Critical domain interactions for type A RNase P RNA catalysis with and without the specificity domain

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Abstract

The natural trans-acting ribozyme RNase P RNA (RPR) is composed of two domains in which the catalytic (C-) domain mediates cleavage of various substrates. The C-domain alone, after removal of the second specificity (S-) domain, catalyzes this reaction as well, albeit with reduced efficiency. Here we provide experimental evidence indicating that efficient cleavage mediated by the Escherichia coli C-domain (Eco CP RPR) with and without the C5 protein likely depends on an interaction referred to as the "P6-mimic". Moreover, the P18 helix connects the C- and S-domains between its loop and the P8 helix in the S-domain (the P8/ P18-interaction). In contrast to the "P6-mimic", the presence of P18 does not contribute to the catalytic performance by the C-domain lacking the S-domain in cleavage of an all ribo model hairpin loop substrate while deletion or disruption of the P8/ P18-interaction in full-size RPR lowers the catalytic efficiency in cleavage of the same model hairpin loop substrate in keeping with previously reported data using precursor tRNAs. Consistent with that P18 is not required for cleavage mediated by the C-domain we show that the archaeal Pyrococcus furiosus RPR C-domain, which lacks the P18 helix, is catalytically active in trans without the S-domain and any protein. Our data also suggest that the S-domain has a larger impact on catalysis for E. coli RPR compared to P. furiosus RPR. Finally, we provide data indicating that the absence of the S-domain and P18, or the P8/ P18-interaction in full-length RPR influences the charge distribution near the cleavage site in the RPR-substrate complex to a small but reproducible extent.

Introduction

Almost all tRNAs carry a phosphate at their 5' ends due to the action of the endoribonuclease RNase P. Bacterial RNase P consists of one protein (C5), and one RNA subunit [1]. The composition of archaeal and eukaryal RNase P is more complex where the sole RNA subunit binds several proteins [2, 3]. Available data suggest that the catalytic activity resides in the RNA irrespective of origin, and the RNA alone can cleave various substrates in the absence of
protein at high ionic strength [2, 4–7]. However, recent data demonstrate the presence of a protein only RNase P activity (PRORP), for example in human mitochondria and in Arabidopsis thaliana, that possesses the capacity to cut precursor tRNAs at the same site as RNase P [8, 9].

On the basis of secondary structure RNase P RNA (RPR) can be divided into different types. Type A [ancestral type; exemplified by Escherichia coli (Eco) RPR] and type B (Bacillus type) are the two main types among the bacteria [10]. Type A also exists in Archaea, as exemplified by Pyrococcus furiosus RPR, Pfu RPR [6]. Other types such as M and T have also been identified in Archaea [11–13]. Irrespective of type, two major domains can be identified, the specificity (S-) and the catalytic (C-) domains albeit in type T RPR the S-domain is degenerated [12–16]. The S-domain provides the binding site (TBS; TSL-binding site) for the pre-tRNA T-stem/loop region referred to as TSL and the interaction to the TSL/TBS-interaction [16–19]. The catalytic activity is associated with the C-domain and with respect to bacterial RPR substantial activity is retained when the S-domain is deleted [20–23]. It has also been suggested that the catalytic activity of archaeal RPR is associated with the C-domain [6, 24]. Although, a cis construct composed of the Methanocaldococcus jannaschii (archaeal type M) C-domain and pre-tRNA is catalytic [11] no cleavage activity in trans without protein has yet been demonstrated for an archaeal RPR lacking the S-domain. For type A RPR two interactions play important roles, the intra-domain P6-interaction in the C-domain and the P8/18-interaction involved in connecting the S- and the C-domains [18, 25, 26]. However, Pfu RPR lacks P18 and consequently the P8/18-interaction is missing [6]. The P6- and P8/18-interactions are absent in type B RPR, however, the intra-domain contact L5.1/L15.1 may fulfill the role of P6 in type B [10, 27]. Of interest, in this context we note that type T RPRs have degenerated S-domains but modeling suggest that P6 is present while P18 is absent [12, 13].

Disruption of the P6- or P8/18-interactions has been reported to affect the catalytic performance of the Eco RPR and the type A Thermus thermophilus RPR, Tth RPR [18, 28–30]. Compared to the full-size type A Eco RPR the cleavage efficiency of a model hairpin loop substrate by Eco CP RPR_{wt} is reduced almost 500-fold ([23]; see also Refs [20, 22]) while deleting the S-domain in the type B Bacillus subtilis (Bsu) reduces the catalytic performance several thousand-fold compared to full-size Bsu RPR [21]. Within type A RPR the C-domain residues in the P15-17 region interact with the 3' end of the precursor substrate and from the crystal structure it can be inferred that their structural position is influenced by the P6-interaction in full-size RPR [18]. Removing the P15-17 region in full-size Eco RPR_{wt}, which also disrupts the P6-interaction, results in almost a 50000-fold reduction in catalytic efficiency [9]. For Eco CP RPR_{wt} no activity could be detected upon deleting P15-17 [22]. Inspection of the Eco RPR_{wt} structure reveals the possibility that residues involved in formation of P6 in full-size RPR can interact and resulting in an interaction mimicking P6 in Eco CP RPR_{wt}, for convenience referred to as the "P6-mimic". Hence, we were interested in understanding whether the "P6-mimic" is formed and if so does it contribute to Eco CP RPR mediated catalysis. Moreover, deleting the S-domain results in disruption of the P8/ P18-interaction. This enabled us to assess the contribution (if any) of P18 to catalysis in an Eco CP RPR context and thereby get insights to its role during catalysis also for full-size RPR. This is of specific interest since the type A Pfu RPR_{wt} lacks P18 and show lower activity than Eco RPR_{wt} in the RNA alone reaction [6, 31]; see also Refs [24, 32]. We were therefor also interested in whether a Pfu RPR lacking the S-domain retains its catalytic activity even in the absence of protein.

Our data indicate that a "P6-mimic" is likely to be present in Eco CP RPR_{wt}, and that its disruption reduce the cleavage efficiency. We also provide data suggesting that P18 does not influence the catalytic performance of Eco CP RPR_{wt} in cleavage of all ribo model hairpin loop.
substrates when the C5 protein is absent, which is not the case for full-size type A RPR. However, in the presence of the C5 protein a modest reduction in cleavage activity for Eco CP RPR<sub>wt</sub> was detected upon deleting P18. Our data also show that deletion of the S-domain of Pfu RPR resulted in an RPR that is catalytically active in the absence of proteins. We also found that deletion as well as disruption of the P8/ P18-interaction in full-size Eco RPR lowers the cleavage efficiency of a model substrate. On the basis of our data combined with the fact that the P8/ P18-interaction (or P6) is not in direct vicinity of where substrate cleavage occurs we raise the possibility that the P8/ P18-interaction acts as a structural mediator between the TSL/ TBS-interaction site and the active center leading to positioning of chemical groups and Mg<sup>2+</sup> that ensures correct and efficient cleavage.

### Materials and methods

#### Preparation of substrates and RPR

The substrates were purchased from Dharmacon, USA and were purified on a 15% (w/v) denaturing PAGE gel followed by an overnight Bio-Trap extraction (Schleicher and Schuell, GmbH, Germany; Elutrap in USA and Canada) and phenol-chloroform extraction. γ-ATP 5’ end-labeled substrates were generated and gel-purified using standard protocols.

