Protein-only RNase P function in *Escherichia coli*: viability, processing defects and differences between PRORP isoenzymes

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ABSTRACT

The RNase P family comprises structurally diverse endoribonucleases ranging from complex ribonucleoproteins to single polypeptides. We show that the organellar (*At*PRORP1) and the two nuclear (*At*PRORP2,3) single-polypeptide RNase P isoenzymes from *Arabidopsis thaliana* confer viability to *Escherichia coli* cells with a lethal knockdown of its endogenous RNA-based RNase P. RNA-Seq revealed that *At*PRORP1, compared with bacterial RNase P or *At*PRORP3, cleaves several precursor tRNAs (pre-tRNAs) aberrantly in *E. coli*. Aberrant cleavage by *At*PRORP1 was mainly observed for pre-tRNAs that can form short acceptor-stem extensions involving G:C base pairs, including tRNA^Asp^ (GUC), tRNA^Ser^ (CGA) and tRNA^His^. However, both *At*PRORP1 and 3 were defective in processing of *E. coli* pre-tRNA^Sec^ carrying an acceptor stem expanded by three G:C base pairs. Instead, pre-tRNA^Sec^ was degraded, suggesting that tRNA^Sec^ is dispensable for *E. coli* under laboratory conditions. *At*PRORP1 and 3 are also essentially unable to process the primary transcript of 4.5S RNA, a hairpin-like non-tRNA substrate processed by *E. coli* RNase P, indicating that PRORP enzymes have a narrower, more tRNA-centric substrate spectrum than bacterial RNA-based RNase P enzymes. The cells’ viability also suggests that the essential function of the signal recognition particle can be maintained with a 5′-extended 4.5S RNA.

INTRODUCTION

RNase P is the endoribonuclease responsible for the 5′-end maturation of tRNA primary transcripts (1–4). While bacterial RNase P is composed of a catalytic RNA subunit and a single protein cofactor (5), the nuclear enzyme of several major eukaryal groups (metazoans, fungi) has evolved into a complex ribonucleoprotein consisting of an RNA subunit plus up to 10 protein subunits (6,7). Until recently, the presence of a catalytic RNA subunit was thought to be the hallmark of all RNase P enzymes. The first exception from this assumption was reported in 2008 when the human mitochondrial activity was found to consist of three protein subunits and to lack an RNA component (8). Homologs of the endonuclease protein subunit of human mitochondrial RNase P, originally termed MRPP3, were identified also in distant eukaryal groups, such as land plants and kinetoplastida (9). This led to the characterization of three homologs to human MRPP3 in the land plant *Arabidopsis thaliana*, all of which were demonstrated to be genuine RNase P enzymes that are active as single polypeptides (9,10). One of the *A. thaliana* RNase P isoenzymes (termed PRORP1 for proteinaceous RNase P 1), which localizes to the mitochondria and chloroplasts, was demonstrated to sustain growth of an *Escherichia coli* mutant strain with an otherwise lethal depletion of its endogenous ribonucleoprotein RNase P (9). The two other isoenzymes *At*PRORP2 and 3 are found in the nucleus (9). Two PRORP isoenzymes also have RNase P function in the protist *Trypanosoma brucei* (*Tb*PRORP1 and 2) (11). Recently, the different PRORPs from *A. thaliana* and *T. brucei* were shown to be able to replace the nuclear RNase P ribonucleoprotein complex in yeast (11,12). An in-depth characterization of *At*PRORP3-dependent yeast strains moreover showed that the RNase P replacement did not compromise the fitness of these cells (12). Overall, these findings lead to the conclusion that protein-based RNase P enzymes are capable of carrying out the basic functions of an RNA-containing RNase P enzyme in Bacteria and Eukarya, but they raise the question why so many organisms have retained a ribonucleoprotein (RNP) RNase P enzyme.
The capability to substitute for E. coli RNase P in vivo was extended here to the nuclear isoenzymes (AtPRORP2 and 3) from A. thaliana. This was motivated by the idea that their complementation capacity in the E. coli host may differ from that of the organellar enzyme (AtPRORP1), as (i) the latter has to act on a different set of substrates in a different cellular environment relative to the nuclear isoenzymes, (ii) recent evidence suggested differences in cleavage fidelity compared to AtPRORP2 and 3 (13) and (iii) AtPRORP1-dependent yeast strains grew slower than AtPRORP2- or 3-dependent ones (12). Principally, all three AtPRORP variants supported growth of a conditional E. coli RNase P mutant strain under non-permissive conditions, complementation capacity in the (Supplementary Table S1) were introduced into vivo in rnpB Expression of the rnpB Bacterial strains and complementation studies MATERIALS AND METHODS

MATERIALS AND METHODS

Bacterial strains and complementation studies

Expression of the rnpB gene (encoding the RNase P RNA subunit) in E. coli strain BW strictly depends on arabinose supplementation (14). In the absence of arabinose and the presence of glucose the chromosomal rnpB expression in strain BW is repressed and survival of the cells depends on plasmid-encoded RNase P activities. For in vivo studies, the different pDG148-based expression vectors (Supplementary Table S1) were introduced into E. coli BW cells by electroporation. The genotypes of the strains used here are specified in Supplementary Table S2 and more methodological details are provided in the Supplementary Material.

