

NIH Public Access

Author Manuscript

J Mol Biol. Author manuscript; available in PMC 2010 November 13.

Published in final edited form as:

J Mol Biol. 2009 November 13; 393(5): 1043–1055. doi:10.1016/j.jmb.2009.08.068.

Solution Structure of an Archaeal RNase P Binary Protein Complex. Formation of the 30-kDa Complex Between *Pyrococcus furiosus* RPP21 and RPP29 is Accompanied by Coupled Protein Folding, and Highlights Critical Features for Protein-Protein and Protein-RNA Interactions

Yiren Xu^{1,2}, Carlos D. Amero^{†,2,3}, Dileep K. Pulukkunat^{†,1,2}, Venkat Gopalan^{*,1,2,3}, and Mark P. Foster^{*,1,2,3}

¹Ohio State Biochemistry Program, The Ohio State University, Columbus, OH 43210, USA

²Department of Biochemistry, The Ohio State University, Columbus, OH 43210, USA

³Biophysics Graduate Program, Center for RNA Biology, The Ohio State University, Columbus, OH 43210, USA

Abstract

RNase P is a ribonucleoprotein (RNP) enzyme that catalyzes the Mg²⁺-dependent 5' maturation of precursor tRNAs. In all domains of life, it is a ribozyme: the RNase P RNA (RPR) component has been demonstrated to be responsible for catalysis. However, the number of RNase P protein subunits (RPPs) varies from one in bacteria to nine or ten in eukarya. The archaeal RPR is associated with at least four RPPs, which function in pairs (RPP21–RPP29 and RPP30-POP5). We used solution NMR spectroscopy to determine the three-dimensional structure of the protein-protein complex comprising *Pyrococcus furiosus (Pfu)* RPP21 and RPP29. We found that the protein-protein interaction is characterized by coupled folding of secondary structural elements that participate in interface formation. In addition to detailing the intermolecular contacts that stabilize this 30-kDa binary complex, the structure identifies surfaces rich in conserved basic residues likely vital for recognition of the RPR and/or precursor tRNA. Furthermore, enzymatic footprinting experiments allowed us to localize the RPP21–RPP29 complex to the specificity domain of the RPR. These findings provide valuable new insights into mechanisms of RNP assembly and serve as important steps towards a three-dimensional model of this ancient RNP enzyme.

Accession numbers:

^{© 2009} Elsevier Ltd. All rights reserved.

^{*}Corresponding authors. Contact: 484 West 12th Ave., Columbus, OH 43210, gopalan.5@osu.edu, foster.281@osu.edu. Current addresses: Institut de Biologie Structurale, CNRS, Grenoble, France (CDA) and Department of Chemistry, Columbia University, New York, NY, USA (DKP)

[†]Equal contributions

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

Coordinates and restraints have been deposited in the Protein Data Bank with accession number 2KI7. NMR resonance assignments have been deposited in the BioMagResBank with accession number 16266.

INTRODUCTION

RNase P, a ribonucleoprotein complex (RNP), catalyzes removal of the 5' leader sequence during tRNA maturation ¹2^{; 3}. Across the three domains of life, it is composed of one RNA subunit and a varying number of protein subunits: one in bacteria; at least four in archaea and nine in eukarya. The RNase P RNA (RPR) from each domain of life has been shown to be catalytic on its own in vitro under elevated monovalent and divalent ion concentrations ^{4; 5; 6}. However, RNase P protein(s) (RPP) enhance catalysis under near-physiological conditions by facilitating RPR folding, substrate recognition and decrease in the Mg²⁺ requirement ^{7; 8; 9; 10}. Interestingly, when comparing the enzyme from the three domains of life, an increase in protein content is associated with the loss of some RPR elements and a concomitant decrease in RPR activity ^{1; 11; 12}. Thus, RNase P is an appealing model to address how protein cofactors might have taken over the structural and functional attributes of RNAs during evolution from a putative RNA-centric world to the present protein-centric one. Towards the goal of understanding this progression, we have focused our efforts on the biochemically tractable and thermostable archaeal version of the RNase P enzyme from the hyperthermophilic archaeon *Pyrococcus furiosus (Pfu)*, whose RPPs are homologous to their eukaryal counterparts ^{13; 14}.

Bacterial RNase P is the best understood form of the enzyme ¹⁵. Based on the primary sequence and secondary structure of the RPRs, bacterial RNase P can be classified into two distinct types: A and B ¹⁶. In both types, the RPR is composed of two independently folding domains, termed the specificity and catalytic domains (S domain and C domain, respectively) held together by interdomain RNA-RNA contacts ¹⁷. To date, crystallographic structures have been reported for two S domains and two full-length RPRs, from bacterial RNase P of types A and B ^{18; 19; ^{20; 21}. The structures of three homologous bacterial RPPs have been solved by crystallography or NMR spectroscopy as well ^{22; 23; 24}. Although the structure of the bacterial RNase P holoenzyme complex has not yet been determined, models of A- and B-type RNPs have been built using a wealth of information from biochemical studies ^{25; 26; 27}.}

Phylogenetic and biochemical studies have revealed that archaeal RNase P is a compositional intermediate between the bacterial and eukaryotic counterparts. Euryarchaeal RNase P can also be categorized into two groups based on their RPR sequences: A and M ^{16; 28}. Euryarchaeal A-type RPR (e.g., *Pfu*) is similar to the bacterial A-type in terms of secondary structures and reported *in vitro* catalytic activity ^{5; 13}; in contrast, the M-type RPR more closely resembles the eukaryotic RPR and has shown no catalytic activity on its own, although it can cleave a substrate tethered in cis ²⁹. Though no high-resolution structure is available of the archaeal RPR, secondary structure similarities and the presence of universally conserved nucleotides in the active site suggest that the archaeal RPR fold (especially the C domain) might resemble that of the bacterial RPR; however, the increased RPP content in archaeal RNase P suggests that RNA-protein interactions in the enzyme are likely to have replaced some of the intra-molecular RNA-RNA interactions present in the bacterial RPR ¹².

Although the archaeal and eukaryotic RPRs share conserved structural features with their bacterial counterparts, none of the RPPs from archaeal or eukaryotic RNase P share sequence similarity with the single bacterial protein. At least four protein subunits are associated with *Pfu* RNase P, and share sequence homology to the human RPPs: RPP21, RPP29, RPP30 and POP5¹⁴. The structures of the four archaeal RPPs have been solved from different archaeal organisms by either NMR spectroscopy, X-ray crystallography, or both. These studies on the isolated proteins revealed the structures of the RPPs to fall within common nucleic acid binding protein families: an Sm-like fold (RPP29) ^{30; 31; 32; 33}, a zinc ribbon (RPP21) ^{34; 35}, an RRM-like fold (Pop5) ³⁶, and a TIM barrel (RPP30) ³⁷

Biochemical data have suggested that at least some of the archaeal and eukaryotic RPPs function in pairs. Yeast two-hybrid studies of proteins from both archaeal and eukaryotic RNase P confirmed the presence of two binary complexes: RPP21–RPP29 and RPP30-POP5 ³⁸; ³⁹; ⁴⁰. Reconstitution assays performed on *Pfu* RNase P have shown that either protein pair is sufficient to activate the RNA enzyme at lower ion concentrations, while no single protein can rescue the RNA enzyme under the same conditions ¹³. Although these studies established a role for RPP binary pairs in enhancing the catalytic activity of archaeal RPRs, the mechanistic basis for their actions is largely unclear without useful structural models of the archaeal RNase P holoenzyme. To that end, high-resolution structure determination of the binary complexes is a necessity.

