Investigation of catalysis by bacterial RNase P via LNA and other modifications at the scissile phosphodiester

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ABSTRACT

We analyzed cleavage of precursor tRNAs with an LNA, 2'-OCH₃, 2'-H or 2'-F modification at the canonical (c₀) site by bacterial RNase P. We infer that the major function of the 2'-substituent at nt -1 during substrate ground state binding is to accept an H-bond. Cleavage of the LNA substrate at the c₀ site by Escherichia coli RNase P RNA demonstrated that the transition state for cleavage can in principle be achieved with a locked C₃'-endo ribose and without the H-bond donor function of the 2'-substituent. LNA and 2'-OCH₃ suppressed processing at the major aberrant m₁ site; instead, the m₁ (nt +1/+2) site was utilized. For the LNA variant, parallel pathways leading to cleavage at the c₀ and m₁ sites had different pH profiles, with a higher Mg²⁺ requirement for c₀ versus m₁ cleavage. The strong catalytic defect for LNA and 2'-OCH₃ supports a model where the extra methylene (LNA) or methyl group (2'-OCH₃) causes a steric interference with a nearby bound catalytic Mg²⁺ during its recoordination on the way to the transition state for cleavage. The presence of the protein cofactor suppressed the ground state binding defects, but not the catalytic defects.

INTRODUCTION

Endonucleolytic 5'-end maturation of tRNA primary transcripts is catalyzed by the ribonucleoprotein enzyme Ribonuclease P (RNase P) in all three domains of life (Archaea, Bacteria and Eukarya) as well as in mitochondria and chloroplasts (1–4). An exception is the human mitochondrial tRNA 5'-end maturation enzyme that lacks an RNA subunit, with its three protein subunits recruited from related and unrelated biochemical pathways (5). Bacterial RNase P enzymes are composed of a catalytic RNA subunit (6), ~400 nucleotides (nt) in length, and a single small protein of typically 120 amino acids (7). The protein is essential in vivo, but in vitro its absence can be compensated by increased mono- and particularly di-valent cations (6). Studies with RNase P RNA ribozymes (P RNAs) from Escherichia coli and Bacillus subtilis have indicated a specific role for at least two metal ions in productive enzyme–substrate complex formation and cleavage chemistry (8–15). Since processing of precursor tRNAs (ptRNAs) by RNase P results in cleavage products with 3'-OH and 5'-phosphate termini, an inherent feature of RNase P catalysis must be to prevent the 2'-hydroxyl group at the cleavage site from attacking the scissile phosphodiester, which would otherwise result in 5'-OH and 2',3'-cyclic phosphate ends. The 2'-hydroxyl group of the ribose at the cleavage site has indeed been identified as being crucial in the catalytic process. A 2'-deoxy or 2'-amino substitution at the canonical RNase P cleavage site (nt -1) affected substrate ground state binding, cleavage site recognition, binding of catalytically important Mg²⁺ and substantially reduced the cleavage rate by E. coli or B. subtilis P RNA (8,16–23). Initial evidence for the 2'-hydroxyl group at nt -1 acting as a ligand for a catalytic Mg²⁺ ion came from the observation that a 2'-deoxy modification at this location reduced the Hill coefficient for the Mg²⁺ dependence of substrate cleavage from 3 to 2 and increased the apparent K₅₀ for Mg²⁺ binding 10⁻³-fold (8). A 2'-amino modification at this position resulted in misincleavage at lower pH, consistent with the idea that protonation of the 2'-amino group at lower pH causes electrostatic repulsion of a metal at this location (20,21). Mn²⁺ rescued cleavage at the canonical c₀ site for ptRNA with a 2'-amino substitution at nt -1, which would be consistent with a direct metal ion coordination to the 2'-substituent at this location; however, a 2'-deoxy modification at the same site was rescued to a similar extent by Mn²⁺, thus questioning the possibility of a direct metal
contact (20). We rather favor the concept of the 2'-OH at nt –1 interacting with a water molecule from the hydration shell of this metal ion (8,20). Pan and coworkers found that a 2'-deoxy substitution at the c0 cleavage site had a large effect on ptRNA cleavage by B. subtilis P RNA, whereas two in vitro selected non-tRNA substrates were little affected by such a substitution at the cleavage site; rather, the 2'-OH one nucleotide upstream was identified to be critical for cleavage of these non-tRNA substrates which lack a T stem-loop module (24). They inferred a model according to which a unique 2'-endo conformational rearrangement or docking step of initial E•S complexes ([E•S] → (E•S)*, Scheme 1) that precedes the catalytic step. This unique 2'-OH is usually the one at the c0 site in the case of ptRNAs that are able to form the T stem-loop interaction with the P RNA S domain, but another 2'-OH in the vicinity may be utilized to fulfill this role in the absence of the T stem-loop interaction. In addition, the P protein shifts the equilibrium towards (E•S)*, thus often mitigating defects of the P RNA-alone reaction which are due to impaired (E•S)* formation. Based on this model, the 2'-OH at the c0 site in ptRNA substrates likely affects $k_{\text{conf}}$ of the docking step as well as catalysis ($k_c$; Scheme 1).

NMR experiments have indicated that the metal ion coordinated with the help of the 2'-OH group at nt –1 is actually ‘prebound’ to ptRNA before complexation with P RNA (25). In a related study, the ribose at nt –1, if not involved in base pairing, was found to be predominantly 2'-endo puckered (26), raising the possibility that a C2'-endo sugar pucker may be the favored conformation in the transition state after potential –1/73 base pairs have been disrupted through formation of the +73/C15 interaction with P RNA (27). To address the role of sugar conformation at the cleavage site for the catalytic process, we analyzed a ptRNA with a locked nucleic acid (LNA) substitution at nt –1, which arrests the ribose in a C3'-endo pucker (28,29) (Figure 1). For better evaluation of effects caused by the locked ribose conformation, the loss of the H-bond donor function at the 2'-oxygen and sterical impediments due to the presence of an additional methylene group, the ptRNA variant with a 2'-OCH3 modification at this position was included in our analysis. As LNA, the 2'-OCH3 group can accept but not donate H-bonds. Unlike LNA, the 2'-OCH3 modified ribose has a flexible conformation (Table 1) and the 2'-OCH3 group can freely rotate around the C2'-O2' bond, whereas the position of the corresponding methylene is fixed in LNA. Other variants with 2'-fluoro (2'-F) or 2'-deoxy (2'-H) substitutions at nt –1 that affect the H-bonding pattern, the free energy of solvation and the electronegativity of the 2'-substituent were included as well. We then characterized the modified ptRNA substrates in reactions catalyzed by type A (E. coli) and type B (B. subtilis) RNase P (RNA) for (i) substrate ground state binding, (ii) cleavage site selection and (iii) cleavage kinetics. Our results indicate that the major function of the 2'-substituent at nt –1 during substrate ground state binding is to accept an H-bond. We demonstrate that cleavage at the canonical site by E. coli P RNA can occur in the presence of a locked C3'-endo ribose and without the H-bond donor function of the 2'-substituent, however, with very low efficiency. The strong catalytic defect in the presence of LNA and 2'-OCH3 modifications at the c0 site supports a model where the extra methylene (LNA) or methyl group (2'-OCH3) causes a steric interference with a nearby bound catalytic Mg2+ during its recoordination on the way to the transition state for cleavage. LNA and 2'-OCH3 at the c0 site also suppressed processing at the major aberrant m +1 site, directing cleavage to the aberrant m +1 site (nt +1/+2). The protein cofactor, although suppressing the ground state binding defects, was unable to substantially relieve the severe catalytic defects. Differences between E. coli and B. subtilis enzymes included a stronger defect caused by 2'-H at nt –1 on the B. subtilis versus E. coli holoenzyme, and different cleavage site selection on the 2'-OCH3 substrate.

MATERIALS AND METHODS

RNA synthesis, 5'-phosphorylation of RNA, 5'-end-labeling of RNA and assembly of ptRNA variants

Chemical and enzymatic RNA synthesis, purification, 5'-phosphorylation and 5'-32P-end-labeling of RNA, as well as the assembly of ptRNA variants with single-site modifications were performed exactly as described (20,30).

Preparation of recombinant RNase P proteins

E. coli and B. subtilis RNase P proteins carrying an N-terminal His-tag were overexpressed and purified as described (31).

