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PPR proteins shed a new light on RNase P biology

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A fast growing number of studies identify pentatricopeptide repeat (PPR) proteins as major players in gene expression processes. Among them, a subset of PPR proteins called PRORP possesses RNase P activity in several eukaryotes, both in nuclei and organelles. RNase P is the endonucleolytic activity that removes 5' leader sequences from tRNA precursors and is thus essential for translation. Before the characterization of PRORP, RNase P enzymes were thought to occur universally as ribonucleoproteins, although some evidence implied that some eukaryotes or cellular compartments did not use RNA for RNase P activity. The characterization of PRORP reveals a two-domain enzyme, with an N-terminal domain containing multiple PPR motifs and assumed to achieve target specificity and a C-terminal domain holding catalytic activity. The nature of PRORP interactions with tRNAs suggests that ribonucleoprotein and protein-only RNase P enzymes share a similar substrate binding process.

Introduction

Pentatricopeptide repeat (PPR) proteins compose a family of RNA binding proteins specific to eukaryotes and mostly involved in gene expression processes in organelles. PPR proteins are particularly numerous in land plants with up to 450 representatives in Arabidopsis thaliana.1 They are composed of tandem arrays of PPR motifs whose primary sequence is very degenerate,2,3 although their tertiary structure seems to be conserved, with each repeat folding into two antiparallel α helices.1-6 A succession of PPR motifs would thus make a superhelix that could act as a platform to bind RNA.7 The combinatorial nature of PPR proteins allows substrate specificity because individual PPR motifs appear to ensure the selection for individual nucleotides.6,7 Since their discovery over a decade ago, functional studies of PPR proteins have helped to answer many persistent questions regarding organelar gene expression processes.1 For example, studies are beginning to unravel how sequence specificity is achieved for hundreds of C to U RNA editing sites in transcripts from higher plant organelles.8 The characterization of PPR proteins has also helped to settle the long-standing debate over the existence of protein-only RNase P enzymes in eukaryotes.9 RNase P is a key enzyme of tRNA maturation. It was initially described as the endonuclease activity that removes the 5' leader sequences of tRNA precursors. It is therefore essential for producing functional tRNAs and, hence, indispensable for translation.10,11 RNase P was first characterized on a molecular level in Escherichia coli, where it is composed of an RNA molecule together with a single protein.12 The discovery that RNase P RNA held the actual catalytic activity of the enzyme13 won Sidney Altman the Nobel prize in 1989 and helped to establish the “RNA world” theory proposing that one stage in prebiotic evolution consisted of RNA molecules that were able both to catalyze biochemical reactions and to store genetic information.14 Subsequently, RNase P enzymes were characterized as similar ribonucleoprotein (RNP) enzymes in numerous other organisms and organelles including bacteria, archaea, yeast nuclei and mitochondria and animal cell nuclei.15,16 Isolated RNA subunits from Bacteria, Archaea, and Eukarya demonstrate catalytic activity only under extreme ionic conditions, whereas the corresponding RNA•protein holoenzymes are maximally active under physiological conditions.13 Apart from tRNAs, RNP RNase P enzymes are involved in the maturation of a wide array of substrates including rRNAs, protein-coding mRNAs, tmRNA, riboswitches, viral RNA, and snoRNA.10,16,17 From a mechanistic point of view, RNP RNases P interact with tRNA mainly in the horizontal stacking domain consisting of the T stem-loop and acceptor stem; they utilize two catalytic metal ions and conserved RNA residues for RNA cleavage.18,19 The structures of RNase P enzymes differ greatly, each containing an RNA molecule (whose structure is considerably reduced in size in some instances20) bound by a variable number of protein subunits ranging from one in bacteria to at least nine in eukaryotes.10,21-23 Still, the central point remained that all RNase P enzymes contained an RNA moiety responsible for catalytic activity, so that the ribonucleoprotein nature of RNase P became a dogma. RNase P, together with the ribosome, was viewed as one of the ultimate universally conserved vestiges of the RNA world.15 Nevertheless, long before the discovery of the PPR protein family, some experimental evidence contradicted the prevailing dogma and suggested that some eukaryotes could use a different kind of enzyme, devoid of RNA, for RNase P activity. Here we review both the early evidence for the existence of protein-only RNase P and the studies describing the actual identification and
Origins and expectations. The earliest reports of protein-only RNase P came from eukaryotic organelles—chloroplasts and mitochondria—that typically encode some or, in plant chloroplasts and vertebrate mitochondria, all of the tRNAs needed for translation of organelar-encoded proteins. In animals, mitochondrial tRNA genes are interspersed among protein-coding genes, such that production of functional mRNA species requires excision of mature tRNAs by precise 5′- and 3′-terminal endonucleolytic cleavages. In chloroplasts, most tRNA genes are transcribed into end-extended precursors bearing 5′- and 3′-terminal extensions that must be removed to yield mature tRNA.