Here we report the NMR-derived solution structure of the 30 kDa complex between two *Pfu* RPPs: the Sm-like RPP29 and the zinc-ribbon RPP21 proteins. This study complements recently reported crystal structures of the *Pyrococcus horikoshii* (*Pho*) POP5-RPP30 and RPP21–RPP29 complexes ^{41; 42}, reveals dynamic features of the proteins and binding-coupled protein folding events, as well as identify additional features important for protein-protein and protein-RNA interactions. Furthermore, footprinting studies allow us to map the RPP21– RPP29 complex onto the S domain of the RPR. Together with biochemical studies on the C domain and RPP30-POP5 complex from *Pyrococcus furiosus* and *Methanocaldococcus jannaschii* (*Mja*) ¹³29, this work represents an important step towards understanding the architecture and function of archaeal and eukaryal RNase P.

RESULTS AND DISCUSSION

NMR spectroscopy of the RPP29-RPP21 complex

At lower temperatures (25, 40°C), NMR spectra of free Pfu RPP21 and RPP29 were of poor quality, with broad lines and highly variable peak intensities, suggesting that the proteins were poorly folded and/or aggregated at those temperatures. Spectra recorded at 55°C yielded generally narrow lines and uniform peak intensities for both proteins, and thus data were recorded at 55°C for both the free proteins and the protein-protein complexes. The ¹⁵N-edited HSQC spectrum of free Pfu RPP29 contains only 62 of 124 expected backbone resonances (Supporting Information), suggesting that only its Sm-like core is folded in solution in the monomer state. This observation is consistent with the previously reported solution structures of RPP29 from Archaeoglobus fulgidus (Afu)³¹ and Methanothermobacter thermautotrophicus (Mth) ³⁰. In free RPP21, 78 backbone amides could be assigned in the free protein, revealing a structured core comprising residues 19 to 105, out of 123 total ³⁵. As unlabeled RPP21 is titrated into ¹⁵N-labeled RPP29, signals from free RPP29 disappear, with concomitant appearance of new set of resonances corresponding to RPP21-bound RPP29, until the samples reach a molar ratio of 1:1 of RPP29 and RPP21, beyond which no further change in the spectrum is observed. This behavior corresponds to the slow exchange regime for the equilibrium between the free and bound states, and is indicative of tight binding and a 1:1 stoichiometry.

Strikingly, 52 new RPP29 backbone amide resonances are observed in ¹⁵N-edited spectra of the RPP21-bound RPP29. This number corresponds overall to 42% of the RPP29 primary sequence and accounts for nearly all of the resonances that were not observed in spectra of the free protein. Among these 52 new signals, 41 arise from residues at the N-terminus and 8 from the C-terminus, implying that the termini, which are disordered in free RPP29, fold upon binding to RPP21 and play an important role in forming the RPP21 binding interface (Figure 1 and Supporting Information).

Spectra of bound RPP21 also revealed signals not observed in the free protein. Upon binding to unlabeled RPP29, nine new backbone amide resonances could be assigned to residues from

the N-terminal helical bundle of the protein. In addition, the amides that exhibited the largest chemical shift perturbations (CSPs) upon binding RPP29 are primarily from residues clustered at the N-terminal helical bundle of RPP21, including D19, I20, L21, L24, A25, R27, V28, S32, R38, L42, and V46 (Figure 1 and Supporting Information). These data highlight the RPP29binding region of RPP21³⁵ and indicate that the interaction with RPP29 stabilizes the secondary structure in this region of RPP21.

Mutations in the unstructured helical bundle of free RPP21 are deleterious to RPP29 binding. We initially used an RPP21 Ala14Val variant (denoted RPP21V14) in our structural studies due to inadvertent selection of a clone with this spurious mutation. The ¹H-¹⁵N correlated spectra of RPP21V14 ³⁵ and of the wild-type protein are nearly identical, indicating that the N-terminal region containing this residue is unstructured in both proteins (data not shown), although helix α_1 was observed to extend through this region in crystallographic studies of *Pho* RPP21³⁴. Ala14 of in RPP21 is highly conserved through archaea and eukarya (Supporting Information), suggesting that the residue may play an important role in maintaining overall structural integrity or interaction with its partner RPP29. Notably, we found that wild-type RPP21 binds to RPP29 three-fold tighter than the mutant (not shown), and no new amide signals appeared when [U-¹⁵N]-RPP21V14 is saturated with RPP29, in contrast to the nine new N-terminal resonances observed when wild type RPP21 binds its partner. This implies that like RPP29, RPP21 experiences binding-coupled folding, and that an A14V mutation in RPP21 interferes with this aspect of the interaction.

Chemical shift perturbations allowed preliminary identification of the binding interface between RPP29 and RPP21. Detailed characterization of the binding interface was achieved by recording and assigning inter-molecular NOEs, which were obtained from 3D ¹³C-filtered/ edited NOESY spectra recorded in 99.8% D₂O ^{43; 44}. Crosspeaks in these spectra arise from NOEs between protons not attached to ¹³C nuclei (the filter step) to protons attached to ¹³C nuclei (the editing step), thereby providing exclusively intermolecular NOEs when one of the proteins is uniformly ¹³C labeled, and the other is unlabeled. We recorded ¹³C-filtered/edited NOESY spectra on both [U-¹³C,¹⁵N]- RPP29 in complex with unlabeled RPP21, and on [U-¹³C,¹⁵N]-RPP21 in complex with unlabeled RPP29 (Figure 2). These spectra yielded a total of 284 intermolecular NOEs (153 and 131 NOEs from each spectrum; Supporting Information).

Solution Structure of Pfu RPP29-RPP21 complex

The solution structure of *Pfu* RPP29-RPP21 complex (Figure 3) was determined by iterative torsion angle refinement using distance restraints derived from inter- and intra-molecular NOEs, hydrogen bond restraints inferred from secondary structure information and torsion angle restraints from analysis of chemical shifts. The ensemble is well defined for the assigned residues (17–123 of RPP29 and 9–54, 57–81, 86–104 of RPP21) with a mean root-mean-square deviation (RMSD) of 0.58 Å and 0.87 Å for backbone and heavy atoms, respectively. The stereochemical quality of the structures was high, with 98.1% of the residues adopting φ and ψ angles falling in the most favored and the additionally favored regions of the Ramachandran plot (Table 1). The extreme N-and C-termini of both proteins (residues 1–16 and 124–127 of RPP29; residues 1–8 and 105–122 of RPP21) are flexible in the complex, as confirmed by {¹H}-¹⁵N heteronuclear NOE data (Supporting Information), indicating these regions are not involved in the interaction with the protein partner.

