup>3</sup> The *S. meliloti* genome has a multipartite architecture consisting of a single chromosome (3.65 Mbp) and two large plasmids termed pSymA (1.35 Mbp) and pSymB (1.68 Mbp). Megaplasmid pSymA harbors the clusters of genes specifying symbiotic functions, among others, but is dispensable for bacterial free-living growth whereas pSymB exhibits chromosome-like features e.g., it accommodates the essential tRNA-Arg encoding gene.<sup>4</sup> This composite arrangement is common to the

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**Abbreviations:** sRNA, bacterial small non-coding RNA; IGR, intergenic region; RACE, rapid amplification of cDNA ends; Smr, *S. meliloti* sRNA; TAP, tobacco acid pyrophosphatase; TSS, transcription start site; CM, covariance model; RBS, ribosome binding site; aSD, anti Shine-Dalgarno; 5'-UTR, 5' untranslated region; GABA,  $\gamma$ -amino butyric acid

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genomes of many bacterial species of the order of the Rhizobiales in which second chromosomes have been proposed to evolve from an early-acquired ancestral plasmid.<sup>5</sup> Similarly to *S. meliloti*, many  $\alpha$ -proteobacteria interacting with plants usually host a variable number of accessory extrachromosomal replicons besides the ancestral set of primary chromosome and megaplasmid. These non-essential plasmids most likely have a mosaic origin and contribute to the adaptive flexibility demanded by the transition of bacteria from a free-living to an intracellular state. At the regulatory level, these adaptations require the coordinated expression of complex gene networks in which sRNAs are also expected to participate. Several recent computational comparative genomics and deep-sequencing approaches have identified more than thousand non-coding RNA elements in the *S. meliloti* genome.<sup>6-9</sup> Nearly two hundred of these molecules have been cataloged as putative *trans*-encoded sRNAs.<sup>9</sup> This is an abundant class of bacterial riboregulators, which mostly target mRNAs via discontinuous nucleotide stretches of sequence complementarity to control the translation and/or stability of the message, many of them in an Hfq-dependent manner.<sup>1,10</sup>

Rhizobial RNomics has been pioneered by a genome-wide comparative genomics screen conducted in our laboratory, which, in combination with the experimental verification of predictions, identified eight sRNA genes in the intergenic regions (IGRs) of the reference strain *S. meliloti* 1021.<sup>6</sup> Northern hybridization experiments and RACE mapping revealed that these sRNAs are differentially expressed from independent transcription units in free-living and endosymbiotic bacteria, thus supporting their putative role as riboregulators in *S. meliloti*. In this work we have combined new experimental data with an extensive in silico structural comparative analysis to further characterize these sRNAs and assess their conservation across  $\alpha$ -proteobacteria.

## Results

### Generation of Smr sRNA family models.

The starting point of this study was the set of eight non-coding transcripts identified

previously in our laboratory on the basis of structure conservation and experimental verification. These sRNAs were initially termed Smr7C, Smr9C, Smr14C, Smr15C, Smr16C Smr22C, Smr35B and Smr45C for *S. meliloti* RNA, where the suffix indicates their respective positions in the output table of candidates along with the genomic location of each *locus* on pSymA (A), pSymB (B) or chromosome (C) (Table 1). TAP-based 5'-RACE experiments mapped the transcription start sites (TSS) of each sRNA to defined positions in the *S. meliloti* genome. Their 3'-ends were assumed to map to the last residue of the consecutive stretches of Us of Rho-independent terminators predicted for most of the transcripts, except for Smr22C and Smr45C which 3'-ends have been inferred from published experimental data<sup>6,9</sup> (Table 1). Recent RNA-Seq based characterization of the small RNA fraction (50–350 nt) of the closely related strain *S. meliloti* 2011 mapped the full-length Smr transcripts in the *S. meliloti* 1021 genome to essentially the same positions reported earlier.<sup>9</sup>

The nucleotide sequences of the full-length Smr transcripts were first used to query the Rfam database v. 10.0 ([www.sanger.ac.uk/Software/Rfam](http://www.sanger.ac.uk/Software/Rfam)).<sup>11</sup> This search revealed full homology of Smr22C to the well characterized 6S RNA family and therefore, this sRNA was not further considered in this study. Of the remaining 7 RNAs, Smr15C/16C and Smr45C exhibited partial structural homology to the *suhB* (RF00519) and *speF* (RF00518) RNA families respectively, whereas the remaining query transcripts did not match