The earliest expectations for the nature of RNase P from these organelles were based on their established bacterial origins: mitochondria descended from the α-proteobacteria and chloroplasts arose from within the cyanobacteria. Members of both bacterial phyla possess “conventional” (E. coli-like) ribonucleoprotein forms of RNase P. In particular, bacterial-like RNase P RNA has been identified in all sequenced red algae chloroplasts and in many green algae in the Prasinophyte lineage. For RNA has been identified in all sequenced red algae chloroplasts and vertebrate mitochondria related to the RNA subunit of bacterial RNase P33 and a nuclear-chondrial RNase P RNA that typically encodes some or, in plant chloroplasts and vertebrate mitochondria, all of the tRNAs needed for translation of organelar-encoded proteins. In animals, mitochondrial RNase P activity could be attributed entirely to a catalytic component of a proteobacterial-type RNase P RNA31 that is dependent upon a cyanobacterial protein subunit.30 (The equivalent protein subunit in C. paradoxa is presumably nuclear-encoded and imported into the cyanelle.) Likewise, the mitochondrion of the early-branching protozoan Reclinomonas americana encodes a proteobacterial-type RNase P RNA46 that is dependent upon a proteobacterial protein subunit for activity.

Initial evidence for an RNA component. Early support for a bacterial-like composition of mitochondrial RNase P was provided by genetic and biochemical determinations that in mitochondria of budding yeast (Saccharomyces cerevisiae), RNase P contained an essential mitochondrial encoded, RNA distantly related to the RNA subunit of bacterial RNase P30 and a nuclear-encoded protein unrelated to the bacterial protein subunit.34

The earliest characterizations of a putative vertebrate mitochondrial RNase P (from rat liver35 or human cells36), did not directly test for the presence of an RNA component. Further efforts by one group, however, led to a claim that human mitochondrial RNase P activity could be attributed entirely to a small amount of nuclear RNase P imported into mitochondria. Because these investigations employed a precursor to E. coli tRNA50—subIII—a substrate for nuclear but not for vertebrate mitochondrial RNase P—the enzyme described is now thought to be the abundant nuclear RNase P present in the starting cytosolic extracts.35,39

A critical assay: The substrate unmasked. Meanwhile, in the plant kingdom, transcription and processing of chloroplast tRNAs had been demonstrated in 1983 by Gruissem and Hallick. Further investigation of chloroplast RNase P was initiated with the expectation that it, too, would resemble the bacterial enzyme. Preliminary evidence accumulated by 1986–87 suggested that crude preparations of RNase P from both spinach and tobacco chloroplasts were sensitive to treatment both with protease and with nuclease, consistent with activity residing in an RNA-protein complex (ref. 40 and Wang et al., poster presentation, 1986 Cold Spring Harbor RNA Processing Meeting). An apparent inhibition of chloroplast RNase P by S. aureus micrococcal nuclease (MN) is shown in Figure 1A, lanes 3–6.

At the time, three primary criteria were used to confirm the presence of an essential RNA component in an RNA processing activity: (1) sensitivity to pre-treatment with nucleases having little or no specificity for RNA sequence or structure, (2) buoyant density in Cs salts, and (3) presence of co-fractionating RNA species of appropriate size (150–400 nucleotide length). In the most common nuclease sensitivity protocol, an enzyme fraction is incubated with micrococcal nuclease (MN) in presence of its catalytic cofactor Ca2+. The nuclease is then inactivated by addition of EGTA, which chelates most divalent cations much more strongly than it does Mg2+, a required cofactor for all RNase Ps. Remaining RNase P activity is then assayed by addition of substrate directly to the treated enzyme fraction. All three assays are, however, susceptible to artifacts or misinterpretation. In particular, nuclease treatment is complicated by the fact that most suitable nucleases are difficult to inhibit cleanly, but residual activity will destroy the reaction substrate. EGTA-inactivated MN often displayed some inhibition of RNA processing.40,41 Moreover, inhibition by active MN of non-RNA-containing enzymes had been observed (e.g., refs. 40 and 42). This was interpreted as resulting from degradation of bulk RNA, present in a crude extract, which was thought to stabilize the RNA processing complex under investigation.42,43

In order to conclusively show whether MN treatment was specifically inactivating chloroplast RNase P, Wang et al. asked whether MN-inhibited RNase P activity could be recovered by addition of non-specific RNA. The dramatic result, as shown in Figure 1A, lanes 7–10, was that addition of yeast RNA or of synthetic polynucleotides completely reversed the apparent inhibition by MN.44 Further work41 showed, not surprisingly, that Ca2+-depleted MN retains substrate binding ability, reversibly binding pre-tRNA with an apparent Kd of 1.35 μM. Polyanions such as heparin or synthetic polynucleotides compete with pre-tRNA for binding MN. The final picture is that binding of catalytically inactive MN to RNA substrate sterically blocks access to the cleavage site. Addition of excess non-specific RNA sequesters the inactive nuclease and frees the pre-tRNA substrate for productive cleavage by the processing enzyme. This phenomenon is referred to as “substrate occlusion” or “substrate masking.”44

Chloroplasts. With a reliable assay in hand, progress was rapid, and Wang et al. determined that chloroplast RNase P is completely insensitive to digestion with concentrations of micrococcal nuclease 20- to 50-fold greater than those required to inactivate E. coli RNase P.44 Furthermore, as shown in Figure 1B, the chloroplast activity has a buoyant density in CsCl (1.28 g/cm3) that is precisely centered within the density distribution.
of bulk protein. In this context, it is essential to note that because observed buoyant densities are a function of the density medium, and for values determined by refractometry, are also influenced by solvent composition, they cannot be directly compared between experiments. (In CsCl gradients, buoyant densities for pure protein, E. coli RNase P, and pure RNA are 1.28, 1.7, and -2.0 g/cm³; in CsSO₄ these are 1.23, 1.55, and 1.65 g/cm³. The most stringent test for presence or absence of an RNA is the extent to which enzyme activity co-fractionates with bulk protein or with a known protein-only enzyme. The coincidence of protein and enzyme densities for plant chloroplast and human mitochondrial RNase P indicates that neither enzyme could possess more than one copy of a 10- to 20-nt long RNA.