In the RPP29-RPP21 complex, additional structural elements from each protein could be resolved that were disordered in their free proteins. For RPP29, in addition to the signals from the β -barrel core, binding RPP21 allowed definition of residues from three helices (helix α_1 , residues 19–23; helix α_2 , residues 27–31; and helix 3, residues 40–44), an extended strand connecting helix α_2 and helix α_3 at the N-terminus, and a C-terminal helix (helix α_4 , residues 117–122). In the context of the RPP21 complex, the packing of several N-terminal residues of

RPP29 within the Sm-like core of the protein (I24, T27, R31, H34, V38, K40 and L44) is stabilized, although these intramolecular interactions were insufficient to stabilize the interactions in the free protein. In contrast to the extensive intra-molecular contacts made by the binding-stabilized N-terminal residues, the RPP29 C-terminal helix observed in the complex appears to be entirely stabilized by inter-molecular contacts to RPP21, as few intra-molecular NOEs are observed from this region of the protein to other parts of RPP29. The Sm-like core of RPP29 is essentially unchanged by RPP21 binding as the RMSD between the free and RPP21-bound structures is within the precision of the ensemble (~1.1 Å for backbone atoms). Thus, it is evident that binding of RPP29 to RPP21 involves binding-coupled folding and stabilization of interfacial structures in RPP29.

When bound to its partner, RPP21 adopts the same overall L-shaped structure observed in the free protein: a long arm containing the two N-terminal α -helices, a short-arm made up of the C-terminal β -sheet comprising the zinc ribbon, and a central linker connecting the two domains. However, in the complex, helix α_1 of RPP21 extends through residues 9–17, indicating that binding is associated with induced fit in RPP21 as well. The RPP29-binding interface of RPP21 is clearly mapped to one face of the helix bundle by the cluster of residues identified with the largest CSPs. Backbone atoms of free and RPP29-bound RPP21 superpose with an overall RMSD of 2.5 Å. However, when superposing the N-terminal helices and C-terminal sheet individually, the agreement is much better, with RMSD of 1.2 Å and 1.1 Å, respectively, reflecting the relatively poorly defined central linker in free RPP21, and corresponding uncertainty in interdomain orientation ³⁵. The N-terminal helices of *Pfu* RPP21 that form the binding interface adopt the same overall structure as observed in crystals of the *Pho* RPP21–RPP29 complex, with a backbone RMSD of 0.7 Å.

Interface between RPP29 and RPP21

The structure of the *Pfu* RPP21–RPP29 intermolecular interface identified from chemical shift perturbations (Figure 1) was defined by 284 unique intermolecular NOEs, many of which arise between methyl bearing and aromatic residues (Supporting Information). These NOEs highlight an extensive interface that buries approximately 2400 Å² of surface on both proteins (RPP29, 1100 Å²; RPP21, 1300 Å²) and is composed of hydrophobic, polar and ionic interactions between residues provided in three separate structural elements: 1) the N-terminal region of RPP29, which extends in an anti-parallel fashion along RPP21 helix α_1 ; 2) RPP29 β 2 interacts with both helices of RPP21 in the center of the interface; and 3) the C-terminal helix of RPP29 stabilizes the end of RPP21 helix α_2 (Figure 4).

In addition to hydrophobic interactions, conserved polar contacts likely play an important role in stabilizing the complex. In an unusual feature, $I71^{RPP29}$ in strand $\beta 2$ adopts backbone ϕ/ψ torsion angles (140 \pm 3 and -53 \pm 12, respectively) that allow sidechains of both I71^{RPP29} and D72^{RPP29} to participate in the RPP21 interface, through hydrophobic interactions and an intermolecular salt-bridge, respectively. Moreover, this unusual backbone configuration orients the backbone carbonyl of I71^{RPP29} into the interface, where it is available for a stabilizing interaction with the sidechain hydroxyl of Y39^{RPP21}. Although intermolecular NOEs were observed between I71^{RPP29} and Y39^{RPP21} (Supporting Information), these NOEs were not sufficient to constrain the sidechain of Y39^{RPP21} such that I71^{RPP29} O and $Y39^{RPP21}$ O^{η} are in hydrogen bonding distance in each member of the ensemble (3.3 ± 0.4 Å). This interaction was also observed in the crystal structure of the Pho RPP21-RPP29 complex 42 , and mutation of Y39^{RPP21} to alanine was shown to be strongly deleterious to activity in an in vitro reconstitution assay ³⁴. Given the sequence conservation in this region of both proteins, this is likely to be a conserved interaction across archaea and eukarya. Adjacent to this interaction, prominent intermolecular salt bridges appear to be formed between E47RPP29 and R17^{RPP21}, and between D72^{RPP29} and R38^{RPP21} (Figure 4d). These charge pairs are highly

conserved in the thermophilic and hyperthermophilic RPPs in archaea (Supporting Information). Many of the residues in the binding surfaces of both proteins are highly conserved or invariant (Supporting Information), suggesting a conserved binding interface between RPP29 and RPP21 in archaeal and eukaryotic RNase P.

RNA Binding Surface

RNA binding studies of *Mth* and *S. cerevisiae* RPPs using the yeast three-hybrid system showed that RPP29 binds the RPR in Mth and yeast RNase P, and both RPP21 and RPP29 interact directly with H1 RNA (RPR) in human RNase P^{40; 45}. Given that RPP29 and RPP21 form a tight and intimate complex, it is reasonable to imagine that this complex forms prior to assembly with the RPR, and thus may constitute an RNA binding "unit" with one or more contiguous RNA binding surfaces. The electrostatic potential map of the Pfu RPP29-RPP21 complex identifies two surface patches with positive electrostatic potential on difference faces of the complex (Figure 5). The larger of these two surfaces (Site 1) indeed spans both proteins, including the RPP21 central linker and helix α_3 at the N-terminus, and $\beta 6$ and the C-terminal α -helix of RPP29. Twenty highly conserved Arg and Lys residues are located on this surface patch; among them, R116, R120, K123 of in the C-terminal helix of RPP29, and K51, K53, K59, R60, R61 and K64 in the central linker of RPP21. The smaller site 2 is localized to the RPP21 β-sheet, with R77, R79, R81, K83, R84, K91, R100 and two His (H87 and H97), all extending on one face of the β -sheet. Of these, R100 is invariant and R77, R79, R81, and K91 are highly conserved in archaea and eukarya (Supporting information). This analysis draws attention to two distinct faces of the binary complex as potential RNA binding interfaces for the RPR catalyst and pre-tRNA substrate.

