

***In vivo* stabilization of endogenous chloroplast RNAs by customized artificial pentatricopeptide repeat proteins**

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Received November 09, 2020; Revised April 05, 2021; Editorial Decision April 27, 2021; Accepted April 28, 2021

ABSTRACT

Pentatricopeptide repeat (PPR) proteins are helical repeat-proteins that bind RNA in a modular fashion with a sequence-specificity that can be manipulated by the use of an amino acid code. As such, PPR repeats are promising scaffolds for the design of RNA binding proteins for synthetic biology applications. However, the *in vivo* functional capabilities of artificial PPR proteins built from consensus PPR motifs are just starting to be explored. Here, we report *in vivo* functions of an artificial PPR protein, dPPR^{*rbcl*}, made of consensus PPR motifs that were designed to bind a sequence near the 5' end of *rbcl* transcripts in *Arabidopsis* chloroplasts. We used a functional complementation assay to demonstrate that this protein bound its intended RNA target with specificity *in vivo* and that it substituted for a natural PPR protein by stabilizing processed *rbcl* mRNA. We targeted a second protein of analogous design to the *petL* 5' UTR, where it substituted for the native stabilizing PPR protein PGR3, albeit inefficiently. These results showed that artificial PPR proteins can be engineered to functionally mimic the class of native PPR proteins that serve as physical barriers against exoribonucleases.

INTRODUCTION

The manipulation of gene expression represents a major challenge for both basic and applied biology. Progress in this field has been made possible by the discovery of natural products holding the potential to be tailored to powerful synthetic tools for genetic engineering. Posttranscriptional mechanisms play a prominent role in the control of

gene expression and RNA binding proteins mediate these processes. Thus, the possibility to engineer RNA binding proteins with desired RNA binding specificity has attracted considerable attention (reviewed in 1,2). Pentatricopeptide repeat (PPR) proteins constitute one of the largest families of RNA binding proteins in eukaryotes comprising more than 400 members in higher plants (3). PPR proteins are nucleus-encoded proteins but they function almost exclusively in mitochondria or chloroplasts where they hold various biological activities: protein barriers to RNA degradation, translational activation, recruitment of effectors to specific RNA sites, regulation of important RNA *cis*-element and remodeling of local RNA structures (reviewed in 4). PPR proteins are characterized by a variable number of tandem repeats (from 2 to over 30) of a degenerate 35 amino acid motif that forms a helix-loop-helix structure (5,6). The consecutive repeats stack to form a right-handed super helix that binds RNA with sequence specificity in a one-repeat/one-nucleotide mechanism (7). The RNA base bound by a PPR repeat is primarily determined by the identity of amino acids at two positions, 5 and 35 (8–11). The modular architecture of PPR repeats and the existence of an amino acid code for RNA base recognition make them an attractive scaffold for the rational design of RNA binding proteins with desired sequence specificity and therefore, the control of RNA functions *in vivo*. In fact, the PPR code has been successfully used to reprogram natural PPR proteins in plants to bind new mRNA sequences *in vivo* that are different from their native ones. For example, the PPR code was used to reprogram the sequence specificity and *in vivo* function of the mitochondrial PPR protein RPF2 in *Arabidopsis* plants (12). RPF2 possesses 16 PPR repeats and targets two RNA sites sharing a strong sequence identity that are located within the 5'-UTRs of *cox3* and *nad9* genes to define the 5' end processing of these transcripts by promoting a likely 5'-3' endonucleolytic activity (13). Colas

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des Francs-Small *et al.* modified the amino acid composition of the RPF2 PPR tract to reprogram its *in vivo* specificity and bind a new RNA target within the mitochondrial *nad6* ORF which induced its subsequent cleavage. Despite its relative success, the assay highlighted a major *ab initio* limitation for the engineering of natural PPR proteins: the nucleotide specificity of only two of the 16 PPR motifs in RPF2 could be manipulated, which greatly restricted the choice of the RNA target to a sequence sharing high identity with RPF2's native targets in mitochondria. The difficulty to freely reprogram the binding specificity of natural PPR proteins was additionally illustrated by a recent study that exploited the maize RNA stabilizer and translation enhancer PPR10 and its cognate chloroplast *atpH* binding site to build an inducible switch for the expression of plastid transgenes in tobacco (14). In this study, a variant of PPR10 was successfully expressed from the tobacco nuclear genome to stimulate the expression of a chloroplast transgene whose mRNA stability and translation were under control of a modified version of the native PPR10 binding site. As for RPF2, however, the modification of PPR10 sequence specificity did not go further than 2 nucleotides.

Thus, in these two instances, the relative success of manipulating the specificity of natural PPR proteins *in vivo* is overshadowed by the inability to fully customize all of their PPR repeats to bind any chosen RNA sequence *in vivo*. The incapacity to recode the specificity of some PPR motifs in natural PPR proteins lies in their amino acid inconsistencies at positions that determine base specificity in relation to the PPR code and their unpredictable contribution to RNA binding (6,15). In order to circumvent the limitation of natural PPR proteins, several synthetic PPR scaffolds have been created (16–18). These artificial PPR proteins (called as well designer PPRs, dPPRs) derived from consensus PPR motifs that offer predictable and reliable sequence specificity *in vitro* (19). Several *in vivo* applications have been envisioned for dPPRs and each of these applications derived from two main functions that are naturally occupied by PPR proteins in organelles: the sequestration of RNA from interaction with other proteins or RNA, or the targeting of effectors to specific RNA sites (reviewed in 20). Therefore, artificial PPR proteins must fulfill these two activities *in vivo* in order to be implementable as tools for the manipulation of RNA functions in living organisms. Currently, there is only one example for the *in vivo* application of dPPRs (21). In this study, a dPPR protein was successfully engineered in transgenic Arabidopsis plants to capture a specific mRNA in chloroplasts, demonstrating that the artificial dPPR scaffold can selectively and reliably bind designated RNA *in vivo*. However, this study did not reveal whether dPPRs hold functional potentialities similar to that of natural PPRs *in vivo*.

Many natural PPR proteins are involved in the control of RNA stability in plant organelles where they specifically bind to intergenic regions of RNA precursors to stabilize and protect the adjacent RNA sequences from degradation by 5'→3' or 3'→5' exoribonucleases, thereby defining the 5' or 3' ends of the processed RNAs. To test whether dPPRs can fulfill these properties *in vivo*, we conducted functional complementation assays in Arabidopsis of nat-

ural PPR proteins that stabilize the 5' end of mature chloroplast mRNAs by an RNase blockade mechanism. We showed that a dPPR made of 13 PPR repeats and programmed to bind the *in vivo* 5' end of a particular mRNA efficiently substituted for the biological function of the endogenous PPR protein. Furthermore, we demonstrated that positioning the RNA binding site of the dPPR to a different genomic location than that of the natural PPR protein creates a new and functional 5' end of the processed mRNA *in vivo*. In conclusion, these results showed that dPPRs hold functional capacities similar to those of natural PPRs by sequestering specific RNA sequences and preventing their access to exoribonucleases. This study provides an additional application of artificial dPPRs in the targeted control of RNA stability *in vivo*.

