Mechanistic comparison of *Bacillus subtilis* 6S-1 and 6S-2 RNAs—commonalities and differences

OLGA Y. BURENINA, PHILIPP G. HOCH, KATRIN DAMM, MARGARITA SALAS, TIMOFEI S. ZATSEPIN, MARCUS LECHNER, TATIANA S. ORETSKAYA, ELENA A. KUBAREVA, and ROLAND K. HARTMANN

1Chemistry Department and A.N. Belozersky Institute of Physico-Chemical Biology, M.V. Lomonosov Moscow State University, 119991 Moscow, Russia
2Institut für Pharmazeutische Chemie, Philipps-Universität Marburg, 35037 Marburg, Germany
3Centro de Biología Molecular “Severo Ochoa” (CSIC-UAM), Universidad Autónoma de Madrid, Cantoblanco, 28049 Madrid, Spain

ABSTRACT

Bacterial 6S RNAs bind to the housekeeping RNA polymerase (σA-RNAP in *Bacillus subtilis*) to regulate transcription in a growth phase-dependent manner. *B. subtilis* expresses two 6S RNAs, 6S-1 and 6S-2 RNA, with different expression profiles. We show in vitro that 6S-2 RNA shares hallmark features with 6S-1 RNA: Both (1) are able to serve as templates for pRNA transcription; (2) bind with comparable affinity to σA-RNAP; (3) are able to specifically inhibit transcription from DNA promoters, and (4) can form stable 6S RNA:pRNA hybrid structures that (5) abolish binding to σA-RNAP. However, pRNAs of equal length dissociate faster from 6S-2 than 6S-1 RNA, owing to the higher A,U-content of 6S-2 pRNAs. This could have two mechanistic implications: (1) Short 6S-2 pRNAs (<10 nt) dissociate faster instead of being elongated to longer pRNAs, which could make it more difficult for 6S-2 RNA-stalled RNAP molecules to escape from the sequestration; and (2) relative to 6S-1 RNA, 6S-2 pRNAs of equal length will dissociate more rapidly from 6S-2 RNA after RNAP release, which could affect pRNA turnover or the kinetics of 6S-2 RNA binding to a new RNAP molecule. As 6S-2 pRNAs have not yet been detected in vivo, we considered that cellular RNAP release from 6S-2 RNA might occur via 6S-1 RNA displacing 6S-2 RNA from the enzyme, either in the absence of pRNA transcription or upon synthesis of very short 6S-2 pRNAs (∼5-mers, which would escape detection by deep sequencing). However, binding competition experiments argued against these possibilities.

Keywords: 6S-1 RNA; *bsrA*; 6S-2 RNA; *bsrB*; pRNA transcripts; affinity for σA-RNAP; 6S RNA:pRNA hybrid stability; 6S-2 RNA release from RNAP

INTRODUCTION

6S RNA is a bacteria-specific noncoding RNA (ncRNA) that primarily binds to RNA polymerase (RNAP) holoenzymes containing the housekeeping σ factor, σ70 in *Escherichia coli* and σA in *Bacillus subtilis* (Wassarman and Storz 2000; Willkomm and Hartmann 2005; Wassarman 2007). Depending on the particular expression profile of a 6S RNA, the RNA sequesters a substantial fraction of the housekeeping RNAP holoenzyme, which then becomes unavailable for transcriptional activity at DNA promoters (Wassarman 2007). This block can be relieved when RNAP synthesizes so-called product RNAs (pRNAs) of a certain length utilizing 6S RNA itself as a transcription template, which then triggers the release of RNAP (Wassarman and Saecker 2006; Gildehaus et al. 2007; Beckmann et al. 2011).

Unlike *E. coli* and many other bacteria encoding a single 6S RNA, the *B. subtilis* genome encodes two 6S RNA homologs. At their discovery, the genes for the two ncRNAs were termed *bsrA* and *bsrB* (Ando et al. 2002; Suzuma et al. 2002), but their nature as 6S RNAs was only unveiled in a later study by the Wassarman group (Trotochaud and Wassarman 2005). Both RNAs were shown to coimmunoprecipitate with the σA housekeeping RNAP holoenzyme using antibodies against σA or the core subunit α (Trotochaud and Wassarman 2005). Recent investigations provided evidence that 6S-1 RNA displays all of the hallmarks of a canonical 6S RNA: (1) It reaches its highest steady-state levels during late exponential/early stationary phase (Barrick et al. 2005; Beckmann et al. 2011); (2) it serves as an RNA template for massive pRNA transcription, particularly during outgrowth from stationary phase upon resupply of nutrients, as inferred from
Mechanistic comparison of 6S-1 and 6S-2 RNAs

Deep sequencing (dRNA-seq) (Beckmann et al. 2011) and Northern blot analyses (Beckmann et al. 2011; Cavanagh et al. 2012); and (3) longer pRNAs (12- to 14-mers), whose fraction increases during outgrowth from stationary phase (Beckmann et al. 2011), form a stable duplex with 6S-1 RNA and induce a structural rearrangement in 6S-1 RNA. The rearranged 6S-1 RNA:pRNA complexes lose their affinity for σA-RNAP to liberate σA-RNAP for transcription at DNA promoters during the newly initiated exponential phase. 6S-1 RNA levels then decrease upon entering a new exponential growth phase and increase again when nutrients become scarce and cells enter another stationary growth phase (Beckmann et al. 2012).

The function of *B. subtilis* 6S-2 RNA is currently a matter of debate. The RNA was observed to reach high levels in the early exponential phase and to decrease toward stationary phase (Ando et al. 2002; Barrick et al. 2005; Beckmann et al. 2011). Deviating from these observations, *B. subtilis* 6S-2 RNA levels were also reported to remain fairly constant in exponential and stationary phase (Trotochaud and Wassarman 2005; Cavanagh et al. 2012).

We have demonstrated in vitro that the σA-RNAP holoenzyme is able to synthesize pRNAs on 6S-2 RNA as a template (Beckmann et al. 2011). So far, deep sequencing analyses have not provided evidence for the synthesis of 6S-2 RNA-derived pRNAs in vivo (Innov et al. 2010; Beckmann et al. 2011). We hypothesized that 6S-2 pRNAs may be rapidly degraded in vivo or substantial 6S-2 RNA synthesis may occur during yet-to-be-defined adaptations to certain environmental or stress conditions (Beckmann et al. 2011). It was recently shown that pRNA synthesis by σA-RNAP is most efficient with GTP as the initiating nucleotide (Cabrera-Ostertag et al. 2013), in line with the finding that a *B. subtilis* strain expressing a chromosomal 6S-1 RNA mutant initiating pRNA transcription with ATP instead of GTP showed reduced pRNA synthesis in *B. subtilis* (Beckmann et al. 2011). Since 6S-2 pRNAs are initiated with ATP (Beckmann et al. 2011) and 6S-1 pRNAs with GTP, it has been posited that 6S-2 RNA may not direct pRNA synthesis under physiological conditions (Cabrera-Ostertag et al. 2013), raising the question of how the sequestration of RNAP by 6S-2 RNA can be overcome.

