Transcription attenuation-derived small RNA rnTrpL regulates tryptophan biosynthesis gene expression in *trans*

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ABSTRACT

Ribosome-mediated transcription attenuation is a basic posttranscriptional regulation mechanism in bacteria. Liberated attenuator RNAs arising in this process are generally considered nonfunctional. In Sinorhizobium meliloti, the tryptophan (Trp) biosynthesis genes are organized into three operons, trpE(G), ppiD-trpDC-moaC-moeA, and trpFBA-accDfolC, of which only the first one, trpE(G), contains a short ORF (trpL) in the 5'-UTR and is regulated by transcription attenuation. Under conditions of Trp sufficiency, transcription is terminated between trpL and trpE(G), and a small attenuator RNA, rnTrpL, is produced. Here, we show that rnTrpL base-pairs with *trpD* and destabilizes the polycistronic *trpDC* mRNA, indicating rnTrpL-mediated downregulation of the *trpDC* operon in *trans*. Although all three *trp* operons are regulated in response to Trp availability, only in the two operons trpE(G) and trpDC the Trp-mediated regulation is controlled by rnTrpL. Together, our data show that the trp attenuator coordinates trpE(G) and trpDC expression posttranscriptionally by two fundamentally different mechanisms: ribosome-mediated transcription attenuation in cis and base-pairing in trans. Also, we present evidence that rnTrpL-mediated regulation of trpDC genes expression in trans is conserved in Agrobacterium and Bradyrhizobium, suggesting that the small attenuator RNAs may have additional conserved functions in the control of bacterial gene expression.

INTRODUCTION

Since the discovery of ribosome-mediated transcription attenuation in bacteria 40 years ago (1,2), it has been presumed that terminated attenuator RNAs have no own functions (3). Here, we show that, in *Sinorhizobium meliloti*, an attenuator RNA that is liberated under conditions of tryptophan (Trp) sufficiency acts in *trans* to destabilize Trp biosynthesis (*trp*) mRNA.

Transcription attenuation sensing specific uncharged tRNA is a sophisticated posttranscriptional mechanism that is used by bacteria to regulate their gene expression. A prime example is the regulation of the *trpEDCBA* operon in Escherichia coli. The structural genes of this operon are preceded by an mRNA leader that is able to form mutually exclusive secondary structures and contains a small open reading frame (sORF), *trpL*, with consecutive Trp codons (4). Ribosome pausing at the Trp codons prevents the formation of a terminator hairpin, resulting in structural genes transcription when cellular Trp concentrations are low. At high Trp levels, the leader peptide TrpL is translated efficiently, resulting in the formation of the terminator hairpin and, consequently, transcription termination, thus leading to the release of a small attenuator RNA. In this way, bacteria respond in a very fast and resource-saving manner to amino acid (aa) availability. Similar ribosomedependent transcription attenuators are widespread (3,5,6). Many Gram-positive bacteria developed different strategies for transcription attenuation of *trp* operons. For example, Bacillus subtilis utilizes a Trp-activated RNA binding protein (TRAP) while other gram-positive bacteria, such as *Streptococcus*, harbor a tRNA^{Trp} sensing structure (T-box) in the leader of trp mRNA (3,7,8).

Thus, at least three different mechanisms for RNA-based regulation of *trp* operons evolved in bacteria, indicating the importance of strict posttrancriptional control of Trp biosynthesis for bacterial survival. Most probably, this is

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due to the fact that Trp is the most costly-to-synthesize amino acid (3). Despite this, many bacteria have split *trp* operons with only one of them being preceded by a transcription attenuator (3). This raises the question of whether the other *trp* operons are posttrancriptionally regulated by alternative mechanisms, for example by small RNAs. Small RNAs (sRNAs) regulate virtually all aspects of bacterial physiology, including metabolism and adaptation to changing environmental conditions. Mechanistically, they can exert their regulatory function by protein binding or by base pairing with mRNAs (9). Bacterial trans-acting and basepairing sRNAs are highly versatile due to their capacity to target multiple mRNAs (10-12). Typically, such sRNAs display limited complementarity to their targets and need an RNA chaperone, such as Hfg or ProQ, for efficient mRNA binding (9,13). In most cases, the interaction between an sRNA and an mRNA prevents translation and/or destabilizes the mRNA target (9, 14, 15).

Most of the functionally characterized sRNAs have orphan genes. Additionally, 3'-UTRs of bacterial mRNAs are considered as established sRNA reservoirs (16,17). In contrast, functional sRNAs originating from 5'-UTRs were reported much less frequently and there are very few examples of riboregulators that act both in *cis* and *trans* (18). These include the S-adenosylmethionine riboswitches SreA and SreB which act as base-pairing sRNAs that control the expression of a virulence gene in *Listeria monocytogenes* (19). Further, in *L. monocytogenes* and *Enterococcus faecalis*, a coenzyme B12 (AdoCbl) riboswitch was shown to work as a protein-binding sRNA that sequesters a response regulator (20,21). To date, small *trans*-acting RNAs derived from ribosome-mediated transcription attenuation have not been described.

We are interested in Sinorhizobium meliloti, a soildwelling alpha-proteobacterium that is capable of fixing molecular nitrogen in symbiosis with some legume plants. In S. meliloti, the structural trp genes are organized into operons trpE(G), trpDC, and trpFBA (3,22), of which only trpE(G) is regulated by transcription attenuation (23). In the past decade, hundreds of sRNA candidates have been detected in S. meliloti (24) and, meanwhile, physiological roles based on direct binding to target mRNAs have been shown for several of them. For example, AbcR1, AbcR2 and NfeR1 were shown to regulate transporter genes (25,26), while RcsR1 was reported to down-regulate the autoinducer synthase encoding mRNA sinI (27), and EcpR1 and GspR were suggested to regulate the cell cycle (28,29). Given that sRNA RcsR1 is transcribed from a position upstream of trpE(G) and is identical to the trpLderived attenuator RNA (23), a possible role of this particular RNA in the (co)regulation of the *trp* operons seemed plausible. In line with this hypothesis, RcsR1 was predicted to base-pair with several mRNAs including trpDC and constitutive ectopic RcsR1 overproduction was shown to result in lower trpC mRNA levels (27). However, it was not clear if this was a direct effect of RcsR1 on trpDC mRNA and if it had any functional implications for bacterial gene regulation in response to Trp availability.

Here, we demonstrate that the liberated attenuator sRNA of the Trp biosynthesis gene trpE(G) base-pairs with trpD and down-regulates trpDC under conditions of Trp suffi-

ciency. Please note that, in this study, the sRNA RcsR1 was re-named rnTrpL to underline its biogenesis (in transcription attenuation) and its role in the posttranscriptional regulation of bacterial *trp* operons. The data obtained in this study provide an interesting example for a multifunctional mRNA leader with regulatory functions in both *cis* and *trans*.

MATERIALS AND METHODS

Cultivation of bacteria

Escherichia coli strains were grown in LB medium (30). Unless stated otherwise, Sinorhizobium (Ensifer) meliloti 2011 (31,32) and Agrobacterium tumefaciens (A. fabrum) NTL4 (pZLR4) (33,34) were grown in TY medium (35) as described (27). For analysis of gene expression at different Trp availability conditions, GMX medium (36) with the indicated Trp concentrations was used. Bradyrhizobium japon*icum (B. diazoefficiens)* 110spc4 (37,38) was grown in PSY medium (39). When appropriate, antibiotics were included at the following concentrations: ampicillin (200 μ g/ml), streptomycin (250 μ g/ml), tetracycline (20 μ g/ml for S. meliloti and A. tumefaciens; B. japonicum was cultivated with 25 μ g/ml Tc in liquid and 50 μ g/ml Tc on plates), gentamycin (10 µg/ml; for S. meliloti on plates, 20 µg/ml were used); kanamycin (25 μ g/ml for *E. coli* and 200 μ g/ml for S. meliloti), spectinomycin (100 µg/ml for B. japonicum). IPTG was used at a final concentration of 1 mM.

Plasmid construction and conjugation

Cloning procedures were performed essentially as described (30). FastDigest Restriction Endonucleases and Phusion polymerase (Thermo Fischer Scientific) were used routinely for cloning in E. coli JM109 (40) or E. coli DH5a. When pJet1.2/blunt (CloneJet PCR Cloning Kit, Thermo Fischer Scientific) was used for cloning, the inserts were subcloned into the conjugative plasmids pRK4352 (41), pSRKGm, pSRKTc (42), or pK18mobsacB (43). Insert sequences were analyzed by Sanger sequencing with plasmidspecific primers (sequencing service by Microsynth Seqlab, Göttingen, Germany) prior to conjugation into S. meliloti. Plasmids used for conjugation in S. meliloti are listed in Supplementary Table S1 and oligonucleotides used in this study are listed in Supplementary Table S2. Oligonucleotides were synthesized by Eurofins Genomics, Ebersberg (Germany) and Microsynth, Balgach (Switzerland).