Footprinting

To better understand how the archaeal RPPs support RPR catalysis, knowledge of the spatial organization of these RPPs on the cognate RPR is necessary. We previously employed footprinting assays and demonstrated that the POP5-RPP30 binary complex, and not RPP21-RPP29, decreased the susceptibility to RNase T1 cleavage of a Pfu RPR deletion derivative containing only the C domain ¹³. As documented before, technical problems complicated footprinting experiments with the full-length Pfu RPR¹³. Therefore, we explored Methanocaldococcus jannaschii (Mja) RNase P as an alternative to its Pfu relative and were able to map the RPP binding sites on this shorter RPR. Full-length Mja RPR was digested with either RNase T1 (which cleaves 3' to unpaired guanines) or RNase V1 (which cleaves basepaired nucleotides in RNA). In the presence of Mja RPP29-RPP21 complex, the paired regions P9, P10/11 and P12 in the S domain are protected from V1 cleavage, while no protection by RPP29-RPP21 complex was observed in the C domain (Figure 6). This observation is also consistent with the finding that addition of RPP21-RPP29 (unlike POP5-RPP30) to the S domain-deleted Pfu and Mja RPRs changes neither the rate nor the NH4⁺/Mg²⁺ requirement, unlike its effects on the full-length archaeal RPRs ^{13; 29}. In contrast to RPP21-RPP29, Mja RPP30-POP5 protects from RNase T1 cleavage the RPR's C domain, especially at or surrounding the universally conserved nucleotides (Figure 6). No additional regions of protection were observed when all four RPPs were present compared to the aggregate of the footprints of each binary pair (compare lanes 4, 6 and 10, Figure 6a, 6b). Collectively, these footprinting studies indicate that RPP30-POP5 and RPP21-RPP29 exclusively interact with the C and S domains, respectively, regardless of whether the other pair is present. Our ongoing studies with another type A RNase P (Mth) reveal a similar binding pattern (data not shown). Considering the conservation of sequence and or secondary structural elements at or near the RPP-binding sites, we can reasonably expect similar RNA-protein interactions in archaeal and eukaryotic RNase P holoenzymes.

CONCLUSION

The NMR data presented here indicate that formation of the *Pfu* RPP29-RPP21 complex involves binding-coupled folding of structural elements in both proteins. The RPP29-RPP21 complex is defined by a combination of conserved interfacial hydrophobic and polar residues, including apparent salt bridges. Outside of the structured cores, residues in the extreme N- and C-termini of both proteins indicate that these remain unstructured in the protein complex. Of particular interest are the long N-terminus of RPP29 (17 residues) and the C-terminus of RPP21 (13 residues), both of which possess an abundance of conserved and basic residues (Lys, Arg) that suggest a role in RNA binding. The surface of the protein-protein complex features two well-defined regions of positive electrostatic potential, highlighting possible RNA binding regions.

The structure of the *Pfu* RPP21–RPP29 complex, together with enzymatic footprinting data, permits informed speculation about the nature of the protein-RNA complex. Enzymatic footprinting of Mja RPP21-RPP29 on the Mja RPR indicates that this protein complex interacts only with the S domain of the RPR, consistent with earlier studies where it was shown to not affect the rate of the phosphodiester bond-breaking step ²⁹ (Figure 6). Since presently there is no three-dimensional model of any archaeal RPR, the crystal structures of the bacterial RPRs serve as the best frame of reference. The crystal structure of the T. thermophilus RPR S domain highlights intramolecular RNA-RNA interactions (especially between stems P13 and P12, and between P14 and P8), that stabilize the tertiary structure necessary for RPR-alone catalysis ¹⁸. However, when compared to the bacterial type-A RPR, P13 and P14 are absent from the S domain of *Pfu* and *Mja* RPRs, which instead exhibit an extended P12. It is tempting to propose that in archaeal RNase P the three-dimensional fold of the RPR is maintained, and that the stabilizing RNA-RNA interactions in the bacterial enzyme are replaced by RNA-protein interactions in archaeal RNase P 46. Thus, given the location of RNA binding by the RPP21-RPP29 complex, the function of this complex might be to compensate for the absence of the P13/P14 structural elements by mediating tertiary contacts between different parts of the RPR.

The presence of a second potential RNA-binding surface localized to the RPP21 β-sheet suggests a function unrelated to stabilizing RPR tertiary structure. Since human RPP21 has been reported to bind precursor tRNA ⁴⁷, it is conceivable that archaeal RPP21 might also be involved in substrate recognition. The S domain in bacterial RPR has been shown to recognize the T stem-loop (TSL) region in the ptRNA. The TSL-S domain interaction triggers a conformational change that aids catalysis by positioning the chemical groups and catalytically important Mg²⁺ near the cleavage site in the C domain ¹⁸; ⁴⁸; ⁴⁹. Thus binding of RPP21– RPP29 to the archaeal RPR's S domain might not only promote intra- and/or inter-domain RPR cooperation but also directly (or indirectly via the RPR) mediate recognition of the TSL in the precursor tRNA. Further experimentation is required to evaluate if indeed the two potential RNA-binding sites identified in the electrostatic potential map of the RPP21–RPP29 complex play distinctive or overlapping roles in RPR and ptRNA binding.

In summary, we have determined the solution structure of the *Pfu* RPP29-RPP21 complex and found that poorly structured elements in each protein fold to form the intermolecular interface in the complex. Analysis of the electrostatic potential surface of this complex revealed two potential RNA-binding surfaces, the larger of which is composed of surface elements from both proteins. Finally, we have localized the RPP29-RPP21 complex to the S domain of archaeal RPR by enzymatic footprinting. These findings provide valuable new insights into mechanisms of RNP assembly and serve as important steps towards a three-dimensional model of this ancient RNP enzyme.

MATERIALS AND METHODS

Protein Expression and Purification

Pfu RPP29—The Pfu RPP29/pET-33b plasmid ¹³ was transformed into Escherichia coli BL21(DE3) Rosetta cells (Novagen). The cells were grown in 2 L flasks in a shaker-incubator at 37°C in LB, or minimal M9 media containing 1 g/L of ¹⁵NH₄Cl and 2 g/L of ¹³C-glucose as the sole nitrogen and carbon sources, supplemented with $30 \mu g/L$ of kanamycin and $34 \mu g/L$ L of chloramphenicol. Production of the recombinant proteins was induced by addition of 0.5 mM isopropyl β -D-thiogalactopyranoside (IPTG) at an A₆₀₀ of 0.6, and harvested after 4 h by centrifugation. The cell pellet from 1 L of culture was resuspended in 30 mL of buffer R (25 mM Tris-HCl, pH 7, 1 mM ethylene diamine tetraacetate (EDTA), 0.1 mM phenylmethylsulphonyl fluoride (PMSF), 50 mM dithiothreitol (DTT)), lysed on ice by sonication, centrifuged (15 min, $8,000 \times g$), and the pellet was resuspended in 30 mL of buffer R containing 7 M urea. The solution was sonicated on ice again and the cell debris was removed by centrifugation (15 min, $8,000 \times g$). The supernatant was filtered (0.45 μ m) and loaded onto a 5-mL HiTrap SP column. Pfu RPP29 was eluted using a 50 mL linear 10-25% gradient of 25 mM to 2 M KCl under denaturing conditions (25 mM Tris-HCl, pH 7, 0.1 mM PMSF, 7 M urea), refolded by dialyzing into NMR buffer [10 mM Tris-HCl, pH 6.7, 10 mM KCl, 0.3 mM ZnCl₂, 0.02% (w/v) NaN₃] and concentrated by ulfiltration with a 5-kDa molecular weight cutoff membrane (Ultra-4, Amicon).