Here, we investigated the mechanistic properties of *B. subtilis* 6S-2 RNA in direct comparison with those of 6S-1 RNA. In vitro at a concentration of 200 µM each NTP, the most abundant pRNA transcripts synthesized by σA-RNAP are 13- to 16-mers in the case of 6S-2 RNA and 14-mers with 6S-1 RNA. Since competition experiments between DNA promoters and 6S RNA have only been done for *E. coli* (Wassarman and Saecher 2006; Gildehaus et al. 2007) and not for *B. subtilis*, we have performed first experiments in this direction. We show that both 6S RNAs bind to σA-RNAP with very similar affinity, and both RNAs are able to inhibit transcription at model DNA promoters in vitro. Transcription by σA-RNAP on 6S-2 RNA as a template results in 6S-2:pRNA hybrids. Their formation reduces the amount of complex with RNAP, but this reduction appears to be less efficient for 6S-2 RNA relative to 6S-1 RNA. An explanation is that pRNAs of equal length dissociate faster from 6S-2 than 6S-1 RNA, thus favoring rebinding of 6S-2 RNA to RNAP within the time frame of the applied experimental conditions. Annealing of longer synthetic 6S-2 pRNAs (20 nt) reduced the rate of pRNA dissociation to a negligible extent, comparable to the stability of 6S-1 RNA:pRNA 14-mer complexes. We also demonstrate that 6S-2 RNA with a bound pRNA is unable to bind to σA-RNAP. Overall, pRNA synthesis from 6S-2 RNA is more sensitive to lowering the NTP concentration than that from 6S-1 RNA as a template, in line with a previous study (Cabrera-Ostertag et al. 2013). We further tested the possibility that 6S-1 RNA may be able to displace 6S-2 RNA from its complex with σA-RNAP.

**RESULTS**

**6S-1 and 6S-2 RNA structures**

The solution 2D structure of *B. subtilis* 6S-1 RNA has been determined with good confidence by probing experiments (Beckmann et al. 2012). The solution structure of 6S-2 RNA is less clear and RNAfold or mfold predictions for 6S-2 RNA are, in contrast to 6S-1 RNA, highly ambiguous (see Supplemental Fig. S1). To improve the confidence of 6S-2 RNA structure prediction, we calculated a consensus secondary structure based on a sequence/structure alignment of closely related homologs to *B. subtilis* 6S-2 RNA from other bacteria of the Firmicutes branch found in the RFAM database. For comparison, the same was done for 6S-1 RNA (Fig. 1A). The predicted 6S-2 RNA consensus secondary structure (Fig. 1B) has all of the hallmarks of a typical 6S RNA (Barrick et al. 2005; Willkomm and Hartmann 2005). Likewise, the in vitro-determined start site for pRNA transcription on 6S-2 RNA as a template (Beckmann et al. 2011) is located in the “central bulge” region, as for 6S-1 RNA (Fig. 1A,B). According to the 6S-2 RNA consensus prediction, the central bulge region closes with three consecutive G-C bp (nt 55–57/124–126) (Fig. 1B). When those 3 bp are constrained in the individual folding prediction for *B. subtilis* 6S-2 RNA, the structure shown in Supplemental Figure S1C is calculated as the minimum free energy structure (MFE) by RNAfold or mfold (the free energy being 7% less optimal than that of the unconstrained MFE structure) (Supplemental Fig. S1B). The structure shown in Supplemental Figure S1C is basically consistent with the 2D structure originally proposed (Trotochaud and Wassarman 2005), except for some weak base-pairing interactions in the central bulge that may form transiently.

**pRNA transcription pattern for 6S-1 versus 6S-2 RNA**

As a next step, the pattern of pRNA transcription by the *B. subtilis* σA-RNAP at increasing 6S-1 or 6S-2 RNA...
FIGURE 1. Consensus secondary structures of (A) 14 different 6S-1 and (B) 16 different 6S-2 RNAs according to miLocarna (Will et al. 2007) and RNAalifold (Bernhart et al. 2008) using default parameters. The 6S-1 and 6S-2 RNA species selected for the prediction are derived from Firmicutes, for which two 6S RNAs were identified in their genomes, one falling into the 6S-1 and one into the 6S-2 RNA cluster (for further details, see text and Supplemental Fig. S1). The black arrows depict the experimentally determined starting points for pRNA transcription in B. subtilis (Beckmann et al. 2011). The minimum free energy of the 6S-1 RNA consensus structure is −95.08 kcal/mol and that of 6S-2 RNA, −92.77 kcal/mol. Circle outline colors depict the probability of being unpaired or paired; residues marked in red have a high probability, i.e., their single-stranded or paired state is well-defined. Letters represent the most frequent base at the corresponding alignment position. Red letters in uppercase indicate highly conserved bases with an occurrence of >90%, or >75% indicated by red lowercase letters. Black letters in uppercase indicate nucleotide identities conserved in >50% of the sequences. A small black letter is used for positions with lower or no conservation in the alignment. Empty circles indicate nucleotide insertions in a minor fraction of sequences.

Conclusions and in the presence of 200 µM each NTP was compared. While 6S-1 RNA gave rise to the 14-mer pRNA as the main “longer” product under these conditions, in line with previous findings (Beckmann et al. 2011, 2012), the prevailing pRNA products were 13–16 nt in length when 6S-2 RNA served as the template (Fig. 2, lanes 10–14), the 14- and 15-mers being most prominent. Our findings demonstrate that 6S-2 RNA shares the capacity with 6S-1 RNA to serve as a template for pRNA transcription in vitro. A difference between the two is the broader pRNA length spectrum seen in the case of 6S-2 RNA-derived pRNA transcription (Fig. 2).

Affinity of 6S-1/2 RNAs for B. subtilis σ^A-RNAP

An electrophoretic mobility shift assay (EMSA) was employed to study 6S-1/2 RNA affinity for the B. subtilis σ^A-RNAP. For this purpose, 5′-^32P-labeled 6S-1 or 6S-2 RNA was incubated with increasing amounts of σ^A-RNAP holoenzyme in the absence of NTPs, followed by 7.5% nondenaturing PAGE analysis (Fig. 3). This revealed similar K_d values for 6S-1 RNA (K_d ≈ 100 nM) and 6S-2 RNA (K_d ≈ 135 nM) (see Fig. 3). Approximately 85% of the σ^A-RNAP holoenzyme batch used in Figure 3 was capable of 65 RNA binding (Supplemental Fig. S2). We conclude that 6S-1 and 6S-2 RNAs have very similar affinities for the B. subtilis σ^A-RNAP holoenzyme.

Competitiveness of 6S RNAs with DNA promoters

Based on studies performed so far only in the Escherichia coli system, an inherent function of 6S RNAs is to compete with DNA promoters for binding to the housekeeping RNAP holoenzyme, σ^70-RNAP in E. coli (Wassarman and Saecker 2006). Thus, an approach to compare 6S-1 and 6S-2 RNA function is inhibition of DNA transcription in the presence of increasing amounts of 6S RNA. For this purpose, we selected two well-studied σ^A-dependent B. subtilis DNA promoters, the veg promoter, which is constitutively expressed during vegetative growth (Fukushima et al. 2003), and the ribosomal rrnB P1 promoter (Fig. 4A). B. subtilis RNase P RNA was employed as a control RNA for which no specific binding to σ^A-RNAP was expected. In such competition assays, the DNA promoter and σ^A-RNAP (each 100 nM) were preincubated with increasing concentrations of 6S-1, 6S-2, or RNase P RNA (0.1–2 µM) before transcription was initiated by addition of NTPs (Fig. 4B,C). Both 6S RNAs showed considerable and comparable inhibition already at 100 nM in reactions with the veg promoter. In contrast, the control RNA (RNase P RNA) had very little effect on transcription, with possibly some nonspecific inhibition at its highest concentration (2 µM) (Fig. 4B). A similar outcome was observed for transcription driven by the rrnB P1 promoter, although 6S RNA-mediated inhibition effects appeared somewhat weaker (Fig. 4C) than for the veg promoter (Fig. 4B). We additionally analyzed four B. subtilis promoters (rrnO, argC, appD, cspB) and one B. subtilis phage ϕ29 promoter (C24ϕ29), which showed similar trends as the veg and rrnB P1 promoters (data not shown). Although we are aware that inhibition in vitro at a few DNA promoters does not allow conclusions about which genes are affected by 6S-1/2 RNAs in vivo on a global scale (Cavanagh et al. 2008; Neusser et al. 2010), our findings provide first evidence that 6S-1 and 6S-2 RNAs have comparable capacities to effectively compete with DNA promoters for binding to σ^A-RNAP, in line with their very similar affinities for the enzyme (Fig. 3).