Reconstitution of RNase P holoenzymes

(i) For cleavage experiments, a solution containing 133 nM E. coli or B. subtilis P RNA, 107 mM KCl, 53 mM MES, pH 6.0 and 10.7 mM MgCl2 was incubated for 5 min at 55°C and 50 min at 37°C. To 15 µl of this solution, 1 µl of P protein solution (fresher thawed and diluted) was added to adjust the solution to 0.5 µM B. subtilis or 1.2 µM E. coli P protein and to 125 nM P RNA, 100 mM KCl, 50 mM MES and 10 mM MgCl2, pH 6.0. This mixture was incubated for another 5 min at 37°C;
16 μl thereof were then combined with 4 μl ptRNA solution (preincubated in the same buffer for 5 min at 55°C and 25 min at 37°C) to start the processing reaction.

(ii) For binding experiments, a solution containing 1.18 μM *E. coli* or *B. subtilis* P RNA, 235 mM NH₄OAc, 59 mM Tris–acetate, pH 7.1, 17.7 mM CaCl₂ and 0.059 (w/v) Nonidet P40 was incubated for 1 h at 37°C. To 8.5 μl of this mixture, 1.5 μl of P protein solution was added to adjust the solution to 5 μM *B. subtilis* or 12 μM *E. coli* P protein (freshly thawed and diluted), 1 μM P RNA, 200 mM NH₄OAc, 50 mM Tris–acetate, pH 7.1, 15 mM CaCl₂ and 0.05 (w/v) Nonidet P40. This holoenzyme stock solution was incubated for 15 min at 37°C and then used to prepare serial dilutions in the same buffer. The dilute solutions were incubated for another 5 min at 37°C; 10 μl of individual holoenzyme solutions were combined with an equal volume of 5'32P-labeled ptRNA solution (<1 nM ptRNA, 80 000 Cerenkov cpm), which had been preincubated for 30 min at 37°C in the same buffer, followed by a final incubation step for 5 min at 37°C before conduction of the spin column assay (see below). For cleavage and binding experiments, the final RNase P holoenzyme concentration was considered equal to the final P RNA concentration. This represents an upper estimate of the enzyme concentration, since 100% ribonucleoprotein complex formation may not have been reached.

Nuclease P1 hydrolysis

A solution of 5'-32P-end-labeled 24-mer RNA (10⁴ Cerenkov cpm/μl), 0.1 μg/μl carrier RNA, 40 mM NH₄OAc, 0.4 mM ZnSO₄, pH 5.3 and 0.01 ng/μl nuclease P1 was incubated for 20 s at 70°C in a heating block. After incubation the reaction tube was placed on ice, the RNA was concentrated and desalted by ethanol precipitation and analyzed by denaturing PAGE.

Kinetics

The ptRNA variants were processed under single turnover conditions at saturating enzyme concentrations (5 μM *E. coli* or *B. subtilis* P RNA or 100 nM of reconstituted *E. coli* or *B. subtilis* RNase P holoenzyme), 1 M NH₄OAc (for RNA-alone reactions) or 0.1 M KCl (for holoenzyme reactions), and at Mg²⁺ concentrations and pH conditions as specified. The pH of 7.0 was chosen to provide conditions where substantial cleavage of all variants occurred, even at low Mg²⁺ concentrations; pH values of 5.5 or 6.0 allowed us to analyze cleavage at rate-limiting chemistry for all substrates (9). All processing reactions were performed at 37°C; P RNA solutions were preincubated for 1 h at 37°C and ptRNA solutions for 5 min at 55°C and 25 min at 37°C before mixing the two. For holoenzyme kinetics, an additional enzyme reconstitution procedure was included (see above). Aliquots withdrawn at different time points from enzyme–substrate mixtures were subjected to ethanol precipitation in the presence of 20 μg of glycogen before analysis by 20% PAGE/8 M urea. Data analysis and calculation of single turnover rate constants of cleavage (kobs) was performed as described (32). For the Mg²⁺ titration experiments in Figure 4E, data were analyzed by non-linear regression analysis using the Hill equation

$$ k_{obs} = k_{obs,max} \cdot [\text{Mg}^{2+}]/(K_c + [\text{Mg}^{2+}]) $$

ptRNA binding by spin column assays

Spin column assays for the determination of equilibrium dissociation constants (Kd) of enzyme–substrate complexes were performed as described (10). Binding of ptRNAs to P RNA was analyzed in a buffer containing 50 mM MES, 1 M NH₄OAc, CaCl₂ (15 mM) or SrCl₂ (5 or 80 mM), 0.1 % (w/v) SDS, 0.05 % (w/v) Nonidet P40, pH 6.0. Binding of ptRNAs to holoenzymes was analyzed in a buffer containing 50 mM Tris–acetate, 0.2 M NH₄OAc, 15 mM CaCl₂, 0.05 % (w/v) Nonidet P40, pH 7.1.

RESULTS

The bacterial ptRNA Gly variants with single modifications at nt -1 were constructed by ligation in the D loop after annealing of a synthetic 24-mer to a 5'-truncated tRNA transcript (Figure 1A). The different 2'-ribose modifications at nt -1 are illustrated in the boxes on the left. Several physico-chemical properties of the 2'-substituents investigated here are summarized in Table 1. The locked C3'-endo conformation of an LNA residue is shown in comparison to that of regular nucleotides in Figure 1B. To put our data into a broader perspective, we analyzed the substrates in reactions catalyzed by *E. coli* and *B. subtilis* RNase P (RNA), the main model systems for the two structurally distinct classes of bacterial P RNAs, type A (*for ‘ancestral’) and type B (*for ‘Bacillus’) (33–37).

Effect of modifications at nt -1 on substrate binding by *E. coli* P RNA

Two sets of variants of ptRNA Gly were investigated. The first set carried ribothymidines (rT) at nt -1 and -2 and a 2'-OH, 2'-OCH₃ or LNA modification at nt -1 [referred to as 2'-OH (T-1), 2'-OCH₃ (T-1) and LNA (T-1)] or LNA at nt -2 [LNA (T-2)]. The ribothymidines at positions -1 and -2 were introduced for reasons of comparability with variants carrying LNA substitutions at nt -1 or -2, because LNA analogs for uridine were not available. The second set of substrates carried a C at nt -1 and U at nt -2, and a 2'-OH, 2'-fluoro (2'-F) or 2'-deoxy (2'-H) substituent at position -1 [referred to as 2'-OH (C-1), 2'-F (C-1) and 2'-H (C-1)]. We determined equilibrium dissociation constants (Kd) for binding of ptRNA Gly variants to *E. coli* P RNA at two Sr²⁺ (5 and 80 mM) and one Ca²⁺ (15 mM) concentration (Table 2). Both metal ions support substrate binding to *E. coli* P RNA with similar efficiency as Mg²⁺, but are inactive (Sr²⁺) or very inefficient (Ca²⁺) as catalytic cofactors [30 and references therein]. The ptRNA with rT residues at -2 and -1, variant 2'-OH (T-1), bound to *E. coli* P RNA with the same affinity (within 2-fold) as the variant with U at -2 and C at -1,
variant 2′-OH (C-1) (Table 2), indicating that the extra methyl groups on the bases as well as the pyrimidine identity at nt −1 had no or very marginal effects on substrate binding in the ground state. This essentially pertains to the LNA (T-2) substrate as well, which differed from the control substrate 2′-OH (T-1) only by a 3–4-fold increase in $K_d$ at the lowest divalent metal ion concentration (5 mM Sr$^{2+}$). Thus, an LNA modification at nucleotide −2 had little effect on substrate binding by *E. coli* P RNA.

At all three tested metal ion conditions, binding defects were seen for the 2′-F (C-1) (2-4-fold), LNA (T-1) (2-13-fold), 2′-H (C-1) (9-60-fold) and 2′-OCH$_3$ (T-1) substrate (30-50-fold). Remarkably, the binding defect for LNA (T-1) was essentially suppressed (within 3-fold) at 15 mM Ca$^{2+}$ or 80 mM Sr$^{2+}$ (Table 2). For comparison, the 2′-H (C-1) ptRNA was still bound with 9–11-fold reduced affinity under the same conditions, thus even exceeding the defect of a 2′-F substitution (4-fold effect on $K_d$). In summary, the LNA substitution at position −1 had only a very minor impact on ptRNA ground state binding, despite its bulky extra methylene group and a sugar pucker arrested in C3′-endo. In contrast, the structurally related 2′-OCH$_3$ modification caused a >30-fold affinity decrease at all tested metal ion concentrations. The 2′-deoxy substitution caused the largest affinity decrease, 60-fold at 5 mM Sr$^{2+}$, which could be mitigated to 10-fold at the higher divalent metal ion concentrations. From the comparison of an LNA versus 2′-H and 2′-F modification at nt −1 we infer that the presence of a 2′-substituent at nt −1 with good H-bond acceptor capacity, such as the 2′-oxygen, is a major determinant for metal-dependent formation of E–S ground state complexes. Such information could not be extracted from the results obtained with the related 2′-OCH$_3$ modification, because the steric effect dominates for 2′-OCH$_3$.