Plasmids pDrive-RcsR1 and pRK-SmelRcsR1 (from now on pDrive-rnTrpL and pRK-rnTrpL, respectively; see Supplementary Table S1) were described previously (27). In *S. meliloti*, pRK-rnTrpL leads to constitutive transcription of rnTrpL from a heterologous ribosomal RNA (*rrn*) promoter originating from the alpha-proteobacterium *Rhodobacter shpaeroides* (41). For constitutive overexpression of the *B. japonicum* rnTrpL homolog Bja-rnTrpL, the corresponding sequence was cloned in the chromosome integration plasmid pRJ-MCS (44) that was cleaved with KpnI and SpeI. Plasmid pRK-rnTrpL-AU1,2UA, which allows for constitutive transcription of an *S. meliloti* rnTrpL derivative that lacks a functional sORF (replacement of the *trpL* AUG codon with UAG), was constructed as follows. The rnTrpL-sequence was amplified with primers RcsR1-ATG/TAG-fw and pRKSmelRcsR1re and cloned into pRK4352. To construct pRK-rnTrpL-AU1,2UA-G44C, the same primers were used in a PCR with pRK-rnTrpL-G44C as the template and the amplicon was cloned into pRK4352. To construct pRK-rnTrpL-CG40,41GC and pRK-rnTrpL-GG46,47CC, site-directed mutagenesis of the rnTrpL-sequence in pDrive-rnTrpL was performed by inverse PCR with Phusion polymerase, followed by treatment with DpnI (Thermo Fischer Scientific). The mutated inserts were excised with BamHI and EcoRI, and cloned into pRK4352. To construct plasmid pRK-trpL-egfp, egfp was amplified with primers Bam-RcsR1-egfp-fw and Ecoegfp-re using pLK64 (45) as template, and the amplicon was cloned into pRK4352 using its BamHI and EcoRI restriction sites. The resulting translational rnTrpL::egfp fusion contains the first six codons of *trpL* fused to the third codon of egfp. Plasmids pSRKTc-rnTrpL and pSRKGmrnTrpL for IPTG-inducible transcription of lacZ'-rnTrpL were constructed as follows. The rnTrpL-sequence of S. meliloti was amplified and ligated with NdeI/SpeI-digested pSRKTc or pSRKGm, resulting in an in-frame insertion of the *trpL* sORF to the ATG of NdeI. Similarly, plasmids pSRKTc-Atu-rnTrpL and pSRKTc-Ec-rnTrpL were constructed by cloning the rnTrpL homologs of A. tumefaciens and E. coli into pSRKTc. To construct pSRKTc-rnTrpL-CG40,41GC and pSRKTc-rnTrpL-GG46,47CC, mutated rnTrpL-sequences were amplified using as template pDrivernTrpL-CG40,42GC and pDrive-RcsR1-GG46,47CC, respectively, and cloned into pSRKTc cleaved with NdeI and SpeI. To construct pSRKGm-trpDC'-egfp, the trpDC' sequence of S. meliloti was amplified with primers NdeItrpD-fw and 5'-egfp-trpC-re. In parallel, egfp was amplified using pLK64 (45) as template and primers trpC-egfpfw and XbaI-egfp-re. The two PCR products were mixed and used for overlap-PCR with the primers NdeI-trpDfw and XbaI-egfp-re. The resulting amplicon was ligated with pJet1.2/blunt, resulting in pJet-trpDC'-egfp. The insert was subcloned into pSRKGm using NdeI and XbaI. The resulting plasmid pSRKGm-trpDC'-egfp allows for IPTG-inducible transcription of a bicistronic *trpDC*'::*egfp* mRNA encoding TrpD and the fusion protein TrpC'-EGFP. The fusion construct contains the first 16 trpCcodons fused in frame to the third codon of egfp. To introduce compensatory mutations into the *trpDC*[']::*egfp* reporter, pJet-trpDC'-egfp was subjected to site-directed mutagenesis as described above. The mutated inserts were subcloned into pSRKGm using NdeI and XbaI, resulting in pSRKGm-trpD-CG985,986GC-trpC'-egfp and pSRKGmtrpD-CC977,978GG-trpC'-egfp. Transconjugants containing one of the pSRK-plasmids were stored at -80°C. Double transconjugants (containing pSRK-plasmids with different antibiotic resistance genes) were used directly after the transfer of the second plasmid to S. meliloti. To construct plasmid pK18mobsacB-∆trpL for marker-less deletion of the rnTrpL-sequence in the chromosome of S. meliloti, regions located upstream and downstream of trpL in the chromosome were amplified separately. Overlap PCR was performed and the resulting amplicon was cloned into pK18mobsacB (43), which was cleaved with EcoRI and PstI. Plasmid pK18mobsacB-∆trpL was used to delete the chromosomal sequence ranging from the third nucleotide after the rnTrpL transcription start site to the last T in the TTTT stretch of the rnTrpL transcription terminator. Similarly, plasmid pK18mobsacB- Δ trpC for markerless in-frame deletion in gene *trpC* of *S. meliloti* was constructed. The in-frame deletion construct contains the first ten codons fused to the last nine codons of *trpC*.

Plasmid constructs were transferred to *S. meliloti*, *A. tumefaciens* or *B. japonicum* by diparental conjugation with *E. coli* S17–1 as the donor (46). Bacteria were mixed, washed in saline and spotted onto a sterile membrane filter, which was placed onto a TY plate without antibiotics. After incubation for at least 4 h at 30°C, serial dilutions were plated on TY agar with selective antibiotics.

Construction of S. meliloti 2011 deletion mutants

The deletion mutants $2011 \Delta trpL$, $2011 \Delta trpC$ and $2011 \Delta trpL \Delta trpC$ were constructed as described (47), using TY solid media. For in-frame deletion of trpC, it was necessary to supplement the TY medium with $20 \ \mu g/ml$ L-tryptophan. Deletion mutant candidates were confirmed by PCR analysis of cell lysates using primers which correspond to sequences located outside the chromosomal region that was cloned in the pK18mobsacB- Δ trpL and pK18mobsacB- Δ trpC, respectively. The $\Delta trpL$ deletion was additionally confirmed by Northern blot analysis in comparison to the parental strain.

TrpC'-EGFP fluorescence measurement

Double *S. meliloti* transconjugants containing pSRKGmand pSRKTc- constructs for IPTG-induced expression of *egfp* reporter fusions and the sRNA rnTrpL of *S. meliloti*, respectively, were grown in TY with Gm (gentamycin) and Tc (tetracycline), but without IPTG, to an $OD_{600nm} = 0.5$. Then, 1 mM IPTG was added to induce simultaneous transcription of the bicistronic *egfp*-fusion mRNA and the regulatory sRNA. Induced cultures were incubated for 20 min at 30°C with shaking. Thereafter, 300 µl of the cultures were transferred to a 96-well microtiter plate and fluorescence of the fusion protein TrpC'-EGFP was measured on a Tecan Infinite M200 reader. Values were normalized to the ODs measured on the Tecan.

RNA purification, Northern blot analysis, qRT-PCR and RNA half-life measurements

RNA was isolated from bacterial cultures grown to an $OD_{600nm} = 0.5$. Unless stated otherwise, 15 ml *S. meliloti* or *A. tumefaciens* culture was filled into a tube with ice rocks and centrifuged. The pellet was resuspended in 250 ml TRIzol and glass beads (with a volume corresponding to the volume of the pellet) were added. Cells were lysed in a shacking mill (4°C), two times for 15 min, interrupted by a 10 min incubation at 65°C. Then 750 µl TRIzol was added and RNA was isolated according to the manufacturer instructions. After precipitation with isopropanol, the RNA was additionally purified by extraction with hot phenol and chloroform-isoamylalcohol to remove residual RNases. RNA from *B. japonicum* or *E. coli* DH5 α was isolated using hot phenol. For RNA half-life measurements,

S. meliloti RNA was isolated using RNeasy columns (Qiagen) as described (27). RNA concentration and purity was determined by measuring the absorbance at 260 and 280 nm. RNA integrity was controlled in a 10% polyacrylamideurea gels stained with ethidium bromide.