6S-2 pRNA transcription changes 6S-2 RNA structure and decreases affinity for RNAP

It was shown that pRNA ~14-mers transcribed by B. subtilis σ^A-RNAP from 6S-1 RNA rearrange the 6S-1 RNA structure.
which a synthetic 32P-labeled pRNA 15-mer was annealed to 6S-2 RNA. The gel mobility of the putative 6S-2 RNA:pRNA complex after pRNA annealing and before RNAP addition (Fig. 5A, cf. lanes 3 and 7). The gel mobility of the putative stable 6S-2 RNA:pRNA complex, which is a control in the absence of RNAP. A pRNA6S-2 15-mer is less effective than a pRNA6S-1 14-mer in decreasing the fraction of 6S RNAs bound to RNAP.

We then analyzed whether pRNAs of comparable length, either derived from 6S-1 or 6S-2 RNA, are similarly effective in decreasing the fraction of the respective 6S RNA bound to RNAP. In the set of experiments now described, we used synthetic pRNAs that were pre-annealed to 6S RNA. Based on the pRNA transcription profiles illustrated in Figure 2, we initially annealed a 6S-1 RNA-specific 14-mer (pRNA6S-1 14-mer) and a 6S-2 RNA-specific 15-mer (pRNA6S-2 15-mer) to the cognate 32P-labeled 6S RNA (10-fold molar excess of pRNA over 6S RNA), followed by addition of α32P-RNA in the absence of NTPs. Nondenaturing PAGE analysis revealed that both 6S RNAs were entirely shifted into the 6S RNA:pRNA complex after pRNA annealing and before RNAP addition (Fig. 6, lanes 4 and 10). While 6S-1 RNA: pRNA6S-1 14-mer complexes almost completely abolished 6S-1 RNA binding to RNAP (Fig. 6A, cf. lane 3 and 2; see also Supplemental Fig. S3), reduced but still substantial complexation with RNAP was observed for samples containing 6S-2 RNA. In the gel image of panel A, the RNAP:6S-1 RNA complex (lanes 1–8) indicated by dashes migrates faster than the neighboring band in lane 1, which likely represents a 6S-1 RNA oligomer.

A pRNA6S-2 15-mer is less effective than a pRNA6S-1 14-mer in decreasing the fraction of 6S RNAs bound to RNAP.

We then analyzed whether pRNAs of comparable length, either derived from 6S-1 or 6S-2 RNA, are similarly effective in decreasing the fraction of the respective 6S RNA bound to RNAP. In the set of experiments now described, we used synthetic pRNAs that were pre-annealed to 6S RNA. Based on the pRNA transcription profiles illustrated in Figure 2, we initially annealed a 6S-1 RNA-specific 14-mer (pRNA6S-1 14-mer) and a 6S-2 RNA-specific 15-mer (pRNA6S-2 15-mer) to the cognate 32P-labeled 6S RNA (10-fold molar excess of pRNA over 6S RNA), followed by addition of α32P-RNA in the absence of NTPs. Nondenaturing PAGE analysis revealed that both 6S RNAs were entirely shifted into the 6S RNA:pRNA complex after pRNA annealing and before RNAP addition (Fig. 6, lanes 4 and 10). While 6S-1 RNA: pRNA6S-1 14-mer complexes almost completely abolished 6S-1 RNA binding to RNAP (Fig. 6A, cf. lane 3 and 2; see also Supplemental Fig. S3), reduced but still substantial complexation with RNAP was observed for samples containing 6S-2 RNA. In the gel image of panel A, the RNAP:6S-1 RNA complex (lanes 1–8) indicated by dashes migrates faster than the neighboring band in lane 1, which likely represents a 6S-1 RNA oligomer.
6S-2 RNA and the pRNA_{6S-2} 15-mer (Fig. 6B, cf. lane 9 and 8). This observation suggested that a fraction of 6S-2 RNA: pRNA complexes underwent dissociation during incubation with RNAP, allowing RNAP molecules to capture transiently free 6S-2 RNAs, which then precludes rebinding of a pRNA.

**Thermodynamics of 6S RNA:pRNA hybrids**

A faster dissociation of 6S-2 pRNAs (relative to 6S-1 pRNAs of equal length) was, indeed, suggested by thermodynamic analysis using RNAcofold (Bernhart et al. 2006), which predicted that 6S-2 RNA:pRNA_{6S-2} 20-mer hybrids are roughly isoenergetic to 6S-1 RNA: pRNA_{6S-1} 14-mer hybrids owing to the A,U-richness of 6S-2 pRNAs (Supplemental Fig. S4).

**Analysis of longer pRNA_{6S-2} variants (16- and 20-mer)**

We thus extended our analysis to pRNA_{6S-2} 16- and 20-mers, using the same setup as in Figure 6. This revealed that the fraction of 6S-2 RNA appearing in the complex with RNAP decreased with increasing pRNA length. With pre-formed 6S-2 RNA:pRNA_{6S-2} 20-mer complexes, essentially no residual complex formation with RNAP was observed (Fig. 7A and B, cf. lanes 2 and 3 vs. 9 and 10 vs. 16 and 17). The same trend was seen when we tested pRNA_{6S-2} 13-, 14-, 15-, 16-, and 20-mers next to each other (Supplemental Fig. S5). Note that the pRNA_{6S-2} 16-mer was still among the prevailing pRNA in vitro transcripts, while the 20-mer was among the pRNA transcripts with lower abundance (Fig. 2, lanes 10–14; see also Fig. 9, below).