MATERIALS AND METHODS

Plant material

The *mr11* Arabidopsis T-DNA homozygote line (SALK_072806, Col-0 background) seeds were a generous donation of Olivier Vallon (22). Arabidopsis plants were grown on soil under controlled conditions: 12-h light (21°C): 12-h dark (18°C); 160–175 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$. *Nicotiana benthamiana* plants were grown on soil under standard greenhouse conditions: 16-h light (22°C): 8-h dark (18°C) cycles; 200–250 $\mu\text{mol photons m}^{-2} \text{s}^{-1}$. Seeds for *pgr3-4* homozygote line (FLAG_086B06, Was-0 background) were a gift of Toshiharu Shikanai (Kyoto University) and is the same allele previously analyzed by ribosome profiling (23). Seeds were germinated on Murashige and Skoog (MS) medium supplemented with 2% (w/v) sucrose and grown at 22°C s in 10-h light: 14-h dark (light intensities as stated in Figure legends) before being transferred to soil.

Complementation of *mr11* and *pgr3* mutants with dPPR-encoding transgenes

The DNA sequences of Arabidopsis chloroplast transit peptide RecA and dPPR^{*rbcl*} were codon-optimized for Arabidopsis and synthesized by GeneCust company. The two sequences were assembled using overlapping PCRs. RecA and dPPR^{*rbcl*} were PCR-amplified using primers K450 Fw/K628 Rev and K627 Fw/K465 Rev, respectively, and assembled by overlapping PCR. Cloning was performed using Gateway technology (Thermo Fisher Scientific, Waltham, MA, USA) following the manufacturer's protocols. The *RecA-dPPR^{rbcl}* gene was first cloned into pDONR207 entry vector using BP clonase II and subsequently cloned into binary vectors pAUL1 (for C-terminal protein fusion with 3xHA tag) (24) or pMDC83 (for C-terminal protein fusion with GFP) (25) using LR clonase II. GV3101 Agrobacteria carrying the *RecA-dPPR^{rbcl}:pAUL1* construct were used for floral dip transformation of homozygous *mr11* plants and Agrobacteria carrying the *RecA-dPPR^{rbcl}:pMDC83* construct were used for tobacco leaf agroinfiltration and protein subcellular localization. Seeds obtained from T₀ Arabidopsis plants were sown on soil and the transgenic seedlings were selected for resistance

to BASTA® (Bayer AG, Leverkusen, Germany). 30 confirmed transgenic plants were screened for dPPR^{rbcl} expression by immunoblot on leaf total protein extracts using HA antibodies (H9658 clone, Sigma-Aldrich) and three lines expressing gradual levels of dPPR^{rbcl} protein were chosen for phenotypic analyses.

The dPPR^{petL} sequence was codon optimized for Arabidopsis, synthesized by Genewiz (South Plainfield, New Jersey) and cloned into a modified version of pCambia1300 that allows the C-terminal fusion of the protein with 3xFLAG, as described previously (21). This plasmid was used to transform Arabidopsis *pgr3-4* plants by the floral dip method. Transgenic seedlings were selected for resistance to hygromycin on MS medium. Plants were grown for two weeks and pools of 6-to-8 seedlings were harvested for protein and RNA analysis.

Information about the primer sequences can be found in Supplementary Table S1. The DNA and protein sequences used in this study are provided in Supplementary Figure S1.

Immunoblot analysis

Total protein extraction was achieved by homogenizing leaf discs (9 mm diameter) in 150 µl of 2× Laemmli sample buffer (120 mM Tris-HCl, pH 6.8, 4% SDS, 20% glycerol, 2.5% β-mercaptoethanol, 0.01% bromophenol blue). Samples were centrifuged at 18 000 g for 5 min at room temperature and 10 µl of the supernatant were analyzed by SDS-polyacrylamide gel electrophoresis (10% polyacrylamide). Proteins were transferred onto PVDF membrane by wet western transfer using the Mini Trans-Blot® Cell Assembly (Bio-Rad) in buffer containing 25 mM Tris, 192 mM glycine, 20% ethanol.

PsaD, AtpA, AtpB and PetD antibodies were described previously (26). The NdhL antibody was a gift of Toshiharu Shikanai (Kyoto University). PsbE and PsbA (D1) antibodies were purchased from Agrisera. Monoclonal anti-HA (H9658 clone), anti-FLAG (M2 clone) and anti-Myc (9E10 clone) antibodies were purchased from Sigma-Aldrich.

Subcellular protein localization

A. tumefaciens strain GV3101 carrying the *RecA-dPPR^{rbcl}:pMDC83* construct were cultured overnight, pelleted for 5 min at 3200 g and resuspended in 10 mM MES pH 5.6, 10 mM MgCl₂, 150 µM acetosyringone to OD₆₀₀ of 0.3. The bacterial suspension was incubated for 2 h at room temperature in the dark and used for infiltration of leaves of 3-week-old *N. benthamina* plants. Two days after infiltration, leaves were digested in protoplast extraction medium (0.01% Macerozyme R10, 0.1% Driselase, 0.2% Cellulase ‘Onozuka’, 4.3 g/l MS salt mix, 0.5 g/l MES pH 5.6, 20 g/l sucrose, 80 g/l mannitol) in the dark for 5 h at 30°C with gentle shaking (50 rpm). Protoplasts were examined under a Zeiss LSM 780 confocal microscope. GFP was excited with a 488 nm laser and emission was acquired between 493 and 556 nm. RFP and chlorophyll were excited with a 561 nm laser line and emissions were acquired between 588–641 and 671–754 nm, respectively.

Purification of recombinant dPPR proteins

DNA encoding dPPR^{rbcl} without the transit peptide sequence was PCR-amplified (K461 Fw/K462 Rev) and cloned into the pMAL-TEV vector using BamHI/SalI restriction sites. DNA encoding dPPR^{petL} without the transit peptide sequence was PCR-amplified (K1245 Fw/K1227 Rev) and cloned in the destination vector pHMGWA (27) by a two-step Gateway® cloning following the manufacturer’s instructions (Thermo Fisher). This destination vector expresses an N-terminal six-histidine in addition to the Maltose binding protein tag. Rosetta 2 E. coli cells containing the protein expression vector were grown to OD₆₀₀ of 0.5 and protein expression was induced with 1 mM IPTG for 3.5 h at 20°C and 220 rpm shaking. Bacteria were pelleted and resuspended in lysis buffer (30 mM Tris pH 7.5, 450 mM NaCl, 5 mM β-mercaptoethanol, cOmplete™ EDTA-free Protease Inhibitor Cocktail (Roche) with the addition of 5 mM of imidazole for rdPPR^{petL}) and lysed by sonication. Recombinant dPPR^{rbcl} and dPPR^{petL} were purified by affinity chromatography to an amylose or Ni-IMAC resin.

