SUPPLEMENTARY MATERIAL

Dissecting functional cooperation among protein subunits in

archaeal RNase P, a catalytic ribonucleoprotein complex

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MATERIALS AND METHODS

Abbreviations. *Eco*, *Escherichia coli*; *Mja*, *Methanocaldococcus jannaschii*; *Mth*, *Methanothermobacter thermautotrophicus*; *Pfu*, *Pyrococcus furiosus*; pre-tRNA, precursor tRNA; RPR, RNase P RNA; RPP, RNase P protein

Cloning of *Mja* **RPR**!**S and** *Mja* **RPR**!**S Min.** The gene encoding *Mja* RPR!S was generated using a PCR-based deletion mutagenesis approach with pBT7-*Mja* RPR (1) as the template. Briefly, T4 polynucleotide kinase was used to phosphorylate the primers Mja RPRAS-F and Mja $RPRAS-R$ (Table S1); these primers flank the region to be deleted and are oriented outward to ensure amplification of the entire plasmid without the nucleotides 67-192 in the Mia RPR-coding region. The resulting PCR products were circularized by ligation to generate pBT7-Mja RPRAS, in which the ΔS derivative is controlled by a T7 RNA polymerase promoter.

A two-step PCR-based approach was adopted to generate a clone encoding *Mja* RPRAS Min. Using pBT7-*Mja* RPRAS as the template and Mja RPRAS Min-F2 and Mja RPRAS Min-R as primers, we first generated a 97-bp product containing the coding sequence corresponding to nts 26 to 112 of *Mja* RPRAS Min. Using this 97-bp product as a template and M ja RPR Δ S Min-F1 and Mja RPR Δ S Min-R as primers, we then generated a 121-bp product containing the entire coding sequence for *Mja* RPRAS Min. This PCR product was digested with *BamHI*, whose recognition site was included in Mja RPRAS Min-R, and then cloned into pBT7 (2) that had been digested with *Stu*I (which generates a blunt end) and *Bam*HI. The resulting plasmid, pBT7-*Mja* RPRAS Min, has the *Mja* RPRAS Min gene under the control of a T7 RNA polymerase promoter.

Cloning of *Reclinomonas americana* **mitochondrial (***Ram* **mt) RPR.** The gene encoding *Ram* mt RPR was amplified by PCR with the primers Ram mt RPR-F and Ram mt RPR-R, using as template a plasmid containing the *Ram* mt RPR gene (kind gift from Drs. Elias Seif and Franz Lang, University of Montreal, Canada) (3). The PCR product was digested with *Eco*RI, whose recognition site was included in the reverse primer, and then cloned into pBT7 (2) that had been digested with *Stu*I and *Eco*RI. The resulting plasmid, pBT7-*Ram* mt RPR, has the *Ram* mt RPR gene under the control of a T7 RNA polymerase promoter.

In vitro **transcription of various RPRs used in this study.** Preparation of *Afu*, *Eco*, *Mja*, *Mth* and *Pfu* RPRs are described elsewhere (1,4,5). *Mja* RPR mutant derivatives were generated using the corresponding *Bsm*AI-linearized pBT7 plasmids as the template DNA for T7 RNA polymerase-mediated run-off transcription (6). For *Ram* mt RPR, the template was *Eco*RI-

linearized pBT7-*Ram*-mt RPR. The RNA transcripts thus generated were subsequently subjected to extensive dialysis to remove unincorporated rNTPs and their concentrations determined from Abs₂₆₀ measurements and their respective extinction coefficients.

Cloning the genes encoding the archaeal RPPs. The genes encoding *Mth or Mja* POP5, RPP21, RPP29 and RPP30 (Table S2) were amplified by PCR from the respective genomic DNA using the corresponding gene-specific primers listed in Table S1. All PCR products were digested with *Bam*HI, whose recognition site was included in the various reverse primers. The *Mth*/*Mja* POP5- and RPP29-encoding ORFs were cloned into pLANT-2b (7) that had been digested with *Nde*I, filled in with Klenow, and subsequently digested with *Bam*HI. The *Mth*/*Mja* RPP21- and RPP30-encoding ORFs were cloned into pET-15b that had been digested with *Nco*I, filled in with Klenow, and subsequently digested with *Bam*HI. These cloning approaches ensured that no additional amino acid residues were introduced in the four encoded proteins.