For real-time RT-PCR (qRT-PCR), 10 µg RNA was treated with 1 µl TURBO-DNase for 30 min and DNA removal was confirmed by PCR with *rpoB*-specific primers for each sample. The gRT-PCR analysis was performed with Brilliant III Ultra Fast SYBR® Green QRT-PCR Mastermix (Agilent) according to the manufacturer manual. Each 10 µl reaction mixture contained 5 µl Master Mix (supplied), 0.1 µl DTT (100 mM, supplied), 0.5 µl Ribo-Block solution (supplied), 0.4 μ l water, 1 μ l of each primer (10 pmol/ μ l), and 2 μ l RNA (20 ng/ μ l). For Figures 1D and 2C, both primers were added simultaneously before the cDNA synthesis step. For other Figures, strand-specific qRT-PCR was performed. For this, first only the primer needed for cDNA synthesis was added to the reaction mixture. After cDNA synthesis and incubation for 10 min at 96°C to inactivate the reverse transcriptase, the probes were cooled to 4°C, the second primer was added, and PCR was performed starting with 5 min incubation at 96°C. Used primer pairs and their efficiencies (as determined by PCR using serial two-fold dilutions of RNA) are listed in Supplementary Table S2. As a reference mRNA, rpoB was used when mRNA steady-state levels were compared. For halflife measurements, the stable 16S rRNA was used as a reference. For the real-time RT-PCR of this reference molecule, the RNA samples were additionally diluted till the quantification cycle (Cq) of mRNA and 16S rRNA was similar. Routinely, 2 μ l RNA with a concentration of 0.002 ng/ μ l was used in a 10 µl reaction for real-time RT-PCR of 16S rRNA. The reactions were performed in a spectrofluorometric thermal cycle (Biorad). The Cq was set to a cycle at which the curvature of the amplification is maximal (48)using BioRad CFX Manager 3.0. When outlier were identified (e. g. technical duplicate range > 0.5 Cq), the qRT-PCR was repeated. No template-controls were always included. qPCR product specificity was validated by a melting curve after the qPCR-reaction and by gel electrophoresis. Fold changes of mRNA amounts were calculated with the Pfaffl formula (49). All qRT-PCR graphs show means and standard deviations from three independent experiments, each performed in technical duplicates. Northern blot hybridization, signal detection and sRNA bands quantification was performed as described (27, 36).

The measurement of mRNA half-lives was performed as described (50) with modifications (27). In brief, transcription was terminated by adding rifampicin (600 μ g/ml final concentration; stock concentration 60 mg/ml in methanol) to the *S. meliloti* culture at an OD₆₀₀ = 0.5. Aliquots of 2 ml were withdrawn at time points 0, 2, 4 and 6 min and mixed with 4 ml RNAprotect Bacteria Reagent (Qiagen). After centrifugation, RNA was purified from the bacterial pellet and used in qRT-PCR analysis with 16S rRNA as a reference. The half-life of rnTrpL was determined as described (50). In brief, 15 μ g total RNA was separated in 10% denaturing polyacrylamide gel and analyzed by Northern blot hybridization with the radioactively labeled oligonucleotide SmelRcsR1 as the probe. 5S rRNA was used as the loading

control. Half-lives were calculated from linear-log graphs. The time after rifampicin addition was plotted against relative mRNA amounts.

In vitro transcription

A DNA template suitable for subsequent in vitro transcription of rnTrpL was obtained by PCR using the forward primer ivTr_RcsR1_f_GGGA, the reverse primer Random_RcsR1_Rev (Supplementary Table S2), and plasmid pRK-rnTrpL as a template. The primer Random_RcsR1_Rev contained an additional sequence at the 5'-end (cgattgtcgttatccattctcgttcatc), generating an artificial 3'-tail in the in vitro transcribed rnTrpL. This tail sequence did not affect the folding of SL1 and SL3 in the transcript and served as a binding site for the oligonucleotide used for primer extension. In this way, modifications in the entire rnTrpL sequence could be probed. To avoid transcription termination by T7 polymerase, the U-stretch of the SL3 transcription terminator was limited to three uridines (51).

The PCR product was purified using HiBind DNA spin-column of the E.Z.N.A DNA probe purification kit (Omega) and eluted in ultrapure water. A total amount of 500 ng of each purified PCR product was used for in vitro transcription using the MEGAshortscript T7 kit (Ambion) or the HiScribeTM T7 High Yield RNA Synthesis Kit (NEB). The reactions were performed according to the manufacturer's instructions (1x T7-polymerase buffer, 7.5 mM ATP, 7.5 mM CTP, 7.5 mM GTP, 7.5 mM UTP, 25 U T7-enzyme mix) for at least 5 h at 37°C. The DNA template was removed by digestion with 1 μ l TURBO-DNase for 1 h at 37°C. The *in vitro* transcript was phenol-extracted, ethanol-precipitated and analyzed by electrophoresis in a 10% polyacrylamide-urea gel and subsequent staining with ethidium bromide.

rnTrpL structure probing

To analyze the structure of rnTrpL, RNA structure probing experiments were done as described (52-54). Typically, 0.6 µg of in vitro-synthesized RNA was heat denatured at 90°C for 1 min and then put on ice for 5 min. The RNA was renatured either in AN buffer (DMS probing, 50 mM sodium cacodylate, pH 7.5, 5 mM MgCl₂, 60 mM KCl) or BN buffer (CMCT probing, 50 mM sodium borate, pH 8.0, 5 mM MgCl₂, 60 mM KCl) for 20 min at room temperature. Next, the samples (total volume of 8 µl) were mixed with 1 µl of yeast tRNA (2 mg/ml, Ambion) and 1 µl of dimethyl sulfate (DMS; Sigma Aldrich) solution (diluted to 1/2, 1/5, 1/10 and 1/20, respectively, in 20% ethanol) or 1 μl of 1-cyclohexyl-3-(2-morpholinoethyl)-carbodiimide metho-p-toluenesulfonate (CMCT; Sigma Aldrich) solution (diluted to 2.5 mg/ml, 5 mg/ml, 10 mg/ml, 20 mg/ml and 50 mg/ml, respectively, in RNase/DNase-free water). Control reactions were done under equal conditions in the absence of DMS and CMCT. Following incubation for 5 min at room temperature, the reactions were terminated by ethanol precipitation in the presence of 1/10 volume 3 M sodium acetate (pH 5.2). Chemical modifications of specific nucleotides by DMS and CMCT, respectively, were determined by primer extension analysis. Briefly, reverse tran-

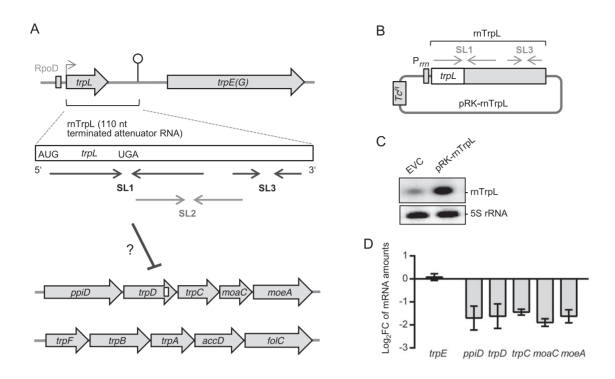


Figure 1. Ectopic overproduction of the attenuator sRNA rnTrpL decreases the level of the polycistronic *trpDC* mRNA in *S. meliloti*. (A) Schematic representation of the three *trp* operons and the attenuator RNA rnTrpL. ORFs are depicted as gray arrows, the RpoD-like promoter is shown as a gray rectangle, a transcription start site is indicated by a flexed arrow and a transcription terminator by a hairpin (according to 23 and 59). The terminated attenuator sRNA rnTrpL (110 nt; synonym RcsR1) is shown as a white box with start and stop codons indicated. Below this box, regions corresponding to stem-loops (SL) 1, 2 and 3 are indicated by convergent arrows. SL2 corresponds to the antiterminator and SL3 to the terminator of transcription. SL2 is preferentially formed in the presence of translating ribosomes that pause upon tRNA^{Trp} shortage. For the rnTrpL secondary structure, see Figure 2. The proposed binding site of rnTrpL in the *trpD* ORF is indicated by a white box. (B) Schematic representation of the plasmid pRK-rnTrpL. Transcription of rnTrpL from this plasmid is expected to be terminated at SL3 in TY medium (see Supplementary Figure S1). (C) Northern blot analysis of RNA isolated from the EVC strain 2011 (pRK4352) and the overexpressing strain 2011 (pRK-rnTrpL), grown in TY. The membrane was first hybridized with a 5S rRNA-specific probe (loading control). (D) Results of qRT-PCR analyses of the indicated mRNAs of the *trpDC* operon and *trpE* (control mRNA). Strain 2011 (pRK-rnTrpL) was compared to the EVC and log₂(fold change) (log₂FC) in mRNA standy-state levels were calculated. Shown are the results from three independent experiments, each performed in technical duplicates (mean values and standard deviations are indicated).