### Effect of modifications at nt -1 on substrate binding by *B. subtilis* P RNA

*B. subtilis* P RNA bound the control all-ribose substrates at 15 mM Ca$^{2+}$ with 19–26-fold lower affinity than *E. coli* P RNA (Table 2), consistent with previous findings (37). Again, whereas the LNA (T-1) substrate showed little perturbation of substrate binding relative to the 2′-OH (T-1) substrate (2.6-fold increase in $K_d$), the defect was most pronounced for the substrate with 2′-OCH$_3$ at −1 (11-fold increase in $K_d$). Here, the 2′-H substitution reduced affinity only 2-fold and a 2′-F substituent was bound with roughly equal affinity as the 2′-OH (C-1) control substrate. These results are qualitatively similar to those obtained with *E. coli*

### Table 1. Some physico-chemical properties of ribose moieties with different 2′-substituents

| 2′-substituent | H-bond donor | H-bond acceptor | Flexible ribose conformation | Preferentially adopted sugar pucker$^a$ | Relative solubility$^b$ | Electronegativity of 2′-substituent | Sterical impediments |
|---------------|--------------|----------------|-----------------------------|-----------------------------------------|------------------------|----------------------------------|------------------|
| 2′-OH         | Yes          | Yes            | Yes                         | C3′-endo                               | Highest                | High                             | No               |
| LNA           | No           | Yes            | No                          | Locked C3′-endo                         | Intermediate           | High                             | No               |
| 2′-OCH$_3$    | No           | Yes            | Yes                         | C3′-endo                               | Intermediate           | Low                              | No               |
| 2′-H          | No           | No             | Yes                         | C2′-endo                               | Lowest                 | Low                              | Yes              |
| 2′-F          | No           | Yes            | Yes                         | C3′-endo                               | Low                    | Highest                           | No               |

$^a$Information for LNA taken from Petersen *et al.* (52) for conformational equilibria of ribose and 2′-substituted riboses (2′-OH, 2′-H and 2′-OCH$_3$), see Guschlbauer and Jankowski (53).

$^b$According to Cramer and Truhlar (54).
P RNA, except for the generally weaker substrate affinity of \textit{B. subtilis} P RNA and its relatively low apparent sensitivity to the 2'-H substitution at C-1.

**Effect of modifications at nt −1 on substrate binding by RNase P holoenzymes**

We further investigated binding of the modified substrates to \textit{E. coli} and \textit{B. subtilis} RNase P holoenzymes at 15 mM \textit{Ca}^{2+} but in the presence of a 5-fold lower monovalent salt concentration (0.2 M) than in RNA-alone assays. The presence of the protein cofactor increased and equalized substrate affinity (K\textsubscript{d} values between 1 and 13 nM; Table 2) and abolished the substrate affinity differences seen for \textit{E. coli} relative to \textit{B. subtilis} P RNA. The 2'-H (C-1) ptRNA was exceptional, as it was bound with 6-fold lower affinity than its control substrate by the \textit{B. subtilis} holoenzyme, thus even exceeding the defect of the 2'-OCH\textsubscript{3} (T-1) substrate (3.5-fold for \textit{E. coli}, 4.1-fold for \textit{B. subtilis} RNase P). Thus, a pronounced defect of 2'-H, not evident in the \textit{B. subtilis} P RNA-alone affinity assay, emerged in the holoenzyme context. LNA at nt −1 affected ground state binding to the holoenzymes insignificantly.

**Cleavage site selection**

\textit{Escherichia coli} P RNA is known to form alternative E–S complexes in response to structural changes in the enzyme and/or substrate RNA, which lead to aberrant cleavage usually one nucleotide upstream (m-1 site, between −2 and −1) of the canonical cleavage site (c0 site, between −1 and +1) (see e.g. 14,27). More rarely, aberrant cleavage at the m\textsubscript{+1} site (between nt +1 and +2) has also been observed in reactions catalyzed by \textit{E. coli} and \textit{B. subtilis} P RNA (Figure 2A) (8,19,22).

Except for the LNA (T-1) and 2'-OCH\textsubscript{3} (T-1) substrates, all ptRNA variants listed in Table 2 were processed exclusively at the c0 site by \textit{E. coli} P RNA (data not shown). The cleavage sites for ptRNA with LNA or 2'-OCH\textsubscript{3} at −1 were analyzed by co-electrophoresis of nuclease P1 hydrolysis ladders (P1 like RNase P generates 5'-phosphate termini) along with the products derived from \textit{E. coli} P RNA cleavage. This revealed cleavage of the LNA (T-1) ptRNA at the c0 and m\textsubscript{+1} sites (Figure 2B and C, lanes 2 and 9, respectively), whereas the 2'-OCH\textsubscript{3} (T-1) ptRNA was cleaved exclusively at the m\textsubscript{+1} site (Figure 2B and C, lanes 4 and 2, respectively). The 7-nt 5'-cleavage product with the 3'-terminal LNA sugar generated by cleavage at the c0 site (Figure 2B, lane 2) migrated somewhat faster than the corresponding all-ribose counterpart (Figure 2B, lane 6). Changes in the mobility of short oligonucleotides with 3'-terminal ribose modifications in high-resolution gels have been documented before (20,22).

The \textit{E. coli} RNAse P holoenzyme cleaved the LNA (T-1) and 2'-OCH\textsubscript{3} (T-1) ptRNAs with very low efficiency relative to the unmodified substrate (Figure 2C, lanes 5

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Table 2. Substrate ground state binding to \textit{E. coli} and \textit{B. subtilis} P RNAs and holoenzymes

| [Me\textsuperscript{2+}] | ptRNA | \textit{E. coli} P RNA | \textit{B. subtilis} P RNA |
|----------------------|-----------------|----------------------|----------------------|
|                      | K\textsubscript{d} (nM) | K\textsubscript{d,rel} | K\textsubscript{d} (nM) | K\textsubscript{d,rel} |
| 5 mM \textit{Sr}^{2+} | 2'-OH (T-1)     | 86 ± 22              | 1.0                   | –                     |
|                      | LNA (T-1)       | 1143 ± 232           | 13.3                  | –                     |
|                      | 2'-OCH\textsubscript{3} (T-1) | 4168 ± 733       | 48.5                  | –                     |
|                      | LNA (T-2)       | 295 ± 70             | 3.4                   | –                     |
|                      | 2'-OH (C-1)     | 48 ± 20              | 1.0                   | –                     |
|                      | 2'-F (C-1)      | 103 ± 49             | 2.1                   | –                     |
|                      | 2'-H (C-1)      | 2887 ± 874           | 60.1                  | –                     |
| 15 mM \textit{Ca}^{2+} | 2'-OH (T-1)     | 6.1 ± 2.3            | 1.0                   | 1.1 ± 0.11 |
|                      | LNA (T-1)       | 15.5 ± 3.7           | 2.5                   | 1.1 ± 0.34 |
|                      | 2'-OCH\textsubscript{3} (T-1) | 191.4 ± 5 | 31.4                  | 3.8 ± 0.64 |
|                      | LNA (T-2)       | 6.7 ± 3              | 1.1                   | –                     |
|                      | 2'-OH (C-1)     | 7.6 ± 1.3            | 1.0                   | 0.92 ± 0.21 |
|                      | 2'-F (C-1)      | 27.1 ± 3.7           | 3.6                   | 1.24 ± 0.35 |
|                      | 2'-H (C-1)      | 82.5 ± 11.3          | 10.9                  | 1.9 ± 0.77 |
| 80 mM \textit{Sr}^{2+} | 2'-OH (T-1)     | 5.9 ± 2.9            | 1.0                   | –                     |
|                      | LNA (T-1)       | 11.5 ± 4.2           | 1.9                   | –                     |
|                      | 2'-OCH\textsubscript{3} (T-1) | 211 ± 61             | 35.8                  | –                     |
|                      | LNA (T-2)       | 7.4 ± 2.4            | 1.25                  | –                     |
|                      | 2'-OH (C-1)     | 7.4 ± 2              | 1.0                   | –                     |
|                      | 2'-F (C-1)      | 29.9 ± 8.3           | 4.0                   | –                     |
|                      | 2'-H (C-1)      | 67.9 ± 16.4          | 9.2                   | –                     |