Automated DNA sequencing at the OSU Plant-Microbe Genomics Facility was used to ascertain the sequence of the clones obtained using the procedures described above.

Construction of expression plasmids containing two archaeal RPP genes in tandem.

Mja RPPTC1: Digestion of pET-15b-*Mja*RPP30 with *Bgl*II and *Blp*I released the *Mja* RPP30 ORF together with its upstream T7 RNA polymerase promoter and downstream T7 terminator. This fragment was then inserted into pLANT-2b-*Mja* POP5 that had been digested with *Bam*HI and *Blp*I to generate *Mja* RPPTC1, a construct in which *Mja* POP5 and RPP30 are present in tandem, each with its own T7 RNA polymerase promoter and control elements as originally conceived in the pET system (8).

Mja RPPTC2: Digestion of pLANT-2b-*Mja* RPP29 with *Bgl*II and *Eco*RI released the *Mja*RPP29 ORF together with its upstream T7 RNA polymerase promoter and downstream T7 terminator as well as the RIL tRNA gene cluster present in pLANT-2b. This fragment was inserted into pET-15b-*Mja* RPP21 that had been digested with *Bgl*II and *Eco*RI to generate *Mja* RPPTC2, a construct in which *Mja* RPP21 and RPP29 are present in tandem.

Mth RPPTC1: The same approach as *Mja* RPPTC1 was used (except with *Mth* RPP-encoding plasmids) to generate a construct in which *Mth* POP5 and RPP30 are present in tandem.

Mth RPPTC2: Digestion of pET-15b-*Mth* RPP21 with *Bgl*II and *Blp*I released the *Mth* RPP21 ORF together with its upstream T7 RNA polymerase promoter and downstream T7 terminator. This fragment was inserted into pLANT-2b-*Mth* RPP29 that had been digested with *Bam*HI and *Blp*I to generate *Mth* RPPTC2, a construct in which *Mth* RPP21 and RPP29 are present in tandem.

Pfu RPPTC1: Digestion of pET-33b-*Pfu* RPP30 (5) with *Bgl*II, followed by fill-in with Klenow and digestion with *Blp*I released the *Pfu* RPP30 ORF with its upstream T7 RNA polymerase promoter and downstream T7 terminator. This fragment was inserted into pET-33b-*Pfu* POP5 (5) that had been digested with *Eco*RI, filled in with Klenow, and then digested with *Blp*I, to generate *Pfu* RPPTC1, a construct in which *Pfu* POP5 and RPP30 are present in tandem.

Pfu RPPTC2: Same approach as *Pfu* RPPTC1, except that the insert was derived from digestion of pET-33b-*Pfu* RPP21 and the vector recipient was pET-33b-*Pfu* RPP29. *Pfu* RPPTC2 is a construct in which *Pfu* RPP29 and RPP21 are present in tandem.

Overexpression of archaeal RPPs. For co-overexpression of archaeal binary RPPs in *E. coli* BL21(DE3) cells, we used plasmids in which the two ORFs were present in tandem. The induction procedures were identical for *Mja*, *Mth* and *Pfu* RPPs. We use the *Mja* RPPs as a representative in the following description. *E. coli* BL21(DE3) cells were freshly transformed with pLANT-2b-*Mja* RPPTC1 or pET-15b-*Mja* RPPTC2. A single colony was used to inoculate 5 ml LB medium containing 35 µg/mL kanamycin (Mja POP5.RPP30) or 100 µg/mL carbenicillin (*Mja* RPP21 • RPP29), respectively, and grown overnight at 37°C with shaking. These overnight cultures were used to inoculate 500 ml of fresh LB media containing the appropriate antibiotics. The cells were grown at 37° C with shaking and induced with 1 mM isopropyl- β -D-thiogalactoside (IPTG) at Abs₆₀₀ ~0.6 to 0.8. To the RPP21 • RPP29-expressing cultures, 1 mM $ZnCl₂$ was also added at the time of induction since RPP21 requires $Zn²⁺$ to fold properly (9,10). Subsequent to IPTG addition, the cells were were grown either at 37°C for 3 h (*Mja* RPP21 • RPP29) or at 24°C for 15 h (*Mja* POP5 • RPP30) and harvested by centrifugation. The cell pellets were stored at -80°C until further use.