scription reactions were performed using primer RcsR1-Random_Rev_PE. Aliquots of chemically modified RNAs were hybridized with 1–3 pmol of 5'-labeled primer (1–2 \times 10^5 c.p.m.). Following a brief heating step (90°C, 2 min), the reaction was cooled slowly (5 min at 75°C, 10 min at 50°C, 5 min at 37°C, 10 min at room temperature). Next, the primer annealing mixture was used to set up a 20-µl reverse transcription reaction in $1 \times \text{SuperScript}^{\textcircled{R}}$ III RTase reaction buffer supplemented with 170 units of SuperScript[®] III RTase (Invitrogen), 20 units RNaseOUT (Invitrogen) and 1 mM of each dNTP. The reaction was performed at 42°C for 50 min and then at 55°C for 60 min. Reactions were terminated by the addition of 1/10 vol of 3 M sodium acetate, pH 5.2, and 10 volumes of ice-cold ethanol. Following centrifugation, the pellets were washed with 70% ethanol. The dried pellets were resuspended in water and treated with DNase-free RNase A for 20 min at 37°C (0.2 mg/ml, Invitrogen). Next, PCR-grade proteinase K (Invitrogen) was added to a final concentration of 1 mg/ml and the reaction was incubated for another 15 min at 55°C. Reactions were stopped by adding Fu-mix (6 M urea, 80% deionized formamide, $1 \times \text{TBE}$, 0.1% (w/v) Bromophenol blue, and 0.1% (w/v) Xylene cyanol). Reaction products were separated in TBE-buffered 10% polyacrylamide gels containing 7 M urea. Signals were visualized using a Typhoon 9200 imager (GE Healthcare) and analysed using Quantity One software (BioRad).

Bioinformatic methods

RNA structures were predicted by Mfold (55) and RNAfold (56) and visualized using VARNA (57). RNA-RNA interactions were predicted by IntaRNA (58) using the following input seed parameters: minimal number of base-pairs in seed, 6; maximal number of mismatches in seed, 2; maximal energy, 20; minimal unpaired probability 8 (each), 0; seed positions were not provided. For other parameters, default settings were used. For alignment of DNA and RNA sequences, T-Coffee (59) and LocaRNA (58) were used, respectively.

RESULTS

Overproduction of the attenuator sRNA rnTrpL decreases the level of the polycistronic *trpDC* mRNA

It was shown previously that, in RNA from *S. meliloti* 2011 cultures grown in the rich TY medium, the terminated at-

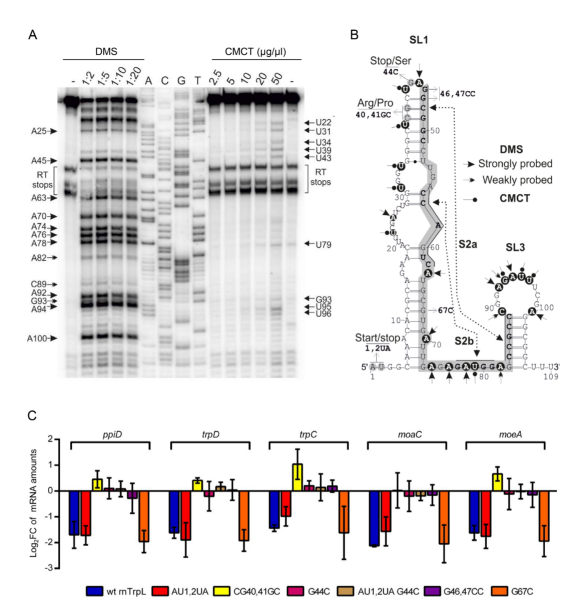


Figure 2. The apical part of rnTrpL stem-loop 1 is critically involved in controlling the mRNA levels of the *trpDC* operon. (A) Chemical probing with DMS or CMCT of *in vitro*-transcribed RNA representing the rnTrpL of *S. meliloti*. DMS and CMCT modifications were identified by primer extension analysis. Shown is the autoradiogram of a representative denaturing polyacrylamide gel. Lanes: -, reaction performed in the absence of DMS; 1:2, 1:5, 1:10 and 1:20, reactions performed with the indicated dilutions of DMS; 2.5, 5, 10, 20 and 50 mg/ml, reactions performed with the indicated concentrations of CMCT; T, G, C, A sequencing reactions using the indicated dideoxynucleotides. (**B**) RNA secondary structure model of rnTrpL. Mutations characterized in this study as well as the SL1 and SL3 structures are indicated. The nucleotides highlighted by a gray box indicate the antitermination structure (SL2) reported previously (23). Nucleotides shown in boldface within the gray box indicate the structural analyses shown here. Chemical modifications of specific nucleotides (highlighted by black circles) that were reproducibly detected in repeated experiments are indicated by arrowheads (DMS) and pinned arrowheads (CMCT), respectively. (C) qRT-PCR analysis of the *trpDC* operon in strains overexpressing wt rnTrpL or rnTrpL variants with the indicated mutations. The mRNA levels of the indicated ORFs in the overexpressing strains were compared to the levels in the EVC. Shown are results from three independent experiments, each performed in technical duplicates (mean values and standard deviations are indicated). For other details, see Figure 1.

tenuator sRNA rnTrpL can be detected in polyacrylamide gels as a 110-nt band (23,27). Since in *E. coli*, in medium containing an excess of Trp, approximately 15% of the transcription events proceed into the structural *trp* genes (60), we tested whether there is a *trpLE(G)* co-transcription in *S. meliloti* 2011 grown in TY. qRT-PCR analyses revealed co-transcription and suggested an approximately 45-fold higher steady-state level of rnTrpL when compared to *trpLE(G)* or *trpE(G)* (Supplementary Figure S1). Thus, in the TY medium routinely used in this work, transcription was efficiently terminated at the attenuator. As a result, the sRNA rnTrpL was accumulated, and was potentially available for regulation in *trans*, while the readthrough transcript was present at a low level only. To further characterize rn-TrpL, we determined its stability in exponentially growing cultures in TY (Supplementary Figure S2). With a half-life of 7.6 ± 0.2 min, the decay of rnTrpL was faster than that of other *S. meliloti* sRNAs under comparable conditions (50).

Previously, constitutive rnTrpL overproduction from plasmid pRK-rnTrpL led to a decrease in the steady-state level of trpC mRNA (27). Here, we asked the question of whether or not rnTrpL overproduction affects the whole polycistronic operon ppiD-trpDC-moaC-moeA (from now on referred to as *trpDC* operon) (see Figure 1A). First, we confirmed higher levels for the sRNA rnTrpL in the overexpressing strain S. meliloti 2011 (pRK-rnTrpL) in comparison to the empty vector control strain (EVC) by Northern blot analysis (see Figure 1B and C). The length of the ectopically overproduced sRNA was similar to that of the native sRNA rnTrpL, which was transcribed from the chromosome. Then, we compared the mRNA levels of the *trpDC* operon in the overexpressing strain and the EVC by gRT-PCR and found that the levels of RT-PCR products covering the entire polycistronic mRNA were decreased (Figure 1D).

The apical part of rnTrpL stem-loop 1 is critically involved in controlling *trpDC* mRNA levels

Previous genetic analyses of the S. meliloti trp attenuator strongly suggested that it is able to form mutually exclusive RNA stem-loop structures: under conditions of efficient trpL translation or SL1 formation in the absence of translation, formation of the antiterminator SL2 is prevented, thus facilitating the formation of the terminator SL3 (23,61). However, there is currently no direct experimental support for the rnTrpL structure. We performed in vitro structural probing to determine experimentally the SL1 and SL3 structures that are expected to be formed in solution (Figure 2A). Based on the probing data, we provide a structure model (Figure 2B) with structural details for the terminator stem-loop. In addition, the model suggests that, with few structural rearrangements involving flexible regions, an alternative (anti-terminator) structure may be formed. According to this model, the sequence predicted to interact with *trpD* (from nt 21 to nt 55, see also Figure 4A) includes the apical region of SL1. In order to investigate the mechanism by which rnTrpL contributes to reducing trpDC mRNA levels, several mutations were introduced in this rn-TrpL region.