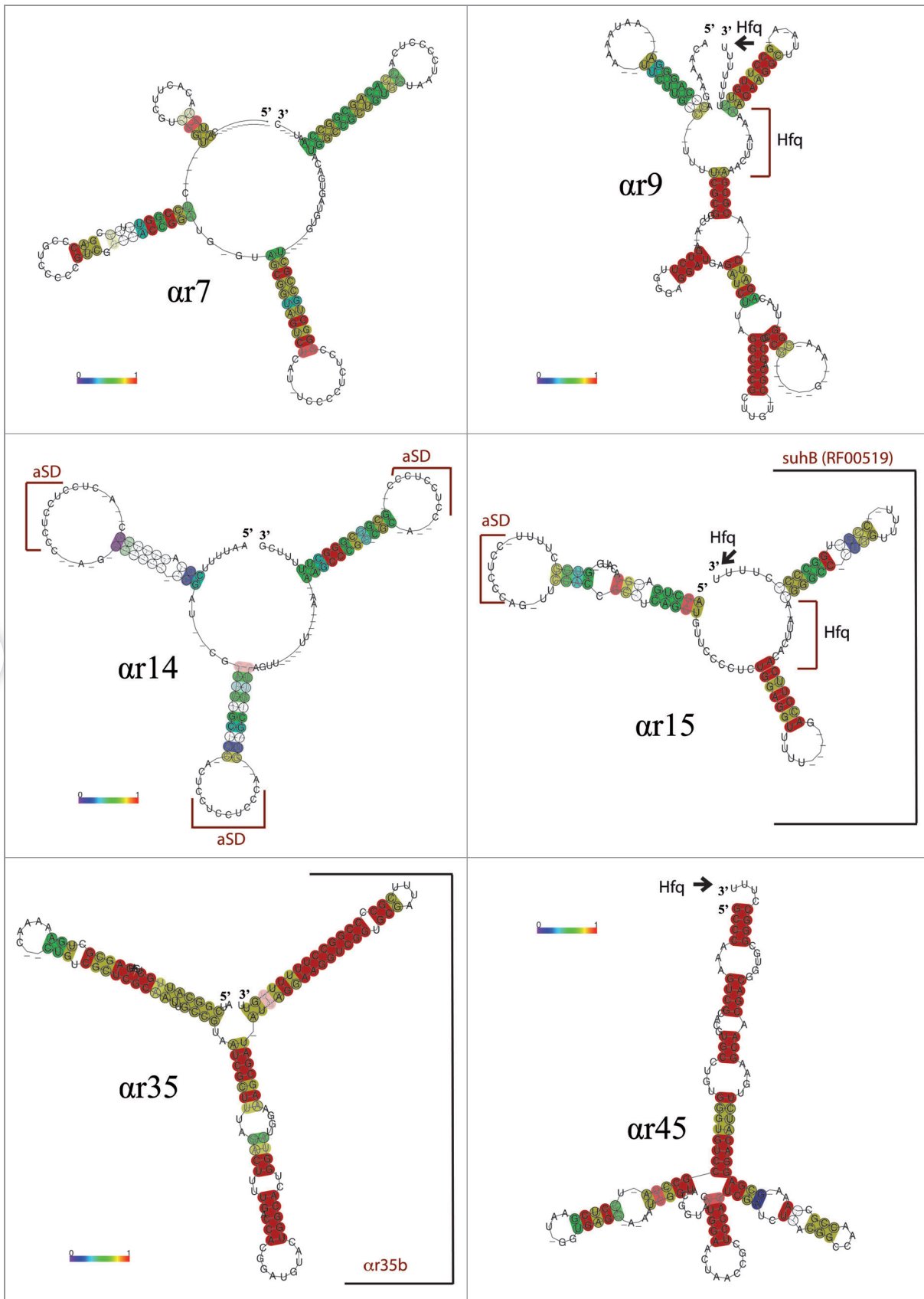
any Rfam entry. These seven sRNA sequences, likely representing previously unknown bacterial sRNA families, were next BLASTed with default parameters against all available bacterial genomes (1,615 sequences at 20th April 2011; [www.ncbi.nlm.nih.gov](http://www.ncbi.nlm.nih.gov)). The genomic regions exhibiting significant degree of homology to the query sequences (78–89% similarity) were collected to generate initial alignments for each RNA that were manually curated to construct an Infernal Model (covariance model; CM) for each sRNA. As expected from their primary nucleotide sequence similarity, this analysis merged the tandemly-encoded Smr15C and Smr16C transcripts into the same RNA family and they were renamed accordingly as Smr15C1 and Smr15C2, respectively. The six RNA families resulting from this study have homologies limited to species of the order of the Rhizobiales within the  $\alpha$ -subgroup of proteobacteria. Consistent with the naming scheme of the query sRNAs, their family models have been referred to as  $\alpha$ rn for  $\alpha$ -proteobacteria RNA, where the suffix identifies the family according to the query sequence. Stockholm formatted alignments for each family is provided at [en.wikipedia.org/wiki/Small\\_non\\_coding\\_RNAs\\_in\\_the\\_endosymbiotic\\_diazotroph\\_%CE%B1-proteobacterium\\_Sinorhizobium\\_meliloti](http://en.wikipedia.org/wiki/Small_non_coding_RNAs_in_the_endosymbiotic_diazotroph_%CE%B1-proteobacterium_Sinorhizobium_meliloti).

**Structural features of the  $\alpha$ r sRNA families.** The inferred consensus secondary structures for each  $\alpha$ r family model are shown in Figure 1.<sup>12</sup> All six RNA families presented the typical sRNA arrangement in sub-structural domains with three to

**Table 1.** Query *S. meliloti* sRNA sequences

Name	Alternative names <sup>a</sup>	5'-end <sup>b,c</sup>	3'-end <sup>b</sup>	Length (nt)
Smr7C	Sra03/Sm13/SmelC023	201,679	201,828	150
Smr9C	Sra32/Sm10/SmelC289	1398,425	1398,277	149
Smr14C	Sm7/SmelC397	1,667,613	1,667,491	123
Smr15C	Sra41/Sm3/SmelC411	1,698,731	1,698,617	115
Smr16C	Sra41/Sm3'/SmelC412	1,698,937	1,698,817	121
Smr35B	SmB6/SmelC053	577,730	577,868	139
Smr45C	SmelC706	3,105,445	3,105,298 <sup>d</sup>	148
Smr22C	Sra56/Sm1/SmelC667/65	2,972,251	2,972,091 <sup>c</sup>	161

<sup>a</sup>Alternative reported names for the Smr transcripts;<sup>7-9</sup> <sup>b</sup>Coordinates according to the *S. meliloti* 1021 genome database at <http://iant.toulouse.inra.fr/bacteria/annotation/cgi/thime.cgi> or [www.rhizogate.de](http://www.rhizogate.de); <sup>c</sup>RACE-based mapping\_ENREF\_32;<sup>6</sup> <sup>d</sup>Deep-sequencing data<sup>9</sup>



**Figure 1.** Consensus secondary structures of the  $\alpha$  sRNA families. Color code for base pairs is that of the Vienna RNA web suite.<sup>12</sup> Putative Hfq-binding sites (Hfq) i.e., free 3'-hydroxyl end of an oligo-U stretch or internal single-stranded A/U-rich regions (Hfq) are marked with an arrowhead and a bracket, respectively. Ultraconserved anti Shine-Dalgarno sequences are indicated as (aSD), and identified sub-structural modules, *suhB* (RF00519) and  $\alpha$ 35b, are marked with brackets.

five main hairpin loops generally interrupted by internal stem-loops and/or single stranded sequence stretches. These structures are supported by a variable degree of nucleotide covariance that was particularly high in the three stem-loops of the  $\alpha r7$ ,  $\alpha r14$  and  $\alpha r15$  family members and the 5' domains of  $\alpha r9$  and  $\alpha r35$  families. In most cases, the 3' domain consists of a GC rich hairpin followed by tails of uridine residues, thus matching the main structural feature of the Rho-independent terminators of transcription. The exception was  $\alpha r45$  which last hairpin is supported by a strong conservation of the primary nucleotide sequences but does not resemble a bona fide Rho-independent terminator.

A remarkable and complex structural situation was found in the  $\alpha r15$  and  $\alpha r35$  families. Members of the  $\alpha r15$  family showed partial homology to the Rfam model RF00519 known as *suhB*. In all cases this structural homology to the full-length *suhB* transcripts was restricted to the second hairpin and the Rho-independent terminator. *SuhB*-like genes have been computationally predicted to occur in multiple copies in a wide range of  $\alpha$ -proteobacterial genomes and some meta-genomes.<sup>13</sup>

Similarly,  $\alpha r35$  sRNAs have three well-defined hairpin loops. The second and third structural motifs are maintained by extensive primary nucleotide sequence conservation and define a sequence stretch with wider occurrence in the genomes of the Rhizobiales (40 sequences) outside the full-length  $\alpha r35$  sRNAs (not shown). Therefore, *suhB* and this newly identified  $\alpha r35$  sub-structural domain ( $\alpha r35b$ ) likely represent widely distributed variants of the  $\alpha r15$  and  $\alpha r35$  sRNA families with a highly variable or even missing 5' stem loops characteristic of the later transcripts.