Gel mobility shift assays

Synthetic RNA oligos were 5′ end-labeled with [γ-³²P]-ATP. Binding was performed for 3 h at 25°C in 20 µl reactions containing increasing amounts of rdPPR, 20 pM of RNA, 100 mM NaCl, 40 mM Tris pH 7.5, 4 mM DTT, 0.5 mg/ml Heparin, 10% glycerol and 1 U RNasin Plus RNase Inhibitor (Promega). Reactions were separated on native 5% polyacrylamide gels in 0.5× TBE buffer at 4°C and gels were dried after electrophoresis and exposed to a phosphorimaging plate.

In vivo protein labeling

In vivo labeling was performed on leaf discs from 4-week-old plants as described (28). ³⁵S-methionine pulse labeling was carried out for 15 min and total proteins were extracted as described previously. 100 000 cpm per sample were resolved on SDS-PAGE. After electrophoresis, the gel was stained in 50% methanol, 10% glacial acetic acid, 0.5g/l Coomassie brilliant blue R-250 and vacuum dried before being exposed to a phosphorimager plate.

RNA gel blot hybridization, cRT-PCR and primer extension assays

Total RNA was extracted from 2-week-old plants using Tri-Reagent (Molecular Research Center, Inc.) according to the TRIzol protocol (Thermo Fisher Scientific, Waltham, MA, USA). RNA was further purified by an additional phenol/chloroform extraction and ethanol precipitation. Five micrograms of total leaf RNA were resolved on denaturing formaldehyde gels as described (29). RNA was blotted onto Hybond-N⁺ membrane (GE Healthcare) by capillary transfer overnight using 20× SSC as blotting buffer. 60-mer DNA oligonucleotides were 5′ end-labeled using [γ-³²P]-ATP and T4 Polynucleotide Kinase (PNK) (Thermo Fisher Scientific, Waltham, MA, USA) according to the manufacturer’s instructions. Blots were hybridized overnight at 50°C in Church buffer (7% SDS, 0.5 M

NaPhosphate pH 7.0, 1 mM EDTA) and washed twice with washing buffer (1× SSC, 0.1% SDS) for 5 min at the hybridization temperature. Results were visualized on an Amersham Typhoon imager. Analysis of *petL* RNA by RNA gel blot hybridization used 6 μg leaf RNA and a 5'-end labeled synthetic DNA probe complementary to the *petL* open reading frame. The blots were hybridized overnight at 48°C in Church buffer and washed five times for 10 min in 0.2% SDS and 5× SSC at 48°C.

cRT-PCR was carried out as described (30) except that SSI V RT (Thermo Fisher Scientific, Waltham, MA, USA) and GoTaq (Promega) enzymes were used. K801 Rev primer was used for cDNA synthesis and K801 Rev/K803 Fw primers for the subsequent PCR. To identify the primary transcript ends, RNA was treated with RNA 5' Pyrophosphohydrolase (NEB) according to the manufacturer's instructions prior to RNA circularization. cRT-PCR products were cloned into pGEM®-T Easy vector (Promega) and more than 10 clones/cRT-PCR product were sequenced.

For primer extension assays of *rbcL*, 10 μg of total RNA were incubated in the presence of [γ -³²P]-ATP 5' end-labeled K626 Rev primer and 0.5 mM dNTPs at 75°C for 5 min. Temperature was then reduced to 50°C and SSI V RT enzyme mix (Thermo Fisher Scientific, Waltham, MA, USA) was added according to the manufacturer's protocol for First-Strand cDNA Synthesis Reaction. Reactions were incubated for 30 min at 50°C and stopped by adding one volume of RNA loading buffer (90% deionized formamide, 20 mM Tris pH 8.0, 20 mM EDTA, traces of bromophenol blue and xylene cyanole) and heating for 5 min at 95°C. Extension products were loaded onto an 8% polyacrylamide gel containing 8 M urea and run in 1× TBE buffer. The DynaMarker® Prestain Marker for Small RNA Plus (BioDynamics Laboratory Inc.) was run in parallel on the gel.

Primer extension analysis of the *petL* 5' end was performed with leaf RNA (2 μg of the wild-type and 7 μg for the *pgr3* mutant) and a 5'-end labeled synthetic DNA oligonucleotide as described previously (31). Products were resolved in an 8% denaturing polyacrylamide gel, and imaged with a Storm phosphorimager.

RIP-Seq and slot blot analysis

RIP-Seq analysis was performed as described previously (32). Briefly, stromal extracts (1 mg protein) were isolated from 2-week-old dPPR^{*rbcL*-1} or Col-0 plants and incubated with monoclonal anti-HA antibodies (Sigma). IgGs were captured with Protein A DynaBeads (Thermo Fisher Scientific, Waltham, MA, USA) and co-precipitated RNA was recovered by Trizol extraction followed by phenol/chloroform extraction and ethanol precipitation. The RNA was purified with the Monarch® RNA Cleanup Kit (NEB). 50 ng of RNA from each experiment were used for library generation with the NEBNext® Ultra™ II RNA Library Prep Kit according to the manufacturer's instructions. Deep sequencing (2 × 250 bp, v3 chemistry) was performed on a MiSeq sequencer (Illumina, San Diego, CA, USA) yielding 13.0 and 14.7 Mio (dPPR^{*rbcL*-1} replicate 1 and 2, respectively) as well as 15.5 and 13.1 Mio (Col-0 replicate 1 and 2, respectively) trimmed paired reads. To

determine depth of coverage (reads/nucleotides), the primary reads were aligned to the Arabidopsis chloroplast genome (accession number NC.000932.1) using CLC Genomics Workbench 6.5.1 (Qiagen, Valencia, CA, USA) with the following parameters: *mismatch cost* = 2, *insertion cost* = 3, *deletion cost* = 3, *length fraction* = 0.5, *similarity fraction* = 0.8, *global alignment* = no, *auto-detect paired distances* = yes. Aligned reads were extracted as reads per nucleotide and the mean value of the two replicates was displayed across the entire chloroplast genome.

For slot blot analysis, the co-immunoprecipitated RNA was recovered as described above. One-fourth of the pellet RNA and 1/40th of the supernatants were heated at 70°C for 10 min in 2× SSPE buffer (300 mM NaCl, 20 mM NaH₂PO₄, and 2 mM EDTA at pH 7.4) and blotted onto Hybond-N⁺ membrane (GE Healthcare) using a Minifold I slot blot system (GE Healthcare). Probe labeling and hybridization were as described for RNA gel blot analysis. Some of the blots were stripped to be rehybridized with another gene probe. Results were visualized on an Amersham Typhoon imager and data quantification was performed with ImageQuant TL (GE Healthcare).