Mechanistic differences between the chloroplast enzyme and the ribozyme-type RNase P affirmed that the chloroplast enzyme could not have an RNA subunit like that of bacterial or yeast nuclear RNase P discussed in detail in the section on structural mimicry. Further studies of the 1000–2000-fold purified chloroplast activity indicated that it does not co-purify with any RNAs that can be 3'-end labeled, and that its hydrodynamic size, determined by gel filtration corresponds to a ~70 kDa globular protein.

Plant nuclei. Knowing that most soluble plastid proteins are encoded in the nucleus, translated in the cytoplasm, and imported into the organelle, Wang et al. suggested that chloroplast RNase P or related peptidyl peptides could have been recruited to process pre-tRNAs encoded in the nucleus and mitochondrion. To investigate this possibility, Oommen used the techniques successful for chloroplasts to demonstrate that wheat embryo extracts contained an authentic RNase P activity with properties essentially identical to those of chloroplast RNase P. (The reaction requirements and substrate specificity of this activity [ref. 48 and unpublished observations] suggested that it was localized to the nucleus). This activity is resistant to amounts of micrococcal nuclease at least 5-fold greater than required to fully inactivate E. coli RNase P. In CsCl gradients, the distribution of wheat RNase P activity is absolutely coincident with the distribution of bulk protein (1.28–1.29 g/cm³). Active fractions across the final ion-exchange column contained no RNA molecules whose abundance was correlated with RNase P activity; trace RNAs larger than tRNA present in the active fractions could be removed without reducing RNase P activity. Finally, gel filtration chromatography in the absence of urea indicated a hydrodynamic size corresponding to a ~120 kDa globular protein or protein complex.

Somewhat later, another group presented essentially identical results: a buoyant density identical with bulk protein and complete resistance to MN treatment. On the basis of its reaction requirements, this activity could be identified with nuclear rather than mitochondrial RNase P. At the time, these data were interpreted as consistent with wheat nuclear RNase P containing an RNA subunit associated with a large number of proteins that conferred a protein-like buoyant density and protected the RNA from nuclease attack. Other researchers separated two RNase P activities, possibly nuclear and mitochondrial, from carrot cell suspension culture. Presence or absence of RNA components was not established: buoyant densities were not determined and results of MN treatment were inconclusive because controls for substrate masking were not included and reaction products were not characterized. Of the two activities, one was inhibited only partially by a 10-fold excess of MN; the second was completely inhibited by either active or inactive MN at all concentrations tested, indicative of unresolved substrate masking.

Plant mitochondria. In 1990, two groups reported processing in vitro of plant mitochondrial pre-tRNAs with homologous mitochondrial extracts. Marchfelder et al. showed that RNase P-like activity in Oenothera mitochondrial lysates was completely
Rossmanith and colleagues in 1995. Using a fully homologous mitochondrial RNase P from vertebrates was reported by were not seen as compelling.

Physical characterization, these observations, though intriguing, were not seen as compelling.

Human mitochondria. The first purification of an authentic mitochondrial RNase P from vertebrates was reported by Rossmannith and colleagues in 1995. Using a fully homologous system with a mitochondrial-specific substrate, they achieved a clean separation of human mitochondrial RNase P from the nuclear enzyme, which was by then known to be an RNA•protein complex. Using an approach similar to that of Wang et al., Rossmannith then made a rigorous finding that this RNase P activity did not require any RNA component. First, activity was fully resistant to digestion with a 10-fold excess of MN. Second, in Cs2SO4 gradients, the buoyant density of RNase P was unaffected by digestion with a 10-fold excess of MN. Some RNAs larger than tRNA were present in the absence of poly(A), consistent with substrate masking. Hanic-Joyce and Gray, on the other hand, stated that the activity in wheat mitochondria was insensitive to MN digestion when assayed in the presence of poly(A). In the absence of further physical characterization, these observations, though intriguing, were not seen as compelling.

Kinetoplastid mitochondria. Mitochondria of the kinetoplastid parasite *Trypanosoma brucei* encode no tRNAs. Instead, all tRNAs are encoded in the nucleus and imported into the mitochondrion. Although it is uncertain whether any tRNAs are imported as 5'-extended precursors, it is known that kinetoplastid mitochondria do possess an active RNase P. In 2001, Salvati used the “masking-free” MN assay to demonstrate that highly-purified *T. brucei* RNase P was unaffected by digestion with a 10-fold excess of MN. Some RNAs larger than tRNA were present in active fractions but could be degraded without effect on RNase P activity. Notably, the hydrodynamic size estimated by gel filtration chromatography was about 70 kDa, the same size as chloroplast RNase P. In the absence of buoyant density or mechanistic data, however, these results were not considered definitive.