RNA-Seq—Illumina sequencing and data evaluation

For the RNA-Seq analysis presented in Figure 3, single colonies of BW[pEcRNPB], BW[pATPRORP1] or BW[pATPRORP3] bacteria (three biological replicates of each strain) picked from agarose plates were used to inoculate 3 ml LB medium supplemented with 10 mM arabinose. After 8 h of growth at 37°C/200 rpm, cells from such pre-cultures were transferred into 50 ml fresh LB medium supplemented with 10 mM glucose to a starting OD_{600} of 0.01. Cultures were grown for 14 h at 37°C/200 rpm into stationary phase. Cells were then transferred into 200 ml of fresh glucose-containing LB medium (starting OD_{600} = 0.1) to initiate exponential growth. At OD_{600} ~ 1.0 (= outgrowth phase), cells were harvested and total RNA was prepared according to Method 1 (‘Extracting RNA three times with hot phenol’) described by Damm et al. (15). Library generation was performed at vertis Biotechnologie AG (Freising-Weihenstephan, Germany). Total RNAs were depleted of rRNA using the Ribo-Zero rRNA Removal Kit for bacteria (Epicenter). The fraction of small RNAs (<200 nt) was separated using the RNeasy MinElute Cleanup Kit (Qiagen). Oligonucleotide adapters were ligated to the 5’- and 3’-ends. First-strand cDNA synthesis was performed using M-MLV reverse transcriptase and the 3’-adapter as primer. The resulting cDNAs were PCR-amplified using a high-fidelity DNA polymerase. The cDNA was purified using the Agencourt AMPure XP kit (Beckman Coulter Genomics). The 3’- and 5’-adapters and the primers used for PCR amplification were designed for TruSeq sequencing according to the instructions of Illumina. The cDNA pool was sequenced on an Illumina NextSeq 500 system with a read length of 75 nt.

Details regarding the preparation of the initial RNA-Seq analysis that led to the detection of the 4.5S RNA processing defect in BW[ATPRORP1] bacteria can be found in the Supplementary Material.

Reads were processed in two steps. First, nucleotides with a read quality lower than Q30 were trimmed from the end of the sequence using the FASTX Toolkit. Then homo-polymeric sequences (≥12 identical nucleotides) were trimmed using custom scripts. Mapping to E. coli str. K-12 substr. MG1655 (GenBank CP012868.1) was done using segemehl (16) with an e-value of 0.01 and a minimal read length of 10 nt. The overall statistics of the three biological replicate libraries for each strain can be found in the Supplementary Table S3.

For the data presentation in Figure 3, only tRNA reads with their 5’-end mapping to position –2, –1 or +1 (according to the RefSeq annotation GenBank CP012868.1) and with a minimal number of 10 reads (sum of reads corresponding to 5’-ends at –2, –1 and +1) of the respective tRNA in at least two of the biological replicates were included in the evaluation. Reads starting further upstream than position –2 are shown in the alignments of the Online Supplement (Results, ‘Categorized mapping alignments for all replicates’) but were not considered further. Reads starting at the 5’-end position +1 were classified as ‘canonically cleaved’ and reads starting at –2 or –1 were classified as ‘miscleaved’. Note that bacterial tRNA^His and tRNA^Sec are naturally ‘miscleaved’ between nt –2/–1. For each tRNA meeting the aforementioned criteria, the canonically cleaved percentage of cDNA reads (5’-end at position +1) was plotted according to the equation \( \frac{N_{+1}}{N_{-2}+N_{-1}+N_{+1}} \times 100\% \), where \( N_{-2}, N_{-1} \) and \( N_{+1} \) correspond to the number of tRNA-specific cDNA reads starting at the 5’-end position –2,
–1 and +1, respectively. Two falsely annotated tRNA^Met molecules were assigned as tRNA^His^2 (17). We furthermore merged sequences with identical tRNA bodies and identical nucleotides at position –1 and –2. Details can be found in the Online Supplement (http://bioinf.pharmazie.uni-marburg.de/supplements/prorp_2017/) under ‘Sequences’. Whenever reads mapped to multiple tRNAs we applied the abundance factors reported by Dong et al. (18). Here, we adjusted the molar ratios to our growth rates of ~2. Hence, we averaged the values given by Dong et al. (18) for growth rates of 1.6 and 2.5. In cases where tRNAs were not differentiated there, we distributed the reads proportionally. For example, the number of mature tRNA reads mapping to both, tRNA^Gln(UUG) 1/2 and tRNA^Gln(UUG) 2/2, were assigned to each tRNA after multiplying with 0.5. The scores can be viewed in the respective alignments in the Online Supplement (Results, ‘Categorized mapping alignments for all replicates’), which also includes the raw and scored read numbers. While this consideration of multimap reads is technically necessary for obtaining an unbiased picture, the influence of these adjustments to the processing fractions (next paragraph) was marginal and did not change the overall picture.

For details on cell growth, plasmid preparations for in vivo studies, the initial RNA-Seq analysis, primer extension, Western blotting, northern blot analysis (for the used probes, see Supplementary Table S4) and in vitro RNase P cleavage assays (conducted essentially as described (19)), see the Supplementary Material.

RESULTS

Complementation of E. coli RNase P RNA deficiency by AtPRORP1, 2 and 3

We compared the organellar AtPRORP1 isoenzyme and the nuclear isoenzymes AtPRORP2 and 3 with respect to their ability to functionally replace E. coli RNase P in vivo (for similarity of the three PRORP isoenzymes, see Supplementary Figure S1A and B). In the test strain BW, a derivative of E. coli MG1655 (Supplementary Table S2), the endogenous RNase P RNA gene (rnpB) is expressed from a chromosome-borne P_bld promoter in the presence of the inducer arabinose, whereas expression is silenced when cells grow in the presence of glucose as the carbon source. As a result, RNase P RNA is depleted (Supplementary Figure S2A) and cells stop dividing (14). Bacteria expressing AtPRORP1 and 3 were able to rescue growth at 37 °C under non-permissive conditions on LB-glucose plates (Supplementary Figure S2B). In contrast, bacteria expressing AtPRORP2 or harboring an empty plasmid failed to rescue growth under these standard growth conditions. We previously observed that recombinant AtPRORP2 is inactive in precursor tRNA (pre-tRNA) processing assays at 37 °C, but displays vivid activity at 28 °C (19). In line with this finding, AtPRORP2 was able to sustain growth of BW bacteria in the absence of arabinose at 28 °C (Supplementary Figure S2B). These findings demonstrate that all three PRORP isoenzymes from A. thaliana can execute tRNA 5′-end maturation to an extent that permits E. coli cell proliferation, consistent with their similar overall structure (Supplementary Figure S1C).