The mutated rnTrpL derivatives were constitutively overproduced from pRK-plasmids in strain 2011. Mutations CG40,41GC and GG46,47CC located at or near the apical loop of SL1 (see Figure 2B) were not predicted to drastically change the secondary structure of rnTrpL, but were expected to weaken the predicted base-pairing interaction of SL1 with *trpD* (see Figure 4A). Also, we produced a mutant (AU1,2UA) in which the small ORF trpL (encoding the leader peptide) was eliminated. Using this mutant, we sought to address the question of whether translation of the liberated rnTrpL has any effect on the sRNA-mediated reduction of *trpDC* levels (see above) or whether the resulting 14-aa peptide translated from this sRNA (independently) affects trpDC mRNA levels (translation of the ectopically overproduced wild type (wt) rnTrpL was confirmed, see Supplementary Figure S3). A double mutant (AU1,2UA-G44C), in which the start and the stop codon of *trpL* were mutated, was also constructed. In addition, two mutant rn-TrpL derivatives constructed previously (G44C and G67C,

respectively) (27) were included in our analysis. All mutated rnTrpL derivatives and wt rnTrpL were confirmed to be overexpressed to similar levels (Supplementary Figure S3).

Next, we analyzed the effect of the mutated rnTrpL sR-NAs on the steady-state level of the polycistronic *trpDC* mRNA using qRT-PCR. Figure 2C shows that all genes of the *trpDC* operon were affected in a similar manner. The sRNA rnTrpL-AU1,2UA and wt rnTrpL diminished the mRNA levels to a similar extent, suggesting that (under the conditions used in this experiment) ribosome occupancy of rnTrpL and/or the leader peptide were not important for the rnTrpL-mediated effects on *trpDC* mRNA. In contrast, all mutations that were predicted to weaken the basepairing between rnTrpL and *trpD* (see Figure 4A), either abolished the sRNA-mediated downregulation or even led to increased mRNA levels (see the effect of the CG40,41GC mutation in Figure 2C). Further, rnTrpL-G67C, carrying a mutation in a region confirmed to specifically interact with sinI mRNA (27), had a readily detectable negative effect on *trpC* levels, very similar to that observed for the wt rnTrpL (Figure 2C), suggesting that this mutation did not impede the sRNA's capability of affecting *trpC* levels. Altogether, these results support the idea that base-pairing interactions between the apical part of rnTrpL SL1 and *trpDC* mRNA are involved in mediating the observed decrease in the trpDC mRNA steady-state level upon rnTrpL overproduction

Short-term overproduction of rnTrpL in a $\Delta trpL$ background suggests a direct effect on trpDC

For further investigations in the absence of native rnTrpL RNA being transcribed from the chromosome, we constructed the deletion mutant $2011 \Delta trpL$, in which the original transcription start site (TSS) of trpLE(G) including the first two nucleotides (AT) was preserved (Figure 3A). To avoid secondary effects due to constitutive sRNA overproduction, we constructed a plasmid, pSRKGm-rnTrpL, for IPTG-inducible transcription of recombinant rnTrpL. In this plasmid, the trpL ORF of rnTrpL was fused in frame to a 38-nt *lacZ* mRNA leader harboring a ribosome-binding site, and transcription was terminated at SL3 of rnTrpL (Figure 3B). Figure 3C shows the induction of this rnTrpL (pSRKGm-rnTrpL).

We also considered the possibility that the sRNA function of lacZ'-rnTrpL may be impaired due to an increased ribosome load compared to the leaderless rnTrpL transcribed from its native locus on the chromosome (23) or from pRK-rnTrpL (see Figure 1B and Supplementary Figure S3). However, we found that the level of *trpDC* mRNA was decreased in strain 2011 $\Delta trpL$ (pSRKGm-rnTrpL) already 10 min post induction, although the lacZ'-rnTrpL level was much lower than the rnTrpL level in the parental strain 2011 (Figure 3C and D). At 20 min post induction, the effect of lacZ'-rnTrpL was similar to the effect of constitutively overproduced rnTrpL. The induction of lacZ'rnTrpL production had no detectable effect on the control mRNA trpE(G) (Figure 3D). This specific, short-term effect on *trpDC* provides additional support to the idea that trpDC mRNA is a direct target of rnTrpL.

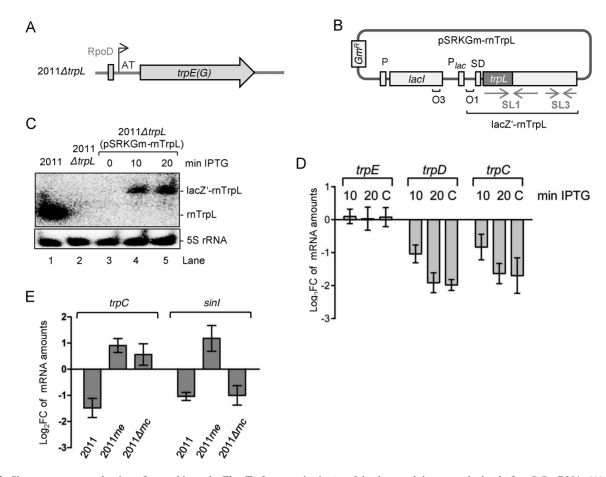


Figure 3. Short-term overproduction of recombinant lacZ'-rnTrpL transcript in $\Delta trpL$ background decreases the level of trpDC mRNA. (A) Schematic representation of the trpE(G) locus in the deletion mutant $2011\Delta trpL$. The original TSS including the first two nucleotides (AT) was preserved. (B) Scheme of pSRKGm-rnTrpL. The pSRK-based plasmids harbor the *lac* repressor gene *lac1* with its own promoter, the *lac1-lacZ* intergenic region, the *lac*-promoter P_{lac} , and the 38-nt *lacZ* leader containing the Shine-Dalgarno sequence (SD), followed by an NdeI restriction site containing the ATG translation start codon. Positions of the LacI-binding operators O1 and O3, the proper spacing of which ensures tight regulation at P_{lac} , are indicated (42). The rnTrpL sequence was cloned in frame in the NdeI site of pSRKGm. The resulting sRNA lacZ'-rnTrpL contains the 38-nt *lacZ*-leader. The scheme shows the plasmid conferring resistance to gentamycin (Gm) (pSRKGm-rnTrpL). A similar plasmid conferring resistance to tetracycline (Tc) was also constructed. (C) Northern blot hybridization showing IPTG-induced lacZ'-rnTrpL transcription at 10 and 20 min post induction in strain 2011 $\Delta trpL$ (pSRKGm-rnTrpL) (lanes 3, 4 and 5). Data for control strains are shown in lanes 1 and 2. Also shown are the hybridization data using a 5S rRNA-specific probe (loading control). (D) qRT-PCR analysis of trpD, trpC and trpE (control) expression. At 10 min and 20 min post induction with IPTG, mRNA levels of the respective mRNA in 2011 $\Delta trpL$ (pSRKGm-rnTrpL) compared to those of the EVC. (E) qRT-PCR analysis of trpD, trpC and $2011 \Delta trpL$ (mRNA strain 2011 (pRK-rnTrpL) compared to those of the EVC. (E) qRT-PCR analysis of trpC and the control mRNA *sinI* in strain 2011 and the RNase mutant strains 2011*rne* and 2011 $\Delta trpC$ (indicated). Each strain contains the plasmid pSRKTc-rnTrpL. Changes in the levels of the respective mRNA were determined 10 min post induction with IPTG. The control mRNA *sinI* in strain 2011

To address the role of ribonucleases in the rnTrpLmediated decrease of *trpDC* mRNA levels, the lacZ'-rnTrpL production was induced for 10 min in the RNase mutants 2011rne (27,62) and 2011 Δ rnc (63) using plasmid pSRKTcrnTrpL, and changes in the levels of trpC mRNA were analyzed. In contrast to the parental strain 2011, in which trpC was confirmed to be decreased, the trpC levels were increased in both mutants (Figure 3E). As a control, we also analyzed the changes in sinI (Figure 3E). As expected, the sinI mRNA level was decreased upon rnTrpL induction in strain 2011 (27). A similar decrease was observed in the RNase III mutant 2011 Arnc, while an increase was detected in the RNase E mutant 2011rne (Figure 3E). The latter result is in line with the RNase E-dependent downregulation of sinI by rnTrpL (27). Together, these results support the importance of RNase E for posttranscriptional gene regulation by rnTrpL in *trans*, and suggest that RNase III may also participate in the regulation of *trpDC*, although it is dispensable for the regulation of *sinI* by rnTrpL.

sRNA rnTrpL base-pairs with *trpD* to destabilize the polycistronic *trpDC* mRNA

The predicted binding of rnTrpL to a region in the *trpD* coding sequence is shown in Figure 4A. To provide experimental evidence for the predicted base-pairing interaction, we performed *in vivo* assays in strain $2011 \Delta trpL$ using bicistronic *trpDC'::egfp* reporter constructs and lacZ'rnTrpL derivatives. The *trpDC'::egfp* fusions (Figure 4B) were expressed from pSRKGm (conferring Gm^r) and challenged with wt or mutated lacZ'rnTrpL transcribed from pSRKTc (conferring Tc^r) (see Figure 3B). To avoid long-