Pfu **RPP21**—The *Pfu* RPP21 protein was overexpressed and purified as previously described ³⁵, except that at each refolding step, the protein was refolded from denaturant solutions (8 M urea) by dialysis, instead of by rapid buffer exchange in a size exclusion column: 1) the protein eluted from the nickel-loaded Hi-Trap chelating column (GE Healthcare) was refolded into NMR buffer by dialysis before thrombin cleavage; 2) the lyophilized protein following HPLC purification was resuspended in denaturing buffer, and refolded by dialyzing against NMR buffer.

NMR Spectroscopy and Resonance Assignments

The *Pfu* RPP21–RPP29 complex was formed by combining each [U-¹⁵N, ¹³C]-labeled protein (~1 mM) with an unlabeled protein partner at a 1:1.2 molar ratio to promote full saturation of the labeled protein ³⁵. Taking advantage of the thermostability of *Pfu* proteins, NMR spectra were recorded at 55°C on 600 and 800 MHz Bruker Avance DRX spectrometers (Billerica, MA) equipped with triple resonance pulse-field gradient probes. NMR spectra were processed and analyzed by NMRPipe ⁵⁰, NMRView ⁵¹ and CARA ⁵².

Assignments of ¹H, ¹⁵N, and ¹³C resonances for each labeled protein in complex with its unlabeled partner were obtained by using data from the following experiments at 600 MHz: ¹⁵N-¹H HSQC, ¹³C-1H HSQC, HNCO, HNCACB, CBCA(CO)NH, C(CO)NH-TOCSY (τ_m 12 ms) and H(C)(CO)NH-TOCSY (τ_m 12 ms) ⁵³. Distance restraints were obtained from 3D ¹⁵N-separated NOESY-HSQC (τ_m = 100 ms) and ¹³C-separated NOESY-HSQC (τ_m = 100 ms) spectra recorded at 800 MHz on samples dissolved in 10 and 99.8% D₂O, respectively. Intermolecular distance restraints were assigned initially from ¹³C-filtered/edited NOESY-HSQC spectra (τ_m 100 ms, 800 MHz, 99.8% D₂O) with the two ¹J_{CH} filter elements tuned to 120 and 160 Hz ⁴³; ⁴⁴.

A heteronuclear $\{^{1}H\}^{-15}N$ NOE experiment was recorded at 600 MHz on $[U^{-15}N, {}^{13}C]$ -RPP29 bound to unlabeled RPP21 in an interleaved manner with (NOE) and without (no NOE) ${}^{1}H$ saturation 54 . Heteronuclear NOE values were determined from the ratios of the peak intensities between the two spectra.

Structure Determination

Distance restraints were derived from the NOE peak intensities, which were calibrated by assigning the median intensity to an interproton distance, r, of 2.7 Å, and scaling the remaining intensities by $1/r^651$. Restraints for methyl groups and geminal protons were adjusted by adding 0.5 Å to the upper bound. Backbone torsion angle restraints were obtained from analysis of the backbone chemical shifts via TALOS ⁵⁵. Hydrogen bonds were identified based on the secondary structural information and characteristic NOE patterns.

Structure calculations were performed using simulated annealing protocols within the XPLOR-NIH software suite ⁵⁶. Preliminary analyses of the ¹⁵N, ¹³C-edited and ¹³C-filtered/edited NOE spectra identified a total of 323 unique (non-redundant) unambiguous NOEs, including 105 intramolecular NOEs in RPP21, 124 intramolecular NOEs in RPP29, and 284 intermolecular NOEs in the binding interface. An initial set of structures of each individual protein was generated using the unique intramolecular NOEs, in addition to short-range NOE restraints and dihedral angles restraints. These initial ensembles (RPP21 and RPP29, respectively) were then used as templates for iterative computer-aided structure-based NOE assignment (SANE) ⁵⁷ of the isolated proteins, resulting in assignment of additional intramolecular NOEs. Then, using the NOE and dihedral angle restraints obtained for the refined structure ensembles of the individual proteins, and the 284 intermolecular NOEs, an initial ensemble of the protein-protein complex was generated from an extended template. This initial docked ensemble of the complex served as the template for iterative SANE, yielding an additional 191 intermolecular NOEs. One-hundred trials of restraint-driven refinement resulted in an ensemble of ~70 structures with similar restraint energies, from which a final set of 10 structures was selected for analysis and evaluation with XPLOR-NIH and PROCHECK-NMR ⁵⁸. Surface burial was calculated using the program STC ⁵⁹.

RNase T1- and RNase V1-based footprinting to map RNA-protein interactions in Methanocaldococcus jannaschii (Mja) RNase P

Details regarding the purification of RPPs and reconstitution of Mja RNase P are described elsewhere ²⁹. The footprinting studies were conducted as follows. Each 40-µL footprinting reaction contained a trace amount of end-labeled (100,000 dpm, ~0.5 nM) and unlabeled (125 nM) RPR either alone or complexed with RPP21–RPP29 or RPP30-POP5 (1.25 μ M) in 50 mM Tris-acetate (pH 7.5), 120 mM Mg(OAc)₂, 400 mM NH₄(OAc). For the reactions with all four RPPs, the reconstitution was performed in 50 mM Tris-acetate (pH 7.5), 30 mM Mg $(OAc)_2$, 800 mM NH₄(OAc). In all cases, the reconstitutions involved two sequential incubations for 10 min at 37°C and 55°C. Subsequently, 1 µL RNase T1 [0.1 U/µL, diluted 10-fold in water from commercial stock (Ambion)] was added to the reconstitution mix and the incubation was continued at 55°C for an additional 1 min for the RNA-alone reaction and 5 min for the RNP reactions. Alternatively, 1 μ L RNase V1 [0.02 U/ μ L, diluted 5-fold in water from commercial stock (Ambion)] was added to the reconstitution mix and the incubation was continued at 55°C for an additional 1 min for the RNA-alone reaction and 8 min for the RNP reactions. Both the T1 and V1 reactions were terminated by adding 10 µL of buffer-saturated phenol (pH 8) followed by extraction with phenol/chloroform. The RNAs were precipitated by adding two volumes of ethanol in the presence of 0.3 M sodium acetate (pH 5.2) and 20 μ g/ml glycogen. The RNAs were then pelleted at 18,000 × g for 15 min and the pellets washed twice with 70% (v/v) ethanol. The air-dried RNA samples were then resuspended in 10 μ L loading dye [9 M urea, 0.9 mM EDTA, 0.05% (w/v) bromophenol blue, 0.05% (w/v) xylene cyanol and 10% (v/v) phenol], separated by 8% (w/v) polyacrylamide/7 M urea gel electrophoresis, and visualized by using a Typhoon PhosphorImager (GE Healthcare). Size markers were generated by using both a partial hydrolysis of the RPR in alkali and digestion of the RPR with RNase T1 under denaturing conditions ¹³. Compression-related artifacts during denaturing PAGE occasionally skews ladder assignments by a few nucleotides.