Growth properties of AtPRORP-depending E. coli cells

To examine how efficiently the AtPRORP isoenzymes fulfill their RNase P function in the E. coli host, growth of the AtPRORP-dependent BW bacteria in liquid medium was monitored (Figure 1). At 28 °C (Figure 1A), all three AtPRORP isoenzymes supported growth of E. coli BW bacteria under non-permissive conditions, reaching comparable cell densities in stationary phase. The growth behavior of the strain harboring the expression vector for AtPRORP3 (BW[pATPRORP3]) was almost identical to that of BW cells harboring the plasmid-encoded E. coli P RNA gene (BW[pEcnpB]). However, cells depending on the RNase P activity of AtPRORP2, or AtPRORP1 in particular, had an extended lag phase before entering exponential growth (Figure 1A; for generation times, see Supplementary Table S5). When growth in liquid culture was monitored at 37 °C, AtPRORP2 was unable to sustain growth of BW cells (Figure 1B), in line with the failure to
grow on agar plates (Supplementary Figure S2B). Again, bacteria expressing AtPRORP3 reproducibly showed a faster transition from log to exponential phase than those depending on AtPRORP1, and strains BW[AtPRORP1] and BW[AtPRORP3] grew to a lower final density at 37°C than BW[pEcnrpB] bacteria (Figure 1B). As shown by Western blot analysis, the expression levels of all three C-terminally His-tagged AtPRORP isoenzymes were comparable in E. coli BW cells grown at 28°C. Likewise, expression levels of AtPRORP1-His and AtPRORP3-His were similar in BW cells grown at 37°C (Supplementary Figure S3). However, compared to BW cells grown at 28°C, the fraction of insoluble AtPRORP1-His and AtPRORP3-His fraction were increased in BW cells grown at 37°C (Supplementary Figure S3A, compare lanes 4 versus 12, and 8 versus 14). Thus, the quantity of functional, soluble PRORP enzymes seems to be decreased at higher growth temperatures and E. coli cells have to cope with larger amounts of insoluble protein. This may have contributed to the lower final density of BW[AtPRORP1] and BW[AtPRORP3] cultures at 37°C relative to BW[pEcnrpB] bacteria (Figure 1B).

RNA-Seq analysis of AtPRORP1-depending E. coli cells

To detect potential processing defects in E. coli cells depending on PRORP, we pursued a deep sequencing (RNA-Seq) approach using total RNA from E. coli BW[AtPRORP1] cells grown for 70 min under non-permissive conditions in arabinose-free medium. The relatively short period of 70 min was chosen to focus on primary effects in the transcriptome upon depletion of endogenous RNase P. The BW[AtPRORP1] strain was chosen at the beginning of this study because it had a longer lag phase than the BW[AtPRORP3] strain (Figure 1), thus increasing the likelihood to detect potential processing defects. As reference, we used E. coli MG1655, the parental wild-type strain of the BW mutant strains. E. coli MG1655 was considered to be equivalent to strain BW[EcnrpB] (Figure 1), since both strains express E. coli wild type RNase P. For strains MG1655 and BW[EcnrpB] we obtained very similar growth curves in LB medium supplemented with 10 mM glucose at 37°C (14), so the potentially higher gene dosage of BW[EcnrpB] expressing E. coli rnpB from a plasmid seems not to impact on growth behavior.

A major outcome of the RNA-Seq analysis was the finding of an increased proportion of reads representing 5'-precursor 4.5S RNA (pre-4.5S RNA) in total RNA isolated from BW[AtPRORP1] compared with MG1655 bacteria (Figure 2A and Supplementary Figure S4). In E. coli, the non-coding RNA 4.5S RNA is a verified substrate of bacterial RNase P (20,21) and plays an important role as part of the signal recognition particle that targets ribosomes to the plasma membrane (22,23).

Accumulation of pre-4.5S RNA in AtPRORP1-depending E. coli cells

To further validate this potential 4.5S RNA processing defect in BW[AtPRORP1] bacteria by an independent approach, we analyzed total RNA from BW[AtPRORP1] cells grown in arabinose-free glucose medium by primer extension using a 4.5S RNA-specific primer. Reverse transcriptase extended the 5’-32P-end-labeled primer (18 nt) by 2 nt on mature 4.5S RNA. The 2-nt extension product corresponding to 5'-mature 4.5S RNA was the only signal obtained with total RNA from the parental wild type strain grown for 6 h in the presence of glucose or with RNA from BW[vector] cells (strain BW harboring the empty expression vector) grown for 6 h in the presence of arabinose (Figure 2B, lanes 1 and 3). However, primer extension using total RNA from BW[AtPRORP1] or BW[vector] grown in the presence of glucose for the same time period resulted in much less extension product corresponding to 5'-mature 4.5S RNA and substantial amounts of longer extension products (Figure 2B, lanes 2 and 4). The identical product pattern in lanes 2 and 4 indicated that AtPRORP1 is essentially unable to contribute to the 5'-end maturation of 4.5S RNA in the E. coli host.

The apparent lengths of primer extension products obtained on pre-4.5S RNA (Figure 2B, lanes 2 and 4) suggested the presence of longer precursor molecules than those reported previously (22 nt) (20). This difference to the previous study may be explained by the fact that the authors used an RNase P mutant strain (A49) under heat shock conditions, which may have caused changes in RNA processing. A search for putative promoters upstream of the 4.5S RNA gene (http://regulondb.ccg.unam.mx; Softberry, BPROM algorithm; http://www.softberry.com) gave predictions for transcription start points of σ70 RNA polymerase at positions −160, −130 and −26 relative to the mature 5'-end (nt +1) of 4.5S RNA. This corresponds to RNA-Seq reads with 5'-ends primarily at −26 and some with 5'-ends at about −153 and −127. However, at present the exact lengths of the 4.5S RNA precursor signals in Figure 2B remain unclear.