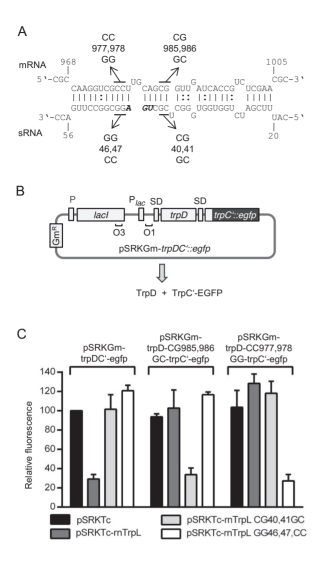


Figure 4. The attenuator sRNA rnTrpL base-pairs with trpD to downregulate trpDC expression. (A) Scheme of the duplex structure predicted to be formed between *trpD* (mRNA) and rnTrpL (sRNA) ($\Delta G = -13.51$ kcal/mol). rnTrpL mutations characterized in this experiment are given below the sRNA sequence. The *trpL* stop codon is shown in bold and italics. Compensatory *trpD* mutations used to restore base-pairing interactions (see C) are shown above the mRNA sequence. Nucleotide numbering starts at the translation start codon of the trpD mRNA and the sRNA harboring the trpL sORF, respectively. (B) Scheme of pSRKGm-trpDC'-egfp. The tr*pDC*[:]::*egfp* fusion contains the first 16 *trpC* codons fused to the third *egfp* codon. For more details, see Figure 3. (C) Analysis of possible base-pairing interactions between lacZ'-rnTrpL and the fusion mRNA trpDC'::egfp in strain $2011 \Delta trpL$. Plasmids used in this experiment are indicated. Fluorescence was measured at 20 min after induction with IPTG; the fluorescence obtained for strain 2011 \(\Delta trpL(pSRKGm-trpDC'-egfp, pSRKTc))\) was set to 100% and used for normalization. Shown are the results from three independent experiments, each performed in duplicates (means and standard deviations are indicated).

term effects, fluorescence was measured at 20 min after simultaneous induction of transcription of the reporter fusion construct and the sRNA.

Figure 4C shows that TrpC'-EGFP fluorescence derived from plasmid pSRKGm-trpDC'-egfp was strongly decreased if lacZ'-rnTrpL was coexpressed, supporting the idea that the sRNA binds to *trpD* and thereby induces a reduction of *trpDC*'::*egfp* mRNA levels. In contrast, if sRNA derivatives carrying CG40.41GC and CC46.47GG mutations, respectively, were co-expressed, no decrease in fluorescence was observed. To test whether this was due to a significantly reduced (or lack of) binding of the mutated sRNAs to *trpD* in the bicistronic reporter mRNA, appropriate mutations were introduced into the *trpD* binding site to restore the presumed base-pairing interactions (see Figure 4A). Indeed, a decrease in fluorescence produced by the pSRKGm-trpD-CG985,986GC-trpC'-egfp construct was (only) observed if the corresponding basepairing sRNA lacZ'-rnTrpL-CG40,41GC was coexpressed. Consistent with this, the fluorescence produced by the pSRKGm-trpD-CC977,978GG-trpC'-egfp construct was (only) decreased if lacZ'-rnTrpL-GG46,47CC sRNA was coexpressed (Figure 4C). These results validate the basepairing between rnTrpL and *trpD* and show that this interaction is responsible for the negative effect of rnTrpL on trpDC.

The negative effect of the sRNA rnTrpL on *trpDC* (a decrease in the steady-state level of the mRNA) can be explained by a destabilization of *trpDC* mRNA. To test whether rnTrpL affects the decay of this mRNA, halflife measurements were performed with strain $2011 \Delta trpL$ (pSRKTc-rnTrpL). The stabilities of the polycistronic trpDC mRNA and the control mRNA trpE were measured without IPTG addition and 10 min post induction of lacZ'rnTrpL overproduction. Figure 5 shows that, with primers specific for trpD and trpC, respectively, a decrease in the mRNA half-life upon rnTrpL induction was detectable, while the half-life of *trpE* mRNA remained unchanged. Similarly, mRNA destabilization upon rnTrpL induction was detected with primers specific for ppiD, moaC and moeA (Supplementary Figure S4). The ppiD- and trpD-specific primers bind upstream of the rnTrpL-binding site in the polycistronic mRNA, while the trpC-, moaC- and moeAspecific primers target downstream regions. Shorter halflives of all transcript parts upon rnTrpL overproduction is consistent with the proposed destabilization of the polycistronic *trpDC* mRNA after base-pairing between rnTrpL and *trpD*.

sRNA rnTrpL down-regulates *trpDC* expression under Trp sufficiency conditions

Finally, we addressed the question of whether the attenuator sRNA rnTrpL is involved in trpDC regulation in response to changes in cellular Trp availability. To assess transcription attenuation between trpL and trpE(G), we constructed a $2011 \Delta trpC$ mutant to prevent cellular Trp production and used minimal medium with defined Trp concentrations. According to previous data (23), during growth in a medium supplemented with 20 µg/ml Trp (Trp sufficiency condition), transcription is regularly terminated between *trpL* and *trpEG*, and the sRNA rnTrpL is liberated. In contrast, during growth at 2 µg/ml Trp (Trp insufficiency), transcription termination is abolished and trpL is cotranscribed with trpE(G). Indeed, Figure 6A shows that the sRNA rnTrpL was readily detectable by Northern blot analysis when $2011 \Delta trpC$ was grown at Trp sufficiency. Under these conditions, readthrough transcripts were present

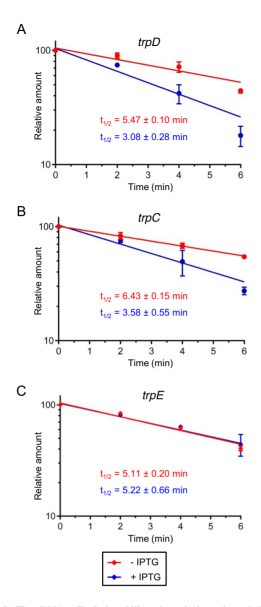


Figure 5. The sRNA rnTrpL destabilizes the polycistronic *trpDC* mRNA. At an OD_{600nm} of 0.5, one half of a liquid culture of strain 2011 $\Delta trpL$ (pSRKTc-rnTrpL) was supplemented with IPTG for 10 min to induce lacZ'-rnTrpL production, while the other half was incubated without IPTG. Then, rifampicin was added to both cultures and RNA was isolated at time points 0, 2, 4 and 6 min after rifampicin addition. Relative mRNA levels in induced (blue) and uninduced cultures (red) were determined by qRT-PCR analysis and half-lives ($t_{1/2}$, in min) were calculated from three independent experiments, each with two technical replicates. The analysis was performed with *ppiD-*, *moaC-* and *moeA-*specific primers is shown in Supplementary Figure S4.

at a very low level (Supplementary Figure S1). Large trpLE(G) transcripts could not be detected because the protocol used in this experiment was optimized for efficient separation and blotting of small RNAs. When the cells were washed in medium without Trp and subsequently grown for 4 h in medium supplemented with 2 µg/ml Trp, the level of rnTrpL was steadily decreased. 10 min after restoring Trp sufficiency, the sRNA rnTrpL level was increased as expected (Figure 6A). Under these conditions, accumulation of a small degradation product of the trpLE(G) transcript was detected (see also Supplementary Figure S5).

To analyze the role of rnTrpL in *trans* in the context of transcription attenuation, we additionally constructed a control strain, the double mutant $2011 \Delta trp C \Delta trp L$. Strains $2011 \Delta trpC$ and $2011 \Delta trpC \Delta trpL$ were used to detect changes in the levels of the polycistronic *ppiD-trpD*- $\Delta trpC-moaC-moeA$ mRNA and trpE(G) mRNA in response to changes in cellular Trp availability. The experimental setup was as described above (transfer from minimal medium containing 20 µg/ml Trp to medium containing 2 $\mu g/\mu l$ Trp, incubation for 4 h with sample collection at onehour intervals and, at the end, incubation with 20 μ g/ml Trp for another 10 min). Upon Trp insufficiency, when the level of the released sRNA rnTrpL was reduced, the level of the polycistronic *ppiD-trpD-ΔtrpC-moaC-moeA* mRNA was found to be increased in strain $2011 \Delta trpC$ (Figure 6B and C). Importantly, 10 min after restoring Trp sufficiency, when the level of the sRNA rnTrpL was increased again, the level of the polycistronic mRNA was decreased (Figure 6B and C). Similar changes were observed for *trpE* (Figure 6B). Notably, such changes were not observed in the double mutant $2011 \Delta trp C \Delta trp L$ (Figure 6B and C), suggesting a posttranscriptional downregulation of the *trpDC* operon by the liberated sRNA rnTrpL in response to Trp availability. In addition, these data lead us to conclude that, similarly to trpE(G) (23), the mRNA level of the trpDC operon is not regulated at the level of transcription initiation in response to Trp availability, but posttranscriptionally by rnTrpL.