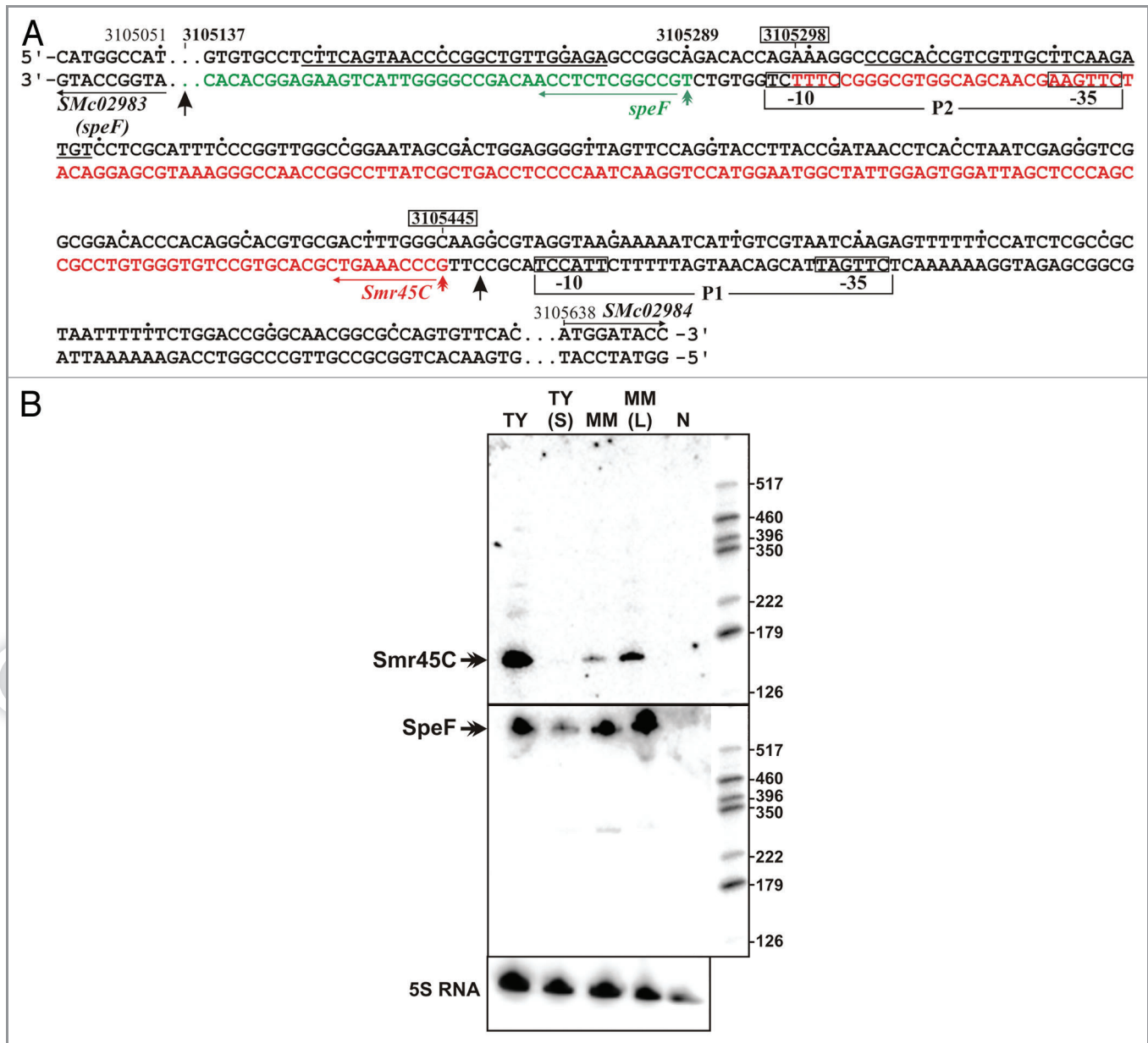
The  $\alpha r$  sRNA families mostly include putative trans-encoded transcripts, which are expected to influence translation of target mRNAs through short base-pairing interactions that usually occlude the ribosome-binding site (RBS). Interestingly, the loop anti Shine-Dalgarno sequence "CUCCUCCC" was found to be conserved in all the three hairpin loops of the  $\alpha r14$  family members as well as in the 5' hairpin loop of  $\alpha r15$  sRNAs.

Nonetheless, paired nucleotide stretches could also bind mRNA sequences if they are released and exposed to the target with the aid of proteins. The RNA chaperone Hfq has been shown to fulfill this function in most of the sRNA-mRNA target interactions documented to date. Internal single-stranded A/U-rich regions as well as a free 3'-hydroxyl end of an oligo-U stretch (e.g., of Rho-independent terminators) have been proposed as preferential sRNA interaction sites for Hfq.<sup>14-16</sup> Both Hfq-binding signatures coexist in the  $\alpha r9$  and  $\alpha r15$  sRNAs, whereas exposed 3'-end poly-U tails of different lengths are also evident in  $\alpha r45$  transcripts. However, the terminal uridines of the Rho-independent terminators predicted for  $\alpha r7$ ,  $\alpha r14$  and  $\alpha r35$  family members are mostly base-paired to upstream sequences and hence could not be easily available for Hfq binding. In good correlation with these observations, the *S. meliloti* Smr9C ( $\alpha r9$ ), Smr15C1, Smr15C2 (both  $\alpha r15$ ) and Smr45C ( $\alpha r45$ ) sRNAs have been detected in the sub-population of transcripts co-immunoprecipitated with a chromosomally-encoded epitope-tagged Hfq protein in lysates of free-living bacteria.<sup>17</sup>

**Smr45C and speF are likely expressed as independent RNA elements in *S. meliloti*.** The  $\alpha r45$  RNA family partially matched the Rfam model *speF* (FR00518), a family of *cis*-acting RNA elements likely involved in the regulation of polyamine biosynthesis that have been identified in several  $\alpha$ -proteobacterial species.<sup>13</sup> Consistent with its proposed role, *speF* RNAs are mostly leader sequences of orthologs of ornithine decarboxylase-encoding genes.<sup>13</sup> The *S. meliloti* *speF* structural homolog has been predicted to map between positions 3,105,448 and 3,105,137 in the chromosome of the reference strain 1021, upstream the *SMc02983* gene which encodes a putative ornithine/arginine decarboxylase (Fig. 2) ([rfam.sanger.ac.uk/genome/266834#tabview=tab1](http://rfam.sanger.ac.uk/genome/266834#tabview=tab1)).<sup>13</sup> Therefore, the 148 nt-long sequence of Smr45C, deduced from experimental mapping,<sup>6,9</sup> would entirely match the 5' region of *speF* (Fig. 2A). To solve this apparent inconsistency in the annotation of *S. meliloti* *speF*, the transcriptional output of this genomic region was further