We further asked the question if the primer extension product for mature 4.5S RNA in BW[AtPRORP1] further decreases after extended growth in glucose medium, and whether BW bacteria complemented with AtPRORP1 or 3 differ in their 4.5S RNA processing defect. However, after ~14 h (stationary phase) as well as following an additional outgrowth from stationary phase (in glucose medium), essentially no mature 4.5S RNA was detectable in BW[AtPRORP1] as well as BW[AtPRORP3] bacteria (Figure 2B, lanes 8–11).

Extended RNA-Seq analysis

To obtain statistically relevant data on tRNA processing defects in E. coli cells depending on PRORP isoenzymes, we performed a second RNA-Seq analysis based on three biological replicates. This included the strains BW[AtPRORP1] and BW[AtPRORP3] and E. coli BW[EcnrpB] as reference strain for direct comparability with the results of our growth analysis (Figure 1). All cDNA reads of E. coli tRNAs were analyzed for their 5'-end position to evaluate cleavage site selection by the different RNase P enzymes. In the libraries of E. coli BW[EcnrpB], BW[AtPRORP1] and BW[AtPRORP3], only tRNA reads starting either at the 5'-end position...
bacterial pre-tRNAHis species analyzed so far are cleaved
pairs or one G:U pair plus two G:C pairs, and pre-tRNAs
see Materials and Methods). In Figure 3, the percentage
–2, –1 or +1 were included in the evaluation (for details,
from stationary phase; under 'Initial Mapping'. (Figure 2.
S6): pre-tRNA Ser(CGA) can form an additional C–1 :
4.5S RNA (m4.5S RNA). Total RNAs of
each form two extra pairs (U:A and G:C).
(26,27), and a G residue at –1 act as positive determinants for selection of the
combination of several structural determinants, as studied
in depth for E. coli RNase P: a U at –2 and a G
terminus at G+1 (corresponding to miscleavage at nt –1/+1) were negligible or only slightly increased in BW[EcrnpB]
and BW[AtPRORP3], respectively, roughly half of the
reads corresponded to miscleaved rather than correctly
processed (5′-end at G+1, cleavage at nt –2/–1) tRNAHis
in BW[AtPRORP1] cells (Figures 3 and 4A). We further
analyzed by northern blotting if the overall levels of
tRNAHis may be altered in BW[AtPRORP1]. However,
tRNAHis levels were equal in all tested strains grown in the presence of glucose (Figure 4B). Of note, the northern blot
-2, –1 or +1 were included in the evaluation (for details,
see Materials and Methods). In Figure 3, the percentage
of canonically processed cDNA reads (5′-end at nt +1) is plotted for all evaluated tRNAs (for read number
cutoff, see Materials and Methods). Whereas cleavage site
selection was overall similar in cDNA libraries derived
from BW[EcrnpB] and BW[AtPRORP3] bacteria, our data
suggested reduced processing at the canonical –1/+1 site for a set of tRNAs in BW[AtPRORP1] (Figure 3). Generally,
aberrant processing correlated well with the propensity of
pre-tRNAs to form acceptor stems extended by one or two
(N-1 : N+73, N-2 : C+74) base pairs (Supplementary Table
S6): pre-tRNA Ser(CGA) can form an additional C-1 : G+73
pair, the three Asp(GUC) variants two extra G:C
pairs or one G:U pair plus two G:C pairs, and pre-tRNAs
Pro(GGG), Thr(GGU) 2/2, Val(UAC) 1/3 and Leu(GAG)
can each form two extra pairs (U:A and G:C).
With the exception of some α-proteobacteria (24,25), all
bacterial pre-tRNAHis species analyzed so far are cleaved
by their cognate RNase P enzymes between nt –2 and
–1 to release a mature functional tRNAHis with an 8-
bp acceptor stem including the extra G–1 :C+N+73 base
pairs (Supplementary Table
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7445

Figure 3. Percentage of reads corresponding to canonical cleavage between nt –1 and +1 (5′-end at position +1) in BW[Ec rnpB], BW[At PRORP1] and BW[At PRORP3] cells. Only tRNA reads with 5′-ends at position –2, –1 or +1 were included in the analysis (for details, see Material and Methods). Note that tRNAHis is naturally miscleaved at nt –2/–1 (5′-end at nt –1) in bacteria, explaining why hardly any 5′-ends at position +1 were identified in BW[Ec rnpB] cells. RNA was prepared from cells outgrown from stationary to exponential growth phase (OD 600 ∼ 0.7) in arabinose-free LB medium and analyzed by RNA-Seq (three biological replicates of each strain).

Northern blot analyses of pre-tRNA Sec processing in AtPRORP-dependent E. coli cells

Reads for tRNA Sec, the second bacterial tRNA species containing an 8-bp acceptor stem generated by non-canonical 5′-end maturation (33), fell below the threshold for minimum read numbers and were thus not represented in Figure 3. Noteworthy, pre-tRNA Sec transcripts have the potential to extend their acceptor stems by two additional G:C base pairs, resulting in pre-tRNAs with a 10-bp acceptor stem (see Figure 5). We then analyzed tRNA Sec processing by northern blotting using total RNA from exponentially grown cells. We surprisingly observed a strong 5′-maturation defect in E. coli cells depending on At PRORP1 or At PRORP3 (Figure 4C, lanes 3 and 4). In BW[At PRORP1] bacteria mature tRNA Sec levels were largely decreased, while a precursor molecule accumulated (a 24-nt leader according to (33), 20–25 nt according to our RNA-Seq data). In BW[At PRORP3] cells grown in the presence of glucose, the pattern looked similar, but overall signal strength was largely diminished (Figure 4C, lane 4), suggesting that unprocessed tRNA Sec transcripts are degraded. In contrast, mature tRNA Sec was detected in E. coli BW[At PRORP1] and BW[At PRORP3] grown in medium supplemented with arabinose (Figure 4C, lanes 5 and 6). All total RNA preparations were quantitatively and qualitatively comparable as shown by the 5S RNA control (Figure 4C, bottom panel). Identical results were obtained with RNA preparations from stationary phase cells (data not shown).