Hindsight. In retrospect, the ability to recognize the existence of protein-only RNase Ps was hindered by (1) justifiable expectations that organelles would have bacterial-type RNase P; most likely containing an organelle-encoded RNA subunit and an imported, nuclear-encoded polypeptide; (2) knowledge that yeast mitochondrial RNase P conformed to this model; (3) indications that RNase P in *C. paradoxa* cyanelles and *R. americana* mitochondria would follow the bacterial paradigm; and (4) evidence that yeast and human nuclear RNase Ps contained an RNA subunit related to the bacterial prototype. On the other hand, there was no obvious reason to doubt the validity of experimental work supporting a protein-only composition for RNase P in animal mitochondria, plant chloroplasts, or plant nuclei, nor was there convincing experimental evidence supporting other interpretations. Nevertheless, these conclusions remained controversial until isolated polypeptides, overexpressed from cloned cDNAs corresponding to defined genetic loci, were shown to possess RNase P activity.

**Identification at the Molecular Level of Protein-Only RNase P**

Characterization of the RNase P enzyme in human mitochondria. The concept of protein-only RNase P was definitely accepted only when the core components responsible for RNase P activity in human mitochondria were identified at the molecular level. In that study, Rossmannith and coworkers confirmed that this RNase P activity did not require any RNA component. Using an elegant approach combining proteomic identification of human mitochondrial RNase P (mtRNase P) complexes, in vitro mtRNase P activity assay and reverse genetics, the authors’ work led to the conclusion that only three individual polypeptide subunits were strictly required for the reconstitution of mtRNase P activity and that their mode of action was concerted. These three polypeptides composing the mtRNase P holoenzyme are nuclear-encoded and were named respectively MRPP1, 2, and 3 (for Mitochondrial RNase P Proteins). MRPP1 (or TRMT10C) encodes a putative tRNA:m1G9-methyltransferase whereas
MRPP2 (or SDR5C1) encodes a 3-hydroxyacyl-CoA dehydrogenase and MRPP3 encodes a protein containing a metallonuclease domain as well as a PPR domain.\textsuperscript{5,55}

MRPP1 catalyzes the methylation of specific bases (G9 or A9) in mitochondrial tRNAs and interacts with tRNAs in vitro,\textsuperscript{56} although its methyltransferase activity is not required for tRNA cleavage by the mtRNase P holoenzyme.\textsuperscript{56} Little is known about the involvement of MRPP2 in mtRNase P activity. Binding to MRPP2 is critical for MRPP1 to perform mitochondrial tRNA methylation, although MRPP2’s dehydrogenase activity seems to be dispensable.\textsuperscript{56} Reciprocally, although MRPP1 and MRPP2 are essential components of the mtRNase P holoenzyme, neither the methyltransferase nor the dehydrogenase activity, respectively, is required for tRNA processing.\textsuperscript{56}

MRPP3 was the only identified subunit of mtRNase P harboring a predicted nuclease domain. Hence, it was hypothesized from the start that the involvement of MRPP3 in mtRNase P activity would be to perform the actual phosphodiester bond hydrolysis.\textsuperscript{55} MRPP3 also features PPR motifs. These elements are helical-repeat motifs considered to bind with specificity to single-stranded RNA stretches; they are found in eukaryotic proteins, predominantly those involved in organelar RNA metabolism.\textsuperscript{1,2,57} Even though the precise role of MRPP3’s PPR motifs in mtRNase P is still unexplored, a tempting proposal is that these repeats contribute to tRNA binding and/or confer base-specific recognition of tRNAs.

Apart from the protein-only RNase P, it was also proposed that RNase P RNA could be imported into human mitochondria,
MRPP3 orthologs could be identified in many eukaryotic organisms and define a new protein family that was named PRORP (for PROteinaceous RNase P). Hence, MRPP3 is now also called human PRORP. These proteins are characterized by the presence of a number of PPR and/or PPR-like motifs, a CXXC Zn finger-like motif and a metallo/nuclease domain belonging to the NYN family.60 The function of putative PRORPs identified by sequence similarities could account for the catalytic activity of RNase P. MRPP3 orthologs could be identified in many eukaryotic organisms and define a new protein family that was named PRORP (for PROteinaceous RNase P). Hence, MRPP3 is now also called human PRORP. These proteins are characterized by the presence of a number of PPR and/or PPR-like motifs, a CXXC Zn finger-like motif and a metallo/nuclease domain belonging to the NYN family.60 The function of putative PRORPs identified by sequence similarities thus leading to the potential cohabitation of both RNP and protein-only RNase P in this organelle.58 The occurrence of RNase P RNA in human mitochondria remains controversial and has been discussed in detail by Rossmanith in 2012.59

The catalytic subunit of protein-only RNase P is the PPR protein. While the three polypeptides that compose the human mitochondrial RNase P enzyme have some RNA-binding potential, only MRPP3 possesses the features of a metallo/nuclease that could account for the catalytic activity of RNase P. MRPP3 orthologs could be identified in many eukaryotic organisms and define a new protein family that was named PRORP (for PROteinaceous RNase P). Hence, MRPP3 is now also called human PRORP. These proteins are characterized by the presence of a number of PPR and/or PPR-like motifs, a CXXC Zn finger-like motif and a metallo/nuclease domain belonging to the NYN family.60 The function of putative PRORPs identified by sequence similarities