The genes encoding full-size Eco RPR<sub>wt</sub> (M1 RNA), Eco CP RPR<sub>wt</sub> and Pfu RPR<sub>wt</sub> have previously been described [22, 31, 33]. The genes encoding the variants Eco RPR<sub>P18CUUG</sub>, Eco RPR<sub>G235</sub>, Eco CP RPR<sub>C83C84</sub>, Eco CP RPR<sub>G278G279</sub>, Eco CP RPR<sub>delP18</sub>, Eco CP RPR<sub>delP18P3Mini</sub>, Eco CP RPR<sub>31</sub>, Eco CP RPR<sub>31delP18</sub> and Pfu CP RPR<sub>wt</sub> have been generated following the same procedure as outlined elsewhere [22, 31, 33] using the Eco CP RPR<sub>wt</sub> and Pfu CP RPR<sub>wt</sub> genes as template and appropriate oligonucleotides. Eco RPR<sub>delP18</sub> was generated by replacing the 3′ half of Eco RPR<sub>wt</sub> with the 3′ half of Eco CP RPR<sub>delP18</sub> using appropriate restriction enzymes. The different RPRs were generated as run-off transcripts using T7 DNA-dependent RNA polymerase and PCR-amplified templates [34, 35]. The C5 protein was purified as described in [34, 36].

#### Assay conditions

The cleavage reactions without the C5 protein were conducted in buffer C [50 mM 4-morpholineethanesulfonic acid (MES) and 0.8 M NH₄Cl (pH 6.1)] at 37˚C and 800 mM Mg(OAc)<sub>2</sub> or as otherwise indicated (see Supporting information S1 Fig). The RPRs were pre-incubated at 37˚C in buffer C and 800 mM Mg(OAc)<sub>2</sub> for at least 10 min to allow proper folding before mixing with pre-heated (37˚C) substrate. In all the experiments the concentrations of substrates were <0.02 μM while the concentrations of the different RPR variants were as indicated in Table and Figure legends.

Reactions with the C5 protein were done in buffer A [50 mM Tris-HCl (final pH 7.2), 5% (w/v) PEG 6000, 100 mM NH₄Cl] and 10 mM Mg(OAc)<sub>2</sub> as described in [34].

In the RPR alone reactions the k<sub>app</sub> values were determined from experiments done under single turnover conditions where we measured the percentage of cleavage as a function of time (with C5 the RPR concentration varied between 0.004 and 0.009 μM). For the calculations we used the 5’ cleavage fragment. To be able to compare with our previously reported data we refer to the so obtained rates as k<sub>app</sub> values. The concentration of substrate was ≤0.02 μM while the RPR concentration varied dependent on RPR variant (see Table 1).

Cleavage of pATSerU<sub>am</sub>G at 37˚C was performed in buffer C, 0.8 M NH₄Cl and 800 mM Mg(OAc)<sub>2</sub> at pH 5.2, pH 6.1 and pH 7.2 [37, 38].
The cleavage reactions were terminated by adding double volumes of stop solution (10 M urea, 100 mM EDTA) and the products were separated on 25% (w/v) denaturing polyacrylamide gels.

### Structural probing

Structural probing of the Eco RPR variants, labeled at the 3'-end with \[^{32}\text{P}\]pCp, was conducted using Pb\(^{2+}\)-induced cleavage and limited RNase T1 digestion under native conditions as described elsewhere [34, 39, 40, 41]. Approximately 2 pmols of labeled RPR in 10 μl was pre-incubated for 10 min at 37°C in 50 mM Tris-HCl (pH 7.5), 100 mM NH₄Cl and 10 mM MgCl₂ together with 4 μM unlabeled tRNA. Cleavage was initiated by adding freshly prepared Pb(OAc)₂ to a final concentration of 0.5 mM (or as indicated in Fig 4 legend) and the reaction was stopped after 10 min. In the digestion with RNase T1, the RPR was pre-incubated as described above. One unit RNase T1 was added followed by incubation on ice for 10 min. The reactions were stopped after 10 min by adding two volumes of stop solution (see above) and the products were analyzed on an 8% (w/v) denaturing polyacrylamide gel.

### RNase H cleavage

Approximately one μg of 3'\[^{32}\text{P}\]\(\text{pCp}\) labeled RPR was re-suspended in H₂O and incubated for 3 min at 95°C. Following this the RPR was re-natured prior to the reaction at 55°C for 5 min in the buffer supplied by the company (20 mM Tris-HCl, 20 mM KCl, 10 mM MgCl₂, 0.1 mM EDTA, 0.1 mM DTT and final pH 7.5; ThermoFisher Scientific) followed by incubation at room temperature. The RPR was mixed with 120 pmols of DNA oligonucleotides 1 (5' TGCCCT) or 2 (5' TGCGCT) and incubated in reaction buffer (see above) for 15 min at 28°C (similar results were obtained using 12 pmols of DNA oligonucleotides 1 or 2). The reaction was initiated by adding one unit RNase H (ThermoFisher Scientific) and the reaction was terminated after 30 min by adding double volumes of stop solution (see above). The reaction

| RPR variant                  | Structural consequence          | without C5 | with C5       |
|-----------------------------|--------------------------------|------------|---------------|
| Eco RPR<sub>wt</sub>        |                                | 12±0.5     | 2161±223      |
| Eco RPR<sub>P18CUUG</sub>  | P8/P18 disrupted G<sub>314</sub>CGA<sub>317</sub> changed to C<sub>314</sub>UGA<sub>317</sub> | 0.58±0.075 | ND            |
| Eco RPR<sub>delP18</sub>   | P18 deleted                    | 0.018±0.0039 | ND          |
| Eco CP RPR<sub>wt</sub>     | S-domain removed               | 0.095±0.002 | 18±1.2        |
| Eco CP RPR<sub>delP18</sub>|                                | 0.12±0.0006 | ND            |
| Eco CP RPR<sub>delP18P3Mini</sub> | P18 deleted P3 size reduced   | 0.003±0.0004 | ND          |
| Eco CP RPR<sub>delP18P3Mini</sub> |                                | 0.002±0.0003 | ND          |
| Eco CP RPR<sub>delP18P3Mini</sub> |                                | 0.002±0.0003 | ND          |
| Eco CP RPR<sub>delP18P3Mini</sub> |                                | 0.004±0.015  | ND          |
| Eco CP RPR<sub>delP18P3Mini</sub> |                                | 0.0047±0.0011 | ND          |
| Pfu RPR<sub>wt</sub>       | S-domain removed               |           |               |
| Pfu RPR<sub>delP18</sub>   |                                |           |               |

Expressed as % of cleavage per min per pmol of RPR. The values are averages of at least three independent time-course experiments ± the maximum deviation from the average value; for Eco RPR<sub>wt</sub> the value is based on two independent experiments. The concentrations of RPR (without C5) varied between 0.8 and 11 μM dependent on RPR variant while with C5 the concentration varied between 0.004 and 0.009 μM. The substrate concentration was ≤0.02 μM. ND = not determined.

https://doi.org/10.1371/journal.pone.0192873.t001

The cleavage reactions were terminated by adding double volumes of stop solution (10 M urea, 100 mM EDTA) and the products were separated on 25% (w/v) denaturing polyacrylamide gels.

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PLOS ONE | https://doi.org/10.1371/journal.pone.0192873 | March 6, 2018 | 4 / 23
products were separated on 10% (w/v) denaturing polyacrylamide gels (see also Fig 3 legend and Ref [39]).