In vitro cleavage of E. coli pre-tRNA His, pre-tRNA Sec and pre-4.5S RNA by AtPRORPs

To further substantiate the results of the in vivo analyses we investigated in vitro processing of 5′-32P-end-labeled pre-tRNA His, pre-tRNA Sec and the non-tRNA substrate 4.5S RNA by AtPRORP1, 2 and 3 (Figure 5). As reference substrate we used pre-tRNA Gly from Thermus thermophilus, a standard class I tRNA (structurally very similar to its E. coli counterpart, but with slightly G:C-richer stem regions). Cleavage assays were performed under conditions of enzyme excess to exclude any rate limitations caused by product release (120 nM enzyme, < 1 nM substrate) for 20 min at 28°C (At PRORP2) or 37°C (At PRORP1 and 3). In parallel, each substrate was incubated with the bacterial RNase P holoenzyme from E. coli under the same reaction conditions at 37°C. Whereas all three AtPRORP enzymes
Figure 4. (A) Reads corresponding to tRNA^{His} canonically processed at nt –2/–1 and aberrantly at nt –1/+1 in *E. coli* BW[Ec\textit{rnpB}], BW[At\textit{PRORP1}] and BW[At\textit{PRORP3}] based on the RNA-Seq data presented in Figure 3. (B, C) Northern blot analysis of tRNA^{His} and tRNA^{Sec} levels in *E. coli* MG1655 (lanes 1), BW[Ec\textit{rnpB}] (lanes 2), BW[At\textit{PRORP1}] (lanes 3 and 5) and BW[At\textit{PRORP3}] (lanes 4 and 6). Total RNA was prepared from outgrowth cultures grown until OD_{600} ∼ 1.0 at 37 °C in glucose-containing (lanes 1–4) or arabinose-containing (lanes 5 and 6) LB medium. After separation on 10% denaturing PAGE (9 μg RNA loaded per lane) the RNA was blotted onto a nylon membrane; tRNA^{His} and tRNA^{Sec} were hybridized with specific digoxigenin-labeled antisense RNA probes (for details, see Supplementary Table S4); signals indicated by filled arrowheads were assigned to mature tRNA^{His} and tRNA^{Sec}, respectively, while signals marked by open arrowheads were interpreted as pre-tRNAs with a few nucleotides at the 5′- or 3′-end in the case of tRNA^{His} and with long 5′-extensions in the case of tRNA^{Sec}. (C) After detection of tRNA^{Sec} levels, the nylon membrane was treated with stripping solution and was reprobed with a 5S RNA (\textit{rrfH})-specific digoxigenin-labeled antisense RNA probe to control for RNA loading and quality (bottom panel).

processed pre-tRNA^{Gly} efficiently and at the canonical site (Figure 5B, lanes 2–5), the other two tRNA substrates were cleaved unequally by the three \textit{AtPRORP} isoenzymes. Only \textit{AtPRORP2} and 3 cleaved *E. coli* pre-tRNA^{His} and pre-tRNA^{Sec} preferentially at the bacteria-typical –2/–1 site producing a 9-nt and 14-nt 5′-flank, respectively (Figure 5B, lanes 8, 9, 13 and 14). With \textit{AtPRORP1}, both substrates were cleaved to >50% at the (for these tRNAs) aberrant –1/+1 site to generate non-functional tRNA^{His} and tRNA^{Sec} moieties (Figure 5B; lanes 7 and 12). Furthermore, all three, and particularly \textit{AtPRORP2} and 3, cleaved pre-tRNA^{Sec} less efficiently than pre-tRNA^{Gly} and pre-tRNA^{His}, as suggested by the residual fraction of uncleaved substrate (Supplementary Table S7).

\textit{AtPRORP1}, 2 or 3 were unable to convert *E. coli* pre-4.5S RNA to its 5′-mature form (Figure 5B, lanes 17–19). In contrast, this non-tRNA substrate was processed by the bacterial RNase P holoenzyme from *E. coli* (Figure 5B, lanes 16). This finding is in line with the initial RNA-Seq and the primer extension data (Figure 2A and B).

**Northern blot and primer extension analyses of other tRNAs in \textit{AtPRORP}-dependent *E. coli* cells**

The Northern blot analysis was extended to tRNA^{Ser}(CGA) and tRNA^{Asp}(GUC) 2/2 (first and second row in Figure 3); for tRNA^{Asp}(GUC) 2/2, RNA-Seq indicated low levels of canonical cleavage in BW[At\textit{PRORP1}] bacteria, for tRNA^{Ser}(CGA) lower levels of canonical cleavage in the same strain but also reduced levels in the other strains (Figure 3). As controls, we used tRNA^{Arg}(CCU) as a tRNA with the potential to extend the acceptor stem by a single U:A bp (not presented in Figure 3 because of cDNA reads below the cutoff) and tRNA^{Ile1}(GAU) as an example for which the RNA-Seq results indicated predominantly canonical cleavage in all strains (Figure 3 and Supplementary Figure S5). In the case of tRNA^{Ile1}(GAU) and tRNA^{Asp}(CCU), Northern blot analysis did not reveal any evidence for precursor accumulation or aberrant processing (Supplementary Figure S5).