The analysis of ptRNA binding was performed using the spin column assay (10). The dissociation constant (K\textsubscript{d}) was measured in the presence of 50 mM MES/NaOH, pH 6.0, 1 M NH\textsubscript{4}OAc (P RNA) and Ca\textsuperscript{2+} (15 mM) or Sr\textsuperscript{2+} (5 or 80 mM) as the divalent metal ion; ptRNA binding to the holoenzyme was assayed at 50 mM Tris/CH\textsubscript{3}COOH, 0.2 M NH\textsubscript{4}OAc, 15 mM Ca\textsuperscript{2+}, pH 7.1. Relative K\textsubscript{d} values (K\textsubscript{d,rel}): individual K\textsubscript{d} divided by K\textsubscript{d} for the corresponding all-ribose ptRNA. K\textsubscript{d} values for the 2'-F (C-1) and 2'-H (C-1) substrates at 15 mM Ca\textsuperscript{2+} are consistent (within 2-fold) with K\textsubscript{d} values determined in a previous study under identical conditions (20).
Figure 2. Analysis of RNase P cleavage site selection; (A) Secondary structure of ptRNA acceptor stem ends; arrows assign the different RNase P (RNA) cleavage sites: c0, canonical site; m−1, m+1, aberrant sites. (B) Cleavage of ptRNA variants LNA (T-1), 2'-OCH3 (T-1) and 2'-OH (T-1) by E. coli P RNA (lanes 2, 4 and 6) and cleavage of the corresponding 24-mers (Figure 1A) used to construct the ptRNA variants by nuclease P1 (lanes 1, 3 and 5). Numbers on the left side indicate the length (nt) of fragments generated by nuclease P1; the corresponding P RNA cleavage sites are indicated in parentheses. The rectangles mark the missing or reduced P1 fragments due to the absence of a 2'-OH at the corresponding position. Hydrolysis by nuclease P1 was performed as described in the ‘Materials and Methods’ section. E. coli P RNA cleavage was performed in buffer A (1 M NH4OAc, 50 mM PIPES, pH 7 and 20 mM Mg2+) for 8 h (lane 2), 22 h (lane 4) and 10 min (lane 6) at 37°C. (C) Analysis of cleavage site selection by E. coli P RNA and holoenzyme. In lanes 1, 2, 8 and 9, processing by E. coli P RNA was performed in buffer A as in panel B; cleavage by the E. coli holoenzyme (lanes 4, 5, 11 and 12) was conducted in buffer B (50 mM MES, pH 6, 0.1 M KCl and 10 mM MgCl2); we also included control reactions in buffer B in the absence of the E. coli protein subunit (lanes 6, 7, 13 and 14), which illustrate the requirement for the protein cofactor under low salt conditions: the LNA and 2'-OCH3 substrates were cleaved by E. coli P RNA in buffer A, but not in buffer B (lanes 2 and 9 versus 7 and 14). The incubation time (at 37°C) was 10 min (lanes 1 and 8), 1 h (lanes 4, 6, 11 and 13), or 20 h (lanes 2, 5, 7, 9, 12 and 14). In lanes 3 and 10, a 5'-end-labeled 6-mer oligoribonucleotide (pCCCUUU) served as size control to assign cleavage at the m−1 site.

and 12 versus 4 and 11). Under the applied holoenzyme assay conditions (buffer B), essentially no cleavage of the LNA (T-1) and 2'-OCH3 (T-1) substrates was seen in the absence of the protein cofactor (Figure 2C, lanes 7 and 14). Thus, the protein exerted its known effect, i.e. improvement of activity under low salt conditions, but a severe catalytic defect caused by LNA (T-1) and 2'-OCH3 at nt −1 remained despite the presence of the protein. Interestingly, the protein switched the cleavage site for the 2'-OCH3 (T-1) ptRNA such that cleavage occurred predominantly at the m−1 site and not at the m+1 site as in the RNA-alone reaction (Figure 2C, lanes 2 versus 5). As in the RNA-alone reaction, the E. coli holoenzyme cleaved the LNA (T-1) substrate with low efficiency at the c0 and m+1 sites, and overexposure of gels even revealed some cleavage at the m−1 site (Figure 2C, lane 12). Processing inhibition by LNA and 2'-OCH3 at −1 also led to appearance of an aberrant cleavage product in reactions catalyzed by the E. coli holoenzyme (marked by asterisks in Figure 2C, lanes 5 and 12). This aberrant cleavage site, attributable to a minor alternative conformation of ptRNA Gly, was tentatively assigned (data not shown) to the anticodon arm. Indeed, cleavage of such weakly populated aberrant tRNA conformers was observed before by Kikuchi and coworkers and termed hyperprocessing (38). The underlying rearrangement of the tRNA structure apparently results from base pairing between the 3'-strand of the acceptor stem and nucleotides of the variable loop and/or the 3'-strand of the anticodon stem to form a hairpin-like structure (38–40).
for the *B. subtilis* RNA-alone reaction (Figure 3A, lanes 6–15) and the *B. subtilis* holoenzyme (Figure 3B, lanes 15–18). In conclusion, all four tested activities (*E. coli* and *B. subtilis* P RNAs and holoenzymes) cleaved the LNA (T-1) substrate with very low efficiency relative to unmodified ptRNA, and utilized the aberrant m₁ site in addition to the c₀ site. However, the 2'-OCH₃ (T-1) ptRNA was cleaved with very low efficiency at the c₀ site by *B. subtilis* P RNA and holoenzyme, whereas *E. coli* P RNA selected exclusively the m₁ site and the *E. coli* holoenzyme cleaved predominantly at m₋₁ and weakly at m₊₁.

**Rate constants of cleavage by *E. coli* P RNA**

All modifications introduced at nt −1 significantly affected the cleavage reaction catalyzed by *E. coli* P RNA. The rate constants for c₀ cleavage under single turnover conditions (*E >> S*) and rate-limiting chemistry (120 mM Mg²⁺, pH 5.5) decreased in the order 2'-OH (T-1) >> LNA (T-2) >> 2'-OH (C-1) >> 2'-F (C-1) >> 2'-H (C-1) in the ptRNA no cleavage at all was detected at the c₀ site (Table 3). The same hierarchy was seen at 20 mM Mg²⁺, pH 5.5, but at this lower Mg²⁺-concentration cleavage of the LNA (T-1) and 2'-OCH₃ (T-1) substrates was at the detection limit and thus not further analyzed. Rate differences were mitigated at pH 7 and 120 mM Mg²⁺, which was expected because at this higher pH value cleavage rates are generally faster, but another step than cleavage chemistry begins to limit the turnover of ptRNA substrates with 2'-OH at nt −1 (20). At lower Mg²⁺ concentrations (20 mM), the defects owing to ribose modifications at nt −1 were exacerbated, consistent with the documented Mg²⁺-binding defect caused by the absence of the 2'-hydroxy group at this location (8,20). Low efficiency cleavage of ptRNA 2'-OCH₃ (T-1) at the m₁ site was comparable at 20 and 120 mM Mg²⁺ (pH 7), although the endpoint of the reaction was lower at 20 mM Mg²⁺ (0.3 versus 0.55; Table 3).

**Rate constants of cleavage by *B. subtilis* P RNA**

In single turnover reactions (5 μM P RNA) at 120 mM Mg²⁺ and rate-limiting chemistry (pH 5.5) (10), the hierarchy of effects caused by the different ribose modifications equaled that observed for *E. coli* P RNA (cf. Tables 3 and 4), but the decreases in rate constant were more pronounced for the 2'-F (C-1) and particularly the 2'-H (C-1) ptRNA analyzed under the same conditions. Furthermore, the endpoint for c₀ cleavage of the LNA (T-1) substrate by *B. subtilis* P RNA was ~2.5% (Table 4), compared with 65% (fitted endpoint) and 42% (highest substrate conversion measured experimentally; Table 3) for the *E. coli* P RNA reaction. Another peculiarity was seen for *B. subtilis* P RNA-catalyzed cleavage of 2'-OCH₃ (T-1) ptRNA at the c₀ site and LNA (T-1) ptRNA at the m₁ site. Here, cleavage was already at saturation at the first time point (1 h; Table 4 and Figure 3A, lanes 6–15), with very low endpoints (1–4%; Table 4). A possible explanation is that weakly populated E–S conformers gave rise to these cleavages.