Figure 4. The current model of the PRORP/tRNA complex suggests a common mode of RNA binding in RNP and PRORP RNases P. (A) Structure of Thermotoga maritima ribozyme (PDBid 3Q1R18) with the catalytic domain in green, the specificity domain in blue, the RNase P protein subunit in orange, the tRNA product in light blue and the molecular surface of the RNP in gray. (B) The two-domain architecture of At-PRORP1 structure offers a concave surface that can be docked on the tRNA acceptor arm. The protein shown in the same orientation and same color code as the RNP with the catalytic domain in green, with metal ions bound (yellow spheres) close to the RNA cleavage site and the RNA-binding PPR domain in blue interacting with the region of the D-ψC loops. The central region (yellow) stabilized by a zinc ion (orange sphere) connects the two main PRORP domains. (C) A close-up of the PRORP1-tRNA complex model shows conserved catalytic aspartates D474 and D475 (blue) adjacent to tRNA cleavage site at position G+1 (red) as well as U16, G18, G19, and C56 (the nucleotides protected in footprint experiments77 in red) in contact with PPR motifs. Current functional data indicate that PRORP proteins have evolved an RNA recognition process very similar to that of RNP RNase P.
has been explored in depth in Arabidopsis, and in the protist *T. brucei*. Data are also available for *Ostreococcus tauri*, a primitive unicellular green alga. Arabidopsis expresses three PRORP proteins: At-PRORP1 is a 62 kDa protein with a pl of 9 and is localized to both plastids and mitochondria, whereas At-PRORP2 and At-PRORP3 are 59 kDa proteins, with pl of 6 and are localized in the nucleus. RNase P catalytic activity was first assigned unequivocally to the single protein At-PRORP1 in Arabidopsis organelles and later to each of the nuclear proteins on its own. In vitro RNase P activity tests using homologous pre-tRNA substrates were performed with purified recombinant forms of the three Arabidopsis RNase P protein candidates, each carefully verified for the absence of contamination by *E. coli* RNase P. Precise mapping of the cleavage site was achieved by high-resolution urea-PAGE or circular RT-PCR, and characterization of the 5' nucleotide of the mature tRNA products showed that each PRORP is a tRNA-specific endonuclease removing 5' extensions from pre-tRNAs and leaving a phosphate group at the 5' end of mature tRNAs. Abolition of the RNase P activity of recombinant PRORPs mutated in two conserved aspartates (predicted to be part of the catalytic site) confirmed that each of the three Arabidopsis PRORPs possessed RNase P activity as a single polypeptide.

Two PRORP genes were identified in the fully sequenced trypanosomatid genomes. In Trypanosoma brucei, PRORP1 is localized to the nucleus and PRORP2 to the mitochondrion. Using in vitro cleavage assays with purified recombinant proteins, each *T. brucei* PRORP protein appeared to perform the canonical 5' tRNA maturation on its own, similar to Arabidopsis PRORPs. Although studied to a lesser extent, a recombinant PRORP from the green algae *O. tauri* is capable of pre-tRNA 5' processing in vitro. The RNase P activity of these eukaryotic PRORP proteins from distant organisms is most likely shared by other members of this family. The association of a nuclease domain with a PPR domain to create RNase P enzymes represents yet another example of the potential and diversity of functions (i.e., RNA editing, splicing, or translation) acquired by the family of PPR proteins.

Beyond the capacity to perform RNase P activity in vitro, an important testimonial to the generality of PRORP tRNA processing capability came with the observation that Arabidopsis organellar PRORP1 and Trypanosoma nuclear PRORP1 could replace, in vivo, the *E. coli* and yeast nuclear RNase P respectively. Wild-type At-PRORP1, but not a protein mutated in the two conserved catalytic aspartates, rescues the lethal knockdown of RNase P RNA in *E. coli*. Similarly, *T. brucei* nuclear PRORP1 can rescue a deletion of the RNA component of yeast nuclear RNase P. These heterologous complementations led to the remarkable result that a single polypeptide can substitute in vivo for a complex ribonucleoprotein structure. Still, PRORP might not be the exact functional equivalent of RNP RNase P as fitness differences were observed between yeast strains non complemented and complemented by PRORP. Kinetic studies reveal that specificity constants of PRORP are not equivalent, i.e., they are lower than that of RNP RNase P.

Further experiments explored the in vivo roles of the three Arabidopsis PRORPs. The lethality of a single-gene knockout of At-PRORP1 and of the double knockout of At-PRORP2 and At-PRORP3 indicate that both the organellar and the nuclear PRORP enzymes fulfill essential functions in vivo, as expected for the authentic RNase P in cellular compartments encoding tRNA. The role of At-PRORP in tRNA 5' maturation in both organelles and the nucleus in planta was further explored by downregulation using virus-induced gene silencing. A decrease in PRORP1 specifically affects internal structures of chloroplast and mitochondria and reduces the level of mature organellar tRNAs, while nuclear-encoded tRNA levels are unchanged. Conversely, downregulation of PRORP2 in a prorp3 knockout background has no effect on organellar tRNAs, while the level of nuclear-encoded processed tRNA is reduced compared with control plants. Since downregulation of each PRORP protein causes a reduction of RNase P activity in the cellular compartment where that protein is found, it can be concluded that each PRORP protein is required for processing the tRNA pool in its respective compartment. On the other hand, downregulation of PO1 and POP4, two essential protein components of RNase MR (a ribonucleoprotein related to the nuclear RNP RNase P and involved in cytosolic tRNA maturation) affected tRNA maturation but did not reduce nuclear tRNA levels. Altogether, these results are consistent with PRORP proteins being the sole source of RNase P activity in both organelles and the nucleus of plants.