Purification of archaeal binary RPP complexes*.*

Mja RPPs: Frozen cells (from a 125 ml culture) were thawed on ice and re-suspended in 20 ml of buffer A [25 mM Tris-HCl (pH 7.5), 5 mM DTT, 0.1 mM PMSF, 25 mM NaCl]. Subsequently, 5 ml of buffer S [25 mM Tris-HCl (pH 7.5), 5 mM DTT, 0.1 mM PMSF and 5 M NaCl] was added and the cells were lysed by sonication. Cell debris was pelleted by centrifugation (9,000 g, 15 min, 4°C). Polyethylenimine (PEI) was added to the cleared supernatant to a final concentration of 0.05% (v/v) and incubated on ice for 30 min. The PEI-precipitated nucleic acid was pelleted by centrifugation (9,000 g, 15 min, 4°C). To the resulting soluble fraction, finely powdered $(NH_4)_2SO_4$ was added slowly (to a final saturation of 40%) and precipitation performed over a period of 60 min with constant stirring on ice. The precipitated RPPs were recovered by centrifugation (9,000 g, 15 min, 4°C). The precipitate was dissolved in 3 ml of buffer RS [1 M $(NH_4)_2SO_4$, 25 mM Tris-HCl (pH 7.5), 5 mM DTT, 0.1mM PMSF, 25 mM NaCl] and diluted by adding buffer A to ensure a final $(NH_4)_2SO_4$ concentration of 375 mM. The solution was filtered through a 0.4 µm filter and loaded on a pre-equilibrated SP-Sepharose (GE Healthcare) column. The bound proteins were eluted with a 0 - 2 M NaCl gradient in buffer A. Binary complexes typically eluted between 1.2 and 1.5 M NaCl.

Mth RPPs: The purification procedures for *Mth* binary complexes were the same as *Mja* RPPs except that the final $(NH_4)_2SO_4$ concentration was adjusted to 250 mM before loading on SP-Sepharose. The binary complexes typically eluted at ~1.5 M NaCl.

Pfu RPPs: The purification procedures for *Pfu* binary proteins were the same as *Mja* RPPs except that *Pfu* POP5 \bullet RPP30 was precipitated with 80% (NH₄)₂SO₄ and the final (NH₄)₂SO₄ concentration was adjusted to either 175 mM (RPP21 \bullet RPP29) or 250 mM (POP5 \bullet RPP30) before loading on SP-Sepharose. The binary complexes typically eluted at ~1.5 M NaCl.

Other comments: An ÄKTA FPLC purifier (GE Healthcare) was used for all chromatographic procedures. Fractions containing the archaeal RPPs were identified by SDS-PAGE analysis and Coomassie Brilliant blue staining. Using the extinction coefficients of the binary RPP complexes and their Abs₂₈₀ values, the concentration of each binary complex was calculated. The purified RPP complexes were dialyzed against 50 mM Tris-acetate (pH 8), 800 mM NH4OAc and 10 mM $Mg(OAc)_2$, and stored at room temperature until further use.

Estimation of the molar ratio of *Pfu* **POP5**!**RPP30 and RPP21**!**RPP29 binary complexes.** About 2.5 µg of each RPP pair was separated in a 15% (w/v) acrylamide gel (37.5:1) using Tricine-SDS-PAGE (11) and stained with SYPRO Ruby protein gel staining dye (Invitrogen). The gels were imaged with VersaDoc System, which utilizes a CCD camera to capture UVfluorescence images of the gels. The intensities of bands were analyzed using the software PDQuest (Bio-Rad). The relative stoichiometries were calculated from the intensities of the individual RPP bands and the molecular weight of the different RPPs.