**Determination of $k_{app}$ and the kinetic constants $k_{obs}$, $k_{obs}/K_{sto}$ and $K_{sto}$**

The rate constants $k_{obs}$ and $k_{obs}/K_{sto}$ were determined under saturating single-turnover conditions at pH 6.1 (where cleavage is suggested to be rate limiting) and 800 mM Mg$^{2+}$ as described elsewhere [19, 23, 34].

On the basis of the simplified scheme $k_{obs}$ reflects the rate of cleavage (Fig 1). We have argued elsewhere that $K_{sto} \approx K_d$ in the Eco RPR-alone reaction [19, 23, 31, 34, 42, 43]. The final concentrations of the different RPR variants were between 0.4 and 47 μM (depending on the combinations of substrate and RPRs); the concentration of the pATSerUG substrate was 0.02 μM. To ensure that the experiments were done under the single-turnover conditions the lowest concentration of RPR was >10 times higher than the concentration of the substrate. For the calculations we used the 5’ cleavage fragment and the time of cleavage was adjusted to ensure that the velocity measurements were in the linear range (i.e., <40% of the substrate had been consumed). To be able to compare with our previously published data $k_{obs}$ and $k_{obs}/K_{sto}$ were obtained by linear regression from Eadie-Hofstee plots as described elsewhere [19, 23, 31, 34, 44, 45]. Each value was an average of at least three independent experiments and is given as a mean ± the deviation of this value.

**Results**

**Structural probing of Eco CP RPR and full-size Eco RPR variants**

Presence of a "P6-mimic" in *Eco* CP RPR<sub>wt</sub>. The "P6-mimic" might form in *Eco* CP RPR<sub>wt</sub> since residues in the P17-loop that constitute one part of P6 are open to pair with other residues in *Eco* CP RPR as a result of deleting the S-domain (marked in green in Fig 2A). To test for the presence of the "P6-mimic" we generated the following *Eco* CP RPR variants (Fig 2A and Table 1): *Eco* CP RPR<sub>C83C84</sub> ("P6-mimic" disrupted), *Eco* CP RPR<sub>G277G278</sub> ("P6-mimic" disrupted), *Eco* CP RPR<sub>C83C84/G277G278</sub> ("P6-mimic" restored), *Eco* CP RPR<sub>31</sub> ("P6-mimic" disrupted due to replacement of the P15-17 domain with P15 RNA [22, 23]).

These variants (except *Eco* CP RPR<sub>31</sub>) were probed with respect to the accessibility of residues 5′A<sub>81</sub>GGGCA<sub>86</sub> (underlined residues altered in the respective CP RPR variant; Fig 2A) to RNase H in the presence of DNA oligonucleotides (i) 5′<sub>tgccct</sub> (oligo 1), complimentary to residues (underlined) 5′<sub>A<sub>81</sub>GGGCA<sub>86</sub></sub> in *Eco* CP RPR<sub>wt</sub> and *Eco* CP RPR<sub>G277G278</sub> and ii)

![Fig 1. Simplified reaction scheme.](https://doi.org/10.1371/journal.pone.0192873.g001)
5’GGGCT (oligo 2), complimentary to residues 5’ A81 GCCCA86 in Eco CP RPR\textsuperscript{C83C84} and Eco CP RPR\textsuperscript{C83C84/G277G278} (Fig 2A; see also Ref [39]). We expected that disruption of the “P6-mimic” should result in RNase H cleavage of Eco CP RPR in the presence of oligo 1 and oligo 2 in a predictable manner. Subjection to cleavage with RNase H in the presence of either of the two DNA oligonucleotides indeed revealed strong cleavage for Eco CP RPR\textsuperscript{G277G278} (oligo 1), and Eco CP RPR\textsuperscript{C83C84} (oligo 2) as expected if the “P6-mimic” does not form. By contrast, for Eco CP RPR\textsubscript{wt} and Eco CP RPR\textsuperscript{C83C84/G277G278} we did observe significant lower
cleavage (Fig 3). We interpreted these data as an indication that the "P6-mimic" is likely to form in both Eco CP RPR<sub>wt</sub> and Eco CP RPR<sub>C83C84/G277G278</sub>.

Structural impact of the P8/ P18-interaction and P18 on full-size Eco RPR and Eco CP RPR.

To investigate the influence of P18 on the Eco CP RPR structure and its contribution to catalysis (see below) we generated the following variants (Fig 2A) Eco CP RPR<sub>delP18</sub> (P18 deleted), Eco CP RPR<sub>delP18P3Mini</sub> (P18 deleted, P3 size reduced) and Eco CP RPR<sub>31delP18</sub> ("P6 mimic" and P18 deleted). For comparison we also generated two full-size variants, Eco RPR<sub>delP18</sub> and Eco RPR<sub>P18CUUG</sub>, which both disrupt the P8/ P18-interaction albeit in different ways, in the former P18 is deleted while in the latter the P8/ P18-interaction is disrupted (Fig 2B). Comparing Eco CP RPR<sub>wt</sub> and Eco RPR<sub>wt</sub> allowed us also to assess whether removal of the S-domain (and the P8/ P18-interaction) affected the structure of the C-domain. First we studied the impact of the P8/ P18-interaction on full-size Eco RPR.

Structural probing of Eco RPR<sub>wt</sub> and Eco RPR<sub>P18CUUG</sub> with Pb<sup>2+</sup> and RNase T1 (Fig 4A) suggested that disruption of the P8/ P18-interaction affected the P18 structure and the region near Pb<sup>2+</sup>-induced cleavage sites IIb and possibly also IIb* (marked with a dot and absent in Eco RPR<sub>P18CUUG</sub> at 10 mM Pb<sup>2+</sup>, cf. lanes 3 and 4; see also Fig 4A legend; Pb<sup>2+</sup> cleavage sites are marked in Fig 2). We also noted that a weak RNase T1 cleavage product (marked with a dot; Fig 4A, cf. lanes 10 and 11) was absent just upstream of the Pb<sup>2+</sup>-induced cleavage site IIc at A248 in Eco RPR<sub>P18CUUG</sub> (Fig 2B), where A248 is close to the tRNA 5' end in the RNase P- tRNA complex [18]. Of note, a higher concentration of Pb<sup>2+</sup> was needed for Eco RPR<sub>P18CUUG</sub> [10 mM vs. 0.5 mM for Eco RPR<sub>wt</sub>; see Fig 4A, cf. lanes 3 (or 8) and 4], which might indicate an effect on Pb<sup>2+</sup> binding affinities perhaps due to a more flexible structure. Deletion of P18
EcoRPR delP18 also resulted in some changes in the Pb\(^{2+}\) cleavage pattern with the appearance of an extra band in the IIa/IIB region (marked with a dot, Fig 4, cf. lanes 14 and 15). Also, Pb\(^{2+}\) mediated cleavage at IIb in EcoRPR delP18, which is in contrast compared to EcoRPR wt and EcoRPR P18 CUUG (Fig 4A, cf. lanes 4 and 15). Moreover, apart from the P18 region the RNase T1 cleavage patterns were similar comparing EcoRPR wt and EcoRPR delP18 (Fig 4A, cf. lanes 16 and 17). Together these data indicate some influence on the overall RPR structure, apart from the P18 region, when the P8/ P18-interaction is absent (or disrupted), in particular in the region near the Pb\(^{2+}\) cleavage site IIb.
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Impact of the "P6-mimic", P18 and P8/ P18-interaction on the catalytic performance of Eco CP RPR and full-size Eco RPR