tRNA^{Asp}(GUC). *Escherichia coli* encodes three tRNA^{Asp}(GUC) genes, of which \textit{aspT} and \textit{aspU} have also
Figure 5. In vitro cleavage of canonical and non-canonical bacterial RNase P substrates by E. coli RNase P and AtPRORP1, 2 and 3. (A) Substrate secondary structures. *T. thermophilus* pre-tRNA Gly containing its natural 5′-flanking sequence (14 nt). *E. coli* pre-tRNA His and pre-tRNA Sec were equipped with a 5′-flanking sequence of 10 and 14 nt, respectively. The *E. coli* pre-4.5S RNA used here contained a 5′-flanking sequence of 23 nt. The canonical cleavage sites are marked by black arrows. (B) Cleavage of 5′-32P-end labeled pre-tRNA Gly, pre-tRNA His, pre-tRNA Sec and pre-4.5S RNA by *E. coli* RNase P (lanes ‘Ec’), 100 nM RNase P RNA, 800 nM RNase P protein, 2 min incubation at 37°C), AtPRORP1, 2 or 3 (lanes ‘P1’, ‘P2’ or ‘P3’, respectively; 120 nM enzyme, 20 min incubation at 37°C for AtPRORP1 and 3, 20 min incubation at 28°C for AtPRORP2); C, control (incubation of substrate for 20 min in the absence of enzyme). Canonical cleavage sites are marked by black, aberrant cleavage sites by gray arrowheads. Cleavage of pre-tRNA His at site –1/+1 was quantified as 7.5 ± 1.1 (in %, ± standard deviation) for Ec, 63.7 ± 1.6 for P1, 9.7 ± 1.5 for P2 and 9.8 ± 1.9 for P3 (based on at least four independent experiments); cleavage of pre-tRNA Sec at site –1/+1 was 6.1 ± 1.8 (in %, ± standard deviation) for Ec, 72.6 ± 3.6 for P1, 16.0 ± 2.8 for P2 and 12.4 ± 1.9 for P3 (based on at least 5 independent experiments). For more details on the cleavage assay, see the Supplementary Material.
identical 5'-leader sequences up to position -4 (Figure 6A); aspT and aspU, corresponding to Asp(GUC) 1/2 in Figure 3, can form a U1 : G575 wobble plus two extra G:C pairs, whereas aspV transcripts, corresponding to Asp(GUC) 2/2, have the potential to expand the acceptor stem by 2 G:C pairs (Figure 6A); aspV transcripts showed the strongest propensity of miscleavage in BW[AtPRORP] according to Figure 3. Elevated precursor levels were indeed observed in northern blots for tRNA^{Asp}(GUC) in BW[AtPRORP] cells (Figure 6B, lane 3), but hardly for BW[AtPRORP3] cells (lane 4) and not at all for MG1655 and BW[EcnpB] bacteria (lanes 1 and 2). We also noticed that the main tRNA^{Asp} band in RNA preparations from BW[AtPRORP] bacteria (Figure 6B, lane 3) migrated slightly slower than in the other lanes including the one representing BW[AtPRORP3] cells (lane 4).

Since the resolution of the northern blots was relatively low, we further performed a primer extension analysis. Using RNA from exponentially grown cells (Figure 6C, outgrowth phase), we could confirm the northern blot results. In BW[AtPRORP] cells, little tRNA^{Asp}(GUC) cleaved at the canonical site was detected, but substantial amounts of tRNA processed at the aberrant -2/-1 site and precursor signals corresponding to 5'-extensions of about 6 to 8 nt. The latter extension products are consistent with 7-nt 5'-extensions of about 6 to 8 nt. The latter extension products are consistent with 7-nt 5'-extensions of about 6 to 8 nt. Our previous finding that hairpin structures representing tRNA acceptor stem and T arm are poor substrates for PRORP3 (34). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35).

DISCUSSION

Processing of 4.5S RNA in AtPRORP-depending E. coli cells

AtPRORP1, 2 and 3 were not able to cleave 4.5S RNA in vitro under conditions where canonical tRNA substrates are processed efficiently. In accordance with this finding, RNA-Seq and primer extension analyses showed that 5'-precursor transcripts of 4.5S RNA accumulated in E. coli BW cells depending on AtPRORP1 or 3. After prolonged growth of the AtPRORP1 or 3 complementation strains, essentially no residual 5'-mature 4.5S RNA was detectable (Figure 2B, lanes 9 and 11). We thus conclude that AtPRORP1 and 3 are essentially unable to contribute to 4.5S RNA 5'-end maturation in the E. coli host. This is in line with our previous finding that hairpin structures representing tRNA acceptor stem and T arm are poor substrates for PRORP3 (34). Moreover, the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35). Thus, we consider it unlikely that the 3'-terminal CCA is no crucial determinant for cleavage by AtPRORP1 or AtPRORP3 (34,35).

Complementation studies with other PRORPs and in Bacillus subtilis

AtPRORP1 was unable to support growth of B. subtilis RNase P mutant bacteria (Supplementary Figure S6). Western blot analyses (Supplementary Figure S7) suggested that insolubility and degradation of AtPRORP1 in B. subtilis are the major causes for the negative complementation results in this Gram-positive bacterium. Complementation by PRORP2 from T. brucei (TbPRORP2) in our E. coli and B. subtilis RNase P mutant strains was found to be negative as well (Supplementary Figure S8), which is attributable to insolubility of the heterologous protein in the bacterial hosts (Supplementary Figure S9). In comparison, AtPRORP1 was predominantly present in the soluble protein fraction of E. coli cells (Supplementary Figure S9C).
Figure 6. Processing of *E. coli* tRNA*_{Asp}*(GUC) and tRNA*_{Ser}*(CGA) by *At*PRORP enzymes in *E. coli* BW. (A) Secondary structures of the analyzed *E. coli* precursor tRNAs encoded by *aspT*, *aspU*, *aspV* and *serU*; all four have the ability to form extended acceptor stem base pairings involving G:C base pairs.
E. coli MG1655 cell can vary between ~450 and ~5300 molecules in minimal versus rich media (38).