RESULTS

Customization of a dPPR targeting the 5' end of mature *rbcL* mRNA

To test whether artificial PPR proteins built from consensus PPR motifs have similar functional capacities as natural P-type PPR proteins *in vivo*, we used an *in vivo* functional complementation assay of the Arabidopsis PPR protein MRL1. MRL1 is a PPR protein that is targeted to chloroplasts, where it stabilizes a processed isoform of RNA from the *rbcL* gene. *rbcL* encodes the large subunit of Ribulose-1,5-bisphosphate carboxylase-oxygenase (Rubisco) and is transcribed into a primary transcript whose 5' end maps at position -177 from the start codon (Figure 1A) (22,33,34). This mRNA precursor is processed to a transcript possessing a shorter 5'-UTR mapping at position -69. Accumulation of the processed isoform requires MRL1, and processing is presumed to involve the canonical PPR barrier mechanism: 5'-to-3' exonucleolytic degradation back to the MRL1 barrier bound immediately downstream of -69 (22). The RNA fragments bound by PPR proteins that stabilize RNA usually accumulate as small RNA footprints (sRNAs) whose termini coincide with the positions of the 5' or 3' ends of the mRNAs they stabilize *in vivo* (35–37). The MRL1 RNA footprint accumulates as a ~20–30 nt sRNA whose 5'-end coincides with the position of the processed *rbcL* mRNA, due to protection by the bound protein (Figure 1A). Despite the loss of the processed *rbcL* mRNA, *mrl1* mutant plants do not show a particular growth or physiological defect besides a slight reduction of RbcL protein content, suggesting that the primary *rbcL* transcript is translationally competent to produce sufficient RbcL (22). Therefore, *mrl1* serves as an ideal surrogate plant to express an artificial PPR and test its ability to complement the *in vivo* function of a natural PPR in gene-specific RNA stabilization by monitoring the recovery of the mature *rbcL* mRNA.

To this end, we conceived a protein, dPPR^{*rbcL*}, targeting the *in vivo* 5'-end of processed *rbcL* mRNA according to

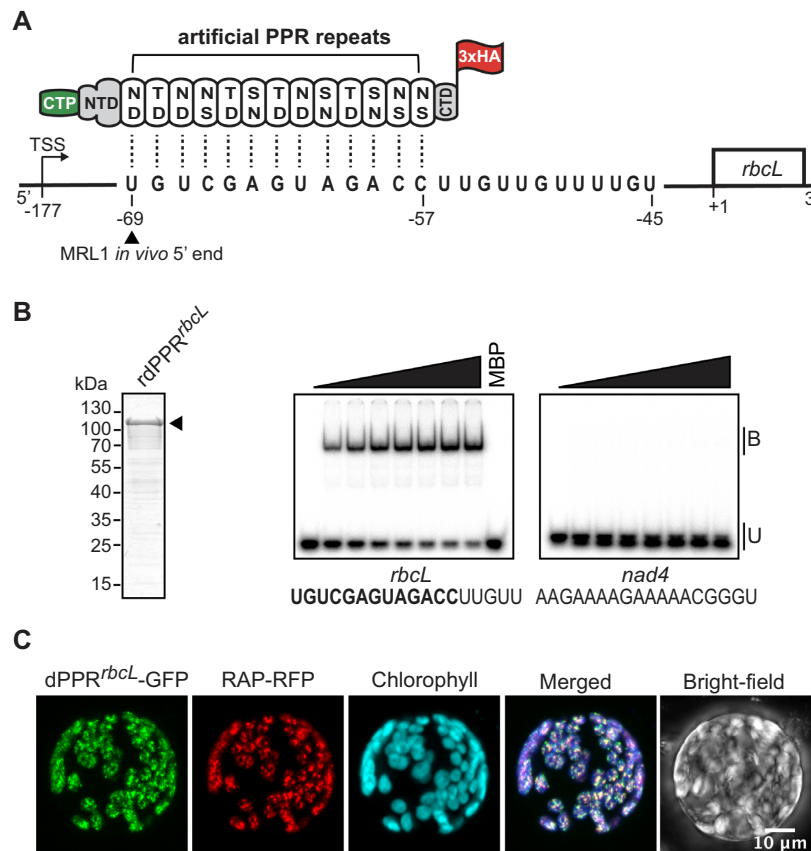


Figure 1. Engineering of an artificial PPR protein (dPPR^{rbcL}) programmed to bind the 5' end of mature *rbcL* mRNA in Arabidopsis chloroplasts. (A) Schematic of dPPR^{rbcL} design. The engineered protein was made of 13 consensus PPR tandem repeats programmed to bind a 13 nucleotide sequence beginning 69 nt and finishing 57 nt upstream of *rbcL* start codon. The dPPR tract was flanked by the N- and C-terminal domains (NTD and CTD) of the maize PPR10 protein. The top and bottom amino-acids featured in each PPR repeat correspond to the amino acid positions 5 and 35 that determine base specificity. An N-terminal chloroplast transit peptide from RecA protein (CTP) and a C-terminal 3xHA epitope tag were added to the protein to allow the chloroplast import and immunodetection of the protein *in vivo*. The *rbcL* transcription start site maps at position -177 and the wild-type (WT) *in vivo* 5' end of processed *rbcL* mRNA at position -69. The MRL1 footprint nucleotide sequence is shown. The genomic positions are given according to *rbcL* start codon. (B) Electrophoretic mobility shift assay (EMSA) showing preferential binding of recombinant dPPR^{rbcL} (rdPPR^{rbcL}) to *rbcL* 5' end. (Left) An aliquot of the purified rdPPR^{rbcL} used for binding assays was analyzed by SDS-PAGE and staining with Coomassie Blue. rdPPR^{rbcL} runs on the gel at a predicted size of ~118 kDa and is indicated by an arrowhead. (Right) rdPPR^{rbcL} was used in EMSAs with radiolabeled RNA oligonucleotides (*rbcL* or *nad4*) whose sequences are shown below. The sequence of the designated dPPR^{rbcL} binding site is highlighted in bold. Protein concentrations were 0, 22, 44, 66, 88, 110, 132 and 154 nM for rdPPR^{rbcL} and 200 nM for the maltose binding protein (MBP). Bound (B) and unbound (U) RNAs are indicated. (C) Chloroplast localization of dPPR^{rbcL} in *N. benthamiana* cells. The dPPR^{rbcL}-GFP fusion protein was transiently co-expressed in tobacco protoplasts with the Arabidopsis chloroplast nucleoid-associated protein RAP fused to RFP. Fluorescence images of GFP, RFP, chlorophyll and a merged image are shown.

the artificial PPR design described by Shen *et al.* (16). The dPPR^{rbcL} was made of 13 consensus PPR repeats flanked by amino- and carboxy-terminal segments of the maize chloroplast-localized protein PPR10 (31) (Supplementary Figure S1), and the PPR tract was programmed to bind a 13 nucleotide sequence matching the processed 5' end of the mature *rbcL* mRNA (Figure 1A). We chose a 13-motif design for dPPR^{rbcL} due to evidence that longer artificial PPR tracts are prone to increased off-target binding (19). For the *in vivo* complementation assay, the dPPR^{rbcL} additionally included the N-terminal chloroplast transit peptide of Arabidopsis RecA (first 68 amino acids) whose efficiency to target exogenous proteins to chloroplasts had already been demonstrated (38) and a C-terminal 3xHA tag for immunodetection (Figure 1A and Supplementary Figure S1). As a preliminary step to the functional complementation assay *in vivo*, we confirmed the RNA sequence binding speci-

ficity of dPPR^{rbcL} by expressing and purifying the mature recombinant protein (lacking the cleaved transit peptide), rdPPR^{rbcL}, in *E. coli* as an N-terminal protein fusion with the maltose binding protein and we used this protein in gel mobility shift assays (Figure 1B). The rdPPR^{rbcL} bound a 18-nt RNA matching the *in vivo* 5' end sequence of processed *rbcL* mRNA but did not bind an unrelated mitochondrial RNA sequence of the same size (*nad4*) demonstrating that, as expected, dPPR^{rbcL} binds with specificity to its designated RNA target *in vitro*. Furthermore, rdPPR^{rbcL} markedly bound its RNA ligand at protein concentrations in the nanomolar range showing that it binds RNA with high affinity.