As substrates we used different well-characterized model hairpin-loop substrates, which are derived from the E. coli tRNA\textsubscript{Ser}Su1 precursor (Fig 5; [34, 39, 50], and references therein). The longer substrates, pATSerUG and pATSerCG, can interact with the TBS-site in the RPR while the shorter, pMini3bpUG and pMini3bpCG, cannot. As we reported elsewhere optimal cleavage of these substrates by Eco RPR\textsubscript{wt}, and Eco CP RPR\textsubscript{delP18} is absent in the Eco CP RPR variants that lack P18 (see also Ref [49]). The bands upstream of IIc (marked with X; Fig 4B) might possibly be the result of Pb\textsuperscript{2+}-induced cleavage near residues constituting the "P6-mimic". RNase T1 cleavage of the Eco CP RPR variants revealed that most of the cleavage sites detected using full-size Eco RPR\textsubscript{wt} were present (Fig 4C, cf. lanes 6–9). However, we noted one apparent difference in the region referred to as the "248-region" (marked with a short vertical grey line, Fig 4C, cf. lanes 8 and 9) where we detected new and stronger cleavage.

Taken together, the P8/ P18-interaction appears to affect the overall structure of full-size RPR to a certain degree (e.g., cf. Fig 4A lanes 3 and 4) while deletion of the S-domain and P18 does not affect the overall structure of the C-domain to any significant extent except for changes, in particular in the P18-region (as expected) and the region referred to as the "248-region". From our data it also appears that the overall structure of the C-domain is not much affected by deleting the S-domain (compare Eco RPR\textsubscript{wt} and Eco CP RPR\textsubscript{wt}).

The "P6-mimic" affects the catalytic performance of Eco CP RPR\textsubscript{wt}. As shown in Fig 6A both Eco CP RPR\textsubscript{C83C84} and Eco CP RPR\textsubscript{G277G278} cleaved pATSerUG with reduced
Fig 5. Secondary structures of the model hairpin loop substrates used in this study. The arrows mark the cleavage sites as indicated: black arrows mark the canonical cleavage sites between residues -1 and +1, and gray arrows mark the alternative site between residues -2 and -1. The differences with respect to the identity of residue -1 are indicated, as is the replacement of the 2’OH to 2’NH$_2$ at the -1 position in pATSerUG. The numbering in the vicinity of the cleavage sites corresponds to that used for tRNA and precursor tRNA [51]. The precursor tRNA (pre-tRNA) is included to
efficiency compared to Eco CP RPR<sub>wt</sub> and Eco CP RPR<sub>C83C84/G277G278</sub>. This was the case in particular in the RNA alone reaction. Determination of the rate of cleavage (k<sub>app</sub>; Table 1) for these Eco CP RPR variants without C5 corroborated these findings with 8- and 28-fold lower k<sub>app</sub> values for Eco CP RPR<sub>C83C84</sub> and Eco CP RPR<sub>G277G278</sub>, respectively, compared to Eco CP RPR<sub>wt</sub>. For Eco CP RPR<sub>31</sub>, which cannot form the "P6-mimic" (Fig 2A; substitution of P15-17 with P15 RNA) we detected an almost 50-fold reduction in k<sub>app</sub> (Table 1; see also Fig 6B and below). The higher impact in response to replacing the P15-17 domain with P15 RNA might reflect a structural effect on establishing the pairing between the 3'ACC in the substrate and the RPR [51].

Together with the structural probing data discussed above suggested that the "P6-mimic" is likely to be present in Eco CP RPR<sub>wt</sub> and that it contributes to its catalytic performance.

**The P8/ P18-interaction influences cleavage of pATSerUG by full-size Eco RPR**. Before analyzing the impact of P18 on catalysis in the Eco CP RPR context we first inquired whether the P8/ P18-interaction influences cleavage of the model hairpin loop substrate pATSerUG in the full-size Eco RPR context. Hence, we used the Eco RPR<sub>delP18</sub> and Eco RPR<sub>P18CUUG</sub> variants (see above), which allowed us also to assess the response upon deleting P18 (Eco RPR<sub>delP18</sub>) and "disruption" of the P8/ P18-interaction (Eco RPR<sub>P18CUUG</sub>). Both these variants cleaved pATSerUG mainly at the +1 site with reduced efficiency (Fig 6B, cf. lane 9; and not shown). Compared to Eco RPR<sub>wt</sub>, the cleavage rates of pATSerUG (k<sub>app</sub>; Table 1) for Eco RPR<sub>P18CUUG</sub> and Eco RPR<sub>delP18</sub> were reduced ≈20-fold and almost 700-fold, respectively. Determination of the kinetic constants under single turnover conditions revealed that "disruption" of the P8/ P18-interaction resulted in a ≈20- and 160-fold decrease in k<sub>obs</sub> and k<sub>obs</sub>/K<sub>d</sub>, respectively, while deleting P18 lowered both k<sub>obs</sub> and k<sub>obs</sub>/K<sub>d</sub> > 3000-fold (Table 2; cf. values for Eco RPR<sub>wt</sub>, Eco RPR<sub>P18CUUG</sub> and Eco RPR<sub>delP18</sub>). The K<sub>obs</sub> values correspond to ≈ K<sub>d</sub> values (see Materials and methods) and no difference was detected comparing Eco RPR<sub>wt</sub> and Eco RPR<sub>delP18</sub> while for Eco RPR<sub>P18CUUG</sub> K<sub>obs</sub> was ≈10-fold higher. These data suggest that the P8/ P18-interaction influence cleavage of pATSerUG, which is consistent with previous findings using pre-tRNAs [28–30]. Also, while deleting P18 (Eco RPR<sub>delP18</sub>) affected k<sub>obs</sub> "disruption" of the P8/ P18-interaction resulted in changes in both k<sub>obs</sub> and K<sub>obs</sub> (see Discussion).

**P18 does not influence the catalytic performance for Eco CP RPR**. On the basis of the data discussed above one prediction was that this might also be the case for Eco CP RPR (see above). However, given that P18 helps to connect the S- and C-domains [18, 46] another possibility was that P18 does not affect catalysis since in Eco CP RPR the S-domain is missing (Fig 2A). To test this, and get insight into the contribution of P18 to catalysis, we studied cleavage of pATSerUG, pATSerCG, pMini3bpUG and pMini3bpCG (Fig 5) followed by determinations of the rate constant k<sub>app</sub> (for pATSerUG) without the C5 protein and for a selected few in its CP RPR context. Hence, we used the Eco CP RPR variants with and without P18 cleaved the four model hairpin loop substrates preferentially at the correct position +1 (see Figs 6B and 7, cleavage with Eco CP RPR<sub>wt</sub>, Eco CP RPR<sub>delP18</sub> and Eco CP RPR<sub>delP18P3Mini</sub> for Eco CP RPR<sub>31</sub> and Eco CP RPR<sub>31delP18</sub> we only tested cleavage of pATSerUG, Fig 6B, cf. lanes 5 and 6; see above). Consistent with our previous data [16, 23, 34] the cleavage efficiencies for pATSerCG and
Fig 6. Cleavage of the model hairpin loop substrate pATSerUG. A. Cleavage with different Eco CP RPR carrying changes that affect the “P6-mimic” (Fig 2A). The experiments were done with and without the C5 protein as indicated. Reactions without the C5 protein were performed in buffer C and 800 mM Mg$^{2+}$ (cf. lanes 1 to 5) while those with the protein were done in buffer A and 10 mM Mg$^{2+}$ (cf. lanes 6 to 10). All the reactions were done at 37°C and black and open circles as defined above. The concentrations of the RPRs were 0.7 nM with C5 and 2.7 μM without. The
reaction times were needed to cleave pATSerCG and pMini3bpCG). Moreover, the cleavage pMini3bpCG were lower than using pATSerUG and pMini3bpUG (Fig 7; note that longer 10.2 μM were 30 min in all cases. Lane 1, pATSerUG alone; lane 2, Eco CP CP RPR variants as indicated. The concentration of pATSerUG was +1 and -1 marks were cleavage had occurred (see Fig 5 and text for details). B. Cleavage with various Eco RPR and CP RPR was deleted (Table 2; cf. values for 31delP18). In the presence of the C5 protein, there was a modest three- to four-fold decrease in kobs of the respective cleavage site differs. For example it has been suggested that residues near the conserved U69 in Eco RPR (Fig 2B) interact with the residue positioned five bases 3’ of the cleavage site [53] and pMini3bp only has a stem of three base pairs (Fig 5).