The *trpFBA* operon is not regulated by the sRNA rnTrpL

In contrast to the *trpDC* operon, *trpFBA* was not predicted to interact with rnTrpL in a previous analysis of putative conserved targets of this sRNA (27). Here, we analyzed *trpB* mRNA and found that its level was not changed 10 min post induction of lacZ'-rnTrpL in strain $2011\Delta trpL$ (log₂FC = 0.12 ± 0.29), suggesting that *trpFBA* mRNA is not a target of rnTrpL.

Next, we asked the question of whether the expression of the *trpFBA* operon is regulated in response to Trp availability. A qRT-PCR analysis of trpF, trpB and trpA was performed with strains $2011 \Delta trpC$ and $2011 \Delta trpC \Delta trpL$ grown in minimal medium containing different Trp concentrations as described above. Growth of strain $2011 \Delta trpC$ in medium containing 2 μ g/ml Trp for 4 h led to increasing trpFBA mRNA levels while, at 10 min after addition of 20 µg/ml Trp to this Trp-starved culture, the level of trpFBA was decreased (Figure 6D). These changes show that trpB is regulated in response to Trp availability. However, in contrast to what was observed for the trpE(G) and *trpDC* operons (Figure 6B and C), similar *trpFBA* changes were also detected in the double mutant $2011 \Delta trp C \Delta trp L$. The *trpF*, *trpB* and *trpA* values obtained for the two strains $2011 \Delta trpC$ and $2011 \Delta trpC \Delta trpL$ were not statistically different (Figure 6D), indicating that regulation of the trpFBA operon in response to Trp availability does not depend on the attenuator sRNA rnTrpL.

In summary, Figure 6 shows that all three *trp* operons are regulated in response to Trp availability. However, while the regulation of the mRNA levels of the two operons trpE(G)

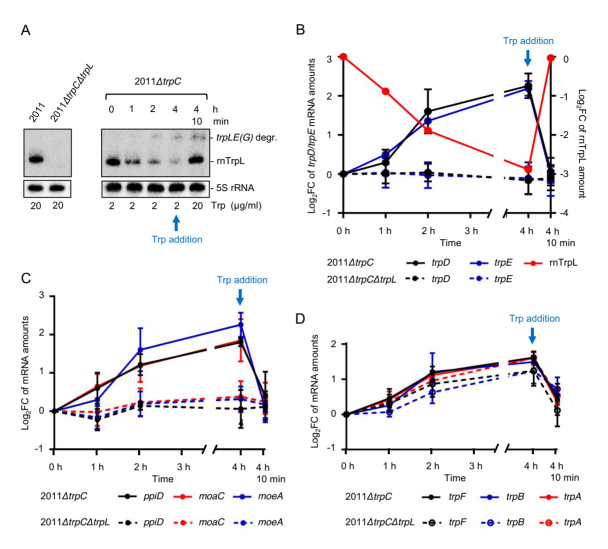


Figure 6. The sRNA rnTrpL down-regulates trpDC in response to Trp availability. (A) Northern blot analysis of rnTrpL, showing transcription attenuation in strain 2011 $\Delta trpC$ which was grown first in minimal medium (MM) containing 20 µg/ml Trp and then transferred to MM containing 2 µg/ml Trp for 0, 1, 2 and 4 h, respectively. Depletion of the cellular Trp pool causes cotranscription of trpL and trpE(G) and a decrease in the level of the sRNA rnTrpL. 4 h 10 min, following incubation with 2 µg/ml Trp for 4 h, the Trp concentration was increased to 20 µg/ml for 10 min. Increased Trp availability led to transcription termination at SL3 of rnTrpL and an immediate increase in the level of the sRNA rnTrpL. The band migrating slower than rnTrpL was identified as a degradation product (degr.) of the trpLE(G) cotranscript (see Supplementary Figure S5). On the left, RNA isolated from the indicated control strains grown in media with 20 µg/ml Trp was loaded. All lanes shown in this panel originate from the same blot. (B) Changes in trpE, trpD and rnTrpL RNA levels in response to Trp availability in the indicated strains. Relative trpD and trpE mRNA levels were measured by qRT-PCR using RNA samples from Trp availability experiments (as shown in panel A). The levels measured at the indicated strains. For details, see B). All mRNA graphs show means and standard deviations from three independent experiments, each performed in duplicates. The rnTrpL graph in B) shows a quantification of the signals from panel A).

and *trpDC* was revealed to critically involve rnTrpL, the regulation of *trpFBA* proved to be independent of rnTrpL.

Conservation of the *trans*-regulating function of rnTrpL on the expression of *trp* genes

Many Alphaproteobacteria have three *trp* operons with gene orders that are identical or very similar to that seen in *S. meliloti* (3). Among them are the plant pathogen *A. tume-faciens* and the soybean symbiont *B. japonicum*. In the three alphaproteobacterial species *S. meliloti*, *A. tumefaciens* and *B. japonicum*, transcription of the attenuator sRNA (and of the bicistronic transcript trpLE(G)) starts with the ATG codon of trpL (64–66) (see Supplementary Figure S6). To

address the question of whether *trp* genes are regulated in *trans* by the rnTrpL homolog in these species, the sequences corresponding to Atu-rnTrpL and Bj-rnTrpL, respectively, were cloned in pSRKTc and pRJ-MCS, respectively. The effect of sRNA overproduction in the corresponding homologous host on the mRNA levels of *trpD* and *trpB* were analyzed by qRT-PCR. We found decreased *trpD* mRNA levels and no change in the levels of *trpB* upon overexpression of the attenuator sRNA in both *A. tumefaciens* and *B. japonicum* (Figure 7A). This indicates that, similarly to what was observed in *S. meliloti, trpDC* is down-regulated by the attenuator sRNA in *trans*, while *trpFBA* is not regulated by this sRNA. To predict base-pairing interactions between *A*.

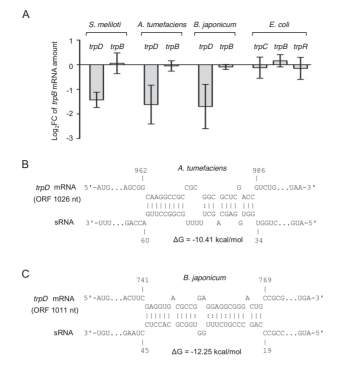


Figure 7. Conservation of *trpDC* regulation by rnTrpL in *trans*. (A) qRT-PCR analysis of the indicated mRNAs and bacterial species. For *S. meliloti* (pSRKTc-rnTrpL), *A. tumefaciens* (pSRKTc-Atu-rnTrpL) and *E. coli* (pSRKTc-Ec-rnTrpL), changes in the levels of the indicated mRNAs were analyzed 10 min post induction. In the constitutively overexpressing *B. japonicum* (pRJ-Bj-rnTrpL), the levels of *trpD* and *trpB* were compared to the EVC. (B) Schematic representations of the duplex structures predicted to be formed between *trpDC* and rnTrpL homologs in *A. tumefaciens* and *B. japonicum*. The numbering starts at the translation start codons of *trpD* mRNA and the *trpL* ORF in the sRNAs, respectively. Predicted secondary structures of the sRNAs Atu-rnTrpL and Bj-rnTrpL are shown in Supplementary Figures S7 and S8.

tumefaciens trpDC and Atu-rnTrpL, as well as between *B. japonicum trpDC* and Bj-rnTrpL, we used IntaRNA (58). Potential interactions were predicted between SL1 of the respective attenuator RNA and the 3'-region of the respective *trpD* (Figure 7B; see also Supplementary Figures S7 and S8). Altogether, these results suggest that *trpDC* regulation by rnTrpL is conserved in the families *Rhizobiaceae* and *Bradyrhizobiaceae*, to which the studied Alphaproteobacteria belong.

To address the conservation of rnTrpL function beyond Alphaproteobacteria, we constructed plasmid pSRKTc-EcrnTrpL and tested whether overproduction of the attenuator RNA Ec-rnTrpL for 10 min affects the level of trpC, trpB and of the separately transcribed Trp repressor mRNA trpR (3) in *E. coli*. The levels of trpC, trpB and trpR were not changed significantly upon Ec-rnTrpL overproduction (Figure 7A), suggesting that this sRNA does not regulate trp genes in *trans*.