investigated. A closer inspection of the *SMc02983/SMc02984* IGR identified two nucleotide sequence stretches that met the consensus CTTGAC-N<sub>17</sub>-CTATAT of  $\sigma^{70}$ -dependent promoters in *S. meliloti* and other  $\alpha$ -proteobacteria.<sup>18</sup> One of these transcription signatures (P1) had been previously identified as the putative promoter of Smr45C and is located immediately upstream the TSS determined for this sRNA, whereas the second one (P2) overlaps the 3' region of the Smr45C coding sequence (Fig. 2A). Transcription initiation from the P2 promoter is predicted to occur at the T residue at 3,105,289 nt position in the *S. meliloti* genome (Fig. 2A). Confirming previously reported data, a probe complementary to the 3' region of Smr45C detected a unique RNA species of the expected size accumulating differentially in free-living microorganisms but not expressed in endosymbiotic bacteria (Fig. 2B). In contrast, a 25-mer oligonucleotide probe targeting a sequence 16 nt downstream the Smr45C 3'-end hybridized to a major RNA molecule visible at top of the gel with an expression profile very similar to that of Smr45C (Fig. 2B). The RNA species detected by this oligonucleotide most likely corresponds to the SMc02983 mRNA with a *speF* leader starting downstream the position previously predicted in silico. This RNA molecule could be originated either by processing of a larger undetectable and hence unstable RNA species transcribed from P1 or, most likely, by transcription from the newly identified promoter P2, independently of Smr45C in the biological conditions tested. In agreement with this observation, a *S. meliloti* map of TSS generated by RNA-Seq of total RNA revealed transcripts with 5'-ends at 3,105,292 and 3,105,166 nt positions in this region of the *S. meliloti* chromosome (A. Becker and J.P. Schlüter, personal communication). Altogether, these new experimental evidences further support classification of Smr45C as a Hfq-binding sRNA, likely unrelated to the *speF* RNA element.

**Distribution of the  $\alpha r$  sRNA families in the Rhizobiales.** The occurrence of the  $\alpha r$  sRNA families in sequenced bacterial species of the Rhizobiales was further

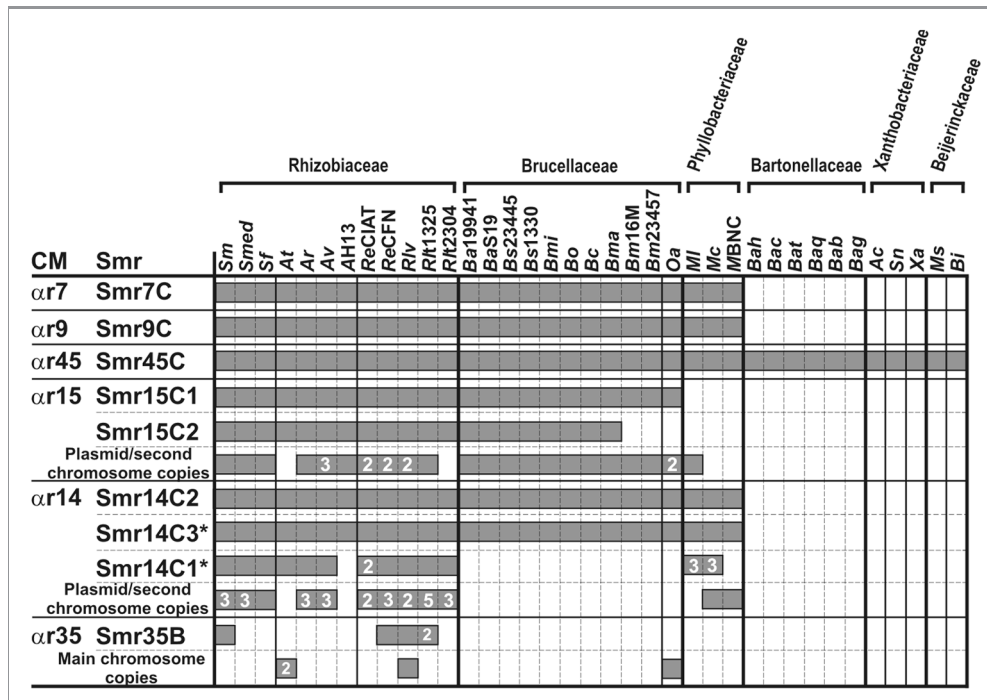


**Figure 2.** Transcription of the *speF* and *Smr45C* RNAs. (A) Nucleotide sequence (both DNA strands) of the *SMc02983-SMc02984* IGR expressing the *speF* and *Smr45C* RNA elements. Numbering indicates coordinates in the *S. meliloti* 1021 genome. The -35 and -10 hexamers of the predicted  $\sigma^{70}$ -dependent promoters (P1 and P2) are boxed. Black arrowheads indicate the predicted start and end of the *speF* RNA as annotated in the Rfam database. Nucleotide positions of 5' and 3' ends previously determined for the *Smr45C* sRNA are boxed and a double arrowhead in red indicates its TSS. A double arrowhead in green indicates the predicted TSS for *speF* from the P2 promoter. The proposed *speF* and *Smr45C* coding sequences are in green and red letters, respectively. (B) Northern analysis of *speF* and *Smr45C* RNAs. Sequences of the 25-mer oligonucleotides used to probe the membranes are underlined in (A). RNA samples were: TY, log TY cultures; TY(S), stationary phase TY cultures; MM, log Minimal Medium cultures; MM(L), luteolin-induced log MM cultures; N, mature alfalfa nodules. Molecular weight markers are shown to the right of the panels. 5S RNA was also probed as RNA loading control.

assessed using the Infernal models (CMs) generated in this work. The results of this comparative analysis are summarized in Figure 3. With the only exception of *Smr35B* ( $\alpha r35$ ), which is encoded in the chromosome-like replicon pSymB, all our query sRNA genes are chromosomally located in *S. meliloti* 1021. Overall,

structure-based clustering of the homologs identified with each of the CMs essentially correlates with the phylogeny of the order ([en.wikipedia.org/wiki/Small\\_non\\_coding\\_RNAs\\_in\\_the\\_endosymbiotic\\_diazotroph\\_%CE%B1-proteobacterium\\_Sinorhizobium\\_meliloti](http://en.wikipedia.org/wiki/Small_non_coding_RNAs_in_the_endosymbiotic_diazotroph_%CE%B1-proteobacterium_Sinorhizobium_meliloti)). The dominant distribution pattern is represented by

$\alpha r7$ ,  $\alpha r9$  and  $\alpha r14$  CMs that identified members in the three taxonomic families of the order that include the bacterial species most closely related to *S. meliloti* i.e., Rhizobiaceae, Brucellaceae and Phyllobacteriaceae. The  $\alpha r15$  family was also found to be widely distributed in the Rhizobiales but lacks chromosomally-encoded relatives