In addition to direct mapping of the cleaved fragments, we also employed primer extension assays as a secondary validation ²⁹. After cleavage by RNase T1, the reaction contents were precipitated, washed with 70% (v/v) ethanol, air-dried and resuspended in water. The oligonucleotide *Mja* RPR-R (5'-GGGGGGATCCGTCTCGGCGGGTATGGGGGGC-3'), which is complementary to the 3'-end of *Mja* RPR was 5'-labeled with [γ -³²P]-ATP and T4 polynucleotide kinase and used to prime the reverse transcription of the partial digestion products of *Mja* RPR. To minimize artificial stops caused by the secondary structure of the RPR, the extension reactions were performed at 50 °C using ThermoScript (Invitrogen) reverse transcriptase as specified by the supplier. The products of the reverse transcription reactions were separated in an 8% (w/v) polyacrylamide/7M urea gel in parallel with a DNA sequencing ladder using *Mja*RPR-R as the primer and the plasmid pBT7-*Mja* RPR ²⁹as the template.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Abbreviations

RNP, ribonucleoprotein (RNA + protein) RNase P, Ribonuclease P RPP, RNase P protein subunit RPR, RNase P RNA subunit tRNA, transfer RNA *Pfu, Pyrococcus furiosus Mja, Methanocaldococcus jannaschii* NMR, nuclear magnetic resonance spectroscopy NOE, nuclear Overhauser effect SANE, structure-assisted NOE evaluation RMSD, root-mean-square deviation

Acknowledgments

We thank C. Yuan and C. Cottrell (CCIC) for assistance with the NMR data collection, and R. C. Wilson, I. R. Kleckner and the members of the V. Gopalan laboratory (especially Lien Lai) for reagents, encouragement, and helpful discussions. This work was supported by a grant from the NIH to M.P.F and V.G. (GM067807).

References

1. Altman S. A view of RNase P. Mol Biosyst 2007;3:604-607. [PubMed: 17700860]

- Gopalan, V.; Altman, S. Ribonuclease P: Structure and Catalysis. In: Gesteland, R.; Cech, T.; Atkins, J., editors. The RNA World. New York: Cold Spring Harbor Laboratory Press; 2006. only online at http://rna.cshl.edu
- Evans D, Marquez SM, Pace NR. RNase P: interface of the RNA and protein worlds. Trends Biochem Sci 2006;31:333–341. [PubMed: 16679018]
- 4. Guerrier-Takada C, Gardiner K, Marsh T, Pace N, Altman S. The RNA moiety of ribonuclease P is the catalytic subunit of the enzyme. Cell 1983;35:849–857. [PubMed: 6197186]
- Pannucci JA, Haas ES, Hall TA, Harris JK, Brown JW. RNase P RNAs from some Archaea are catalytically active. Proc Natl Acad Sci U S A 1999;96:7803–7808. [PubMed: 10393902]
- Kikovska E, Svard SG, Kirsebom LA. Eukaryotic RNase P RNA mediates cleavage in the absence of protein. Proc Natl Acad Sci U S A 2007;104:2062–2067. [PubMed: 17284611]
- 7. Hsieh J, Andrews AJ, Fierke CA. Roles of protein subunits in RNA-protein complexes: lessons from ribonuclease P. Biopolymers 2004;73:79–89. [PubMed: 14691942]

- Sun L, Campbell FE, Zahler NH, Harris ME. Evidence that substrate-specific effects of C5 protein lead to uniformity in binding and catalysis by RNase P. EMBO J 2006;25:3998–4007. [PubMed: 16932744]
- 9. Sun L, Harris ME. Evidence that binding of C5 protein to P RNA enhances ribozyme catalysis by influencing active site metal ion affinity. RNA 2007;13:1505–1515. [PubMed: 17652407]
- Smith JK, Hsieh J, Fierke CA. Importance of RNA-protein interactions in bacterial ribonuclease P structure and catalysis. Biopolymers 2007;87:329–338. [PubMed: 17868095]
- Hartmann E, Hartmann RK. The enigma of ribonuclease P evolution. Trends Genet 2003;19:561– 569. [PubMed: 14550630]
- Gopalan V. Uniformity amid diversity in RNase P. Proc Natl Acad Sci U S A 2007;104:2031–2032. [PubMed: 17287341]
- Tsai HY, Pulukkunat DK, Woznick WK, Gopalan V. Functional reconstitution and characterization of Pyrococcus furiosus RNase P. Proc Natl Acad Sci U S A 2006;103:16147–16152. [PubMed: 17053064]
- Hall TA, Brown JW. Archaeal RNase P has multiple protein subunits homologous to eukaryotic nuclear RNase P proteins. RNA 2002;8:296–306. [PubMed: 12003490]
- Kazantsev AV, Pace NR. Bacterial RNase P: a new view of an ancient enzyme. Nat Rev Microbiol 2006;4:729–740. [PubMed: 16980936]
- 16. Brown JW. The Ribonuclease P Database. Nucleic Acids Res 1999;27:314. [PubMed: 9847214]
- Loria A, Pan T. Domain structure of the ribozyme from eubacterial ribonuclease P. RNA 1996;2:551– 563. [PubMed: 8718684]
- Krasilnikov AS, Xiao Y, Pan T, Mondragon A. Basis for structural diversity in homologous RNAs. Science 2004;306:104–107. [PubMed: 15459389]
- Krasilnikov AS, Yang X, Pan T, Mondragon A. Crystal structure of the specificity domain of ribonuclease P. Nature 2003;421:760–764. [PubMed: 12610630]
- Torres-Larios A, Swinger KK, Krasilnikov AS, Pan T, Mondragon A. Crystal structure of the RNA component of bacterial ribonuclease P. Nature 2005;437:584–587. [PubMed: 16113684]
- Kazantsev AV, Krivenko AA, Harrington DJ, Holbrook SR, Adams PD, Pace NR. Crystal structure of a bacterial ribonuclease P RNA. Proc Natl Acad Sci U S A 2005;102:13392–13397. [PubMed: 16157868]
- 22. Kazantsev AV, Krivenko AA, Harrington DJ, Carter RJ, Holbrook SR, Adams PD, Pace NR. Highresolution structure of RNase P protein from Thermotoga maritima. Proc Natl Acad Sci U S A 2003;100:7497–7502. [PubMed: 12799461]
- Stams T, Niranjanakumari S, Fierke CA, Christianson DW. Ribonuclease P protein structure: evolutionary origins in the translational apparatus. Science 1998;280:752–755. [PubMed: 9563955]
- Spitzfaden C, Nicholson N, Jones JJ, Guth S, Lehr R, Prescott CD, Hegg LA, Eggleston DS. The structure of ribonuclease P protein from Staphylococcus aureus reveals a unique binding site for single-stranded RNA. J Mol Biol 2000;295:105–115. [PubMed: 10623511]
- Tsai HY, Masquida B, Biswas R, Westhof E, Gopalan V. Molecular modeling of the threedimensional structure of the bacterial RNase P holoenzyme. J Mol Biol 2003;325:661–675. [PubMed: 12507471]
- 26. Buck AH, Kazantsev AV, Dalby AB, Pace NR. Structural perspective on the activation of RNAse P RNA by protein. Nat Struct Mol Biol 2005;12:958–964. [PubMed: 16228004]
- Niranjanakumari S, Day-Storms JJ, Ahmed M, Hsieh J, Zahler NH, Venters RA, Fierke CA. Probing the architecture of the B. subtilis RNase P holoenzyme active site by cross-linking and affinity cleavage. RNA 2007;13:521–535. [PubMed: 17299131]
- Harris JK, Haas ES, Williams D, Frank DN, Brown JW. New insight into RNase P RNA structure from comparative analysis of the archaeal RNA. RNA 2001;7:220–232. [PubMed: 11233979]
- Pulukkunat DK, Gopalan V. Studies on Methanocaldococcus jannaschii RNase P reveal insights into the roles of RNA and protein cofactors in RNase P catalysis. Nucleic Acids Res 2008;36:4172–4180. [PubMed: 18558617]