A report by Krehan, et al. has shown that RNase P activity as well as RNase MR RNA are present in a wheat embryo extract immune-precipitated with PO1 antibodies. This result has been interpreted as a clue for the presence of an RNP RNase P enzyme in plant nuclei. Since the downregulation of PO1 in planta resulted in decreased RNase MRP activity and did not affect RNase P activity, we believe that the results instead reflect the presence of both PRORP and RNase MRP in the immune-precipitated fraction, i.e., that the two enzymes might be present in a single complex in planta as also proposed by Krehan, et al. In Trypanosoma, PRORP activity was analyzed after immunodepletion, with anti-PRORP antibodies, of RNase P activity in a whole-cell extract. Depletion of both nuclear PRORP1 and mitochondrial PRORP2 abolishes all activity, suggesting that *T. brucei* contains no other RNase P. More studies are required, however, to understand the function of *T. brucei* PRORP in vivo. Since a complete set of tRNAs is imported from the cytosol into mitochondria in Trypanosoma, it will be particularly interesting to identify the substrates of the mitochondrial PRORP2 in vivo.

Collectively, experimental data obtained in distantly-related eukaryotes has clearly established that RNase P activity can reside in a single polypeptide. Moreover, in plants and Trypanosomes, PRORP proteins provide RNase P activity in vivo in both organelles and in the nucleus.

The substrate spectrum of PRORP, like that of RNP RNase P, goes beyond tRNAs. RNase P was first defined as the activity performing the 5' maturation of tRNA precursors. Still, extensive analyses of ribonucleoprotein RNase P functions have revealed that RNase P can be involved in the maturation of a much
wider variety of substrates in both prokaryotes and eukaryotes. After finding that PRORP proteins could perform the 5’ maturation of tRNA precursors in Arabidopsis, Trypanosoma, and Ostreococcus, it was logical to investigate whether PRORP proteins are entirely tRNA-specific or whether they, like RNase P, are involved in the maturation of other substrates.

The assumption that PRORP enzymes might be involved in the maturation of other RNAs is supported by the fact that numerous tRNA-derived sequences or structures are present in plant genomes. For instance, tRNA-like sequences called “t-elements” are present in transcripts of plant mitochondrial DNA, where they separate individual mRNAs. Similarly, in the nucleus, SINE RNAs are derived from tRNAs, although their canonical cloverleaf structure has apparently been lost. Another argument comes from the observation that Arabidopsis PRORP1 can replace E. coli RNP RNase P in vivo. Bacterial RNP RNase P is responsible for the maturation of many non-tRNA substrates, including the precursor to the 4.5S RNA. Two substrates that contain tRNA-like recognition elements are the precursor to C4 antisense RNA of bacteriophage P1 and P7, which possesses a tRNA-like structure with short D- and T-loops, and the precursor to tRNA, part of whose structure resembles the horizontal stacking domain (acceptor stem plus T-stem and loop) of tRNA, a known minimal substrate for E. coli RNase P. E. coli RNase P is also involved in processing polycistronic mRNAs such as the histidine operon transcript and in cleavage of some riboswitches, including those for the coenzyme B12. It can thus be speculated that Arabidopsis PRORP1 could catalyze the maturation of all these E. coli non-tRNA substrates. Alternatively, it is possible that some of these non-tRNA maturation steps are not essential or that they can be rescued by other enzymatic systems in the absence of ribonucleoprotein RNase P.

Preliminary results, both in vitro and in vivo, have confirmed that Arabidopsis PRORP1 are indeed involved in the maturation of other RNA substrates. In particular, PRORP1 is able to perform in vitro the endonucleolytic cleavage of tRNA-like t-elements present in the mitochondrial transcripts of Arabidopsis. PRORP1 activity is required in vivo to accumulate nad6 mRNA. Similarly, Arabidopsis PRORP2 and 3 are indirectly involved in the maturation of snoRNA. In Arabidopsis, a dicistronic precursor to tRNA and the snoRNA snoR43.1 is processed by both RNase P and the pre-tRNA 3’-processing endonuclease RNase Z, with RNase P cleavage of the pre-tRNA portion being a prerequisite for the cleavage by RNAse Z that separates mature tRNA from mature snoR43.1. In PRORP downregulation mutants, snoR43 failed to accumulate to normal levels whereas tRNA-snoRNA precursor levels increased, showing that nuclear PRORP activity is required for the accumulation of this snoRNA.

An initial investigation of the PRORP/tRNA complex has revealed that minimal tRNA structural features are required for recognition by PRORP alone. For example, and like the bacterial RNP RNase P, the tRNA acceptor stem is essential whereas the anticodon domain is not. Unlike the bacterial enzyme, PRORP cleavage is impaired by the absence of the D domain from tRNAs or t-elements. Thus, maturation of mitochondrial mRNAs by cleavage of some t-elements (such as the one from ccmC mRNA that lacks both D and anticodon domains) might require additional proteins acting as PRORP partners to recognize these structures. Similarly, in humans, the requirement for MRPP1 and 2 might reflect an inability of Hs-PRORP alone to bind the non-canonical tRNA structures characteristic of vertebrate mitochondria. This would also explain why the plant PRORP1 can function in both chloroplasts and mitochondria, since tRNAs from plant chloroplasts and mitochondria closely resemble bacterial tRNAs.