In a related setup, 5'-32P-end-labeled pRNA instead of 6S-2 RNA was used. Since pRNA was present in excess over 6S-2 RNA to saturate 6S-2 RNA:pRNA hybrid formation, only a sub-fraction of labeled pRNA appeared in the complex with 6S-2 RNA (Fig. 7A, lanes 6, 7, 13, 14, 20, and 21). Nevertheless, no 6S-2 RNA:pRNA_{6S-2} hybrids migrating as complexes with RNAP were observed (Fig. 7A, lanes 7, 14, and 21), supporting the notion that 6S RNA:pRNA hybrids have lost the ability to form gel-resolvable complexes with RNAP. Based on these findings, we posit that some 6S-2 RNA:RNAP complex formation seen in lanes 3 and 10 (Fig. 7A,B) is due to significant dissociation of 6S-2 RNA:pRNA_{6S-2} 15/16-mer hybrid duplexes during incubation. The resulting “free” 6S-2 RNAs may then either rebind a pRNA oligonucleotide or bind to...
A recent study has reported evidence that pRNA transcription in vitro is less efficient from 6S-2 than 6S-1 RNA, because *B. subtilis* σ^A^-RNAP prefers a GTP as the initiating nucleotide (iNTP), but 6S-2 pRNAs initiate with ATP (Cabrera-Ostertag et al. 2013). In addition, the authors saw only weak 6S-2 pRNA transcription at 50 µM each NTP, an NTP concentration fourfold lower than the one used here (200 µM). We thus analyzed pRNA transcription at different NTP concentrations. In line with the findings of Cabrera-Ostertag et al. (2013), overall pRNA synthesis from 6S-2 RNA decreased more at lower NTP concentrations (20 and 50 µM) relative to that from 6S-1 RNA as template (Fig. 9). At present, it is unclear how this relates to NTP concentrations in vivo. The ATP concentration was recently reported to be ~60 µM in exponentially growing *B. subtilis* cells (Meyer et al. 2011), whereas 1–3 mM was reported for intracellular GTP (Ratnayake-Lecamwasam et al. 2001). Yet, the ATP concentration was measured to be three- to fourfold higher than the other three NTPs (Lopez et al. 1979). For exponentially growing *E. coli* cells, NTP concentrations were determined to be ~4–10 mM for ATP and ~1–5 mM for GTP (Bennett et al. 2009), suggesting that 1–3 mM GTP (Ratnayake-Lecamwasam et al. 2001) in *B. subtilis* seems more like a realistic estimation. Thus, the intracellular concentration of NTPs available for transcription may well be in the range of 0.2 mM or even higher.

**Analysis of 6S-2 RNA displacement from σ^A^-RNAP by 6S-1 RNA**

Although 6S-2 RNA is basically able to direct pRNA synthesis in vitro, deep sequencing and Northern blot experiments have not provided evidence for 6S-2 RNA-templated pRNA synthesis occurred in the presence of shorter synthetic pRNAs, the more shorter the synthetic pRNA (Fig. 8A, lanes 6–10, and Fig. 8B, lanes 16,17). Titration experiments with synthetic pRNA_{6S-2} 15-, 16-, and 20-mers and 6S-2 RNA as the target confirmed these findings (Supplemental Fig. S6). In conclusion, 6S RNA complexes with shorter pRNAs (≤16 nt for 6S-2 RNA; <14 nt for 6S-1 RNA) (see also Supplemental Fig. S3) underwent measurable dissociation within the experimental time frame.
and 1 end-labeled 6S-1 RNA were subjected to the folding and annealing procedure, either alone (lanes 1 and 2) or in the presence of 25 µM pRNA6S-1 14-mer (lanes 3 and 4) or 25 µM pRNA6S-2 15-mer as a noncomplementary control (lanes 5 and 6) in 4 µL 1×TE buffer. Then, 2 µL 5× activity buffer and 3.5 µL ddH2O were added, and samples were kept at 37°C before adding 0.5 µL RNAP holoenzyme (8 µg/µL) (lanes 2,3,5) or 0.5 µL RNAP storage buffer instead (lanes 1,4,6), followed by incubation for 30 min at 37°C (f.c. σ5-RNAP: 1 µM, f.c. 6S RNA: 1 µM, f.c. pRNA: 10 µM), gel loading, and 7.5% nondenaturing PAGE. (B) The analogous experiment with 6S-2 RNA.

**FIGURE 6.** A pRNA6S-2 15-mer is less effective than a pRNA6S-1 14-mer in promoting release of the cognate 6S RNA from σA-RNAP. (A) 2.5 µM 6S-1 RNA including trace amounts of 5’-[32P]-end-labeled 6S-1 RNA were subjected to the folding and annealing procedure, either alone (lanes 1 and 2) or in the presence of 25 µM pRNA6S-1 14-mer (lanes 3 and 4) or 25 µM pRNA6S-2 15-mer as a noncomplementary control (lanes 5 and 6) in 4 µL 1×TE buffer. Then, 2 µL 5× activity buffer and 3.5 µL ddH2O were added, and samples were kept at 37°C before adding 0.5 µL RNAP holoenzyme (8 µg/µL) (lanes 2,3,5) or 0.5 µL RNAP storage buffer instead (lanes 1,4,6), followed by incubation for 30 min at 37°C (f.c. σ5-RNAP: 1 µM, f.c. 6S RNA: 1 µM, f.c. pRNA: 10 µM), gel loading, and 7.5% nondenaturing PAGE. (B) The analogous experiment with 6S-2 RNA.

synthesis in vivo (Beckmann et al. 2011; Cavanagh et al. 2012). It is thus possible that pRNA synthesis from 6S-2 RNA is negligible, at least under the laboratory growth conditions tested. This raises the question of how σA-RNAP may escape from the block by 6S-2 RNA. We tested a simple “6S-1 RNA displaces 6S-2 RNA” model, prompted by the finding that a B. subtilis 6S-1 knockout strain can be rescued from its retarded outgrowth phenotype by resupplying 6S-1 RNA or by further deleting the 6S-2 RNA gene (Cavanagh et al. 2012). For this purpose, we incubated 0.25 µM σ5-RNAP (approximately twofold above the Kd for 6S RNA binding) (see Fig. 3) with a fivefold excess of radiolabeled 6S-2 RNA (1.25 µM) to allow complex formation, followed by addition of 12.5 µM unlabeled 6S-1 RNA (10-fold molar excess over 6S-2 RNA) and incubation for up to 2 h. However, displacement of radioactive 6S-2 RNA from the complex with RNAP was not detectable within this time frame (Fig. 10A).

We then considered the possibility that very short pRNA6S-2 transcripts, which would escape detection by RNA-seq or Northern blotting, might provide sufficient transcriptional and structural dynamics to permit 6S-2 RNA exchange on RNAP by excess 6S-1 RNA. This was tested by further adding ATP and GTP to the reaction samples of the type shown in Figure 10A, allowing synthesis of pRNA 5-mers (or 6-mers in the case of nontemplated addition of an extra nucleotide) on 6S-2 RNA but essentially preventing any pRNA synthesis retarded gel mobility, and (5) with both, formation of stable 6S RNA:pRNA hybrid structures abolishes binding to RNAP.

Despite these commonalities, differences between 6S-1 and 6S-2 RNA were observed as well, which are related to the different G,C-content of the major pRNA species (6S-2 pRNA 13- to 16-mers: 3× G,C; 6S-1 pRNA 14-mer: 6× G,C). A mechanistic consequence might be that short 6S-2 pRNAs (<10 nt) dissociate faster instead of being elongated to longer pRNAs, which could make it more difficult for 6S-2 RNA-stalled RNAP molecules to escape from the sequestration (see discussion below). Also, compared with 6S-1 RNA, 6S-2 pRNAs of equal length will dissociate more rapidly from 6S-2 RNA after RNAP release, which could accelerate pRNA decay or favor binding of 6S-2 RNA to a new RNAP molecule.