Consistent with our previous data [23] Eco CP RPRwt cleaved pATSerUG with a reduced rate (kapp decreased ≈100-fold) both with and without the C5 protein compared to full-size Eco CP RPRwt (Table 1). In contrast to full-size Eco RPR, where “disruption” (Eco CP RPR wt 31CUG) of the P8/ P18-interaction resulted in a 20-fold (or almost 700-fold upon deleting P18; see above) reduction in kobs, deleting P18 in Eco CP RPR did not affect kapp (if anything, there was a modest ≈two-fold increase for Eco CP RPR delP18). In the presence of the C5 protein, there was a modest three- to four-fold decrease in kapp for the Eco CP RPR variant lacking P18 (Table 1; cf. values for Eco CP RPR wt, Eco CP RPR delP18 and Eco CP RPR delP18P3Mini). Determination of the kinetic constants for cleavage of pATSerUG without C5 corroborated the data presented in Table 1 and revealed no change in either kobs, or ksto when P18 in Eco CP RPR was deleted (Table 2; cf. values for Eco CP RPR wt and Eco CP RPR delP18). These data are in contrast to full-size Eco RPR were deleting and “disrupting” the P8/ P18-interaction affected kobs and kobs/Ksto, respectively (see above: Table 2). Of note, the Ksto values (≈Kd, see above) for Eco CP RPR 31CUG and Eco CP RPR wt (or Eco CP RPR delP18) only differed by a factor of two (Table 2).

Taken together, these data emphasized the importance of the S-domain and the P8/ P18-interaction for catalysis and substrate binding for full-size Eco RPR while P18 does not contribute to the catalytic performance of Eco CP RPR to any significant extent. However, the presence of P18 in full-size Eco RPR that cannot properly interact with P8 does affect pATSerUG binding whereas its absence does not (see Discussion). Also, comparing kapp values (Table 1) for Eco CP RPR delP18 and Eco CP RPR delP18P3Mini suggested that the length of P3 does not appear to influence the catalytic performance in an Eco CP RPR context.

The C-domain derived from Pyrococcus furiosus (Pfu) is catalytically active in the absence of the S-domain and protein. The type A Pfu RPR lacks P18 (Fig 2C and S2 Fig) and it is catalytic also in the absence of the S-domain but only in the presence of proteins ([6]; pMini3bpCG were lower than using pATSerUG and pMini3bpUG (Fig 7; note that longer reaction times were needed to cleave pATSerUG and pMini3bpUG). Moreover, the cleavage pMini3bpUG was ≈0.02 μM while the concentrations of the RPR varied: Eco CP RPRwt, 8.2 μM (lane 3); Eco CP RPR delP18, 9.2 μM (lane 4); Eco CP RPR delP18P3Mini, 10.2 μM (lane 6), and Eco CP RPR wt 31CUG, 3.2 μM (lane 9). The reaction times were 20 min for Eco CP RPR wt and Eco CP RPR wt 31CUG, 60 min for Eco CP RPR wt 31P18delP18 and Eco CP RPR delP18. Controls, incubation of pATSerUG alone without RPR (lane 1), cleavage of pATSerUG (lane 2) and pATSerUG (lane 6) (known to cleave at +1 and -1, see 18 Wu et al. 2011; lane 8) with Eco CP RPR, S = substrate, 5’-L = 5’ cleavage fragments and +1 and -1 marks cleavage sites (see text for details).
see Discussion). Full-size *Pfu* RPR<sub>wt</sub> alone cleaves the model hairpin substrates used above at high Mg<sup>2+</sup> concentration [31]. To test whether *Pfu* RPR<sub>wt</sub> is catalytically active also without the S-domain and protein we generated *Pfu* CP RPR<sub>wt</sub> (Fig 2C). Indeed, *Pfu* CP RPR<sub>wt</sub> cleaved pATSerUG, pATSerCG, pMini3bpUG and pMini3bpCG mainly at the correct position +1 (Fig 7, cf. lanes 9, 13, 17 and 21). In addition, *Pfu* CP RPR<sub>wt</sub> cleaved pMini3bpCG at the alternative site -1 (Fig 7, lane 21) while we could not detect any cleavage of pATSerCG at -1 (Fig 7B, lane 17). However, this could be because *Pfu* CP RPR<sub>wt</sub> cleaved pATSerCG with a very low efficiency such that cleavage at -1 could not be detected and quantified.

| Table 2. Kinetic constants for cleavage of pATSerUG with RPR variants at 800 mM Mg<sup>2+</sup> as indicated. |
|-------------------------------------------------------------|
| **RPR variant** | **k<sub>obs</sub>**<br>(min<sup>-1</sup>) | **k<sub>obs</sub>/K<sub>sto</sub>**<br>(= k<sub>cat</sub>/K<sub>m</sub>)<br>(min<sup>-1</sup> x μM<sup>-1</sup>) | **K<sub>sto</sub>**<br>(μM) | **ΔΔG# (kcal)** |
|-----------------|-----------------|-----------------|-----------------|----------------|
| *Eco* RPR<sub>wt</sub> | 12<sup>a</sup> | 19<sup>a</sup> | 0.63 | 1 |
| *Eco* RPR<sub>P18CUUG</sub> | 0.50±0.09<sup>a</sup> | 0.12±0.01<sup>a</sup> | 4.3 | +3.1 |
| *Eco* RPR<sub>delP18</sub> | 0.0033±0.0018 | 0.066±0.0007 | 0.56 | +5 |
| *Eco* CP RPR<sub>wt</sub> | 0.34<sup>a</sup> | 0.04<sup>a</sup> | 8.3 | +3.8 |
| *Eco* CP RPR<sub>delP18</sub> | 0.32±0.01<sup>a</sup> | 0.036±0.002<sup>a</sup> | 8.9 | +3.9 |
| *Pfu* RPR<sub>wt</sub> | 0.058<sup>a</sup> | 0.03<sup>a</sup> | 1.9 | 1 |
| *Pfu* CP RPR<sub>wt</sub> | 0.016±0.003<sup>a</sup> | 0.0018±0.00035<sup>a</sup> | 8.9 | +1.7 |
| *Thh* RPR<sub>wt</sub> (without C5) | ND | 25<sup>b</sup> | ND | 1 |
| *Thh* RPR<sub>P18(304/27)</sub> (without C5) | ND | 1.4<sup>b</sup> | ND | +1.8 |
| *Eco* RPR<sub>wt</sub> (without C5) | ND | 2.4<sup>c</sup> | ND | 1 |
| *Eco* RPR<sub>P18UUCG(L18m)</sub> (without C5) | ND | 0.2<sup>c</sup> | ND | +1.5 |
| *Eco* RPR<sub>wt</sub> (with C5) | ND | 568<sup>c</sup> | ND | 1 |
| *Eco* RPR<sub>P18UUCG(L18m)</sub> (with C5) | ND | 237<sup>c</sup> | ND | +0.54 |
| *Eco* RPR<sub>wt</sub> (without C5) | ND | 0.012<sup>d</sup> | ND | 1 |
| *Eco* RPR<sub>delP18</sub> (without C5) | ND | 0.0023<sup>d</sup> | ND | +1.0 |