Pulse-chase experiments with *Mth* **RNase P to demonstrate that the dissociation rate (k-1) is greater than the rate of chemical cleavage (k₂) under single-turnover conditions. These** pulse-chase experiments were performed essentially as described elsewhere for studies on the hammerhead ribozyme (12). RPR folding, RNP assembly and assay conditions are elaborated in Materials & Methods (of the main text). *Mth* RPR (16.6 µM) was mixed with a trace amount of internally labeled pre-tRNA^{Tyr} (~2 nM) and incubated at 55°C in a final reaction volume of 20 μ l. From this mixture, two 5-ul aliquots were diluted 100-fold in 55° C pre-warmed assay buffer [50] mM MES (pH 5.8), 2 M NH₄OAc, 500 mM MgCl₂], one at 20 and the other at 30 min after the initiation of the reaction; the remainder proceeded without any dilution. Similarly, for *Mth* RPR + RPP21•RPP29 (3 μ M), two 5- μ l aliquots were diluted 100-fold in pre-warmed assay buffer [50 mM MES (pH 5.8), 800 mM NH₄OAc, 120 mM MgCl₂], one at 5 and the other at 10 min after the initiation of the reaction. From both the undiluted and diluted reactions, aliquots were removed at defined time intervals to measure the product formed. Reactions were terminated by adding stop solution [10 M urea, 5 mM EDTA, 0.05% (w/v) bromophenol blue, 0.05% (w/v) xylene cyanol, 10% (v/v) phenol]. The reaction products were separated on denaturing PAGE and quantitated as described in the Materials & Methods (of the main text).

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Oligonucleotides used in this study

*Note***:** All oligonucleotides were purchased from either Integrated DNA Technologies or Sigma Genosys. Sequences are listed from 5' to 3'. Restriction sites, if incorporated into primers to facilitate subsequent cloning into appropriate vectors, are shown in italic font.

Characteristics of RPPs from different archaea

Note**:** Gene IDs and sequences from the Comprehensive Microbial Resource, J. Craig Venter Institute (http://cmr.jcvi.org).

Quantitation of the stoichiometric ratio of *Pfu* **RPPs in purified binary complexes***

* The values reported are the mean from at least two independent experiments.

** The numbers indicate the molecular ratio of POP5 to RPP30 and RPP21 to RPP29.

Steady-state kinetic parameters for processing of *Eco* **pre-tRNATyr by** *in vitro* **reconstituted** *Pfu* **RNase P***

* The values reported are the mean and standard deviation from at least three independent trials. The curve-fit errors for k_{cat} and K_M were less than 25% in individual experiments. The experiment was performed essentially as described elsewhere (5). Assays were performed at 55°C in 50 mM Tris (pH 7.5), 0.8 M NH₄OAc and 30 mM MgCl₂. The *Pfu* RNase P holoenzyme was assembled using 10 nM folded RPR and 100 nM RPPs. The pre-tRNA^{Tyr} concentration range used for measuring the K_M was 31 - 2000 nM. Turnover numbers were calculated assuming that all of the RPR is assembled into holoenzyme.

Figure S1. Secondary structures of *Eco*, *Pfu*, and *Mth* RPRs (13). Paired helices (e.g., P1, P2) are numbered consecutively from 5' to 3' and according to the *Eco* RPR nomenclature (13). A linker that joins helices P11 and P12 is termed J11/12. Boxes and dotted lines indicate loophelix tertiary interactions.

Figure S2. Purification of recombinant *Pfu* RPPs as binary complexes: (A) POP5•RPP30, (B) RPP21•RPP29 and (C) comparison of RPPs purified either individually or as binary complexes. M indicates size marker. UI and I denote the crude extracts from un-induced and induced cultures, respectively. IP represents the ammonium sulfate-fractionated sample that was subjected to the SP-sepharose column chromatography.