We recently proposed for 6S-1 RNA that its release from RNAP depends on the interplay of rate constants for pRNA dissociation from 6S-1 RNA (koff), for pRNA elongation by 1 nt (kp) and for the structural rearrangement (kcont) required for RNAP release (Beckmann et al. 2012). The observation that initiation of pRNA6S-2 synthesis is more reduced for 6S-2 than 6S-1 RNA at lower NTP concentrations (<200 µM) owing to iATP being less favorable than iGTP (Cabrera-Ostertag et al. 2013) requires to further invoke the rate constant for initiation of pRNA synthesis, kinit. A lower kinit for 6S-2 RNA is expected to decrease the overall pRNA synthesis rate on the 6S-2 RNA template, while the overall kp for pRNA...
elongation (averaged over all pRNA positions starting from addition of nt +2) may be similar or higher/lower than for the 6S-1 RNA template. The rate constant \( k_{\text{off}} \) strongly depends on the length of the nascent pRNA transcript. In the case of 6S-2 RNA, \( k_{\text{off}} \) is predicted to be higher for pRNAs of comparable length (e.g., a 6S-2 pRNA 8-mer forms two, a 6S-1 pRNA 8-mer forms five G-C pairs). However, to what extent this predicted faster dissociation rate of 6S-2 pRNAs takes effect depends on how fast \( k_{\text{on}} \) is at the individual nucleotide addition steps (position+2, +3, +4, etc.).

With respect to the function of 6S-2 RNA in vivo, there are two major possibilities: Either (1) the RNA’s principle capacity to give rise to pRNA synthesis and to rearrange its structure to release \( \sigma^\text{A}-\text{RNAP} \), as demonstrated in the study presented here, may indicate that 6S-2 RNA exerts this function in vivo under some conditions to be identified; or (2) pRNA synthesis from 6S-2 RNA has lost its functional relevance in vivo, and the blockade of \( \sigma^\text{A}-\text{RNAP} \) by 6S-2 RNA may be lifted by other means. The latter possibility would be consistent with the observation that significant amounts of 6S-2 pRNAs have neither been detected by Northern blotting nor deep sequencing (Beckmann et al. 2011; Cavanagh et al. 2012), and that efficient pRNA synthesis in vitro by \( \sigma^\text{A}-\text{RNAP} \) requires a GTP as the initiating nucleotide (Cabrera-Ostertag et al. 2013). According to this scenario, 6S-2 RNA would represent a paralog of 6S-1 RNA that has, on evolutionary time scales, lost its classical biological function only very recently, as in vitro, the RNA still has all the hallmark mechanistic capacities of canonical 6S RNAs.

The latter scenario raises the question about the functional advantage of having a second 6S RNA. The different expression profiles of 6S-1 and 6S-2 RNAs suggest that their functions have, indeed, diversified. It was found in three independent studies (Ando et al. 2002; Barrick et al. 2005; Beckmann et al. 2011) that 6S-2 RNA levels substantially decrease in (extended) stationary phase, indicating that sequestration of RNAP molecules complexed with 6S-2 RNA can be overcome in vivo. The recent finding that a 6S-1 RNA knockout strain displays a retarded outgrowth phenotype, whereas a double knockout of 6S-1/2 does not, suggests that 6S-1 RNA

![FIGURE 7](image-url) Length of pRNAs affects the release of \( \sigma^\text{A}-\text{RNAP} \) from complexes with 6S-2 RNA. (A) 1.67 \( \mu \text{M} \) 6S-2 RNA were subjected to the folding and annealing procedure, either alone (lanes 1, 2, 5, 8, 9, 12, 15, 16, 19) or in the presence of 16.7 \( \mu \text{M} \) pRNA\(_{6S-2} \) 15-mer (lanes 3, 4, 6, 7) or pRNA\(_{6S-2} \) 16-mer (lanes 10, 11, 13, 14) or pRNA\(_{6S-2} \) 20-mer (lanes 17, 18, 20, 21) in 6 \( \mu \text{L} \) 1× TE buffer. In lanes 1–5, 8–12, and 15–19 the 6S-2 RNA was \( 5'\)\(^{\text{32P}} \)-end-labeled, whereas pRNAs were \( 5'\)\(^{\text{32P}} \)-end-labeled in lanes 6, 7, 13, 14, 20, and 21. After annealing, 2 \( \mu \text{L} \) 5× activity buffer and 1.5 \( \mu \text{L} \) ddH\(_2\)O were added, and samples were kept at 37°C. Then, 0.5 \( \mu \text{L} \) RNAP holoenzyme (8 \( \mu \text{g/µL} \)) was added (lanes 2, 3, 7, 9, 10, 14, 16, 17, 21) or 0.5 \( \mu \text{L} \) RNAP storage buffer instead, and samples were incubated for 30 min at 37°C, followed by gel loading (f.c. \( \sigma^\text{A}-\text{RNAP} \); 1 \( \mu \text{M} \), f.c. 6S RNA: 1 \( \mu \text{M} \), f.c. pRNA: 10 \( \mu \text{M} \)) and 15% non-denaturating PAGE analysis. (B) As lanes 1–4, 8–11, and 15–18 in panel A but analyzed by 7.5% non-denaturating PAGE for improved gel resolution.

![FIGURE 8](image-url) The extent of inhibition of pRNA de novo transcription by preformed 6S RNA:pRNA complexes depends on the pRNA length. (A) 6S-2 RNA (2 \( \mu \text{M} \)) was subjected to the folding and annealing procedure in 5 \( \mu \text{L} \) of 1× TE buffer, either alone (lane 4) or in the presence of 40 \( \mu \text{M} \) pRNA\(_{6S-2} \) 20-mer (lane 5), 16-mer (lane 6), 15-mer (lane 7), 14-mer (lane 8), 13-mer (lane 9), 12-mer (lane 10), or pRNA\(_{6S-1} \) 14-mer as control (lane 11, noncomplementary pRNA). After annealing, 0.5 \( \mu \text{L} \) water, 2 \( \mu \text{L} \) 5× activity buffer, and 0.5 \( \mu \text{L} \) \( \sigma^\text{A}-\text{RNAP} \) (8 \( \mu \text{g/µL} \)) were added, and reactions were incubated for 30 min at 37°C followed by addition of 2 \( \mu \text{L} \) of NTP mix (f.c. 200 \( \mu \text{M} \) each NTP, also containing \( \alpha^\text{32P}-\text{ATP} \) and further incubation for 30 min at 37°C. As size markers, \( 5'\)\(^{\text{32P}} \)-end-labeled pRNA\(_{6S-2} \) 20-mer (lane 2) and 15-mer (lane 3) were used. (B) 6S-1 RNA (2 \( \mu \text{M} \)) was subjected to the folding and annealing procedure in 5 \( \mu \text{L} \) of 1× TE buffer, either alone (lane 14) or in the presence of 40 \( \mu \text{M} \) pRNA\(_{6S-1} \) 14-mer (lane 15), 13-mer (lane 16), 12-mer (lane 17), or pRNA\(_{6S-2} \) 15-mer as control (lane 18, noncomplementary pRNA). In vitro transcription by \( \sigma^\text{A}-\text{RNAP} \) was performed as in panel A, except that the NTP mix contained \( \alpha^\text{32P}-\text{UTP} \) instead of \( \alpha^\text{32P}-\text{ATP} \). 5\(^{\text{32P}} \)-end-labeled pRNA\(_{6S-1} \) 14-mer (lane 13) and 8-mer (lane 19) were loaded as size markers. Lanes 1 and 12 are negative transcription controls where 6S RNA was omitted. Transcription products were analyzed by 25% denaturing PAGE.
can, by some unknown mechanism, help overcome or prevent the 6S-2 RNA-mediated block of RNAP. Here, we ruled out a simple displacement of 6S-2 by 6S-1 RNA owing to different binding affinities for RNAP. A 6S-2 RNA replacement with 6S-1 RNA when RNAP is capable of synthesizing very short 6S-2 pRNAs (5- or 6-mers, in the presence of ATP and GTP only) seems possible, but the slow rate of this process makes its biological relevance questionable. One possibility is that nucleolytic degradation of RNAP-bound 6S-2 RNA can occur and trigger the release of RNAP. Nonetheless, at present we think it is premature to exclude that pRNA synthesis from 6S-2 RNA occurs in vivo at some stage and to such an extent that RNAP molecules sequestered by 6S-2 RNA are released by this cotranscriptional mechanism.