In summary, our data show that in bacteria with *trp* genes organized in several operons, the attenuator sRNA rnTrpL acts in *trans* to regulate the *trpDC* operon posttranscriptionally (Figure 8).

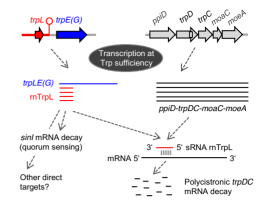


Figure 8. Model of gene regulation in *trans* by the attenuator sRNA rn-TrpL in *S. meliloti*. At TrpL sufficiency, transcription is generally terminated between *trpL* and *trpE(G)*. The liberated attenuator sRNA rn-TrpL binds to *trpD* and destabilizes the polycistronic mRNA *ppiD-trpDCmoaC-moeA*. In this way, the *trp* attenuator posttranscriptionally coordinates the expression of the two *trp* operons trpE(G) and trpDC in response to Trp availability. In addition, the destabilization of the polycistronic *trpDC* mRNA by rnTrpL probably affects protein translocation through the outer membrane (*ppiD*) and MoCo biosynthesis (*moaC-moeA*). Moreover, the sRNA rnTrpL binds to and destabilizes *sinI* (27), thus probably linking quorum sensing to Trp availability.

DISCUSSION

In this study, we demonstrate that a bacterial sRNA derived by ribosome-dependent transcription attenuation acts by base-pairing in *trans*. According to our data, the liberated attenuator RNA of the trpE(G) operon down-regulates the expression of the trpDC operon in *S. meliloti* and other bacteria.

The gene trpE(G) encodes anthranilate synthase that catalyzes the first step of Trp biosynthesis starting with chorismate, while trpD and trpC encode anthranilate phosphoribosyltransferase and indole-3-glycerol phosphate synthase that catalyze the two subsequent reactions. The remaining enzymes of the pathway, phosphoribosylanthranilate isomerase and Trp synthase are encoded by the trpFBA operon. Our data show that ectopic oveproduction of rn-TrpL does not influence trpFBA but decreases the steadystate level of the polycistronic trpDC mRNA, in line with the predicted base-pairing between rnTrpL and trpD. Consistent with a direct effect of rnTrpL on *trpDC* mRNA, a decrease of the *trpC* level was detected as early as 10 min after induction of lacZ'-rnTrpL in the $\Delta trpL$ background (Figure 3). The results shown in Figure 4C and Figure 5 provide strong support for this hypothesis. Thus, for example, a base complementarity between specific rnTrpL and *trpD* sequences proved to be required for *trpDC* downregulation (Figure 4C). Furthermore, parts of the polycistronic mRNA located upstream (in ppiD and trpD) and downstream (in trpC, moaC and moeA) of the rnTrpL binding site were destabilized upon lacZ'-rnTrpL induction (Figure 5 and Supplementary Figure S4). The latter result shows that rnTrpL acts at the level of trpDC mRNA (rather than DNA, for example, by disturbing transcription). Based on the data presented in this study, we conclude that the attenuator sRNA rnTrpL base-pairs with the distal part of the trpD coding region in the polycistronic trpDC mRNA and

thereby promotes degradation of the entire transcript (see the model in Figure 8).

Upon sRNA binding to an mRNA in gram-negative bacteria, an endoribonuclease, such as RNase III or RNase E, is recruited. After an initial endonucleolytic cleavage, RNA fragments are degraded by (other) exo- and endoribonucleases, leading to rapid decay of both the sRNA and the mRNA. Alternatively, inhibition of translation by the sRNA results in unprotected mRNA regions that are endonucleolytically cleaved, mostly by RNase E (9,67), a mechanism suggested for the action of rnTrpL on sinI mRNA (27). According to Figure 3E, both RNase E and RNase III are important for the rnTrpL-mediated downregulation of *trpDC*. Future analyses are needed to elucidate whether these RNases directly participate in the destabilization of the *trpDC* message by rnTrpL. Further, according to Figures 2 and 4, the apical region of SL1 of rnTrpL is critically involved in the base-pairing interaction with *trpD*, suggesting that rnTrpL may need an RNA chaperone for efficient binding to this target. Since rnTrpL was described as an Hfg-independent sRNA (27), and since S. meliloti does not harbor ProQ (13), another RNA-binding protein that interacts with rnTrpL may be identified in the future.

The physiological role of rnTrpL as an attenuator RNA in cis and base-pairing sRNA in trans is an intriguing example for the evolution of bacterial operons and their co-regulation. Although bacteria belonging to Escherichia, Salmonella, Vibrio and most members of Bacillales and Lactobacillales have all the trp genes in a single operon, most other bacteria (Alpha-, Beta-, Gamma-, Delta-, and Epsilonproteobacteria, Spirochaeta and most Actinobacteriadae) carry trp genes that are organized into several operons. It has been proposed that this allows for more flexibility in the regulation of the genes involved in each step of Trp synthesis since, in different bacteria, specific Trp precursors have additional functions and are linked to different metabolic pathways (3). However, as already mentioned in the introduction, the high costs of Trp synthesis could pose a selection pressure for evolving RNA-based mechanisms suitable to mediate an appropriate response to varying Trp concentrations in the cell. This is in an apparent contradiction to the observation that, in bacteria with split *trp* operons, only one of them (usually *trpE*) is equipped with an mRNA leader that mediates transcription attenuation. Our finding that, in Sinorhizobium, Agrobacterium and *Bradyrhizobium*, rnTrpL is capable to regulate *trpDC* in trans provides an example of how bacteria evolved posttranscriptional strategies to coordinate the expression of trp operons according to Trp availability (Figure 8). In contrast, we did not detect posttranscriptional regulation by ectopically overexpressed Ec-rnTrpL in E. coli where all the *trp* biosynthesis genes are co-transcribed (Figure 7A).

Our data show that the sRNA rnTrpL can downregulate the entire *trpDC* operon including *ppiD*, *moaC* and *moeA* (Figure 1, Figure 2, Figure 5 and Supplementary Figure S4) The gene *ppiD* encodes peptidyl-prolyl isomerase required for folding of outer membrane proteins/protein translocation (68,69), while *moaC* and *moeA* encode enzymes required for the first and the last step in the biosynthesis of the molibdenum cofactor MoCo (70). This pterin-based cofactor is the active compound present in the catalytic site of all molybdenum-containing enzymes except for the bacterial molybdenum nitrogenase (71). The MoCo biosynthesis is highly conserved in all domains of life and enzymes requiring this cofactor are necessary for carbon, nitrogen, and sulfur metabolism (72). Because the trpLE(G) and ppiD-trpDC-moaC-moeA operons are conserved in many Alphaproteobacteria and because rnTrpL overexpression downregulates trpD in A. tumefaciens and B. japonicum, we propose that the specific regulatory function of rnTrpL revealed in this study is widely conserved. It is tempting to suggest that, in bacteria with trp genes organized in several operons, the Trp sufficiency status may influence not only the cellular Trp biosynthesis but also other functions, such as protein translocation through the outer membrane and MoCo-dependent metabolic processes.

It is also noteworthy that, in S. meliloti, rnTrpL can downregulate the autoinducer synthase gene sinI (27), thereby probably connecting Trp availability to quorum sensing. According to our data, the sRNA rnTrpL is available for regulation in *trans* under conditions of Trp sufficiency. Under such conditions, destabilization of sinI mRNA by rnTrpL may impede the autoinducer accumulation and, as a consequence, quorum sensing may begin at higher population density than under conditions of Trp insufficiency. In line with this, it was recently shown that quorum sensing begins at higher population density, when S. meliloti cultures are supplied with water-soluble humic materials as additional nutrient sources (73). Indeed, guorum sensing is known to be influenced by environmental factors in addition to the population density (74). Previously, it was shown that, during growth in TY medium, the steady-state level of rnTrpL is influenced by temperature and salt (27). The effects of these factors on rnTrpL generation and/or decay as well as the regulation of the rnTrpL targets remain to be elucidated.

It was predicted that, in addition to *sinI* and *trpD*, mR-NAs of several regulatory genes can base-pair with rn-TrpL (27). Thus, it is possible that this attenuator sRNA is at the center of a regulatory network that responds to Trp availability as a marker indicating the nutrient status of the cell. Under conditions of Trp sufficiency, the signal (charged tRNA-Trp) is the critical regulator of ribosome-mediated transcription attenuation, while the liberated attenuator sRNA may get involved in regulatory functions in *trans* by base-pairing with specific mRNAs in addition to *trpD* (Figure 8).

As one of the rare examples of 5'-UTR-derived sRNAs, the *trans*-acting attenuator RNA rnTrpL characterized in this study highlights the regulatory potential and the versatility of bacterial mRNA leaders.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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