- 30. Boomershine WP, McElroy CA, Tsai HY, Wilson RC, Gopalan V, Foster MP. Structure of Mth11/ Mth Rpp29, an essential protein subunit of archaeal and eukaryotic RNase P. Proc Natl Acad Sci U S A 2003;100:15398–15403. [PubMed: 14673079]
- Sidote DJ, Hoffman DW. NMR structure of an archaeal homologue of ribonuclease P protein Rpp29. Biochemistry 2003;42:13541–13550. [PubMed: 14622001]
- 32. Sidote DJ, Heideker J, Hoffman DW. Crystal structure of archaeal ribonuclease P protein aRpp29 from Archaeoglobus fulgidus. Biochemistry 2004;43:14128–14138. [PubMed: 15518563]
- Numata T, Ishimatsu I, Kakuta Y, Tanaka I, Kimura M. Crystal structure of archaeal ribonuclease P protein Ph1771p from Pyrococcus horikoshii OT3: an archaeal homolog of eukaryotic ribonuclease P protein Rpp29. RNA 2004;10:1423–1432. [PubMed: 15317976]
- Kakuta Y, Ishimatsu I, Numata T, Kimura K, Yao M, Tanaka I, Kimura M. Crystal structure of a ribonuclease P protein Ph1601p from Pyrococcus horikoshii OT3: an archaeal homologue of human nuclear ribonuclease P protein Rpp21. Biochemistry 2005;44:12086–12093. [PubMed: 16142906]
- Amero CD, Boomershine WP, Xu Y, Foster M. Solution structure of Pyrococcus furiosus RPP21, a component of the archaeal RNase P holoenzyme, and interactions with its RPP29 protein partner. Biochemistry 2008;47:11704–11710. [PubMed: 18922021]
- 36. Wilson RC, Bohlen CJ, Foster MP, Bell CE. Structure of Pfu Pop5, an archaeal RNase P protein. Proc Natl Acad Sci U S A 2006;103:873–878. [PubMed: 16418270]
- Takagi H, Watanabe M, Kakuta Y, Kamachi R, Numata T, Tanaka I, Kimura M. Crystal structure of the ribonuclease P protein Ph1877p from hyperthermophilic archaeon Pyrococcus horikoshii OT3. Biochem Biophys Res Commun 2004;319:787–794. [PubMed: 15184052]
- Kifusa M, Fukuhara H, Hayashi T, Kimura M. Protein-protein interactions in the subunits of ribonuclease P in the hyperthermophilic archaeon Pyrococcus horikoshii OT3. Biosci Biotechnol Biochem 2005;69:1209–1212. [PubMed: 15973057]
- Hall TA, Brown JW. Interactions between RNase P protein subunits in archaea. Archaea 2004;1:247– 254. [PubMed: 15810434]
- 40. Houser-Scott F, Xiao S, Millikin CE, Zengel JM, Lindahl L, Engelke DR. Interactions among the protein and RNA subunits of Saccharomyces cerevisiae nuclear RNase P. Proc Natl Acad Sci U S A 2002;99:2684–2689. [PubMed: 11880623]
- Kawano S, Nakashima T, Kakuta Y, Tanaka I, Kimura M. Crystal structure of protein Ph1481p in complex with protein Ph1877p of archaeal RNase P from Pyrococcus horikoshii OT3: implication of dimer formation of the holoenzyme. J Mol Biol 2006;357:583–591. [PubMed: 16430919]
- 42. Honda T, Kakuta Y, Kimura K, Saho J, Kimura M. Structure of an archaeal homolog of the human protein complex Rpp21–Rpp29 that is a key core component for the assembly of active ribonuclease P. J Mol Biol 2008;384:652–662. [PubMed: 18929577]
- 43. Breeze A. Isotope-filtered NMR methods for the study of biomolecular structure and interactions. Progress in Nuclear Magnetic Resonance Spectroscopy 2000;36:323–372.
- 44. Zwahlen CPL, Vincent SJF, Greenblatt J, Konrat R, Kay LE. Methods for Measurement of Intermolecular NOEs by Multinuclear NMR Spectroscopy: Application to a Bacteriophage λ N-Peptide/boxB RNA Complex. J Am Chem Soc 1997;119:6711–6721.
- 45. Jiang T, Guerrier-Takada C, Altman S. Protein-RNA interactions in the subunits of human nuclear RNase P. RNA 2001;7:937–941. [PubMed: 11455963]
- Gopalan V. Uniformity amid diversity in RNase P. Proc. Natl. Acad. Sci. U S A 2007;104:2031– 2032. [PubMed: 17287341]
- 47. Jarrous N, Reiner R, Wesolowski D, Mann H, Guerrier-Takada C, Altman S. Function and subnuclear distribution of Rpp21, a protein subunit of the human ribonucleoprotein ribonuclease P. RNA 2001;7:1153–1164. [PubMed: 11497433]
- Brannvall M, Kikovska E, Wu S, Kirsebom LA. Evidence for induced fit in bacterial RNase P RNAmediated cleavage. J. Mol. Biol 2007;372:1149–1164. [PubMed: 17719605]
- Pan T, Loria A, Zhong K. Probing of tertiary interactions in RNA: 2'-hydroxyl-base contacts between the RNase P RNA and pre-tRNA. Proc. Natl. Acad. Sci. U S A 1995;92:12510–12514. [PubMed: 8618931]
- Delaglio F, Grzesiek S, Vuister GW, Zhu G, Pfeifer J, Bax A. NMRPipe: a multidimensional spectral processing system based on UNIX pipes. J Biomol NMR 1995;6:277–293. [PubMed: 8520220]