**MATERIALS AND METHODS**

**Synthetic RNA oligonucleotides**

6S-1 p12: 5′-GUUCGGUCAAACG-3′ (IDT)
6S-1 p13: 5′-GUUCGGUCAAACG-3′ (IDT)
6S-1 p14: 5′-GUUCGGUCAAACG-3′ (Noxxon)
6S-2 p12: 5′-AAAGGUUUAAAACGU-3′ (Noxxon)
6S-2 p13: 5′-AAAGGUUAAAACGU-3′ (MSU)
6S-2 p14: 5′-AAAGGUUAAAACGU-3′ (MSU)
6S-2 p15: 5′-AAAGGUUAAAACGU-3′ (IDT)
6S-2 p16: 5′-AAAGGUUAAAACGU-3′ (MSU)
6S-2 p20: 5′-AAAGGUUAAAACGU-3′ (MSU)

IDT: Integrated DNA Technologies; Noxxon: Noxxon Pharma GmbH; MSU: Lomonosov Moscow State University (for more synthesis details, see Supplemental Material).

**T7 transcription of 6S-1 and 6S-2 RNA**

Mature 6S-1 RNA (190 nt; gene bsrA) and 6S-2 RNA (203 nt, gene bsrB) from *B. subtilis* were synthesized by T7 transcription as described (Beckmann et al. 2011). The respective pUC18 derivative plasmids were linearized with HindIII for run-off transcription. The transcription reaction (final volume 0.5 mL) contained 80 mM HEPES, pH 8.0, 15 mM DTT, 33 mM MgCl2, 1 mM spermidine, 15 mM each NTP, 80 μg/mL template DNA, and 2 units/mL pyrophosphatase. The reaction mixture was prewarmed to 37°C, and guanosine (30 mM stock solution preheated to 75°C) was added to a final concentration (f.c.) of 9 mM to generate transcripts by this cotranscriptional mechanism. The transcription reaction (final volume 0.5 mL) contained 80 mM HEPES, pH 8.0, 15 mM DTT, 33 mM MgCl2, 1 mM spermidine, 15 mM each NTP, 80 μg/mL template DNA, and 2 units/mL pyrophosphatase. The reaction mixture was prewarmed to 37°C, and guanosine (30 mM stock solution preheated to 75°C) was added to a final concentration (f.c.) of 9 mM to generate transcripts by this cotranscriptional mechanism.

**Preparation of *B. subtilis* σ^A^-RNAP holoenzyme**

The *B. subtilis* σ^A^-RNA polymerase holoenzyme was prepared as described (Sogo et al. 1979). For transcription at DNA promoters, *B. subtilis* RNAP was prepared from the *B. subtilis* strain MH5636 as described (Anthony et al. 2000), except that the only chromatographic step performed was affinity-purification using Ni-NTA agarose (Qiagen). This cruder RNAP was highly active in transcription at DNA promoters located on linear PCR fragments.

**PCR**

Linear DNA fragments harboring the *B. subtilis* rrnB and veg promoters (~240 and ~290 bp, respectively) were amplified from genomic *B. subtilis* RNA 168 DNA by standard PCR procedures using the following primers (Metabion): Bsub_veg_22Fwd: 5′-d(CATAA TTTACGAAACTTGGG)-3′; Bsub_veg_19Rev: 5′-d(GGGTACGTCTCAGC)-3′; Bsub_rrnB_31Fwd: 5′-d(GAACG TTACACGGCTTTAGAAATCATGG)-3′; Bsub_rrnB_32Rev: 5′-d (GACGTGACGTACATTGTCTTCTCTCTAAT)-3′. DNA products were then purified by the Wizard SV Gel and PCR Clean-Up System (Promega).

**In vitro transcription at DNA promoters**

Transcription reactions (final volume 10 μL) were prepared as follows: 1 pmol of linear DNA fragment (f.c. 100 nM) was mixed with 2 μL 5× activity buffer (0.2 M Tris-HCl, pH 8.0, 25 mM MgCl2, 50 mM KCl), 2.5 mM each NTP, 10 μg/mL RNAP, and incubated for 2 h at 37°C. Transcription products were then purified by the Wizard SV Gel and PCR Clean-Up System (Promega).

**FIGURE 9.** 6S-1 or 6S-2 RNA-templated pRNA synthesis at varying NTP concentrations. For details, see Materials and Methods. Lanes 1, 2, 9, 10: 5′-32P-labeled RNA length markers mimicking a 6S-1 RNA-derived pRNA 8-mer (5′-GUUCGGUUC, lanes 1,9) or a 14-mer (5′-GUUCGGUUCACCUAC, lanes 2,10). Samples were analyzed by denaturing 25% PAGE. Asterisk at the right margin: These species appear only in the presence of 6S RNA; their nature is unknown and currently under investigation.
Mechanistic comparison of 6S-1 and 6S-2 RNAs

FIGURE 10. (A) Analysis of 6S-2 RNA displacement from α′-RNAP by 6S-1 RNA. 0.25 µM α′-RNAP was incubated with a fivefold excess of radiolabeled 6S-2 RNA (1.25 µM) in 1× activity buffer for 15 min at 37°C to allow complex formation. Then, 12.5 µM unlabeled 6S-1 RNA were added, and samples were incubated for 5, 10, 30, 60, or 120 min at 37°C before loading onto a 7.5% native PAA. Lane 1: ddH2O instead of 6S-1 RNA was added; lane 7: as lane 6, but without RNAP (for details, see Materials and Methods). The same result was obtained in two additional independent experiments (data not shown). Also, quantification of phosphorimage signals for free 6S-2 RNA and the complex with RNAP did not reveal any decrease in the proportion of 6S-2 RNA:RNAP complexes over time. (B) Experiment as in panel A, but incubation with ATP and GTP (each 200 µM). Lane 8: 6S-2 RNA, no RNAP, no 6S-1 RNA, no NTPs; lane 9: 6S-2 RNA, RNAP, no 6S-1 RNA, no NTPs; lane 10: 6S-2 RNA, RNAP, no 6S-1 RNA, ATP and GTP; lane 11: 6S-2 RNA, no RNAP, 6S-1 RNA, no NTPs; lanes 12–17: as lanes 2–6, but after addition of 6S-1 RNA, samples were incubated for 15 min at 37°C, followed by addition of ATP/GTP (ddH2O instead in lane 12) and incubation for the time period indicated above each lane.