**Figure 3.** Conservation of the *S. meliloti* Smr sRNAs in the Rhizobiales. CMs generated in this work along with the name of the query *S. meliloti* sRNA sequences are listed to the left. The newly predicted chromosomal copies of the Smr14 gene are indicated with an asterisk. All bacterial species with representatives of the  $\alpha$ r RNA families are indicated on top of the panel grouped by taxonomic families i.e., Rhizobiaceae, Brucellaceae, Phyllobacteriaceae, Bartonellaceae, Xanthobacteriaceae and Beijerinckaceae, as follows; Sm, *S. meliloti* 1021; Smed, *S. medicae* WSM419; Sf, *S. fredii* NGR234; At, *Agrobacterium tumefaciens* C58; Ar, *A. radiobacter* K84; Av, *A. vitis* S4; AH13, *A. sp* H13-3; ReCIAT, *Rhizobium etli* CIAT652; ReCFN, *R. etli* CFN42; Rlv, *R. leguminosarum* bv. viceae 3841; Rlt1325, *R. leguminosarum* bv. trifolii WSM1325; Rlt2304, *R. leguminosarum* bv. trifolii WSM2304; Ba19941, *Brucella abortus* bv. One 9-941; BaS19, *B. abortus* S19; Bs23445, *B. suis* ATCC23445; Bs1330, *B. suis* 1330; Bmi, *B. microti* CCM4915; Bo, *B. ovis* ATCC25840; Bc, *B. canis* ATCC 23365; Bma, *B. melitensis* bv. abortus 2308; Bm16M, *B. melitensis* bv. 1 16M; Bm23457, *B. melitensis* ATCC23457; Oa, *Ochrobactrum anthropi* ATCC49188; Ml, *Mesorhizobium loti* MAFF303099; Mc, *M. ciceri* bv. biserrulae WSM1271; MBNC, *M. sp* BNC1; Bah, *Bartonella henselae* Houston-1; Bac, *B. clarridgeae* 73; Bat, *B. tribocorum* CIP105476; Baq, *B. quintana* Toulouse; Bab, *B. bacilliformis* KC583; Bag, *B. grahamii* as4aup; Ac, *Azorhizobium caulinodans* ORS571; Sn, *Starkeya novella* DSM506; Xa, *Xanthobacter autotrophicus* Py2; Ms, *Methylocella silvestris* BL2, Bi, *Beijerinckia indica* subsp indica ATCC9039. Grey bars indicate distribution of each sRNA family in these bacterial species. If more than one, the number of chromosomal and extrachromosomal copies of each sRNA gene is also indicated.

in *Mesorhizobium* species (family Phyllobacteriaceae). The widest distribution corresponded to  $\alpha$ r45 which occurrence extended to species of other three taxonomic families with larger phylogenetic distances to *S. meliloti* i.e., Bartonellaceae, Xanthobacteriaceae and Beijerinckaceae.  $\alpha$ r7,  $\alpha$ r9 and  $\alpha$ r45 members are all encoded by single-copy genes with well-defined promoter regions on the main bacterial chromosomes. Further, with a very few exceptions, complete microsynteny, i.e., conservation of upstream and downstream genes, was observed for representatives of all these three sRNA families in genomes of bacterial species from the same taxonomic family whereas only one of the two flanking genes appears variable across the Rhizobiales (en.wikipedia.org/wiki/%CE%B1r7\_RNA;

en.wikipedia.org/wiki/%CE%B1r9\_RNA; en.wikipedia.org/wiki/%CE%B1r45\_RNA). Thus, the current distribution pattern of the  $\alpha$ r7,  $\alpha$ r9 and  $\alpha$ r45 sRNA families in bacteria is likely the result of the vertical inheritance of their respective sRNA genes located in the ancestral chromosome of the Rhizobiales.

In contrast, sRNA genes of the  $\alpha$ r15 and  $\alpha$ r14 families exist in highly variable copy numbers in the individual genomes; many of them located on extrachromosomal replicons i.e., large accessory plasmids in Rhizobiaceae/Phyllobacteriaceae representatives and the second chromosome in *Brucella* species.  $\alpha$ r15 members occur in two chromosomal copies in 19 genomes of bacteria belonging to the Rhizobiaceae and Brucellaceae families. These two genes are clustered in the same IGR in genomes

from Rhizobiaceae whereas in *Brucella* species map to distant positions on chromosome I. The second chromosomal  $\alpha$ r15 loci were missed by our search in the genomes of *B. melitensis* bv. abortus 2308, *B. melitensis* bv.1 16M and *Ochrobactrum anthropi* ATCC49188. With the exceptions of *A. tumefaciens* C58 and *R. leguminosarum* bv. trifolii 2304, at least a third  $\alpha$ r15 gene is located in extrachromosomal replicons of the host genomes. The  $\alpha$ r14 RNA family showed an even more complex distribution pattern in the Rhizobiales. Two tandem copies of the *S. meliloti* Smr14C2 (formerly Smr14C) and Smr14C3 homologous genes were also identified in *Sinorhizobium* and *Mesorhizobium* species whereas in *O. anthropi* ATCC49188, *Agrobacterium* and *Brucella* species the second chromosomal

gene predicted by the  $\alpha$ r14 CM does not occur in such a syntenic context. A variable number of additional  $\alpha$ r14 copies (up to six more in the genome of *R. leguminosarum* bv. *trifolii* WSM1325) were identified in the main chromosome and accessory plasmids of most of the bacterial species belonging to the Rhizobiaceae and Phyllobacteriaceae families. The  $\alpha$ r15 and  $\alpha$ r14 family members are mostly encoded in IGRs with a few exceptions of genes predicted within or antisense to annotated ORFs. However, these ORFs are frequently small, putatively coding for hypothetical proteins and/or absent from syntenic positions in bacterial genomes, thus representing probable mis-annotations as protein coding regions ([en.wikipedia.org/wiki/%CE%B1r14\\_RNA#Genomic\\_Context](http://en.wikipedia.org/wiki/%CE%B1r14_RNA#Genomic_Context); [en.wikipedia.org/wiki/%CE%B1r15\\_RNA#Genomic\\_Context](http://en.wikipedia.org/wiki/%CE%B1r15_RNA#Genomic_Context)). In general, tandemly-arranged  $\alpha$ r15 and  $\alpha$ r14 genes occur in complete or partial microsynteny with the flanking genes in genomes of Rhizobiaceae and Phyllobacteriaceae as do their homologs on the main chromosome of *O. anthropi* ATCC49188 and *Brucella* species. However, microsynteny is much more fragmented or even absent for many of the remaining chromosomal and plasmidic copies of the  $\alpha$ r14 and  $\alpha$ r15 *loci*. Altogether, these observations suggest that  $\alpha$ r14 and  $\alpha$ r15 constitute families of paralogous sRNA gene copies in the Rhizobiales probably emanated from duplication events of their respective ancestral chromosomal genes over evolutionary time scales. Nonetheless, horizontal transfer events could certainly contribute to the current distribution patterns of some  $\alpha$ r14 and  $\alpha$ r15 gene copies, particularly of those occurring without signs of microsynteny in the accessory plasmids of plant-interacting bacteria. Noteworthy, some of the  $\alpha$ r15 *loci* were flanked by insertion sequences or transposase-encoding genes, among other genetic elements involved in mobility events and genomic rearrangements ([en.wikipedia.org/wiki/%CE%B1r15\\_RNA#Genomic\\_Context](http://en.wikipedia.org/wiki/%CE%B1r15_RNA#Genomic_Context)).