- Johnson BA. Using NMRView to visualize and analyze the NMR spectra of macromolecules. Methods Mol Biol 2004;278:313–352. [PubMed: 15318002]
- 52. Keller R. The Computer Aided Resonance Assignment Tutorial. 2004
- 53. Cavanagh, RPsR. Protein NMR Spectroscopy: Principles and Practice. Vol. 2nd edit. New York: Academic Press; 2006.
- Kay LETD, Bax A. Backbone dynamics of proteins as studied by nitrogen-15 inverse detected heteronuclear NMR spectroscopy: application to staphylococcal nuclease. Biochemistry 1989;28:8972–8979. [PubMed: 2690953]
- 55. Cornilescu G, Delaglio F, Bax A. Protein backbone angle restraints from searching a database for chemical shift and sequence homology. J Biomol NMR 1999;13:289–302. [PubMed: 10212987]
- Schwieters CD, Kuszewski JJ, Tjandra N, Clore GM. The Xplor-NIH NMR molecular structure determination package. J Magn Reson 2003;160:65–73. [PubMed: 12565051]
- Duggan BM, Legge GB, Dyson HJ, Wright PE. SANE (Structure Assisted NOE Evaluation): an automated model-based approach for NOE assignment. J Biomol NMR 2001;19:321–329. [PubMed: 11370778]
- Laskowski RA, Rullmannn JA, MacArthur MW, Kaptein R, Thornton JM. AQUA and PROCHECK-NMR: programs for checking the quality of protein structures solved by NMR. J Biomol NMR 1996;8:477–486. [PubMed: 9008363]
- Lavigne P, Bagu JR, Boyko R, Willard L, Holmes CF, Sykes BD. Structure-based thermodynamic analysis of the dissociation of protein phosphatase-1 catalytic subunit and microcystin-LR docked complexes. Protein Sci 2000;9:252–264. [PubMed: 10716177]



Figure 1.

Binding-coupled folding in Pfu RPP21 and RPP29 as detected by NMR. Overlay of ¹⁵N HSQC spectra of each ¹⁵N-labeled protein in the absence (black) and presence (red) of its partner illustrates the site-specific chemical shift perturbations (CSPs). Weighted average CSP values (c, d) are mapped onto cartoon diagrams of the proteins (e, RPP21 and f, RPP29) using a linear color ramp from grey (no change) to red (maximal CSP). Blue indicates residues whose signals were only observed in the spectra of the complex. Black, undetermined CSP.



Figure 2.

Representative strips from ¹³C-filtered/edited NOE spectra recorded on (a) $[U^{-13}C, {}^{15}N]$ -RPP21 bound to unlabeled *Pfu* RPP29, and (b) $[U^{-13}C, {}^{15}N]$ -RPP29 with unlabeled RPP21. The black (positive) crosspeaks arise from inter-molecular NOEs, while the red (negative) peaks result from incomplete suppression of self- and intra-molecular NOEs. The ¹H and ¹³C shifts of the labeled partner are indicated above and below each strip, while the y-axis corresponds to the shift of ¹Hs from the unlabeled partner to which the labeled proton NOEs.

Xu et al.



Figure 3.

Solution structure of *Pfu* RPP29-RPP21 complex. Ensemble of 10 lowest-energy structures superimposed on the backbone heavy atoms of residues 18–122 of RPP29 and 10–54, 57–81 and 88–104 of RPP21. *Pfu* RPP29 and RPP21 are labeled in red and blue, respectively.

Xu et al.



Figure 4.

Interface of the *Pfu* RPP29-RPP21 complex. Electrostatic potential maps on the surface of RPP29 (a) and RPP21 (b) illustrate that the interface is dominated by hydrophobic interactions surrounding a prominent charge-charge interaction; the binding partner is shown as a ribbon (RPP21 in cyan and RPP29 in green). (c–e) Close-up showing three elements that stabilize the protein-protein complex. The lowest energy structure is chosen as the representative. The structure ensemble of these same views are provided in Supporting Information.

Xu et al.

Page 18



Figure 5.

Electrostatic potential map suggests two RNA binding regions. (a) Electrostatic potential map of *Pfu* the RPP29-RPP21 complex reveals two extended electropositive surface patches. The largest of these surfaces is composed of residues contributed by both proteins; the smaller is localized solely to the RPP21 zinc ribbon. (b) Highly conserved basic residues are shown on the ribbon diagrams of *Pfu* RPP29 (red) and *Pfu* RPP21 (green). The orientation is as in (a).

Xu et al.

Page 19



Figure 6.

Footprinting using RNase V1 and RNase T1 to identify RPP-binding sites in Mja RPR. Mja RPR labeled at the 5'-end (a) or 3'-end (b) was incubated either without (lanes 1, 3, 5, 7 and 9) or with (lanes 2, 4, 6, 8 and 10) RNase V1 (panel a) or RNase T1 (panel b). Mja RPR was present either alone (lanes 1, 2, 7 and 8), with RPP21-RPP29 (lanes 3 and 4), with RPP30-POP5 (lanes 5 and 6) or with both binary complexes (lanes 9 and 10). Since reconstitution of the RPR with each binary RPP complex is performed in a buffer different from that used for reconstitution with both binary complexes together, two different control RNase T1/V1 digestions of the RPR are shown (lanes 1, 2 for binary RPPs and lanes 7, 8 for both binary pairs). "Alk." and "T1" represent molecular size ladders generated by subjecting end-labeled, denatured Mja RPRs to alkaline hydrolysis and partial RNase T1 digestion, respectively. The RNase T1 cleavage sites were also mapped by using primer extension assays (data not shown ²⁹). * Longer electrophoretic runs were used to map protection patterns distal to the labeled termini (see Supporting Information). (c) Summary of the RPP footprinting data depicted on a secondary-structure model of Mja RPR. Circled and boxed nucleotides indicate protection to RNase T1 and RNase V1, respectively; blue and red colors indicate regions of protection by RPP30-POP5 and RPP21-RPP29, respectively. The green arrow indicates an RPR position that showed increased susceptibility to RNase T1 in the presence of either RPP30-POP5 or all four RPPs. RNase V1 cleavages around nucleotides 130-150 suggest that the secondary structure as drawn may need to be revised.

Table 1

Structural Statistics for RPP29-RPP21 complex

NMR constrains	RPP29	RPP21
NOEs	2038	1407
Intraresidue $(i - j = 0)$	833	606
Sequential $(i - j = 1)$	475	364
Short range $(1 < i - j < 5)$	204	203
Long range $(i - j > 5)$	526	234
Intermolecular (RPP29 – RPP21)	472	
Ambiguous	376	328
Hydrogen bonds ^a	38	80
Dihedral angles	188	162
Structure statistics		
Violations		
Distance violations > 0.5 Å	1.50 ± 1.02	
Dihedral angle violations $> 5^{\circ}$	1.63 ± 0.10	
Deviation from idealized geometry		
Bonds, Å	0.0046 ± 0.00008	
Angles, °	0.77 ± 0.02	
Impropers, °	0.56 ± 0.01	
Ramachandran statistics ^b , %		
Favored	77.0	
Additionally allowed	21.1	
Generously allowed	1.4	
Disallowed	0.5	
Precision ^C (RMSD form the mean structure)		
Backbone atoms, Å	0.58	
All heavy atoms, Å	0.87	

 a Hydrogen bonds were applied as upper bound restraints between amide proton and oxygen atoms and between amide nitrogen and oxygen atoms.

 $^{b}{\rm Ramachandran}$ analysis performed with PROCHECK-NMR 58

^CStructure statistics were calculated using the 10 lowest energy structures; RMSDs calculated by superimposing residues 18–122 of RPP29, and 9–54, 57–81 and 86–104 of RPP21.