<sup>a</sup>ΔΔG values (change with respect to the RPR<sub>wt</sub> in each case) were calculated using k<sub>obs</sub>/K<sub>sto</sub> (k<sub>cat</sub>/K<sub>m</sub>) values and ΔΔG = -RTln(k<sub>obs</sub>/K<sub>sto</sub>)<sub>mut</sub>/(k<sub>obs</sub>/K<sub>sto</sub>)<sub>wt</sub> [52]. The experiments were conducted under single-turnover conditions at 800 mM Mg<sup>2+</sup> and pH 6.1 as described in Materials and Methods. The concentration of substrate was ≤0.02 μM while the concentration of the different RPR variants varied dependent on RPR and substrate as stated in Materials and Methods. Numbers are averages of at least three independent experiments ± the maximum deviation of the average value.

Substrates used in the different reports were:
- pATSerUG model hairpin loop substrate;
- pre-tRNA<sup>Gly</sup> from *T. thermophilus*;
- pre-tRNA<sup>Tyr</sup> Su3 from *E. coli*;
- pre-tRNA<sup>Asp</sup> from *B. subtilis*.

Values taken from:
1. Sinapah et al. 2011 [31];
2. Wu et al. 2012 [34];
3. Schlegl et al. 1994 [28];
4. Pomeranz-Krummel and Altman, 1999 [30];
5. Haas et al. 1994 [29], values based on the experiment done at 1M NH<sub>4</sub>Cl, at 3M NH<sub>4</sub>Cl no difference in k<sub>cat</sub>/K<sub>m</sub> indicating that lack of P18 can be compensated for by increasing the ionic strength.
Fig 7. Cleavage of different model hairpin loop substrates as indicated. A. Cleavage of pATSerUG with Eco RPR<sub>wt</sub> (0.8 μM; lane 5), Eco CP RPR<sub>wt</sub> (20.5 μM; lanes 6, 10, 14 and 18), Eco CP RPR<sub>delP18</sub> (23 μM; lanes 7, 11, 15 and 19), Eco CP RPR<sub>delP18P3Mini</sub> (27 μM; lanes 8, 12, 16 and 20) and Pfu CP RPR<sub>wt</sub> (20 μM; lanes 9, 13, 17 and 21). The concentration of substrates was ≤0.02 μM. Reaction times were 15 min (pATSerUG and pMini3bpUG) and 90 min (pATSerCG and pMini3bpCG) irrespective of RPR variant. In the case when full-size Eco RPR<sub>wt</sub> was used the reaction time was 4 sec. The negative controls, incubation without RPR for 90 min (pATSerUG; lane 1), (pMini3bpUG; lane 2), (pATSerCG; lane 3) and (pMini3bpCG; lane 4). 5'-L marks the migration of the 5' cleavage fragments as a result of cleavage at +1 and at -1. Lanes 14–16 and 18–21, the numbers correspond to the frequency of cleavage at +1 expressed in percentage. The numbers are averages of at least three independent experiments with the following errors: 92±0.64 (lane 14), 96±0.07 (lane 15), 95±0.27 (lane 16), 82±1.4 (lane 18), 79±0.5 (lane 19), 84±0.65 (lane 20) and 92±0.5 (lane 21). For experimental details Materials and methods (see also Ref [50]). B. Lanes 14–17 overexposure of the 5' cleavage fragments shown in panel A.

https://doi.org/10.1371/journal.pone.0192873.g007
The rate of cleavage ($k_{\text{app}}$; Table 1) for Pfu CP RPR$_{\text{wt}}$ (without protein) was $\approx$ 7-fold lower than for Pfu RPR$_{\text{wt}}$ while determination of $k_{\text{obs}}$ and $K_{\text{sto}}$ revealed that both were affected $\approx$ four- to five-fold, respectively, resulting in a 17-fold reduction in $k_{\text{obs}}/K_{\text{sto}}$ (Table 2). This is a significant lower reduction compared to the 500-fold drop in $k_{\text{obs}}/K_{\text{sto}}$ in response to deleting the S-domain in the Eco RPR system (Table 2; see also Refs [20, 23]).

We conclude that the S-domain is not essential for cleavage in the Pfu RPR alone reaction and as such supporting that the C-domain is responsible for catalysis also in the case of type A archaeal RNase P (see also Refs [11, 24, 32, 42]). However, the S-domain boosts the catalytic performance but to a lesser extent than for Eco RPR (see Discussion).

The absence of the S-domain or disruption of the P8/ P18-interaction affects the charge distribution at and in the vicinity of the cleavage site. A correct TSL/ TBS-interaction leads to efficient and correct cleavage [16, 19, 23]. Moreover, cleavage of pATSer derivatives in which the 2‘OH at position -1 in the substrate had been replaced with 2‘NH$_2$ showed that the frequency of cleavage at -1 is reduced with increasing pH. Most likely this is because the 2‘NH$_2$ becomes protonated and positively charged with decreasing pH, thereby reducing cleavage at the canonical site +1 [38, 54]. The shift of the cleavage site is also dependent on the structural topography of the +1/+72 base pair in the substrate. We have argued that this is due to a change in the charge distribution at the cleavage site in the RPR-substrate complex ([37]; however, see Ref [55] for an alternative model). Hence, to test whether the absence of the S-domain and disruption of the P8/ P18-interaction influence the charge distribution/ protonation near the cleavage site in the RPR-substrate complex we studied the cleavage pattern of the pATSerUG variant pATSerU$_{\text{am}}$G, in which the 2‘OH was replaced with 2‘NH$_2$ at -1, at different pH (Fig 5; see Materials and methods).

The cleavage frequency of pATSerU$_{\text{am}}$G at +1 increased with increasing pH for all the RPRs variants as expected from our previous data (Fig 8 and S3 Fig). However, compared to Eco RPR$_{\text{wt}}$ and Eco CP RPR$_{\text{wt}}$ the trend was that higher pH was required to reach 50% cleavage at +1 for the other RPR variants (including the Pfu RPR variants). For the all-ribo substrate pATSerUG the site of cleavage did not change with pH irrespective of RPR variant (S3 Fig).

We also inquired if a structural change in the TBS region (in the vicinity of where P18 contact P8; Fig 2B) in the S-domain affects cleavage of pATSerU$_{\text{am}}$G as a function of pH differently compared to Eco RPR$_{\text{wt}}$. Hence, we examined the cleavage pattern for Eco RPR$_{G235}$ at different pH. This change in the RPR is known to influence cleavage site recognition (Fig 2B; [16, 19, 23]). Again a higher pH was needed to give 50% cleavage at the +1 position compared to Eco RPR$_{\text{wt}}$ (Fig 8).