Finally, the  $\alpha$ r35 family exhibits a more restricted and dispersed representation, not only at the species but also at the strain levels. Only seven candidates were identified by the  $\alpha$ r35 Infernal models

in addition to the *S. meliloti* Smr35B sRNA. Three of these predicted Smr35B homologs are encoded on the chromosomes of *A. tumefaciens* C58, *O. anthropi* ATCC49188, and *R. leguminosarum* bv. *viciae* 3841, whereas the remaining four  $\alpha$ r35 genes are extrachromosomal and were identified on the *R. etli* CFN42 plasmid p42f, *R. leguminosarum* bv. *viciae* 3841 plasmid pRL11 and *R. leguminosarum* bv. *trifolii* 1325 plasmids pRL132502 and pRL132504. Again, the majority of the  $\alpha$ r35 genes appeared to be independent transcription units with recognizable promoters with the exceptions of the chromosomal and plasmidic *loci* of *R. leguminosarum* bv. *viciae* 3841 and *R. etli* CFN42, respectively, which putatively overlap to annotated ORFs of unpredicted function. *S. meliloti* 1021 and *O. anthropi* ATCC49188  $\alpha$ r35 genes occur in complete microsynteny with the flanking genes whereas the genomic regions of the other six  $\alpha$ r35 representatives revealed partial or no conservation at all ([en.wikipedia.org/wiki/%CE%B1r35\\_RNA#Genomic\\_Context](http://en.wikipedia.org/wiki/%CE%B1r35_RNA#Genomic_Context)).

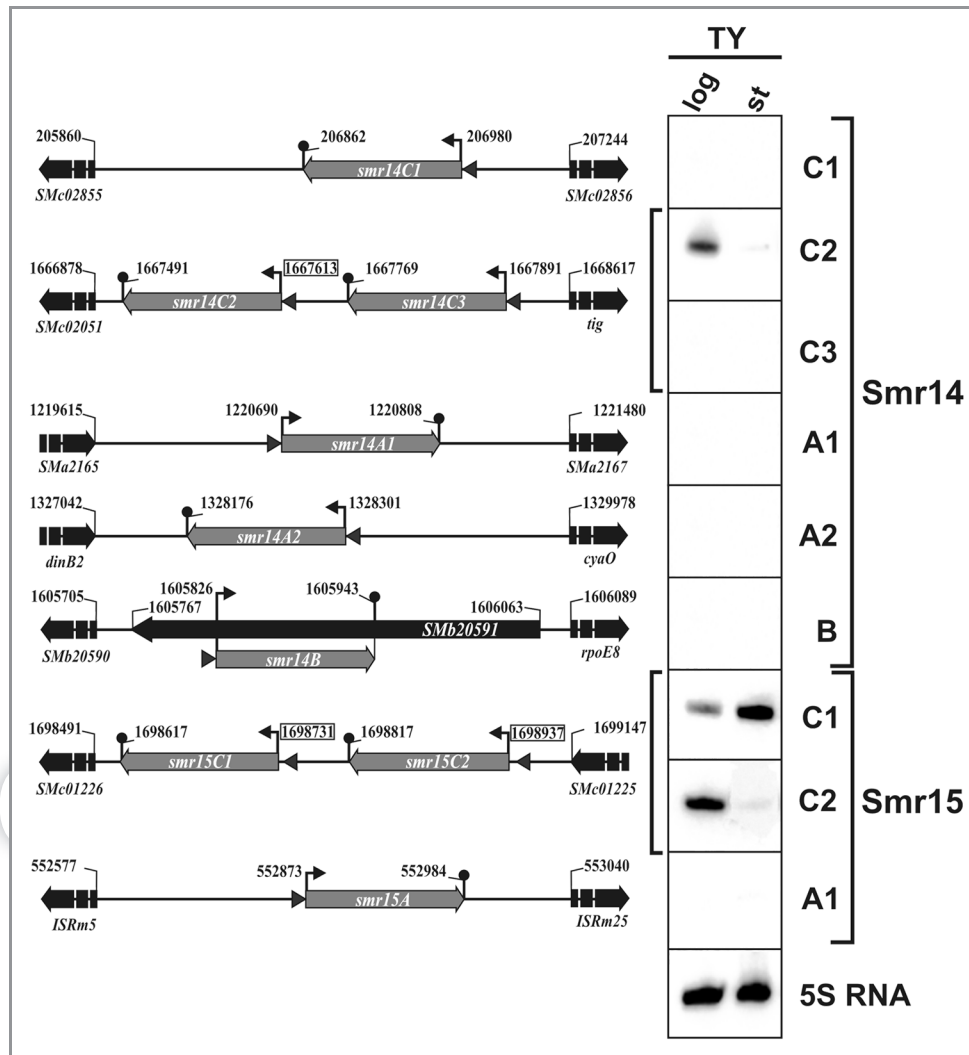
**$\alpha$ r14 and  $\alpha$ r15 representatives are differentially regulated in *S. meliloti*.** The  $\alpha$ r14 and  $\alpha$ r15 CMs also identified several related genes in the *S. meliloti* 1021 genome. A third copy of the Smr15C *locus* was found in the megaplasmid pSymA (Smr15A) and up to 5 additional copies of the query Smr14C2-encoding gene, were also identified; two of them chromosomally located (Smr14C1 and Smr14C3), two in pSymA (Smr14A1 and Smr14A2) and the remaining one in pSymB (Smr14B) (Fig. 4). Similarly to the situation of Smr15C1/Smr15C2, genes arranged in tandem in the same *S. meliloti* 1021 IGR encode Smr14C2 and Smr14C3. All the newly predicted Smr14- and Smr15-like sRNAs in the *S. meliloti* genome are encoded in IGRs, with the exception of Smr14B, which is encoded antisense to the *SMb20591* gene (Fig. 4).