Figure S3. Heterologous reconstitution of RNase P holoenzymes using different archaeal RPRs and RPPs. Archaeal RPRs representing type A (*Pfu* and *Mth*) and M (*Mja* and *Afu*) were reconstituted with all four RPPs of either type A (*Pfu*) or M (*Mja*) RNase P. For these reconstitutions, 50 nM folded RPR was assembled with 250 nM RPPs in 50 mM Tris·HCl (pH 7.5), 800 mM NH₄OAc and 30 mM Mg(OAc)₂. The resulting RNPs were incubated with 500 nM pre-tRNA^{Tyr} for 15 min at 55 °C.

Figure S4. Pulse-chase experiments with *Mth* RNase P to demonstrate that the dissociation rate (k_{-1}) is greater than the rate of chemical cleavage (k_2) under single-turnover conditions. (A) *Mth* RPR (16.6 μ M) and pre-tRNA^{Tyr} (2 nM) assayed in 50 mM MES (pH 5.8), 2 M NH₄OAc and 500 mM MgCl₂: the rate of chemical cleavage was measured without dilution $(•)$ and after dilution at 20 min (\bullet) or 30 min (\bullet). (B) *Mth* RPR + RPP21•RPP29 (3 μ M) and pre-tRNA^{Tyr} (2 nM) assayed in 50 mM MES (pH 5.8), 800 mM NH₄OAc and 120 mM MgCl₂: the rate of chemical cleavage was measured without dilution $(•)$ and after dilution at 5 min $(•)$ or 10 min $(•)$. The green and blue lines indicate the plateau theoretically expected post-dilution when $k_1 \gg k_2$.

Figure S5. Effects of *Mth* RPPs on the single-turnover rate of *Mth* RPR-catalyzed pre-tRNA cleavage. (A) Increased product formation as a function of incubation time is illustrated in a representative time-course of pre-tRNA^{Tyr} processing by the *Mth* RNase P holoenzyme. (B) Examples of the curve fits of the rate of product formation by *Mth* RPR with and without RPPs under their respective optimal conditions (see Table I for details). The concentrations of *Mth* RPR, RPR + POP5•RPP30, RPR + RPP21•RPP29 and RPR + 4 RPPs used were 1.5 μ M, 1.2 μ M, 1.2 μ M and 0.5 μ M, respectively. Most of these ES complexes are productive as indicated by their $\geq 90\%$ amplitudes, except for the reaction with RPR + RPP21•RPP29. Since we use a two-fold stoichiometric excess of RPPs:RPR to facilitate assembly of the respective RNP, it is possible that any RPP21•RPP29 uncomplexed with the RPR engages in RPR-independent

interactions with the pre-tRNA substrate and lowers the amplitude to 40%; however, this problem does not exist when RPP21•RPP29 is used in the context of 4 RPPs, presumably due to differences in assay conditions and/or the presence of POP5•RPP30.

Figure S6. Dependence of k_{obs} on assay pH for *Mth* RPR + 4 RPPs. These single-turnover assays were performed with a vast excess of enzyme $(3 \mu M)$ over the substrate (2 nM) , and in 50 mM MES (indicated pH), 800 mM NH₄OAc and 20 mM MgCl₂. Note that the assays were performed in 20 mM Mg²⁺ and not 30 mM Mg²⁺, which is optimal for the RPR + 4 RPPs; this change lowered the rate by ~6-fold and permitted manual rate measurements over the pH range used in this experiment.

Figure S7. Kinetic scheme as a framework to interpret the contribution of the two binary RPP complexes to archaeal RNase P catalysis. For simplicity, the scheme does not take into account possible thermodynamic coupling in the binding of the RPPs (or pre-tRNA) and Mg²⁺ to the RPR. While the role of RPPs to binding of Mg^{2+} ions is indicated, no distinction is made between metal ions essential for RPR folding and catalysis. Although POP5•RPP30 reduced K_{M(STO)} by two-fold in our single-turnover kinetic studies, in the absence of detailed binding studies, we are unable to comment on the role of this binary complex in substrate binding; this uncertainty is indicated by a question mark.