**Rate constants of cleavage by RNase P holoenzymes**

Rates in the RNase P holoenzymes reactions were measured under single turnover conditions at pH 6.0 and 4 mM Mg²⁺, or at 10 mM Mg²⁺ for the most inhibitory LNA and 2'-OCH₃ modifications. Effects were similar in reactions catalyzed by the *E. coli* and *B. subtilis* holoenzymes, with two major exceptions: (i) similar to the *B. subtilis* RNA-alone reaction, the defect caused by the 2'-H (C-1) ptRNA was much higher in the *B. subtilis*
Table 3. Observed (k_{obs}) and relative (k_{rel}) rate constants for cleavage of modified ptRNAs by E. coli P RNA

| ptRNA                | pH 5.5 | E. coli P RNA | pH 7 |
|----------------------|--------|---------------|------|
|                      |        | 20 mM Mg^{2+} | 120 mM Mg^{2+} | 20 mM Mg^{2+} | 120 mM Mg^{2+} |
|                      | k_{obs} (min^{-1}) | k_{rel} | k_{obs} (min^{-1}) | k_{rel} | k_{obs} (min^{-1}) | k_{rel} | k_{obs} (min^{-1}) | k_{rel} |
| 2'-OH (T-1)          | 0.9 ± 0.25 | 1.0 | 3.2 ± 1.1 | 1.0 | 8.7 ± 3.9 | 1.0 | 14.5 ± 6.5 | 1.0 |
| LNA (T-1) c_{0} site | n.a. | n.a. | (1 ± 0.5) x 10^{-3} | 3200 | (8 ± 5) x 10^{-4} | 10 875 | 0.024 ± 0.004 | 604 |
| LNA (T-1) m_{1} site | n.a. | n.a. | (2.8 ± 0.1) x 10^{-3} | 1143 | (3.1 ± 0.1) x 10^{-3} | 2806 | (2.2 ± 0.1) x 10^{-2} | 659 |
| 2'-OCH_{3} (T-1) c_{0} site | n.a. | n.a. | n.d. | ∞ | n.d. | ∞ | n.d. | ∞ |
| 2'-OCH_{3} (T-1) m_{1} site | n.a. | n.a. | n.d. | ∞ | n.d. | ∞ | n.d. | ∞ |
| LNA (T-2)            | 0.52 ± 0.02^b | 1.7 | 2.34 ± 0.16 | 1.4 | 2.9 ± 1 | 3000 | (3.1 ± 1) x 10^{-3} | 4677 |
| 2'-OH (C-1)          | 1.2 ± 0.1 | 0.8 | 5.2 ± 0.7 | 0.6 | 8.3 ± 0.5^c | 1.0 | 12.0 ± 2.0^b^e | 1.2 |
| 2'-F (C-1)           | (2.3 ± 0.9) x 10^{-2} | 39 | 0.5 ± 0.07 | 6.4 | 1.3 ± 0.08 | 6.7 | 7.0 ± 2.5 | 2.1 |
| 2'-H (C-1)           | (7 ± 3) x 10^{-4} | 1286 | (4.0 ± 0.7) x 10^{-2} | 80 | (2.8 ± 0.6) x 10^{-2} | 311 | 0.6 ± 0.15 | 24 |

n.a.: not analyzed; n.d.: cleavage not detectable. Cleavage experiments were performed under single turnover conditions (5 μM E. coli P RNA, <1 nM ptRNA, 1 M NH_{4}OAc and the indicated pH and Mg^{2+} concentration, essentially as described (30); for low efficiency cleavage of the LNA (T-1) and 2'-OCH_{3} (T-1) substrates, average values for the endpoint (L = limit) of the reaction determined by curve fitting and average values for the highest substrate turnover measured experimentally are given; k_{rel} is the ratio of the respective k_{obs} value to that of the 2'-OH (T-1) ptRNA. For the LNA (T-1) and 2'-OCH_{3} (T-1) substrates, cleavage is differentiated for the c_{0} and m_{1} sites; all other substrates were cleaved exclusively at the c_{0} site.  ^b Determined from the linear range of substrate turnover (no endpoint from curve fitting/ highest turnover experimentally measured).

Table 4. Observed (k_{obs}) and relative (k_{rel}) rate constants for cleavage of modified ptRNAs by B. subtilis P RNA and E. coli and B. subtilis RNase P holoenzymes

| ptRNA                | E. coli holoenzyme | B. subtilis P RNA | B. subtilis holoenzyme |
|----------------------|--------------------|-------------------|------------------------|
|                      | pH 6               | pH 5.5            | pH 6                   |
|                      | k_{obs} (min^{-1}) | k_{rel} | k_{obs} (min^{-1}) | k_{rel} | k_{obs} (min^{-1}) | k_{rel} |
| 2'-OH (T-1)          | 2.5 ± 0.5 | 1.0 | 2.64 ± 0.4 | 1.0 | 0.73 ± 0.07 | 1.0 |
| LNA (T-1) c_{0} site | Extent: 0.03 at 1 h; ≤ 0.2 at 20 h | 3 (1 ± 1) x 10^{-3} | 880 | Extent: 0.03 at 1 h; 0.05 at 20 h | 0.04 at 20 h | 0.05 at 20 h |
| LNA (T-1) m_{1} site | Extent: 0.03 at 1 h; ≤ 0.07 at 20 h | Extent: 0.03 at 1 h; 0.04 at 20 h | 0.04 at 20 h |
| 2'-OCH_{3} (T-1) c_{0} site | n.d. | n.d. | Extent: 0.01 at 1 h; 0.02 at 23 h | n.d. | n.d. | n.d. |
| 2'-OCH_{3} (T-1) m_{1} site | (6.0 ± 0.1) x 10^{-3} | 417 | (6.0 ± 0.1) x 10^{-2} | 33 | (6.0 ± 0.1) x 10^{-2} | 40 |
| 2'-OH (C-1)          | 2.5 ± 0.2 | 1.0 | 5.8 ± 0.1 | 0.5 | 1.57 ± 0.04 | 0.5 |
| 2'-F (C-1)           | 0.39 ± 0.06 | 6.4 | (6.0 ± 0.5) x 10^{-2} | 40 | 0.11 ± 0.02 | 6.6 |
| 2'-H (C-1)           | (7.5 ± 1.5) x 10^{-2} | 33 | 0.001 (L = -0.12)^b | 2640 | (L = -0.08)^b | 1570 |

n.d.: Cleavage not detectable. Cleavage experiments were performed under the following conditions: 5 μM B. subtilis P RNA, <1 nM ptRNA, 1 M NH_{4}OAc, 50 mM MES, pH 6 and 4 or 10 mM Mg^{2+} (E. coli/B. subtilis holoenzyme). Holoenzyme reconstitution and kinetic experiments were performed as described under Materials and Methods. Cleavage is differentiated for the c_{0} and m_{1} site in the case of the 2'-OCH_{3} (T-1) substrate, and for the c_{0} and m_{1} site in the case of ptRNA LNA (T-1); all other substrates were cleaved exclusively at the c_{0} site. For the 2'-OCH_{3} (T-1) and LNA (T-1) substrates, cleavage occurred to very low extents [in most cases with saturation already reached at the first data point (1 h)]; thus, only the extent of cleavage instead of cleavage rate constants is given here.

^b Determined from the linear range of substrate turnover (no endpoint from curve fitting/ highest turnover experimentally measured).
than *E. coli* holoenzyme reaction; (ii) weak cleavage of the 2′-OCH₃ substrate at the c₀ site occurred with *B. subtilis* but not with *E. coli* RNase P, as mentioned before. As observed for *B. subtilis* P RNA (see above), the extent of cleavage for the LNA (T-1) and 2′-OCH₃ (T-1) substrates by both holoenzymes was extremely low (Table 4).