Next, we tested the subcellular localization of the chimeric dPPR^{rbcL} protein in plant cells. To this end, a dPPR^{rbcL}-GFP fusion protein was transiently expressed in *Nicotiana benthamiana* leaves and leaf protoplasts were ex-

amed by confocal microscopy (Figure 1C). The GFP signal overlapped with the chlorophyll fluorescence demonstrating that dPPR^{*rbcl*}-GFP was efficiently imported to chloroplasts. In addition, the protein was found in discrete foci within chloroplasts that colocalized with the fluorescence of the nucleoid marker RAP-RFP (39) as demonstrated by the degree of colocalization measured between the two fluorophores (Supplementary Figure 2). These results indicate that dPPR^{*rbcl*} associates with the nucleoids in chloroplasts. This observation is in agreement with the reported association of MRL1 and several other PPRs controlling mRNA stability with the nucleoids of chloroplasts in maize (40). The results demonstrated that the customized dPPR^{*rbcl*} was expressed in plant cells where it localized to the chloroplasts similarly to its natural PPR counterpart, MRL1.

dPPR^{*rbcl*} can substitute for MRL1 to stabilize the 5' end of *rbcl* mRNA *in vivo*

To test the *in vivo* activity of dPPR^{*rbcl*} and its capacity to complement the biological function of MRL1, we generated transgenic Arabidopsis plants expressing dPPR^{*rbcl*} in the *mrl1* mutant background. Three independent transgenic lines expressing different levels of dPPR^{*rbcl*} protein (Figure 2A) were selected for further phenotypic analyses. The expression of dPPR^{*rbcl*} in the transgenic plants did not cause any growth phenotype as compared to *mrl1* and wild-type (Col-0) plants (Figure 2B), and immunoblot analyses using antibodies against core subunits of the major thylakoid membrane complexes did not reveal any change in chloroplast protein accumulation (Figure 2A). These results suggest that expression of dPPR^{*rbcl*} does not cause deleterious pleiotropic effects in plants.

To determine whether dPPR^{*rbcl*} stabilized the 5' end of processed *rbcl* mRNAs in the *mrl1* mutant, we performed a primer extension assay that allows the identification of both the primary and 5'-end processed *rbcl* transcripts (Figure 3A). As expected, both the primary and processed *rbcl* mRNAs were detected in wild-type plants whereas the processed RNA was absent in *mrl1* mutant. However, we observed that *mrl1* mutant also showed some reduction in the primary *rbcl* mRNA suggesting that the protein might have a secondary binding site upstream, or that its absence might expose an otherwise inaccessible endonuclease-sensitive site. In the three transgenic *mrl1* lines expressing dPPR^{*rbcl*} protein, a processed *rbcl* mRNA isoform accumulated, and the abundance of this isoform correlated with the abundance of the dPPR^{*rbcl*} protein in each line. Altogether, these results demonstrated that the artificial dPPR^{*rbcl*} had the capacity to control the stability of its RNA target *in vivo* and to complement the function of the natural PPR MRL1.

The primer extension analysis revealed that the processed 5' end of *rbcl* mRNA that accumulated in the transgenic plants was several nucleotides longer than that in the wild-type plants. We conducted cRT-PCR to map precisely the ends of the different *rbcl* mRNAs accumulating in the different genotypes (Figure 3B). An RNA treatment with the RNA 5' pyrophosphohydrolase prior to cRT-PCR allowed to distinguish the 5' ends of the primary *rbcl* mRNA from

the processed ones. In agreement with the primer extension results, the cRT-PCR analysis revealed that the 5' end of the primary transcript mapped 177 nt upstream of the *rbcl* start codon in all genotypes. In addition, the results confirmed that the major 5' ends of the processed *rbcl* mRNA in the transgenic plants mapped at position -75/-73 from the *rbcl* start codon, 4 to 6 nucleotides upstream of the processed 5' end in the wild-type. The position of the 5'-end that is stabilized by dPPR^{*rbcl*} is as expected based on the fact that PPR10 (from which the N-terminal amino acid region of this dPPR is derived) protects several nucleotides upstream of the sequences bound by its canonical PPR motifs (41). Since we designated the first PPR repeat of dPPR^{*rbcl*} to bind the immediate 5' nucleotide of the MRL1-dependent processed *rbcl* mRNA, the engineered protein was expected to stabilize a processed *rbcl* mRNA with an end mapping several nucleotides 5' to that in the wild-type.

Consistent with the loss of the mature *rbcl* mRNA in the *mrl1* mutant, examination of the accumulation of *rbcl* transcripts by RNA gel blot hybridization showed that the level of *rbcl* mRNAs was reduced in the mutant compared to the wild-type (Figure 3C). In the complemented *mrl1* lines, *rbcl* mRNA level was restored according to the level of dPPR expression in these plants with a full recovery of mRNA abundance in the strong expressor line that was comparable to the wild-type level.

Despite the reduction of *rbcl* mRNA abundance in *mrl1*, Coomassie staining of total leaf protein resolved by SDS-PAGE did not reveal a noticeable reduction of RbcL protein in the mutant compared to the wild type (Figure 2A). To assess RbcL synthesis rate in the different genotypes, we performed pulse labeling analysis using intact plant leaf tissues fed with the ³⁵S-radiolabeled methionine (Figure 3D). The results showed that the loss of *rbcl* mRNA in *mrl1* was accompanied by a small reduction in the rate of RbcL synthesis. This modest effect of MRL1 on Rubisco synthesis and accumulation is consistent with results reported in the original description of the *mrl1* mutant (22). The level of *rbcl* mRNA correlated well with the abundance of neosynthesized RbcL protein in transgenic, wild-type and *mrl1* mutant plants. These results indicate that the 5' end processed *rbcl* mRNA contributes to the synthesis of RbcL protein in plants and confirmed that the 5' end processed *rbcl* mRNA that was newly defined by dPPR^{*rbcl*} in chloroplasts is functional in the transgenic plants.