**Processing of tRNA^{His} by ATRORP in E. coli**

ATRORP1 mislocalizes *E. coli* pre-tRNA^{His} in vivo and *in vitro* to a substantial extent (Figures 4A and 5). The absence of a G_{-1} and thus the lack of the G_{-1} : C_{+73} bp, in *E. coli* tRNA^{His} was shown to reduce aminoacylation efficiency by the cognate HisRS enzyme more than two orders of magnitude (39, 40). Thus, at least half (Figure 5) of the tRNA^{His} processing products in *E. coli* BW[ATPRORP] bacteria are predicted to be non-functional. This may well have a negative effect on the protein synthesis rate, particularly affecting the translation of mRNAs enriched in histidine codons. We analyzed whether the cells may respond to the depletion of chargeable tRNA^{His} by increasing transcription from its single hisR gene, but northern blot analysis provided no evidence for elevated tRNA^{His} levels in BW[ATPRORP] bacteria grown in the presence of glucose (Figure 4B).

The finding that the nuclear ATRORP2 and 3 isoenzymes processed bacterial pre-tRNA^{His} at the −2/−1 site (34, this study) points to the previously noted flexibility of ATRORP3 to efficiently fit and process substrates extended by a single G:C base pair (34). As previously discussed, the biological significance of this feature is elusive and such acceptor stem extensions appear to be under negative selection in the nuclear tRNA genes of *A. thaliana* (34). Counterintuitively, *A. thaliana* mitochondria and chloroplasts encode bacterial-type tRNA^{His} genes (http://planttna.ibmp.cnrs.fr/; 41), however the organellar ATRORP1 cleaved only <50% of *E. coli* pre-tRNA^{His} at the upstream ‘bacterial-like’ site (−2/−1). These observations, although not understood at present, provide evidence that substrate recognition by organellar ATRORP1 is not identical to that of the nuclear isoenzymes.

**Aberrant processing of tRNA^{Sec} by ATRORP1 in E. coli**

Except for the American cranberry *Vaccinium macrocarpon* (42), genes for tRNA^{Sec} have not been found in flowering plants (41). Eukaryal tRNA^{Sec} molecules, if present, also differ from their bacterial counterparts in their acceptor and T stem structures (34). Thus, a bacterial-like tRNA^{Sec} is absent from the natural substrate spectrum of ATRORP enzymes. *E. coli* tRNA^{Sec}, encoded by the selC gene, carries a variable arm with more base pairs than in any other tRNA. Furthermore, it has an unusual D-arm with a 6-bp stem and a 4-nt loop. Nonetheless, bacterial tRNA^{Sec} assumes an L-shaped structure from which the long variable arm protrudes (44). tRNA^{Sec} is the second tRNA in *E. coli* beside tRNA^{His} that is subject to non-canonical processing by endogenous RNase P. 5′-End maturation leaves an 8-bp acceptor stem with an additional G_{-1} : C_{+73} bp as in tRNA^{His} (33). This is essential for incorporation of selenocysteine into three *E. coli* proteins, the formate dehydrogenases (FdH) Fdh-O, Fdh-N and Fdh-H (45). The three Fdh proteins are involved in anaerobic energy metabolism, where they function in electron transfer from formate to nitrite via cytochrome c552 (46). However, other glycolysis metabolites (NADH, and possibly lactate and ethanol) can also mediate nitrite reduction (46). Considering that *E. coli* cells depending on ATRORP1 or ATRORP3 were (almost) devoid of mature tRNA^{Sec} (Figure 4C; BW[ATPRORP] even more than BW[ATPRORP] bacteria; lane 4 versus 3), this defect can be excluded as a reason for the slower growth of BW[ATPRORP] relative to BW[ATPRORP] cells (Figure 1).

The reduction of precursor and mature tRNA^{Sec} levels in BW[ATPRORP] and BW[ATPRORP] cells grown under non-permissive conditions can be explained by degradation of unprocessed precursor, and inefficient processing of pre-tRNA^{Sec} by ATRORP1 and 3 is attributable to the potential of this precursor to form an acceptor stem extended by three G:C base pairs (Figure 5A). Such G:C-rich extensions are not found in precursor tRNAs of *A. thaliana* (34), indicating that ATRORP enzymes have not been under evolutionary pressure to cope with such substrates. This is also in line with the *in vitro* processing results, demonstrating less efficient ATRORP processing of pre-tRNA^{Sec} relative to pre-tRNA^{His} and pre-tRNA^{Gly} in particular (Supplementary Table S7). We assign the observation of some ATRORP-catalyzed processing of pre-tRNA^{Sec} *in vitro*, but barely in *E. coli* cells, to the use of enzyme excess in the *in vitro* experiments. As observed for pre-tRNA^{His}, ATRORP1 had a higher propensity than the two other isoenzymes to cleave pre-tRNA^{Sec} *in vitro* at the (in this case) aberrant −1/+1 site.