Mg²⁺ dependence of LNA (T-1) ptRNA cleavage by *E. coli* P RNA at sites c₀ and m₊₁

The LNA (T-1) ptRNA was simultaneously processed at the c₀ and m₊₁ site by *E. coli* P RNA (Figure 2B and C). Simultaneous processing at different sites can be attributed to parallel reaction pathways (14,19,22,23,41) as illustrated in Scheme 1. Here we investigated the Mg²⁺ dependence of c₀ and m₊₁ cleavage, with the following outcome: First, at low [Mg²⁺], the fraction of substrate cleaved at the c₀ site (F₀) was substantially lower at early versus late time points (Figure 4A); with increasing [Mg²⁺], these differences largely disappeared (Figure 4B–D). This indicates that the two (EₛS) complexes reacted with different single exponential decay rate constants (kₒbs) at low [Mg²⁺] and similar rate constants at high [Mg²⁺]. Second, F₁ values at experimental endpoints (last measured time point) and theoretical endpoints (derived from the single exponential decay curve fit) were lower at 20 mM Mg²⁺ (F₁ < 0.7) than at e.g. 120 mM Mg²⁺ (F₁ > 0.8). Third, plotting kₒbs as a function of [Mg²⁺] (Figure 4E) and allowing for a cooperative Mg²⁺ dependence at both cleavage sites (see Materials and methods section) revealed a higher Mg²⁺ dependence for cleavage at the c₀ site (K₁/₂ = 184 mM) than at the aberrant m₊₁ site (K₁/₂ = 129 mM), consistent with a severe metal ion binding defect for c₀ cleavage in the presence of LNA at nt −1 (the defect may include contributions from Kₛₑₙₖ or Kₛₑₚ see Scheme 1). A double logarithmic plot of the same data illustrates that the kₒbs for c₀ cleavage was slower than that for m₊₁ cleavage at low [Mg²⁺], whereas this relation was reversed at high [Mg²⁺] (Figure 4F).

pH dependence of c₀ and m₊₁ cleavage pathways

In the hydrolysis reaction catalyzed by RNase P, a linear relationship of log kₒbs and pH with a slope of ~1 is taken as evidence for the catalytic step being rate-limiting (8,9,11,20,22,42). This pH dependence, consistent with a single ionization, is thought to reflect ionization of a metal ion-coordinated water molecule acting as the nucleophile in this phosphodiester hydrolysis reaction (8,11,22). Analysis of the pH dependence of log kₒbs for cleavage of the LNA (T-1) ptRNA at sites c₀ and m₊₁ (Figure 5) revealed a slope of ~0.9 up to pH 7 for c₀ cleavage, after which the curve inflected (Figure 5D), consistent with another step limiting the rate of the reaction above pH 7 (9). In contrast, cleavage at the m₊₁ site was linear over the entire pH range (5.5–8.1) but with a slope of only 0.45 (Figure 5D), indicative of another step than cleavage chemistry limiting the reaction at all tested pH values.

The pH variation experiments were performed at an intermediate Mg²⁺ concentration (60 mM). Interestingly, lowering the pH caused similar effects as lowering the Mg²⁺ concentration at neutral pH (Figure 4): at low pH, m₊₁ cleavage was faster than c₀ cleavage, while this relation was reversed above pH 7 (Figure 5D); and, related to this, at low versus higher pH (Figure 5A–C) the fraction of substrate cleaved at the c₀ site (F₀) was substantially lower at early than late time points.

DISCUSSION

LNA and 2′-OCH₃ modifications at nt −1: blockage of steps after substrate ground state binding

A 2′-OCH₃ modification at position −1 was previously reported as the most inhibitory substrate ribose modification in the reaction catalyzed by *E. coli* P RNA (8). In this previous study, a ptRNA with a single nucleotide as 5′-leader was miscleaved at the m₊₁ site as observed here. The substrate with a 1-nt 5′-leader precluded analysis of m₋₁ cleavage, but as shown here this site appears to be blocked by 2′-OCH₃ at nt −1 in *E. coli* P RNA-alone reactions with ptRNAs carrying longer 5′-leaders (Figure 2). The inhibitory strength of 2′-OCH₃ at −1 exceeded that of a 2′-deoxy modification, consistent with a steric effect caused by the extra methyl group (8).

Our results obtained with *E. coli* P RNA demonstrate that the transition state for canonical cleavage can be achieved in the presence of a fixed C3′-endo conformation at nt −1 (Figure 2B and C), indicating that a C2′-endo conformation prevailing at this position based on NMR analyses (26) is not an absolute requirement for catalysis to occur. It is unclear if the low efficiency of c₀ cleavage with LNA at −1 is primarily due to the steric effect of the extra methylene group or includes substantial contributions from the lack of ribose flexibility at this position. Also, the strong inhibitory effect of LNA might stem from the fact that the free electron pairs of the 2′-oxygen in LNA are orientationally constrained; this may impair the correct positioning of the Mg₂⁺hydrate complex containing the water molecule which we propose to protonate the 3′-oxymion leaving group (Figure 6A). The moderate effect of LNA at −1 on substrate ground state binding (Table 2) relative to 2′-OCH₃ suggests that the extra methylene group causes less steric constraints than the 2′-OCH₃ group; likewise, the LNA (T-1) substrate was cleaved by *E. coli* P RNA at the c₀ site, whereas c₀ cleavage was not seen at all in the presence of the 2′-OCH₃ (T-1) modification (Figure 2). However, in contrast to the mild defect in substrate ground state binding, the high Mg²⁺ requirement for c₀ cleavage of LNA (T-1) ptRNA at saturating enzyme concentrations (Figure 4E) is consistent with a severe block in the binding of catalytically relevant Mg₂⁺. Thus, the defect is on Kₛₑₙₖ and/or kₒbs rather than on Kₛₑₚ (Scheme 1). It should also be noted that the effect of 2′-OCH₃ at −1 on ground state binding (30-fold at 15 mM Ca²⁺; Table 2) was still moderate compared with the complete blockage of cleavage at site c₀ for this modification. Thus, we favor a model according to which the extra methylene or methyl group causes a steric interference with the nearby bound catalytic Mg₂⁺ during its re-coordination on the way to the transition state for
cleavage (Figure 6). To which extent the defect is exerted at the level of $K_{\text{conf}}$ or bond breakage ($k_c$) cannot be discerned on the basis of our data.

Despite their similar chemical nature, the effects of LNA and 2'-OCH$_3$ substitutions at nt $-1$ differed in detail. Relevant towards understanding these differences could be the fact that the 2'-O-methylene group in LNA occupies a position in space that coincides only with one of the many positions that can be adopted by the 2'-OCH$_3$ group. The latter can rotate around the C2'-O2' bond, and
additional orientations result from the equilibrium between C3'- and C2'-endo sugar puckers (Figure 7). The orientation flexibility of the 2'-OCH3 group relative to the methylene group in LNA may explain the differential effects of the two substitutions on substrate ground state binding (Table 2). In addition, we recently showed that LNA at C01 stabilizes base pairing between T-1 and U73 (see Figure 1) (43). Thus, in the ground state the LNA (T-1) substrate is expected to adopt an extended acceptor stem structure to a larger degree than the 2'-OCH3 (T-1) substrate. However, based on our recent results, the breaking of an additional base pair at the end of the acceptor stem leaves the energetic barrier for reaching the transition state of the chemical step for cleavage at the canonical (c0) phosphodiester essentially unaffected (43). An effect on the aberrant m+1 and m-1 cleavage pathways cannot be excluded.

Regarding metal-dependent substrate ground state binding to E. coli P RNA, 2'-H and 2'-OCH3 substituents at nt -1 were most deleterious, whereas LNA and 2'-F at this position had, overall, mild effects. We infer (i) that the absence of an H-bond donor function at the 2'-position of nt -1, as with LNA, is not crucial for substrate ground state binding; (ii) an extra 2'-methyl(ene) group has a mild effect if its position is constrained as in LNA; (iii) the mild effect of 2'-F can be explained in several ways; we favor the idea that the electronegative 2'-fluorine, as 2'-OH or LNA but in contrast to 2'-H, can accept a hydrogen bond from a metal ion-coordinated water molecule (20); this should also be the case for 2'-OCH3, but here steric interference seems to dominate; (iv) a locked C3'-endo sugar pucker at nt -1 is compatible with high affinity ground state binding, indicating that the sugar pucker at this position has not to be constrained in C2'-endo for ground state binding, a possibility that arose from our previous NMR study (26). The observed effects were similar for B. subtilis P RNA, but apart from its lower substrate affinity, this RNA was less sensitive to the 2'-deoxy modification at nt -1 (Table 2). However, sensitivity to the 2'-deoxy substitution in the B. subtilis holoenzyme reaction, where the effect of 2'-H even exceeded that of 2'-OCH3. Possibly, the 2'-deoxy effect was masked in the B. subtilis P RNA-alone context owing to the generally low substrate affinity of this RNA in the absence of the protein cofactor. In summary, our results are consistent with the idea that acceptance of a hydrogen bond from a metal ion-coordinated water molecule is a key feature of the 2'-substituent at nt -1 during substrate ground state binding.