Altogether, the molecular analyses demonstrated that the artificial dPPR^{*rbcl*} protein has the capacity to complement the *in vivo* function of the PPR protein, MRL1 and control RNA stability in chloroplasts.

dPPR^{*rbcl*} binds preferentially to the 5' UTR of *rbcl* mRNA *in vivo*

To get a genome-wide view of dPPR^{*rbcl*} RNA binding specificity *in vivo*, we performed RIP-seq analysis using chloroplast stroma isolated from the dPPR^{*rbcl*} transgenic and wild-type plants in two replicate experiments (Figure 4A). The read values (reads/nucleotide) and read mapping of the two immunoprecipitation replicates for each genotype (wild type and *mrl1*:dPPR^{*rbcl*}) were highly reproducible (Supplementary Figure S3). Therefore, the mean read val-

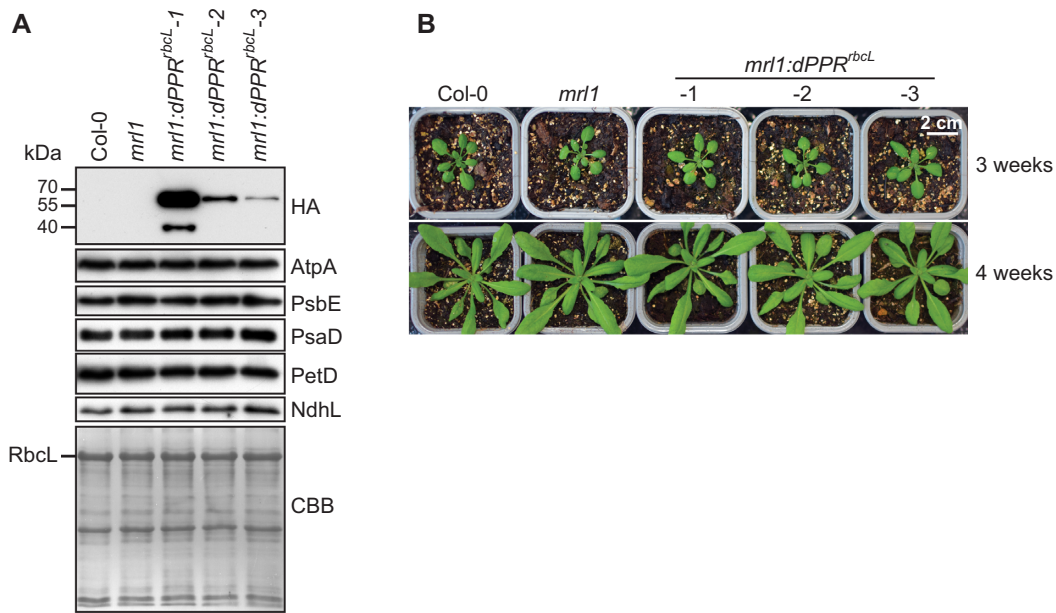


Figure 2. Phenotype of transgenic *mrl1* Arabidopsis plants expressing *dPPR^{rbcL}*. (A) Three *mrl1:dPPR^{rbcL}* independent lines expressing gradual levels of *dPPR^{rbcL}* were analyzed by immunoblots on total leaf protein extracts along with wild type (Col-0) and *mrl1* mutant plants. The *dPPR^{rbcL}* was detected with antibodies against the HA tag. The abundance of the chloroplast photosynthetic enzyme complexes in the different plant genotypes were analyzed with antibodies against core subunits: AtpA (ATP synthase), PsbE (Photosystem II), PsaD (Photosystem I), PetD (Cytochrome *b₆f*) and NdhL (NDH complex). One of the membrane duplicates was stained with Coomassie brilliant blue (CBB) and serves as the protein loading control. The band corresponding to the large subunit of Rubisco (RbcL) is marked. (B) Visible phenotype of 3- (top) and 4-week-old (bottom) Col-0, *mrl1* and transgenic *mrl1:dPPR^{rbcL}* Arabidopsis plants.

ues of the two replicates from the experimental and control immunoprecipitations were aligned to the Arabidopsis chloroplast genome (Figure 4B). The RNA immunoprecipitation using wild-type stroma yielded very low read coverage of the chloroplast genome except for the region covering the rRNA genes (Figure 4B, Supplementary Figure S3) whereas the experiment using the transgenic plants yielded several local peaks in the *psbA*, *atpF*, *psbC*, *psaB*, *rbcL*, *psbE* and *psbT* genes, with the *rbcL* peak being the most prominent (Figure 4B and Supplemental Table S2). Consistent with our experimental design, the genomic position of the prominent *rbcL* peak coincided with the *rbcL* 5' UTR (Supplemental Table S2, Supplementary Figure S4 and Figure 4C). Calculation of the enrichment of RNAs in the experimental versus control immunoprecipitations showed that RNAs in the 5' UTR of *rbcL* were enriched more than 700-fold whereas RNAs mapping within the *rbcL* ORF were more weakly enriched (less than 100-fold) demonstrating that *dPPR^{rbcL}* primarily binds the 5' UTR (Figure 4C). The RNAs from the other loci (*psbA*, *atpF*, *psbC*, *psaB*, *psbE* and *psbT*) were considered as potential off-targets and were examined in an independent RIP experiment. In this experiment, the experimental and control immunoprecipitations were both performed on stroma from the *mrl1:dPPR^{rbcL}* transgenic plants but used different antibodies. The immunoprecipitated RNAs with HA antibodies were compared to those from an immunoprecipitation with Myc antibodies that do not recognize *dPPR^{rbcL}* protein and the recovered RNAs from the pellets and supernatants were subsequently analyzed by slot blot hybridizations (Figure 4D). In agreement with the RIP-seq analysis,

the slot blot data confirmed that *rbcL*, *psbA*, *atpF*, *psbC*, *psaB*, *psbE*, *psbT* RNAs were specifically recovered in the *dPPR^{rbcL}* immunoprecipitation but with different degrees of enrichment. Quantification of the signals in the pellet versus supernatant showed that *rbcL* RNA was strongly enriched in the pellet whereas RNAs from *psbA*, *atpF*, *psbC*, *psaB*, *psbE* and *psbT* loci weakly coimmunoprecipitated with *dPPR^{rbcL}* (Figure 4D). Altogether, the data argue that *dPPR^{rbcL}* strongly associates to its designated *rbcL* RNA target *in vivo* while binding to a few off-targets with less affinity.

***dPPR^{petL}* can partially substitute for PGR3 to stabilize the 5' end of *petL* mRNA *in vivo*.**

To provide a second example, we addressed whether an artificial PPR protein can functionally substitute for the Arabidopsis PPR protein PGR3. PGR3 has three sites of action: (i) it binds the *petL* 5'-UTR, where it protects the downstream RNA from degradation and activates *petL* translation (42–44); (ii) it binds the *ndhG* 5' UTR where it activates *ndhG* translation (45); and (iii) it binds the *rpl14-rps8* intergenic region, where it stabilizes a processed 3'-end and mildly stimulates *rps8* translation (23). The footprint of PGR3 on *petL* mRNA is represented *in vivo* by an abundant sRNA whose 5'-end matches that of the PGR3-dependent processed *petL* 5'-end (35–37) (Figure 5A).