**PRORP isoenzyme-dependent complementation efficacy**

Here, we have shown that all three PRORP isoenzymes from *A. thaliana* are able to support growth of *E. coli* mutant bacteria in rich medium under conditions where endogenous RNase P levels are insufficient to sustain
growth. Thus, all three protein enzymes can take over basic and essential functions in tRNA 5′-end maturation normally catalyzed by the endogenous RNP enzyme. Surprisingly, E. coli bacteria depending on AtPRORP3 reproducibly showed a shorter lag phase than those complemented with AtPRORP1 (Figure 1). 4.5S RNA primer extension data (Figure 2) and northern blot results revealed that both isoenzymes are unable to process pre-tRNA Sec mature molecules and are largely deficient in tRNA 5′-end maturation (AtPRORP3 even more than AtPRORP1; Figure 4) precursor molecules. Thus, the 4.5S RNA and tRNA Sec processing defects cannot be the reason for the shorter lag phase of the BW[AtPRORP3] strain. However, for several tRNAs our RNA-Seq analysis revealed increased levels of aberrant cleavage in BW[AtPRORP1].

For selected tRNAs we were able to validate the RNA-Seq data in a more detailed biochemical analysis. For example, AtPRORP3 cleaved tRNA His at the functional -2/-1 site, whereas AtPRORP1 selected this cleavage site to 50% or less (Figures 3 and 5). Also, the retardation of tRNA Asp(GUC) and tRNA Ser(CGA) 1 nt upstream of the canonical site was only observed in the AtPRORP1 complementation strain (Figure 6C and D). Generally, aberrant processing by AtPRORP1 correlated well with pre-tRNA Asp acceptor stem extensions involving G:C base pairs (Supplementary Table S6). Those substrates lack a T loop equivalent, one of the most conserved tRNA features. The capacity of bacterial RNase P to specifically act on substrates deviate from canonical tRNA structures. The core task of both is the 5′-end maturation of tRNA His carrying an extra G –1 : C73 base pair. In Eukarya, the G –1, not encoded in tRNA His genes, is added by tRNA His guanylyltransferase (THG1) after RNase P processing. In those Archaea, where G –1 is genomically encoded, the RNase P enzymes (one RNA plus five protein cofactors) are thought to be still able to switch the cleavage site on pre-tRNA His. However, in other Archaea with G –1 not encoded in the tRNA His gene, the G –1 has to be added by the respective THG1 enzyme (47). For A. thaliana mitochondria, it has been proposed that both the bacterial and the eukaryal nuclear RNP 5′-maturative pathways for tRNA His coexist (32).

The presence of the eukaryal pathway was inferred from two lines of evidence: (i) expression of larch pre-tRNA His lacking G –1 in potato mitochondria and the subsequent identification of larch tRNA His molecules with G –1 added post-transcriptionally; (ii) transfer of a radioactive G –1 residue onto a 5′-G –1-tRNA His by a guanylyltransferase activity in potato mitochondrial extracts; the identity of the enzyme remained unclear and was shown to be different from the two A. thaliana THG1 homologs that localize to the nucleoplasm (32). Whatever the identity of this mitochondrial guanylyltransferase activity is, it seems to be quite inefficient, as substantial steady-state levels of non-functional tRNA His molecules lacking G –1 are detectable in the mitochondria of potato plants grown under normal conditions. Apparently, plant mitochondria tolerate the presence of a substantial fraction of inactive tRNA His molecules (with 5′-G –1 ) derived from imprecise pre-tRNA His processing by AtPRORP1. The presence of non- aminoacylated tRNA His resulting from ambiguous 5′-end maturation may also hint at a novel and so far unknown regulatory function in plant mitochondria.

**Evolutionary implications**

Catalytic RNA-based bacterial RNase P and PRORP enzymes are apparently the result of convergent evolution. The core task of both is the 5′-end maturation of canonical tRNAs, whose structures have remained essentially unchanged throughout evolution. However, the two types of RNase P differ in terms of specificity for substrates that deviate from canonical tRNA structures. E. coli RNase P, but not PRORP, can act efficiently on hairpin-like substrates. This is demonstrated in the present study for 4.5S RNA, a long-known non-tRNA substrate of E. coli RNase P (20,21), and is also suggested for structurally related substrates including phage φ80-induced M3 RNA (48), CI RNA from satellite phage P4 (49), the C4 repressor RNA of bacteriophages P1/P7 (50) and some other E. coli RNAs (for a review, see (1)). Those substrates lack a T loop equivalent, one of the most conserved tRNA features. The capacity of bacterial RNase P to specifically act on minimalistic substrates, originally discovered by Altman and coworkers (51,52), can be attributed to the enzyme’s recognition of multiple elements near the cleavage site, including the base pairing interaction between substrate 3′-ends and the L15 loop of RNase P RNAs, as well as selective binding of unstructured 5′-precursor fragments through the bacterial RNase P protein. For PRORP enzymes, the only natural, non-canonical substrates...
T. elements are associated with 5'- or 3'-termini of several plant mitochondrial mRNAs and appear to act as cis-signals for endonucleolytic cleavages by RNase P and/or RNase Z (54). T-elements look like degenerated tRNAs, harboring acceptor stem/T-arm mimics but lacking the anticodon arm (A. thaliana nad6) or a canonical D arm (Brassica napus orf138 (9, 10, 54)). However, with their conservation of T loop and the presence of at least a degenerated D arm, t-elements are structurally more closely related to canonical tRNAs than extended hairpin substrates such as 4.5S RNA. In addition, our study has revealed that the organellar AIPRORP1 enzyme has difficulties to act on pre-tRNAs that can form acceptor stem extensions involving G:C base pairs. AIPRORP3 was less sensitive to such extensions, but failed on pre-tRNA sec with its acceptor stem expanded by three G:C pairs, again consistent with previously reported findings on cleavage-site selection by AIPRORP3 (34). Future studies will have to address the molecular basis for the differences we have observed here between AIPRORP1 and AIPRORP3. Despite these differences between isoenzymes, it appears that PRORPs, like nuclear RNP RNase P enzymes (55–57), use different, structure- rather than sequence-based mechanisms of substrate recognition and cleavage-site selection (34) and, as a result, PRORPs have a narrower, more canonical tRNA-centric substrate spectrum than bacterial RNase P enzymes.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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