Figure 5. Kinetics of ptRNA LNA (T-1) processing by E. coli P RNA at different pH values. Cleavage experiments were performed under single turnover conditions (5 μM P RNA, <1 nM ptRNA) in the presence of 1 M NH4OAc, 50 mM MES, PIPES or HEPES buffer and 60 mM Mg2+.

(A–C) Examples of kinetic analyses at low, intermediate and the highest pH tested. For further details, see legend to Figure 4. (D) Plots of log kobs dependence on pH. For cleavage at the c0 site (open circles), the first four data points gave a linear fit with a slope of 0.92 ± 0.08; for cleavage at the m+1 site (filled triangles), all five data points fit to a line with a slope of 0.45 ± 0.04.
Figure 6. Transition state models for E. coli RNase P RNA-catalyzed phosphodiester hydrolysis at the canonical c₀ site with a ribose (A) or LNA (B) at nt −1. The model in (A) [adapted from ref. (55)] predicts that both Mg₁ and Mg₂₈ directly coordinate to the pro-Rp oxygen, with Mg₁ coordinator the OH⁻ nucleophile, and Mg₂₈ interacting with the 2'-oxygen and the 2'-hydroxy group at nt −1 via a water molecule of its hydration shell; we propose this water molecule to donate the proton to the 3'-oxyanion leaving group. The ribose at nt −1 is assumed to mainly populate the 2'-endo sugar pucker based on NMR analyses of acceptor stem mimics (26). (B) Here, the −1 nucleotide is LNA with the sugar moiety fixed in the C3'-endo conformation; the steric interference caused by LNA or 2'-OCH₃ at position −1 (indicated by the dashed curved line) likely disrupts the binding site for Mg₂₈. The model differs from others [e.g. (56)] by assuming that only two instead of three metal ions are essential; also, in our model Mg₂₈ is proposed to simultaneously interact with the pro-Rp oxygen, the 3'-oxygen and the 2'-OH, whereas the latter interaction is assigned to the third metal ion in the model by Kazantsev and Pace (56). Group I intron ribozymes have a similar size as bacterial P RNA and also catalyze phosphoryl transfer reactions yielding products with 3'-OH termini. The two-metal-ion mechanistic model we propose here is essentially identical to that for the second step of group I intron splicing inferred from a crystal structure of the Azoarcus sp. ptRNAαᵦ mRNA group I intron (57), with two differences: (i) the contact of Mg₂₈ to the 2'-OH is outer-sphere in the RNase P reaction, but inner-sphere in the group I intron reaction; (ii) the nucleophile is a hydroxide in the RNase P reaction, but the terminal 3'-OH of the 5'-exon in the second step of the group I intron reaction (57). Despite striking similarities, one mechanistic difference has been observed in phosphodiester hydrolysis reactions catalyzed by the Tetrahymena group I intron-derived ribozyme and E. coli P RNA; whereas a 2'-amino modification at ptRNA position −1 inhibited cleavage at the canonical site with increasing protonation of the 2'-amine in the P RNA reaction (consistent with electrostatic repulsion of a metal ion), the reverse was seen with the Tetrahymena ribozyme; in the latter reaction, protonation of the 2'-amine at equivalent position rather stimulated cleavage at the canonical site (20,58). The molecular basis of this difference is not clear at present.

**Cleavage fidelity**

For simple class I tRNAs, such as the ptRNAαᵦ Gly used here or B. subtilis ptRNAαᵦ (22), it has previously been discussed that interactions between E. coli P RNA and the ptRNA cleavage site are of redundant nature. For example, base identity switches at nt −1 that weaken the interaction with A248 of E. coli P RNA, 2'-substitutions at nt −1 (e.g. 2'-H or 2'-F), an Rp-phosphorothioate substitution at the scissile phosphodiester or a G292C or G293C mutation in the CCA-binding site of P RNA do not by themselves lead to substantial miscleavage (22). However pairwise modification, such as 2'-deoxy modification at nt −1 combined with an A248U or G292C mutation resulted in substantial miscleavage (20,22). In contrast, LNA and 2'-OCH₃ at nt −1 induced severe miscleavage in the absence of any other modification, emphasizing the deleterious effect of the extra methyl(ene) group on cleavage chemistry or a preceding conformational rearrangement of E*S complexes (Scheme 1).

**Differences in the Mg²⁺ and pH dependence of the correct (c₀) and mis-cleavage (m₁) pathways**

Harris and coworkers recently demonstrated communication between a 2'-deoxy modification at nt −1 and weakening of the A248/nt −1 contact. Alterations at both sites, such as combination of 2'-deoxy U-1 ptRNAαᵦ with the A248U mutant E. coli P RNA, resulted in cleavage at the c₀, m₁ and also at the m₁ site (22). When studying the proportion of cleavage events at the c₀ and m₁ site as a function of the Mg²⁺ concentration, the authors observed increased utilization of the m₁ site with increasing Mg²⁺ concentration. They concluded that the pathway leading to cleavage at the c₀ site exhibits a greater apparent affinity for Mg²⁺ than the pathway leading to cleavage at the m₁ site (22). We performed a similar analysis for the m₁ site in the presence of LNA at −1 (Figure 4) and observed a reverse correlation: here the amount of cleavage at the c₀ site increased with increasing Mg²⁺, indicating that the LNA modification lowers the apparent affinity for Mg²⁺ of the canonical cleavage pathway below that of the pathway leading to m₁ cleavage.

The same authors (22) observed that F₉ remained essentially constant over time in E. coli P RNA cleavage of ptRNAs with 2'-deoxy modifications at nt −1 and/or mutations at A248 of P RNA. Thus, they assumed that cleavages at the canonical and aberrant site (m₁ in their case) follow the same single exponential decay rate (kobs) calculated from the sum of the c₀ and m₁ site cleavage products accumulating at individual time points. Using F₉ and kobs, the apparent first-order rate constant for
cleavage ($k_{\text{app}}$) at a particular site was calculated according to $k_{\text{app}}[c_0] = F_c \times k_{\text{obs}}$, and $k_{\text{app}}[m+1] = (1 - F_c) \times k_{\text{obs}}$ [for details, see ref. (22)]. Although this assumption also holds for our data obtained at Mg$^{2+}$ concentrations between 60 and 200 mM (Figure 4F), it is not applicable particularly to the $F_c$ and $k_{\text{obs}}[c_0]$ data obtained under conditions of low [Mg$^{2+}$] and low pH (see Figures 4A, F and 5A). Changes in $F_c$ and $k_{\text{obs}}[c_0]$ may include contributions from the equilibria described by $K_c$ and $K_{\text{conf}}[c_0]$ (and the corresponding forward and back rate constants) as well as $k_{c_0}$ (see Scheme 1). Our findings ($F_c$ increases with [Mg$^{2+}$], $k_{\text{obs}}[c_0] < k_{\text{obs}}[m+1]$ at low [Mg$^{2+}$] and pH, $k_{\text{obs}}[c_0] > k_{\text{obs}}[m+1]$ at high [Mg$^{2+}$] and high pH) can be explained within the following framework (also illustrated in Scheme 2): When enzyme and substrate are mixed, rapid equilibria of (E$\cdots$S)$_{c_0}$ and (E$\cdots$S)$_{m+1}$ form, which commit the complexes to reaction along the different pathways. Partitioning between (E$\cdots$S)$_{c_0}$ and (E$\cdots$S)$_{m+1}$ is sensitive to the Mg$^{2+}$ concentration. At low [Mg$^{2+}$], the ratio of equilibrium binding constants $K_{c_0}/K_{m+1}$ (Scheme 1) exceeds that at higher [Mg$^{2+}$], explaining why $F_c$ increased with [Mg$^{2+}$] (Figure 4A–D). The rate constant for bond breakage at the $c_0$ site ($k_{c_0}$) and/or the preceding conformational step ($K_{\text{conf}}[c_0]$; see Scheme 1) have a higher Mg$^{2+}$ requirement than that for $m+1$ cleavage (Figure 4F). Finally, different steps are rate-limiting for cleavage at sites $c_0$ and $m+1$, as inferred from the different pH dependencies between pH 5.5 and 7.0 (Figure 5D).