Oligonucleotides specific to all the Smr14 and Smr15 *loci* were used to probe *S. meliloti* RNA obtained from log and stationary phase cultures in TY broth (Fig. 4). These experiments confirmed the growth-dependent expression of Smr14C2, Smr15C1 and Smr15C2 transcripts with

preferential accumulation of Smr15C1 upon entry of bacteria into stationary phase (Fig. 4). Despite their sequence and structural similarity Smr15C1 and Smr15C2 displayed opposite expression profiles. Strikingly, this set of Northern hybridizations did not reveal signs of expression of any of the other five Smr14 genes whereas the Smr15A transcript was barely detected on gels (Fig. 4). Multiple nucleotide sequence alignments of the promoter regions of all the genes encoding  $\alpha$ r15 and  $\alpha$ r14 members in species of the Rhizobiales identified diverse conserved motifs that could contribute to the differential expression of these genes in specific biological conditions ([en.wikipedia.org/wiki/%CE%B1r14\\_RNA#Promoter\\_Analysis](http://en.wikipedia.org/wiki/%CE%B1r14_RNA#Promoter_Analysis); [en.wikipedia.org/wiki/%CE%B1r15\\_RNA#Promoter\\_Analysis](http://en.wikipedia.org/wiki/%CE%B1r15_RNA#Promoter_Analysis)). Supporting this prediction, RNA-Seq of the *S. meliloti* sRNAs expressed in a number of stress conditions has rendered variable number of reads for the *S. meliloti*  $\alpha$ r14- and  $\alpha$ r15-like transcripts, possibly correlating with a diversity of accumulation profiles.<sup>9</sup>

## Discussion

The repertoire of non-coding RNAs expressed by the legume endosymbiont *S. meliloti* is one of the best characterized among those of its  $\alpha$ -proteobacterial counterparts.<sup>6,9</sup> However, current information about the function of these transcripts in bacteria is certainly scarce. The first set of sRNAs identified in the reference strain *S. meliloti* 1021 included eight transcripts with genomic boundaries experimentally determined by independent approaches.<sup>6,9</sup> Here, we have performed a comprehensive computational comparative analysis of these eight sRNA sequences to identify conserved structural motifs putatively relevant to their function as well as to assess their conservation patterns in bacterial genomes. CMs derived from alignments of the Smr sRNA homologs first identified Smr22C as the *S. meliloti* ortholog of the ubiquitous 6S sRNA. This RNA constitutes an example of a well-characterized *trans*-acting protein-binding sRNA.<sup>19</sup> The remaining seven transcripts represent structural and functional novel prokaryotic sRNAs and were



**Figure 4.** Northern analysis of the Smr14 and Smr15 sRNAs in *S. meliloti*. Maps of the genomic regions (not drawn to scale) of all the genes predicted by the  $\alpha$ 14 and  $\alpha$ 15 CMs in *S. meliloti* 1021 are shown to the left of the panels. Numbers denote coordinates of the genes in the genome. Name of the oligonucleotide probes used to hybridize each membrane are indicated to the right and their corresponding nucleotide sequences are listed in **Table 2**. RNA samples were obtained from logarithmic (log) and stationary phase (st) *S. meliloti* 1021 cultures in TY broth. 5S RNA was also probed as RNA loading control.

collected into six different Infernal models. These CMs were used to accurately identify new members of each family in available sequenced bacterial genomes. This search revealed conservation of the Smr sRNAs in bacterial species belonging to the order of the Rhizobiales within the  $\alpha$ -subgroup of proteobacteria and, hence these RNA families were accordingly termed  $\alpha$ . Such a distribution pattern, limited to phylogenetically related bacterial species, is a general feature of the Hfq-dependent base-pairing riboregulators.<sup>1</sup> Indeed, the consensus secondary structures deduced from each family model evidenced Hfq-binding and exposed aSD signatures in

$\alpha$ 15 and  $\alpha$ 14 transcripts as recognizable functional motifs involved in the sRNA-target mRNA interaction. In this regard, it is also noteworthy that previously reported pull-down experiments as well as stability assays on a *S. meliloti* *hfq* mutant background independently confirmed the Smr-Hfq interactions predicted by our CMs.<sup>17,20</sup>

Two particular CMs representing the  $\alpha$ 15 and  $\alpha$ 45 families rendered partial hits to the Rfam models corresponding to the *suH*B and *spe*F non-coding RNA elements, respectively. The secondary structure of the  $\alpha$ 15 sRNAs is predicted to consist of three hairpin motifs, in good

agreement with the mapping of the *A. tumefaciens* Smr15C1 homolog (AbcR1) by enzymatic probing.<sup>21</sup> Furthermore, the aSD-containing 5' hairpin loop of *A. tumefaciens* AbcR1 has been shown to be the functional domain of this transcript for targeting the 5'-UTR of the mRNA encoding the GABA-binding protein.<sup>21</sup> Confirming these experimental results preliminary predictions of Smr15C1/C2-mRNA interactions in *S. meliloti* using diverse bioinformatics tools anticipate a major involvement of the 5' hairpin in target recognition (O. Torres-Quesada and J.I. Jiménez-Zurdo, unpublished results). This 5' stem loop is a variable



or missing domain in *suhB*-like transcripts. Our comparative analysis revealed a similar situation for the  $\alpha$ r35 sRNA family and its variant  $\alpha$ r35b. The dispersed occurrence of the  $\alpha$ r35 *loci* in the Rhizobiales points also to the primary hairpin of these molecules as a functional domain, which probably has co-evolved with its target protein or mRNA in these genomes. Some 5' located sRNA domains have been shown to be critical elements for specific pairing-based mRNA target recognition that can act autonomously when fused to unrelated sRNA molecules.<sup>22</sup> Therefore, the structural modules shared by  $\alpha$ r15/*suhB* and  $\alpha$ r35/ $\alpha$ r35b could be regarded as a kind of  $\alpha$ -proteobacteria-specific “structural Legos” which could accommodate autonomous 5' domains to create functionally diverse sRNAs.<sup>23</sup>

We have also shown that the *S. meliloti* Smr45C sRNA and its downstream mRNA containing the *cis*-regulatory element *speF* are detected as different RNA species on Northern membranes under several biological conditions. Nonetheless, our comparative analysis also revealed that Smr45C always occurs in a syntenic context with a downstream ornithine decarboxylase-encoding gene in the Rhizobiales ([en.wikipedia.org/wiki/%CE%B1r45\\_RNA](http://en.wikipedia.org/wiki/%CE%B1r45_RNA)). Therefore, it cannot be ruled out that under not yet tested specific biological conditions, probably relevant to polyamine biosynthesis, *speF* and Smr45C can be transcribed as a single *cis*-acting RNA element likely controlling translation of the ornithine decarboxylase enzyme. Possible processing of sRNAs from riboswitches was first described in *E. coli*.<sup>24</sup> Furthermore, a dual function of a sRNA as *trans*- and *cis*-acting riboregulator has been recently reported for a lysine riboswitch which lies in the 5'-UTR of the lysine transporter gene in *Listeria monocytogenes*.<sup>25</sup>

Chromosomal location and conservation of at least one of the flanking protein-coding genes are also dominant features of the intergenic base pairing sRNA *loci*.<sup>1</sup> The  $\alpha$ r7,  $\alpha$ r9 and  $\alpha$ r45 CMs represent new examples of bacterial sRNAs encoded by conserved unique chromosomal genes that occur in extensive microsynteny across phylogenetically related species.