We designed an artificial PPR protein, *dPPR^{petL}* to bind a sequence within the PGR3 binding site in *petL*, but beginning several nucleotides downstream of the PGR3-dependent 5' end (Figure 5A). The *dPPR^{petL}* design was

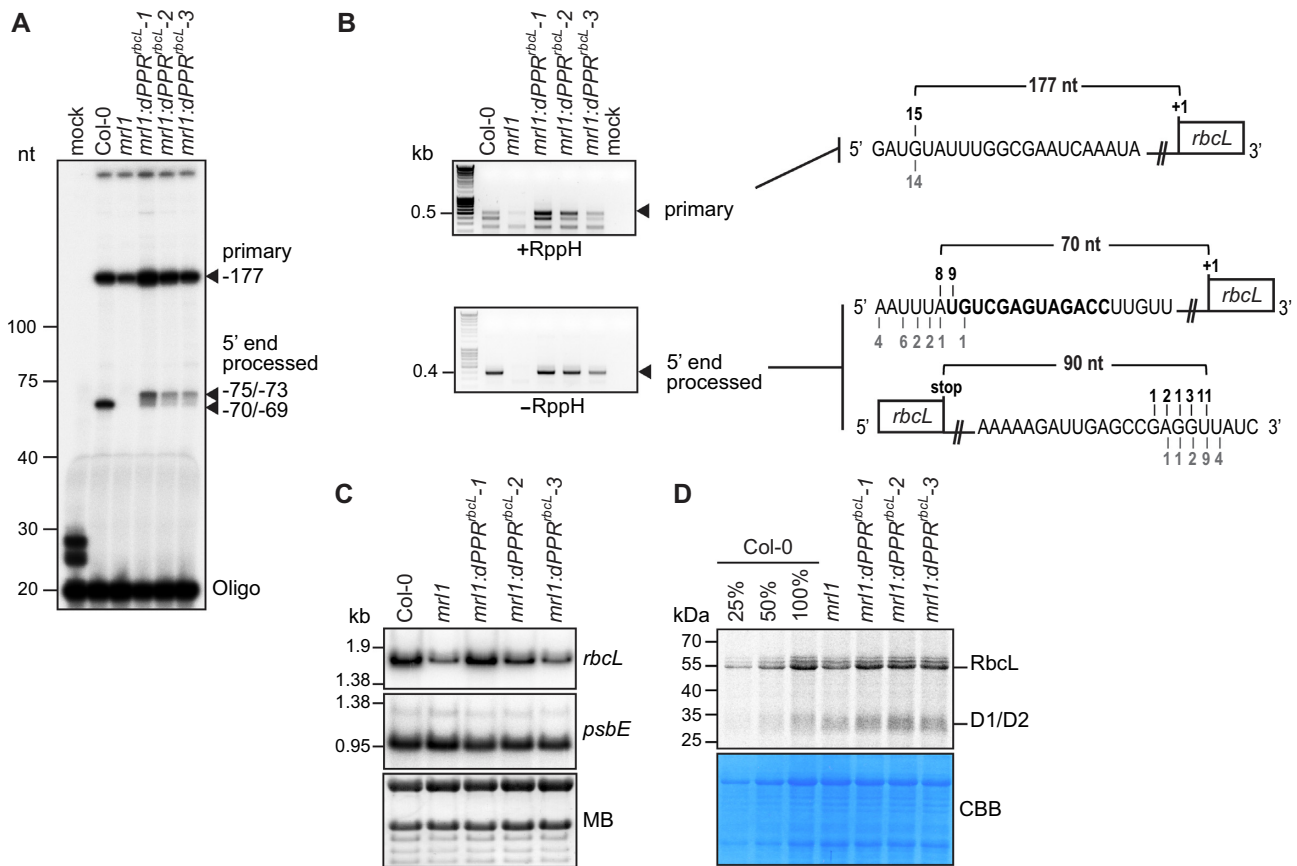


Figure 3. Functional complementation assay of *mrl1* Arabidopsis mutant by dPPR^{*rbcL*}. (A) Primer extension analysis of *rbcL* mRNAs 5' end accumulating in Arabidopsis chloroplasts. The extension products were separated on a 8% denaturing polyacrylamide gel. RNA samples from Col-0, *mrl1* and the transgenic plant lines were analyzed. The arrowheads indicate the 5' end of the primarily transcribed and processed *rbcL* mRNA. Their positions are given according to the *rbcL* start codon and their mapping by cRT-PCR. (B) cRT-PCR mapping of the termini of the primary (+RppH) and processed (-RppH) *rbcL* mRNAs in the Col-0 and complemented *mrl1* plants. The arrowheads indicate the cRT-PCR bands that were excised from the agarose gel for sequencing. The diagrams on the right display the RNA sequences annotated with the 5' or 3' ends determined by cRT-PCR. The dPPR^{*rbcL*} binding site is highlighted in bold and the number of clones mapping to each position is indicated above and underneath the RNA sequence for the Col-0 and *mrl1:dPPR^{rbcL}* genotypes, respectively. RppH: RNA 5' RNA 5' Pyrophosphohydrolase. (C) RNA gel blot hybridizations of leaf RNAs from plants of the indicated genotypes using strand-specific oligonucleotide probes for *rbcL* and a chloroplast control *psbE* gene. A portion of one of the blots stained with methylene blue (MB) is shown to illustrate equal sample loading. (D) *In vivo* chloroplast protein synthesis analysis. Intact leaves from the indicated plant genotypes were ³⁵S pulse labelled in presence of cycloheximide to block cytosolic translation. Leaf proteins were fractionated by SDS-PAGE. The autoradiograph and Coomassie blue stained gel (CBB) pictures are shown. The signal bands corresponding to radiolabeled RbcL and D1/D2 proteins are indicated.

not expected to complement PGR3 functions at *ndhG* and *rpl14-rps16* because those binding sites involve contacts with PGR3's C-terminal repeats (23,45), which are not represented by dPPR motifs in the synthetic protein (see Figure 6). The protein design is analogous to that of dPPR^{*rbcL*} except for the use of a different chloroplast targeting sequence (PPR10 versus RecA) and C-terminal tag (FLAG versus HA) (Supplementary Figure S1). The native PPR10 chloroplast targeting peptide was previously shown to target a dPPR to Arabidopsis chloroplasts *in vivo* (21). As for dPPR^{*rbcL*}, *in vitro* EMSA assays showed that recombinant dPPR^{*petL*} binds to its intended *petL* RNA target *in vitro* with specificity and high affinity (Figure 5B). We introduced the transgene expressing dPPR^{*petL*} into the genome of a null *pgr3* mutant, and used immunoblot analysis to identify lines expressing dPPR^{*petL*} (Figure 5C). The *pgr3* mutant has a barely discernable phenotype under the growth conditions we used, and the expression of dPPR^{*petL*} did not have

an obvious effect on the phenotype (Supplementary Figure S5A). When grown at low light intensity, the *pgr3* mutant exhibited a decrease in subunits of the cytochrome *b₆f* complex (PetD) and NDH complex (NdhL) (Figure 5C), as expected based on results reported for different *pgr3* alleles (42–44). When grown at a higher but still moderate light intensity, core subunits of other complexes were reduced as well (Supplemental Figure S5B). This is consistent with the mild defect in chloroplast translation reported for the same allele (23). Expression of dPPR^{*petL*} had little or no effect on the protein deficiencies. This is consistent with the fact that dPPR^{*petL*} was not expected to bind PGR3's *ndhG* or *rpl14-rps8* sites, together with its modest effects on *petL* mRNA as discussed below.