MgCl2, 0.8 M KCl, 5 mM DTT) supplemented with 0.5 µg/µl heparin, 0.6 µL α′-RNAP holoenzyme (0.7 µg/µL) and ddH2O to yield a volume of 6 µL. This mixture was incubated for 10 min at 37°C to promote DNA:RNAP complex formation. Then, 2 µL containing different amounts of 6S-1 RNA, 6S-2 RNA, or B. subtilis RNase P RNA (409 nt, used as a negative control) were added, followed by incubation for another 10 min at 37°C to allow competing RNA:RNAP and DNA:RNAP complexes to equilibrate. Finally, the transcription reaction was started upon addition of 2 µL NTP mix (250 µM each ATP, CTP, GTP, and 62.5 µM UTP) containing α-[32P]-UTP (~50,000 Cherenkov c.p.m. per gel lane), followed by incubation for 20 min at 37°C. The f.c. of α′-RNAP was ~100 nM and that of 6S-1, 6S-2, or RNase P RNA either 0.1, 0.2, 0.5, 1, or 2 µM. After transcription, samples were mixed with an equal volume of 2× denaturing RNA loading buffer (Thermo Scientific), heated for 5 min at 95°C, followed by immediate transfer onto ice, and then loaded onto a 5% denaturing PAA gel (1× TBE: 89 mM Tris base, 89 mM boric acid, 2 mM EDTA, pH ~8.3).

Folding and annealing procedure

In vitro-transcribed 6S-1 or 6S-2 RNA, either alone or in the presence of chemically synthesized pRNA oligonucleotides, were heated to 95°C for 5 min in a volume of 4–8 µL 1× TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0) and stepwise-cooled (5 min each 80, 70, 60, 50, and 37°C) in a thermocycler (Biometra) to accomplish folding and annealing. Samples also contained trace amounts of either 5′-[32P]-labeled 6S-1/2 RNA or pRNA oligonucleotide; RNA concentrations are specified in the respective figure legends.

Analysis of 6S RNA:RNAP complex formation

Electrophoretic mobility shift assays were conducted as described (Beckmann et al., 2011), using trace amounts of 5′-[32P]-labeled 6S-1 or 6S-2 RNA (~5000 Cherenkov c.p.m. per gel lane) and varying excess amounts of α′-RNAP. After pre-incubation for 30 min at 37°C, samples were mixed with an equal volume of 2× native RNA loading buffer (0.025% bromophenol blue [w/v], 0.025% xylene cyanol blue [w/v], 20% glycerol), followed by 7.5% native PAGE (1× TBE). For data analysis and Kd determination, see Beckmann et al. (2011).

6S RNA:RNAP gel shifts after pRNA annealing

To 4 µL of 6S-1 or 6S-2 RNA that had been subjected to the folding and annealing procedure (see above) in the presence of a complementary pRNA oligonucleotide (controls without pRNA), 2 µL of 5× activity buffer (supplemented with 0.5 µg/µl heparin) and 2.9 or 3.5 µL ddH2O were added, followed by addition of 1.06 µL or 0.5 µL of α′-RNAP holoenzyme (8 µg/µL) to give a final volume of 10 µL. Samples were incubated for 30 min at 37°C to promote 6S RNA:RNAP complex formation (f.c. of α′-RNAP: 2 or ~1 µM; f.c. 6S RNA: 1 or 10 µM); finally, samples were mixed with an equal volume of 2× native RNA loading buffer (or adjusted to 10% glycerol without dyes, giving the same results) and loaded onto a 7.5% nondenaturing PAA gel (1× TBE).

6S RNA:RNAP gel shifts after transcription

This protocol pertains to the experiment shown in Figure 5. To 4 µL 6S-2 RNA (2.5 µM) that had been subjected to the folding and annealing procedure, 2 µL of 5× activity buffer (supplemented with 0.5 µg/µl heparin), 1 µL ddH2O and 1.06 µL α′-RNAP holoenzyme (8 µg/µL) were added, and samples were incubated for 30 min at 37°C. Transcription was started by adding 2 µL NTP mix (1 mM each NTP; f.c. 200 µM each) or 2 µL ddH2O instead (negative control); the f.c. of α′-RNAP was ~2 µM and that of 6S-2 RNA 1 µM. Samples were then loaded onto a 7.5% nondenaturing PAA gel (1× TBE) as described above.

Transcription of [32P]-labeled pRNAs using 6S-1 or 6S-2 RNA as template

Four microliters 6S RNA (2.5 µM; or other concentration if indicated) that had been subjected to the folding and annealing procedure, either alone or in the presence of a synthetic pRNA oligonucleotide
min, followed by cooling on ice for 20 min. Before gel loading, samples were heated to 98°C for 3 min of each sample was mixed with 15 µL of highly denaturing RNA loading buffer (0.02% [w/v] bromophenol blue, 0.02% [w/v] xylene cyanol blue, 8 M urea, 50% [v/v] deionized formamide, 2× TBE, pH 8.0) and loaded onto a 25% denaturing PAA gel.

6S-1 and 6S-2 RNA-templated pRNA synthesis at varying NTP concentrations

Two microliters 6S-1 or 6S-2 RNA (5 µM) in 1× TE buffer were heated to 80°C and cooled down to 50°C in steps of 10°C (2 min each step), with a final incubation step at 37°C for 2 min. Then, 4.6 µL mix containing activity buffer and σA-RNAP were added, followed by incubation for 10 min at 37°C. Reactions were started by adding 3.4 µL NTP mix (f.c. of each NTP: 20, 50, or 200 µM; including ∼250,000 Cherenkov c.p.m. of α-[32P]-UTP per gel lane). The final reaction volume of 10 µL contained 1 µM 6S RNA, 1 µM RNAP, and 1× activity buffer (see above). After 1 h at 37°C, half of each sample was mixed with 15 µL of highly denaturing RNA loading buffer (see above) and analyzed by 25% denaturing PAGE (1× TBE). Before gel loading, samples were heated to 98°C for 3 min, followed by cooling on ice for 20 min.

Prediction of 6S-1 and 6S-2 RNA structures from Bacilli

6S RNA sequences listed for the phylum Firmicutes were taken from the RFAM database (Burge et al. 2012) and aligned to the 6S/SsrS RNA covariance model provided there. The alignment was clustered using the neighbor-joining algorithm implemented in ClustalW (Larkin et al. 2007). Thus, a similarity tree was built that grouped similar sequences (in terms of the RFAM SsrS RNA covariance model). Thus, a similarity tree was built that grouped similar sequences (in terms of the RFAM SsrS RNA covariance model). 6S-1 and 6S-2 RNA from Bacilli from

Analysis of 6S-2 RNA displacement from RNAP in the absence of NTPs

In this experiment, σA-RNAP was pre-incubated with a fivefold molar excess of radiolabeled 6S-2 RNA, followed by addition of a 10-fold molar excess of 6S-1 RNA over 6S-2 RNA and varying times of incubation. 6S-2 RNA-RNAP complexes were then analyzed by native PAGE (1× TBE buffer). In practice, to each of seven reaction tubes, 2 µL of 5× activity buffer, 1 µL heparin (400 ng/µL; f.c. 40 ng/µL), 12.5 pmol unlabeled 6S-2 RNA (f.c. 1.25 µM), trace amounts of radiolabeled 6S-2 RNA (~10,000 Cherenkov c.p.m.), and ddH2O (to 8.1 µL) were added; all tubes were put at 37°C. Immediately, 1.3 µL B. subtilis σA-RNAP holoenzyme (0.8 µg/µL; f.c. ~0.25 µM) was added to tube #6, while 1.3 µL σA-RNAP storage buffer were added to tube #7, followed by incubation for 15 min at 37°C. Then, 0.55 µL 6S-1 RNA (231 µM) (f.c. 12.5 µM) was added to tubes #6 and #7 (time point zero) to reach final volumes of 10 µL, followed by incubation for 120 min at 37°C (Fig. 10A, lanes 6 and 7). After 45 min (relative to the time point zero just mentioned), components were added in the same order (as for tube #6) to tube #5 (addition of RNAP, 15 min at 37°C, then addition of 6S-1 RNA and incubation for 60 min at 37°C). After 75, 95, and 100 min, respectively, the same was done for tubes #2–4 (addition of RNAP, 15 min at 37°C, then addition of 6S-1 RNA and incubation with 6S-1 RNA for 30, 10, or 5 min, respectively). After 105 min, 1.3 µL B. subtilis σA-RNAP were added to tube #1, and the mixture was incubated for 15 min at 37°C, followed by addition of 0.55 µL ddH2O instead of 6S-1 RNA. Then, all samples were instantly mixed with 10 µL of 2× native RNA loading buffer and loaded onto a 7.5% native PAA gel (1× TBE).