However, single-copy genes hardly represent 58% of the total gene content of the *S. meliloti* genome.<sup>4</sup> The genomes of plant-interacting bacteria usually evidence high levels of paralogy suggesting that their expansion through gene duplications has been little constrained during the evolution, facilitating the acquisition of new adaptive functions for life in the soil and within plant cells.<sup>2,4</sup> The  $\alpha$ r14 and  $\alpha$ r15 family members occur in multiple copies in the individual genomes. Multiple sRNA copies are not unusual in bacteria, although the physiological/ecological advantages of these reiterations have been only investigated in a subset of cases.<sup>1</sup> Seemingly homologous sRNAs could act either redundantly, serving as backups in critical pathways, additively sensing different stimuli to integrate diverse environmental signals, independently, regulating different set of genes or hierarchically upon each other.<sup>26-28</sup> In this work we have investigated the expression in free-living bacteria of the Smr14 and Smr15 genes copies identified by the respective covariance models in *S. meliloti* 1021. Northern experiments, promoter predictions and reported RNA-Seq data<sup>9</sup> provide evidences for the differential regulation of these genes. In particular, the opposite expression patterns of Smr15C1 and Smr15C2 contrast with those of their *A. tumefaciens* homologs, which encoding genes are similarly arranged in tandem in the circular chromosome of this bacterium but showed identical expression profiles.<sup>21</sup> Interestingly, Smr15C1 retained its accumulation pattern in a *S. meliloti*  $\Delta$ Smr15C2 derivative and vice versa suggesting that these sRNAs act independently or additively rather than hierarchically as riboregulators in *S. meliloti* (O. Torres-Quesada and J.I. Jiménez-Zurdo, unpublished). On the other hand, the undetectable expression of some transcripts in our assays, particularly of those grouped within the  $\alpha$ r14 sRNA family anticipates that they could be only expressed under not tested specific biological conditions to fulfill different adaptive functions in this bacterium.

In summary, our findings provide a baseline for the forthcoming investigation of the functional plasticity and evolution of the small non-coding RNAs in *S. meliloti* and related plant-interacting bacteria.

## Materials and Methods

**Computational tools and methods.** In a first step the *smr* gene sequences were BLASTed with default parameters against all currently available bacterial genomes (1,615 sequences at 20 April 2011; [www.ncbi.nlm.nih.gov](http://www.ncbi.nlm.nih.gov)). The regions exhibiting significant homologies to the query sequence (78–89% similarity) were used to generate automated infernal alignments<sup>29</sup> for each family. This initial alignment was hand-curated and manually inspected to deduce a consensus secondary structure for each family. The consensus structure was also independently predicted with the program locARNATE<sup>30</sup> in an automatic manner and differences reconciled giving priority to the structural conservation. Given the initial hand-curated structural alignment of close homologs Infernal was used to interrogate the same set of bacterial genomes, searching for new members of the models. The alignment process was repeated during three iterations. The candidates obtained with the Infernal models were selected as members of a given family if their Infernal E-value was  $\leq 10^{-03}$  or lower, or after manual inspection for those with higher Infernal E-values. The hierarchical cluster-tree for each family is derived by WPGMA clustering of the pairwise alignment distances and the optimal number of clusters was calculated from the tree using RNAclust ([www.bioinf.uni-leipzig.de/~kristin/Software/RNAclust/](http://www.bioinf.uni-leipzig.de/~kristin/Software/RNAclust/)). A Stockholm format text file of each family alignment is provided in the links to the family wiki pages at [en.wikipedia.org/wiki/Small\\_noncoding\\_RNAs\\_in\\_the\\_endosymbiotic\\_diazotroph\\_%CE%B1-proteobacterium\\_Sinorhizobium\\_meliloti](http://en.wikipedia.org/wiki/Small_noncoding_RNAs_in_the_endosymbiotic_diazotroph_%CE%B1-proteobacterium_Sinorhizobium_meliloti).

In order to study the microsynteny of each  $\alpha$ r family, we located and extracted the flanking genes of their respective members. Non-annotated ORFs were further annotated using Blast2GO,<sup>31,32</sup> and the high-throughput pipelines ProtSweep, and DomainSweep.<sup>33</sup> The obtained results were later manually inspected in order to annotate and predict a biological function for these ORFs. In the few cases where the predicted sRNAs overlapped ORFs, the same procedure as with the flanking genes was carried on. ORFs

**Table 2.** Oligonucleotide probes used in Northern hybridizations

sRNA	Nucleotide sequence	Target sequence <sup>a</sup>
speF	5'-CTTCAGTAACCCCGCTGTTGGAGA-3'	3,105,282–3,105,258
Smr45C	5'-CCGACCCTCGTTGCTTCAAGATGT-3'	3,105,328–3,105,304
Smr14C1	5'-AACCGACCGAATGCCGGGCGCCGTG-3'	206,954–206,930
Smr14C2	5'-TGCTTGATCTGATTGGCAACCGGA-3'	1,667,552–1,667,528
Smr14C3	5'-ACCGGCGGGCGTCATAAAGCGATT-3'	1,667,818–1,667,794
Smr14A1	5'-AACCGATCGGCTCTTGCCTGG-3'	1,220,715–1,220,739
Smr14A2	5'-GAGGAAAGTGCCTCGCATATCGAA-3'	1,328,303–1,328,279
Smr14B	5'-GTGCGCCGGCTTTCGATCTGACC-3'	1,605,895–1,605,919
Smr15C1	5'-GAGGAGAAAGCCGTAGATGCACCA-3'	1,698,728–1,698,704
Smr15C2	5'-ACTGGGAGGAGAAGCCACCAAGAT-3'	1,698,928–1,698,904
Smr15A	5'-GGAGAAAAGTCCATGCGCATCAA-3'	552,875–552,899

<sup>a</sup>Coordinates of the sequence stretches complementary to each probe in the *S. meliloti* 1021 genome according to [iant.toulouse.inra.fr/bacteria/annotation/cgi/rhime.cgi](http://iant.toulouse.inra.fr/bacteria/annotation/cgi/rhime.cgi).

shorter than 30 aa, that neither showed similarity with any database entry, nor motif or signatures when searched against family and motif databases such as Interpro,<sup>34</sup> PFAM<sup>35</sup> or Smart<sup>36</sup> were considered as miss-annotations and thus not registered in the genomic context graph of the corresponding  $\alpha$ r family.

**Experimental methods.** Growth of *S. meliloti* strain 1021 in TY and MM broths, RNA extraction from free-living and endosymbiotic bacteria and Northern hybridizations were performed as previously described.<sup>6</sup> Sequences of the 25-mer oligonucleotides used to probe Northern membranes are detailed in Table 2.

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