The diversity of substrates identified so far for PRORP remains limited. Other potential RNA substrates will have to be investigated at the transcriptome-wide level, for example, through comparative transcriptome analyses of PRORP downregulation mutants or by global sequencing of RNA partners immune-precipitated in complex with PRORP proteins.

Emergence and distribution of PRORP enzymes in eukaryotes. RNase P is a ubiquitous enzyme, found in all organisms with the exceptions of symbiotic Archaea, such as Nanoarchaea equitans, several species of Pyrobaculum and Aquifex aeolicus in which transcription of tRNAs starts at position +1. The RNase P form of RNase P is widespread as it is present in Bacteria, in Archaea and in Eukarya with characterized activities in both the nucleus and mitochondria (as, for example, in yeast). On the contrary, known PRORP RNase P are limited to eukaryotes (Fig. 2), having been identified in human mitochondria, Arabidopsis thaliana mitochondria, chloroplasts, and nuclei and in Trypanosoma brucei mitochondria and nuclei. In the green alga Ostreococcus tauri, a PRORP protein was found to have RNase P activity but its localization was not determined. However, bacterial-type RNase P ribozymes can be found encoded in both mitochondrial and plastidial genomes along with an RNP RNase P protein in the nucleus. Nonetheless, although all characterized PRORP enzymes are eukaryotic, they are not restricted to endosymbiotic organelles as was previously assumed.

Database analyses confirm that PRORP proteins constitute a eukaryote-specific family of enzymes. Putative PRORP sequences can be found in nearly all major eukaryotic groups (i.e., in Metazoa, Streptophyta, Chlorophyta, Kinetoplastida, Stramenopiles, and Oomycetes) with the notable exceptions of fungi and amoeboida. The appearance of PRORP can essentially be defined by the event that led to the fusion of a PPR domain with an NNY nucleic domain. The precise timing of this event and the evolutionary history of PRORP remain to be established. Still, its occurrence as experimentally shown for Metazoa, Euglenozoa and for both Streptophyta and Chlorophyta in Viridiplantae (Fig. 2), already suggests that PRORP appeared very early in the evolution of eukaryotes.

The emergence of PRORP has been proposed to be related to the acquisition of organelles. Similarly, Howard et al. suggested that the evolutionary drive for RNase P replacement by PRORP might have resided in different substrate specificities between nuclear and organelle RNase P enzymes, in the difficulty of importing a large RNA such as that for RNase P into mitochondria, or in the “vulnerability” of organelle RNA toward
RNP RNase P enzymes. All these propositions assume that PRORP initially arose as an organelle-targeted enzyme, which is not established and not necessarily true. Indeed, PRORP clearly emerged as a nuclear gene by fusion of genes encoding a PPR RNA-binding protein and an NYN metallonuclease domain (discussed in the following section). Because the nuclear RNP RNase P activity of PRORP is found in distantly related eukaryotes, PRORP nuclear activity is most likely ancient. It is thus possible that PRORP might have first functioned as a nuclear enzyme. If so, the evolutionary impetus to replace a RNP complex containing one RNA and up to ten proteins by a protein-only enzyme might have resided in the fact that the simpler enzyme assembles faster, is easier to regulate and requires fewer cellular resources for its biogenesis.

PRORP enzymes are two-domain proteins. Initial structural predictions of PRORP based on sequence analyses indicated the presence of PPR modules in the N terminus and of a NYN-like catalytic domain in the C terminus. This organization into two α-helix-rich domains was supported by biophysical characterization (circular dichroism and small angle X-ray scattering) of At-PRORP in solution and is consistent with the X-ray crystal structure of At-PRORP. Taking together 1-, 2-, and 3-D data available for this enzyme family, comparative models of representative PRORP members are presented in Figure 3A. These models pinpoint the general conservation of the PRORP fold from unicellular algae to humans. Small variations are observed, mainly in peripheral loops. Long insertions are present in plasmodial enzymes, as is often observed in proteins from this parasite family.

The N-terminal PPR domain forms a superhelical structure very similar to those described in TPR (TetratricoPeptide Repeat) domains, an evolutionary-related domain involved in protein–protein interactions. As illustrated in Figure 3B, it contains five PPR and PPR-like motifs: two canonical ones and three displaying remote sequence similarities. Despite their divergent sequences, these PPR modules are structurally similar and superimposable on those found in the only other PPR protein of known three-dimensional structure, i.e., human mitochondrial RNA polymerase. This confirms, as was originally proposed, that the defining feature of PPR family members is a conserved structural fold of PPR motifs rather than of conserved sequence elements.

The catalytic domain of PRORP adopts an α/β/α sandwich fold (Fig. 3C) belonging to the PIN-like nuclease family. A similar architecture is found in the nuclease domain of T4 RNase H and of human SMG6 and SMG5, two essential factors in nonsense-mediated mRNA decay, as well as of a recently characterized MCPIP1 RNase (MCP-1 induced protein 1) that participates in the regulation of immune response by degrading the mRNA of inflammatory cytokines. Among the four aspartate residues involved in the binding of metal ions, two are strictly conserved in PRORPs (D474 and D475 in At-PRORP1) and in other nucleases of the PIN/NYN family and are essential for pre-tRNA cleavage.