Suppression of $m+1$ cleavage

Another idiosyncrasy of LNA or 2'-OCH$_3$ at $-1$ is the complete absence of cleavage by *E. coli* P RNA at the $m+1$ site (Figure 2B and C), which represents the prevailing miscleavage site [e.g. (44)]. Pan and colleagues showed that a 2'-deoxy substitution at $-1$ in ptRNA not only decreased the cleavage rate constant by *B. subtilis* P RNA 240-fold for the $c_0$ site but also 470-fold for the $m+1$ site and 210-fold for another aberrant site (between nt +2 and +3). This finding suggested a contact between the ribozyme and the ptRNA substrate 2'-OH at $-1$ which promotes a conformational rearrangement required for catalysis at both the $c_0$ as well as at aberrant sites, such as the $m+1$ site (19,24). Blockage of cleavage at the unmodified $m+1$ site owing to LNA or 2'-OCH$_3$ at $-1$ would be consistent with the findings of Loria and Pan.
Based on their model, utilization of the m+1 site in E. coli P RNA-catalyzed cleavage of ptRNA LNA (T-1) or 2'-OCH3 (T-1) may suggest that the 2'-OH group at nt +1 took over the role of that at nt −1 in promoting the conformational step preceding catalysis (Scheme 1). Likewise, redirection of cleavage from the m+1 to the m−1 site in the 2'-OCH3 (T-1) substrate when cleaved by the E. coli RNase P holoenzyme (Figure 2C) may be due to the protein cofactor allowing utilization of the 2'-OH at nt −2 as anchor point for the conformational step (24).

**pH dependence of c0 and m+1 cleavage by E. coli P RNA**

The shallow pH dependence of log kobs for cleavage at the m+1 site is consistent with one or more slow steps upstream of the hydrolysis step in the misincorporation pathway, such as conformational rearrangements. Our findings and those of others (22,45) indicate that cleavage at different sites (c0, m−1, m+1 or others) has different pH and metal ion dependencies and involves different contributions from rate-limiting steps. Accordingly, E–S complexes leading to cleavage at the different sites are structurally distinct. Cleavage at the c0 site has usually the highest apparent metal ion affinity, which still pertains to 'mild' modifications, such as 2'-deoxy at nt −1. However, more deleterious modifications, such as LNA or 2'-OCH3 at −1, can break this rule (Figure 4E). Considering that LNA at nt −1 had little effect on substrate binding in the ground state, this modification is a paradigm for a modification that specifically inhibits formation of later steps along the cleavage pathway.

**Cleavage of modified substrates by B. subtilis P RNA**

Loria and Pan (19) have studied processing of yeast ptRNA Phe by B. subtilis P RNA, representing a tRNA with a canonical 7-bp acceptor stem as the ptRNA Gly used in our investigation (Figure 1). They observed substantial miscleavage of ptRNA Phe, particularly at the m−1 site and even in the absence of any ribose modifications. In contrast, we did not observe significant miscleavage by B. subtilis P RNA for any of the ptRNA Gly variants used here, except for the LNA (T-1) substrate which gave rise to cleavage at the m+1 site (Table 4). Even the 2'-OCH3 (T-1) variant was cleaved exclusively at the c0 site (Figure 3), in contrast to E. coli P RNA which entirely miscleaved this substrate. Thus, we observed little propensity of B. subtilis P RNA to miscleave the ptRNA Gly substrate. The discrepancy to the previously observed miscleavage of yeast ptRNA Phe (19) may be explained by the fact that yeast ptRNA Phe has a rather labile acceptor stem with 3A–U and 1G–Ubp in a row in its lower part; in comparison, the acceptor stems of B. subtilis tRNA Phe molecules consist of 6G–C and a single A–Ubp, which is more typical of bacterial tRNAs. Low rigidity of the tRNA body is expected to produce conformational heterogeneity, which would explain increased miscleavage. This further raises the question if miscleavage at a specific site necessarily proceeds along the same mechanistic pathway for ptRNAs with rigid and less rigid tRNA bodies.

**Cleavage of modified substrates by the E. coli RNase P holoenzyme**

In several previous analyses, essentially identical misincorporation patterns were observed in E. coli P RNA-alone and holoenzyme reactions. This pertains, for example, to the disruption of the 3'-CCA interaction by mutation (36,46), or to reactions utilizing a A248U mutant P RNA and a ptRNA with a 2'-deoxy-U at nt −1 (41). However, in a study investigating the effect of a substrate 2'-amino modification at nt −1 on cleavage by E. coli RNase P (RNA), the P protein suppressed misincleavage at the m−1 site and redirected cleavage to the c0 site (21). This suggests that effects of the P protein on cleavage fidelity depend on the specific structural alteration.

The cleavage site of the 2'-OCH3 (T-1) substrate was almost entirely shifted from the m−1 to the m−1 site in the E. coli holoenzyme versus P RNA-alone reaction (Figure 2C, lanes 2 and 5). With LNA at −1, also some cleavage at the m−1 site appeared in the presence of the protein cofactor (Figure 2C, lanes 9 and 12). Evidently, the protein, by providing additional substrate binding energy and by enhancing the affinity of key metal ions, modulates the activation barriers of pathways leading to cleavage at different sites. The differences in mono- and di-valent salt conditions used in RNA-alone and holoenzyme reactions may also have contributed to the observed protein-dependent changes, resulting in differential effects on the individual misincorporation pathways. Specifically, the shift of cleavage site for the 2'-OCH3 (T-1) substrate may be explained by (EoS)m−1 complex formation favored over that of (EoS)m−1 complexes in the context of the holoenzyme reaction, and/or by a differential effect of the protein on the individual equilibrium docking steps [(EoS) → (EoS)*; Scheme 1] preceding catalysis (24). The protein might also affect active site architecture more profoundly than as yet thought.

**Differences between E. coli and B. subtilis RNase P (RNA)**

Architectural differences between bacterial P RNAs of type A (for 'ancestral') and type B (for 'Bacillus') (33) are associated with biochemical differences. The B. subtilis RNase P holoenzyme binds ptRNA with a much higher affinity than mature tRNA, but this discrimination against mature tRNA is much less pronounced for E. coli RNase P (34,35). RNase P enzymes containing the B. subtilis P RNA are less prone to aberrant cleavage than those including E. coli P RNA (36), and B. subtilis P RNA alone binds ptRNA substrates weaker than its E. coli counterpart (37, this study). The E. coli protein reduces the Mg2+ concentration requirement for tertiary structure formation of P RNA and increases the melting temperature at physiologically relevant Mg2+ concentrations (37). In contrast, the B. subtilis protein does not exert such stabilizing effects on its cognate or E. coli P RNA (37). Taking these reported differences between the two RNase P model systems into account, it is not surprising that we saw differences and commonalities (for details, see Results). Differences included a stronger defect caused...
by 2'-H at nt -1 on the *B. subtilis* versus *E. coli* holoenzyme, and deviating cleavage site selection for the 2'-OCH₃ substrate; major commonalities included the severe catalytic block by LNA and 2'-OCH₃ and, for the LNA substrate, the occurrence of low efficiency parallel cleavage pathways leading to hydrolysis at the c₀ and m₀ sites. The differences illustrate the complex interplay of functional groups and of thermodynamic as well as kinetic parameters in this RNA-catalyzed and protein-assisted phosphodiester hydrolysis reaction. Compared with protein enzyme orthologs with rigid and strongly conserved active sites, type A and B RNase P enzymes appear more as homologs that still catalyze the same reactions, but with distinct architectural and non-identical mechanistic concepts.

**The P protein is unable to compensate the catalytic block by LNA**

The protein subunit of bacterial RNase P increases substrate affinity and specificity (47,48), stabilizes the local RNA conformation (37), preorganizes S- and C-domain orientation (24) and increases the affinity of local RNA conformation (37), preorganizes S- and substrate affinity and specificity (47,48), stabilizes the functional groups and of thermodynamic as well as sites. The differences illustrate the complex interplay of functional groups and of thermodynamic as well as kinetic parameters in this RNA-catalyzed and protein-assisted phosphodiester hydrolysis reaction. Compared with protein enzyme orthologs with rigid and strongly conserved active sites, type A and B RNase P enzymes appear more as homologs that still catalyze the same reactions, but with distinct architectural and non-identical mechanistic concepts.

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