To conclude, we interpret these data to suggest that the S-domain, the P8/ P18-interaction and the structural topology of the TBS region influence the pKa of the 2‘NH$_2$ and/or the charge distribution at the cleavage site.

Discussion
Importance of the P6- and P8/ P18-interactions
The P6- and P8/ P18-interactions play important structural roles in folding the RPR where P6 is an intra C-domain interaction while P8/ P18 is involved in connecting the S- and C-domains (Fig 2B). However, information of their impact on the structure and function of RPRs lacking the S-domain, e.g. Eco CP RPR$_{\text{wt}}$, is scarce. In this context, type T archaeal RPRs are equipped with a degenerated S-domain and secondary structure modeling suggests the presence of P6 but its contribution to catalysis has not been studied. As for Pfu RPR, P18 is also absent in type T RPRs [6, 12, 13]. Studying RNase H accessibility and cleavage of the model hairpin loop substrates pATSerUG we provide data suggesting that residues
5'G_{276}C_{277}C_{278}C_{279} are likely engaged in pairing with residues 5'G_{82}G_{83}G_{84}C_{85} in the absence of the S-domain (Fig 2A). We refer to this interaction as the "P6-mimic" and our data indicate that its presence contributes to the catalytic performance by Eco CP RPR. As such, our findings also provide support for the existence and functional importance of P6 in type T RPR. Moreover, deleting the S-domain in the type B B. subtilis RPR decreased the cleavage rate 25000-fold [21]. This is in contrast to the 120-fold reduction in the rate observed for the type A Eco RPR (Table 1; see also Refs [22, 23]). Type B lacks P6, however, recent data indicate that disruption of the intra-domain interaction between L5.1 and L15.1 in the C-domain affects both folding and the catalytic activity in a full-size RPR context [27]. Hence, it will be of interest to understand whether the L5.1/L15.1 interaction (or a mimic) is present in the absence of the S-domain (see also Ref [21]). If this is the case we predict that it contributes to catalysis by the type B RPR lacking the S-domain.
In contrast to disruption of the "P6-mimic", removal of P18 did not result in any significant change in the catalytic performance for Eco CP RPR (Table 2). However, disruption of the P8/ P18-interaction, or deletion of P18, in full-size Eco RPR reduced cleavage of pATSerUG and affected the overall structure to a certain degree. While disrupting the P8/ P18 (P18 still present) influenced both the kinetic constants \( (k_{\text{obs}} \text{ and } K_{\text{sto}}); 24\text{-} \text{and } \approx 10\text{-fold change, respectively, compared to Eco RPR}_{\text{wt}}; \text{ Table 2}) \) deleting P18 resulted in a dramatic decrease in \( k_{\text{obs}} \) (>3000-fold) in cleavage of the model substrate pATSerUG while no change in \( K_{\text{sto}} \) was detected. Given that \( K_{\text{sto}} \approx K_d \) this might indicate that P18 interferes with binding of pATSerUG when its interaction with P8 is disrupted while this is not the case in its absence. Nevertheless, previous multiple turnover kinetic studies using pre-tRNA and pre-4.5S RNA reported that disruption of P8/ P18 or deletion of P18 affects binding affinity \( (K_m) \) and \( k_{\text{cat}} \) in cleavage of pre-tRNAs with type A RPR with and without protein. The levels of change differ comparing our single turnover data and previously reported results \([28–30]; \text{ see below})\). This is likely due to different reaction conditions, choice of substrate and RPR \( (\text{Eco RPR and Thermus} \text{ thermophilus, Tth, RPR}) \). In this context we also note that earlier data suggested that the impact of deleting P18 is suppressed by raising the ammonium concentration to 3 M \([29]\). We conclude that irrespective of substrate presence of P18 and the P8/ P18-interaction have an impact on the catalytic performance by bacterial type A RPR while in the absence of the S-domain (and the P8/ P18 interaction), as in Eco CP RPR, P18 has no significant impact on catalysis.

Calculating the \( \Delta \Delta G \) using \( k_{\text{obs}}/K_{\text{sto}} \) values \([52]\) revealed that the contribution of the P8/ P18-interaction is between 3.1 and 5 kca/mol for full-size Eco RPR while the contribution of the S-domain is approximately 3.8 kca/mol (Table 2). Extracting and using the \( k_{\text{cat}}/K_m \) \( (= k_{\text{obs}}/K_{\text{sto}}) \) values (for type A Eco and Tth RPRs; Table 2) from previous reports \([28–30]\) to calculate the \( \Delta \Delta G \) values suggest that the contribution of P18 and the P8/ P18-interaction varies between 1 and 1.8 kca/mol. In the Eco RPR pre-tRNA-system without the C5 protein disruption of the P8/ P18-interaction resulted in a loss of 1.5 kca/mol \([30]\), which is two-fold lower than the value obtained using pATSerUG (Table 2). The reason for this discrepancy could be due to the difference in reaction conditions \((e.g., \text{here we used higher } [\text{Mg}^{2+}])\) and/or the way the two substrates interact with the RPR where pre-tRNA has a structurally intact TSL-region. The importance of P18 in the full-size RPR context can be rationalized by its structural role in connecting the S- and C-domains and structurally orient these domains in a productive/ correct manner as discussed by Li et al. \([24]; \text{ Figs 2 and 3})\). This together with that a productive TSL/ TBS-interaction in the S-domain affects catalysis \([16, 19, 23, 31, 34]\) opens for the possibility that the P8/ P18-interaction acts as a structural mediator in the "communication" between TSL/ TBS-interaction and the cleavage site leading to positioning of chemical groups and Mg\(^{2+}\) that result in correct and efficient cleavage. Consistent with this is that disruption of the P8/ P18-interaction, removal of the S-domain (and P18) as well as alteration in the vicinity of the structure were P18 connects (as in the Eco RPR\(_{G235}\) variant; \text{Fig 2B}; \text{Ref [19]}) seems to influence the charge distribution in the vicinity of the cleavage site (\text{Fig 8}). That the P8/ P18-interaction influences events at the cleavage site is also supported by data using a derivative of pATSerCG in which the loop had been replaced with a GAAA-tetra loop (pATSerCG\text{GAAA}; \text{Fig 5}). Eco RPR\(_{wt}\) cleaves this substrate preferentially at -1 (81\% \pm 2\%) while absence of the S-domain (and the P8/ P18-interaction) results in cleavage preferentially at +1 \([23]\). Taken together, the type A Eco RPR S-domain and P8/ P18-interaction play important roles for the catalytic performance and site selection. Moreover, since removing P18 in Eco CP RPR did not affect
cleavage (or the structure to any significant extent) it is likely that P18 itself does not influence catalysis but the P8/ P18-interaction does, consistent with that Eco RPR$_{\text{delP18}}$ and Eco RPR$_{\text{P18CUUG}}$ are poor catalysts in cleaving pATSerUG (Table 2). But noteworthy, the presence of P18 that cannot interact with P8 does affect pATSerUG binding and the reason to this is unclear (however see above). In this context, our unpublished structural probing data of full-size Eco RPR variants suggest that substitution of A248, which is positioned close to the cleavage site in the RNase P-tRNA structure [18], influence the structure of P18.