These two functional domains are connected by a split zinc-binding module derived from the central and the C-terminal regions of PRORP (Fig. 3D) and which forms the tip of the overall “V shape” of PRORP, the PPR and catalytic domains being the two arms of the V. The concave surface of the PPR superhelix in one arm thus faces the catalytic groove in the other arm, thereby exposing conserved aspartate residues and metal ions, making the overall architecture look like tweezers.

Is PRORP a structural mimic of ribonucleoprotein RNase P? The bacterial RNP RNase P docks onto the acceptor stem of its pre-tRNA substrate, with an interaction extending from the tRNA corner (T and D loops), which is recognized by the specificity domain (S-domain) to the cleavage point between nucleotides −1 and +1, which is apposed to the catalytic domain (C-domain). In E. coli tRNAs, the 3’ terminal CCA interacts specifically with a complementary sequence in a loop of the RNase P RNA, whereas the pre-tRNA leader interacts with the protein subunit of the holoenzyme.

The bipartite organization of PRORPs (Fig. 3) is reminiscent of that of RNP RNase P, with the PPR domain playing the role of the S-domain to ensure recognition of the pre-tRNA and its orientation in the catalytic domain. In support of this role, removal of the four N-terminal PPR motifs of At-PRORP1 leads to a 34-fold drop of affinity for the substrate and a > 2000-fold loss of enzymatic activity. Similarly, the deletion of the S-domain in the RNP RNase P resulted in 30- to 13 000-fold loss in catalytic performance, depending upon the substrate used. However, the S-domain deletion, surprisingly, led to more accurate cleavage site selection.

On the substrate side, deletions altering the pre-tRNA structure show that for PRORP, just as for the RNP RNase P, the anticodon stem-loop is dispensable, whereas the D and T loops are required. Footprint experiments confirmed that the corner of the tRNA L-fold interacts with At-PRORP1 to give strong protection of residues U16, G18-19, and C56. The PRORP/pre-tRNA complex was modeled based on the At-PRORP1 crystal structure using as geometrical restraints the binding of the T/D loops by the PPR domain and the positioning of the cleavage point in the vicinity of conserved aspartate groups constituting the metal-binding site. Figure 4 illustrates the potential similarity between PRORP and RNP RNase P in the way they bind their pre-tRNA substrates. Another model of PRORP/pre-tRNA complex has been proposed, it shows PRORP interacting on the side rather as on the top of tRNAs. However, the latter does not take in account footprinting and tRNA deletion results that suggested contacts between PRORP and tRNA residues U16, G18, G19 as well as C56, while the anticodon stem is dispensable for recognition.

Despite the overall similarity of their substrate-binding modes, however, the two types of RNase P—employing a protein or an RNA catalytic component—are mechanistically distinct. Both cleave a phosphodiester bond by nucleophilic attack of hydroxide ion apical to O3′ of the upstream ribose, generating products with 3′-hydroxyl and 5′-phosphoryl termini. The presence of metal-binding sites in the structure of At-PRORP1 suggests that the proteinaceous enzymes use a two-metal-ion mechanism to deprotonate water and to stabilize the transition state. However, the tolerance of PRORPs
to an Rp-phosphorothioate modification of the scissile bond in the presence of Mg\(^{2+}\) as cofactor is a striking difference from the RNase P enzyme,\(^{5,33}\) indicating that the metal in PRORP does not directly coordinate the pro-Rp-oxide of the target phosphodiester. Rather, it appears that, whereas the RNase P RNA subunit employs one hydrated divalent cation to provide the attacking hydroxide and a second metal hydrate to protonate the leaving group,\(^{5,7}\) the proteinaceous RNase P utilizes a more conventional mechanism akin to that of known protein metal-lonucleases, in which the metal ions serve primarily to stabilize the charge and structure of the trigonal bipyramidal transition state, and general acid-base chemistry is accomplished by the carboxylate groups of aspartate (and possibly the imidazole nitrogen of histidine). The binding affinities of PRORPs for their pre-tRNA substrate are in the micromolar range.\(^{5,7}\) These values are one or two orders of magnitude lower than for RN P RNases P and may indicate more transient interaction with substrates. Nevertheless, these proteinaceous enzymes are efficient enough to complement \(E.\) coli RN P RNase P.\(^{5,41}\) So the precise functional advantages of the PRORP and RNase P mechanism remain to be identified.

**Concluding Remarks**

Within the PPR family, the characterization of PRORP proteins has finally settled the long-lasting debate over the existence of an alternative system devoid of RNA for RNase P activity in eukaryotes. The discovery of protein-only RNase P indicate that the distribution and evolutionary history of RNase P are more complex than previously thought. The functional and mechanistic comparison of PRORP with RN P RNase P will have important implications for our understanding of the evolution of living systems. Indeed, it will illustrate how convergent evolution has found two independent routes to catalyze the 5′ maturation of tRNAs: either with an RNA-based enzyme or a protein-only enzyme. This mechanistic comparison leads to important questions. For instance, the mechanism by which PPR motifs confer PRORP substrate specificity remains to be elucidated. Future work, in particular determination of the crystal structure of PRORP in complex with tRNA, will establish whether PPR motifs indeed bind conserved residues in the single-stranded D and T loops of tRNAs as was previously suggested,\(^{7,7}\) and thus whether the PRORP mode of RNA recognition is in conformity with the overall mode of RNA recognition recently proposed for PPR proteins.\(^{7,9}\)

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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