Analysis of 6S-2 RNA displacement from RNAP in the presence of ATP and GTP

In this type of experiment, the staggered incubation procedure was identical to that in the absence of NTPs (see above) to the point of 6S-1 RNA addition, after which samples were incubated for another 15 min at 37°C. Then, ATP and GTP were added to final concentrations of 200 µM each (Fig. 10B, lanes 13–17), and samples were incubated for 5, 10, 30, 60, or 120 min at 37°C before gel loading. The sample in lane 12 of Figure 10B (time 0) was the last to receive RNAP 30 min before gel loading, followed by incubation for 15 min at 37°C, addition of 6S-1 RNA, another incubation for 15 min at 37°C, and addition of ddH2O instead of ATP/GTP immediately before gel loading.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

ACKNOWLEDGMENTS

We thank Dominik Helmecke for excellent technical assistance. This work was supported by the Deutsche Forschungsgemeinschaft (GK 1384) to P.G.H., K.D., M.L., and R.K.H., the Russian Foundation for

Burenina et al.
REFERENCES

Ando Y, Asari S, Suzuma S, Yamane K, Nakamura K. 2002. Expression of a small RNA, BS203 RNA, from the yocL-yocF intergenic region of Bacillus subtilis genome. FEMS Microbial Lett 207: 29–33.

Anthony LC, Artsimovich I, Svetlov V, Landic R, Burgess RR. 2000. Rapid purification of His(6)-tagged Bacillus subtilis core RNA polymerase. Protein Expr Purif 19: 350–354.

Barrick JE, Sudarasa N, Weinberg Z, Ruzzo WL, Breaker RR. 2005. 6S RNA is a widespread regulator of eubacterial RNA polymerase that resembles an open promoter. RNA 11: 774–784.

Beckmann BM, Burenina OY, Hoch PG, Marz M, Willkomm DK, Salas M, Hartmann RK. 2012. A pRNA-induced structural rearrangement of bacterial RNA polymerase resembles an open promoter. RNA 11: 599–599.

Bernhart SH, Tafer H, Mückstein U, Flamm C, Stadler PF, Hofacker IL. 2006. Partition function and base pairing probabilities of RNA heterodimers. Algorithms Mol Biol 1: 3.

Bernhart SH, Hofacker IL, Will S, Gruber AR, Stadler PF. 2008. RNAalifold: Improved consensus structure prediction for RNA alignments. BMC Bioinformatics 9: 474.

Burge SW, Daub J, Eberhardt R, Tate J, Barquist L, Nawrocki EP, Eddy SR, Gardner PP, Bateman A. 2012. Rfam 11.0: 10 years of RNA families. Nucleic Acids Res 41: D226–D232.

Cabrera-Ostertag IJ, Cavanagh AT, Wassarman KM. 2013. Initiating nucleotide identity determines efficiency of RNA synthesis from 6S RNA templates in Bacillus subtilis but not Escherichia coli. Nucleic Acids Res 41: 7501–7511.

Cavanagh AT, Klocko AD, Liu X, Wassarman KM. 2008. Promoter specificity for 6S RNA regulation of transcription is determined by core promoter sequences and competition for region 4.2 of D3. Mol Microbiol 67: 1242–1256.

Cavanagh AT, Sperger JM, Wassarman KM. 2012. Regulation of 6S RNA by pRNA synthesis is required for efficient recovery from stationary phase in E. coli and B. subtilis. Nucleic Acids Res 40: 2234–2246.

Fukushima T, Ishikawa S, Yamamoto H, Ogasawara N, Sekiguchi J. 2003. Transcriptional, functional and cytotoxic analyses of the veg gene in Bacillus subtilis. J Biochem 133: 475–483.

Gildehaus N, Neusser T, Wurm R, Wagner R. 2007. Studies on the function of the riboregulator 6S RNA from E. coli: RNA polymerase binding, inhibition of in vitro transcription and synthesis of RNA-directed de novo transcripts. Nucleic Acids Res 35: 1885–1896.

Hofacker IL, Fekete M, Stadler PF. 2002. Secondary structure prediction for aligned RNA sequences. J Mol Biol 319: 1059–1066.

Irrov I, Sharma CM, Vogel J, Winkler WC. 2010. Identification of regulatory RNAs in Bacillus subtilis. Nucleic Acids Res 38: 6637–6651.

Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, et al. 2007. Clustal W and Clustal X version 2. Bioinformatics 23: 2947–2948.

Lopez JM, Marks CL, Freese E. 1979. The decrease of guanine nucleotides initiates sporulation of Bacillus subtilis. Biochem Biophys Acta 587: 238–252.

Meyer FM, Jules M, Mehne FM, Le Coq D, Landmann JJ, Görke B, Aymerich S, Stülke J. 2011. Malate-mediated carbon catabolite repression in Bacillus subtilis involves the HPrK/CcpA pathway. J Bacteriol 193: 6939–6949.

Neusser T, Polen T, Geissen R, Wagner R. 2010. Depletion of the non-coding regulatory 6S RNA in E. coli causes a surprising reduction in the expression of the translation machinery. BMC Genomics 11: 165.

Nakamura K, Yamane K. 2002. Identification and characterization of a small RNA, BS203 RNA, from the yocL-yocF intergenic region of Bacillus subtilis. J Biochem 131: 475–483.

Suzuma S, Asari S, Bunai K, Yoshino K, Ando Y, Kakeshita H, Fujita M, Nakamura K, Yamane K. 2002. Identification and characterization of novel small RNAs in the aspS-yrvM intergenic region of the Bacillus subtilis genome. Microbiology 148: 2591–2598.

Trotchaud AE, Wassarman KM. 2005. A highly conserved 6S RNA structure is required for regulation of transcription. Nat Struct Mol Biol 12: 313–319.

Wassarman KM. 2007. 6S RNA: A small RNA regulator of transcription. Curr Opin Microbiol 10: 164–168.

Wassarman KM, Saecher RM. 2006. Synthesis-mediated release of a small RNA inhibitor of RNA polymerase. Science 314: 1601–1603.

Wassarman KM, Storz G. 2000. 6S RNA regulates E. coli RNA polymerase activity. Cell 101: 613–623.

Will S, Reiche K, Hofacker IL, Stadler PF, Backofen R. 2007. Infering non-coding RNA families and classes by means of genome-scale structure-based clustering. PLoS Comput Biol 3: e65.

www.rnajournal.org 359