Comparing the type A Eco and Pfu RPRs

Our data show that the Pfu RPR retained its catalytic activity upon removing the S-domain. Tsai et al. [6] provided data where they showed that a Pfu CP RPR construct is indeed catalytic however only in the presence of proteins. A rational for that they did not detect any cleavage activity without proteins might be differences in reaction conditions. We used a significantly higher Mg$^{2+}$-concentration, which has been reported to increase low or unnoticed cleavage efficiency by Eco RPR variants [22, 34]. In addition, the choice of substrate differs in these two studies, pre-tRNA vs. pATSer, which might also be a factor. It should also be noted that our data are in accordance with that the activity of the archaeal type A M. thermoautotrophicus RPR substantially increases when its C-domain is linked to the Eco RPR S-domain [24].

Comparison of our current data, where we removed the S-domain of Pfu RPR$_{\text{wt}}$ with our previous data [31] shows an effect on both the kinetic constant $k_{\text{obs}}$ and $K_{\text{sto}}$ in cleaving pATSerUG. This is similar to the situation for Eco RPR but the magnitude of change in $k_{\text{obs}}$ for Eco RPR was higher (cf. 35-fold vs. 3.6-fold in the case of Pfu RPR; Table 2). Using the $k_{\text{obs}}/K_{\text{sto}}$ values for Pfu RPR$_{\text{wt}}$ and Pfu CP RPR and calculating $\Delta\Delta G$ gives a loss of 1.7 kcal/mol as a result of removing the S-domain (Table 2). This should be compared to the 3.8 kcal/mol loss seen for Eco RPR$_{\text{wt}}$ (see above). Full-size Eco RPR$_{\text{wt}}$ can form a productive/ correct interaction with the substrate TSL-region while Pfu RPR$_{\text{wt}}$ in the absence of proteins interacts differently with TSL [31]. Together this suggests that in the RPR alone reaction the Pfu S-domain plays a less important role in cleavage of the model hairpin loop substrate pATSerUG than in the Eco RPR case. Similar reduction in the rate (12-fold) was also observed upon removing the S-domain in a cis RPR-pre-tRNA construct based on the archaeal M. jannaschii type M system [42]. In this context, as reported previously the S-domain of type A archaeal RPR appears to hamper the activity in the RNA alone reaction [24, 32]. These authors also provided data where structural changes improved the activity of the type A archaeal Methanothermobacter thermoautotrophicus RPR and their findings are likely to be applicable to rationalize the difference in activity comparing Eco and Pfu RPRs.

Taken together, for Pfu RPR the influence of the S-domain is perhaps reflected by the fact that two of the five RNase P proteins, Rpp21 and Rpp29, bind to the S-domain and influence the interaction with the T-loop region of pATSer model substrates [31]. Moreover, considering the absence of P18 in archaeal type A RPRs it has been discussed that its role has been taken over by the Pop5 and Rpp30 proteins [24]. In this context we emphasize that removal of P18 in full-size Eco RPR reduced $k_{\text{obs}}$ to a level $\approx$10-fold lower compared to $k_{\text{obs}}$ as determined for Pfu RPR$_{\text{wt}}$ and a loss of 5 kcal/mol relative to Eco RPR$_{\text{wt}}$, one kcal/mol lower compared to Pfu RPR$_{\text{wt}}$ Table 2; both Eco RPR$_{\text{delP18}}$ and Pfu RPR$_{\text{wt}}$ lack P18). To conclude, combined these data raise the question whether the evolution of a more complex RNase P in terms of the number of protein subunits is linked to a decrease in the contribution of the S-domain to catalysis see also Refs [24, 32].
Supporting information

S1 Fig. Cleavage of pATSerUG as a function of Mg^{2+} concentration. Cleavage by Eco CP RPR_{delP18} (A), Eco CP RPR_{delP18P3Mini} (B) and Pfu CP RPR_{wt} (C). Cleavage by Eco CP RPR_{delP18} (A), Eco CP RPR_{delP18P3Mini} (B) and Pfu CP RPR_{wt} (C). The experiment was performed in buffer C, 0.8 M NH₄OAc (pH 6.1) at 37˚C in the presence of indicated amount of Mg(OAc)₂. The concentrations of RPRs ranged between 1 to 1.5 μg per μl and the substrate concentration was ≤0.02 μM. For the calculations, we used the 5’ cleavage fragments and the data are the mean of three independent experiments. The bars indicate the experimental errors. For details see main text.

(TIFF)

S2 Fig. Secondary structure model of the type A Pfu RPR.

(TIF)

S3 Fig. Cleavage of pATSerUG and pATSerU_{am}G by Eco CP RPR_{wt}, Eco CP RPR_{delP18}, Eco CP RPR_{delP18P3Mini}, Pfu CP RPR_{wt}, Eco RPR_{wt} and Eco RPR_{P18CUUG}. The experiment was performed at 37˚C in buffer C, 0.8 M NH₄OAc, 800mM Mg(OAc)₂ at pH 5.2, 6.1 and 7.2 (the black triangles mark the increase in pH). The concentration of substrates were 0.02 μM while the concentration of RPRs were as indicated below: S = substrate, C1 and C2 = controls, no RPR added (1) and cleavage of pATSerUG with Eco RPR_{wt} 0.8 μM for 4 sec (2), 5’-L mark the cleavage 5’ cleavage fragments due to cleavage at +1 and at -1. (A) and (B) longer exposure of selected region shown in A) The concentrations of the RPRs and reaction times (in parenthesis) were: lane marked with C2 (4 sec), control (see above), Eco RPR_{wt} 0.8 μM (10 min), Pfu RPR_{wt} 4.5 μM (60 min), Eco CP RPR_{wt} 11 μM (270 min), Eco CP RPR_{delP18} 12 μM (270 min), Eco CP RPR_{delP18P3Mini} 14.5 μM (270 min) and Pfu CP RPR_{wt} 20 μM (270 min). (C) Cleavage of pATSerUG with RPRs as indicated (only the migration of 5’ cleavage fragments are shown). Concentrations of the RPRs and reaction times (in parenthesis) were: lanes marked with C1 (60 min) and C2 (4 sec) controls (see above), Eco CP RPR_{wt} 13.7 μM (10 min), Eco CP RPR_{delP18} 12 μM (30 min), Eco CP RPR_{delP18P3Mini} 14.5 μM (30 min), Pfu CP RPR_{wt} 20.3 μM (60 min). (D) Cleavage of pATSerUG and pATSerU_{am}G with RPRs as indicated (only the migration of 5’ cleavage fragments are shown). Concentrations of the RPRs and reaction times (in parenthesis) were: control lanes marked with C1 (pATSerUG no RPR, 10 min), C2 (pATSerUG with 0.8 μM Eco RPR_{wt}, 4 sec) and C3 (pATSerU_{am}G no RPR, 10 min), Eco RPR_{wt} 0.8 μM (pATSerUG, 4 sec and pATSerU_{am}G, 10 min), Eco RPR_{P18CUUG} 0.8 μM (pATSerUG, 1 min and pATSerU_{am}G, 10 min) and Pfu RPR_{wt} 4.5 μM (10 min). For details see main text.

(TIFF)

Acknowledgments

We thank our colleagues for discussions throughout this work. Dr V. Gopalan and Ms T. Bergfors are acknowledged for critical reading of the manuscript. We thank Dr A. C. Forster for discussions about the title. This work was supported by the Swedish Research Council (N/T), Uppsala RNA Research Center (Swedish Research Council Linneus support) and the Carl Tryggers Foundation for Research. LAK is a shareholder in Bioimics AB.

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