RNA gel blot hybridization showed the expected loss of *petL* transcripts from the *pgr3* mutant (Figure 5D). Expression of dPPR^{*petL*} resulted in partial restoration of the major PGR3-dependent transcript isoform (a tricistronic *petL*-

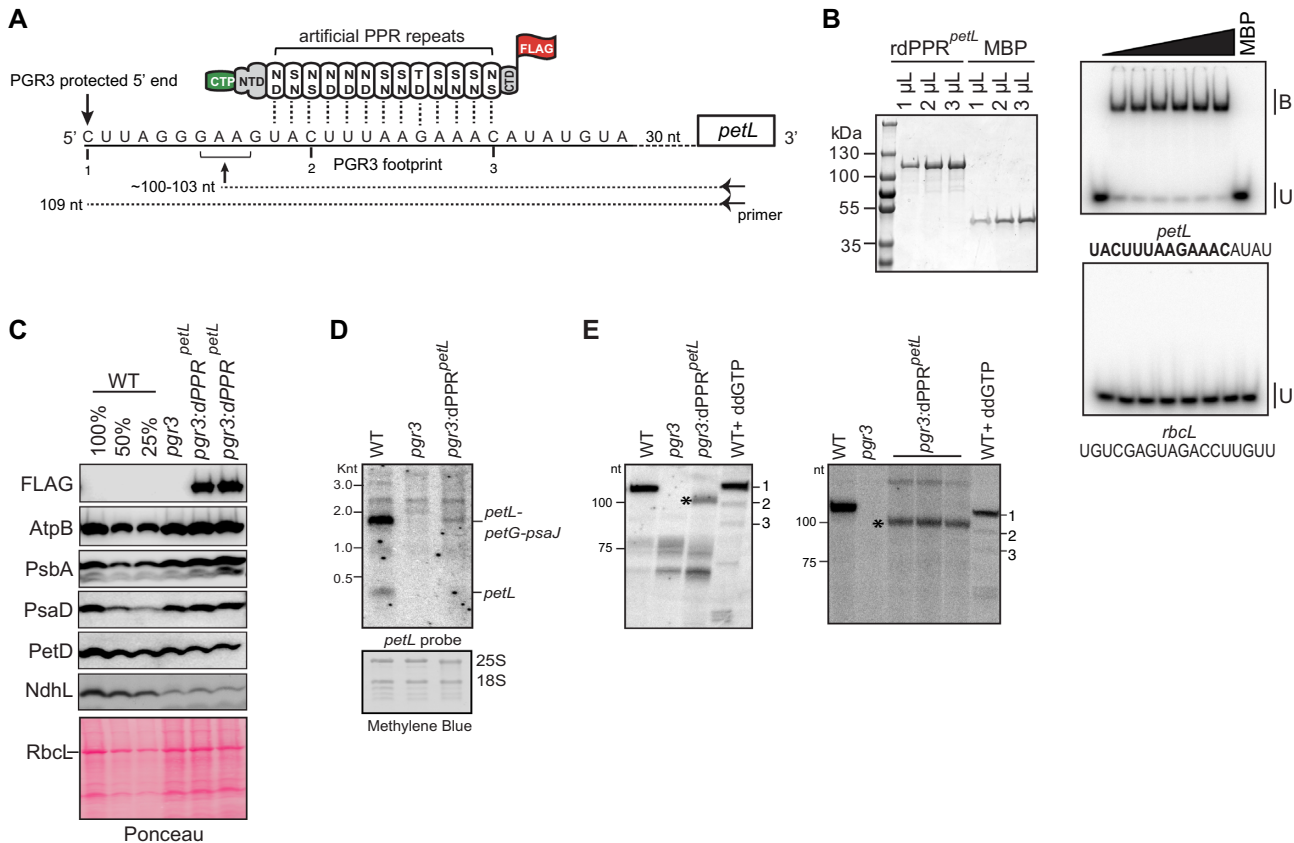


Figure 5. Partial complementation of PGR3's *petL* RNA stabilization function by dPPR^{*petL*}. (A) Schematic of dPPR^{*petL*} design. The protein consists of 13 consensus PPR motifs programmed to bind the indicated sequence in the *petL* 5'UTR. The amino acids at positions 5 (top) and 35 (bottom) in each dPPR motif are indicated. The dPPR tract was flanked by the N- and C-terminal regions (NTD and CTD) of maize PPR10, including PPR10's chloroplast targeting peptide (CTP). A 3xFLAG tag was fused to the C-terminus. The 5' end that is stabilized by PGR3 *in vivo* is marked with a vertical arrow, and the PGR3 footprint is underlined with a solid line. The horizontal arrow denotes the primer used for primer extension and the asterisk marks the 5'-end that is stabilized by dPPR^{*petL*}. The nucleotides marked 1, 2, and 3 are primer extension stops resulting from the use of the chain terminator ddG, which serve as size markers on the primer extension gels in panel E. (B) EMSA showing preferential binding of recombinant dPPR^{*petL*} (rdPPR^{*petL*}) to *petL* 5' end. (Left) An aliquot of the purified rdPPR^{*petL*} and MBP used for binding assays was analyzed by SDS-PAGE and staining with Coomassie Blue. rdPPR^{*petL*} has a predicted size of ~118 kDa. (Right) rdPPR^{*petL*} was used in EMSAs with radiolabeled RNA oligonucleotides (*petL* or *rbcl*) whose sequences are shown below. The sequence of the designated dPPR^{*petL*} binding site is highlighted in bold. Protein concentrations were 0, 5.6, 11.25, 22.5, 45, 90 and 118 nM for rdPPR^{*petL*} and 200 nM for the maltose binding protein (MBP). Bound (B) and unbound (U) RNAs are indicated. (C) Immunoblot analysis of *petL* transcripts in *pgr3* mutants expressing dPPR^{*petL*}. Plants used for this analysis were grown at low light intensity (60 μ E). Results from plants grown at 120 μ E are shown in Supplementary Figure S5B. Protein analyzed in each lane comes from pooled plants that were confirmed to have the indicated genotype by PCR. Replicate blots were probed with antibodies to the indicated proteins. The FLAG antibody detects dPPR^{*petL*}. The Ponceau S-stained blot is shown below to illustrate equal sample loading and the abundance of the large subunit of Rubisco (RbcL). (D) RNA gel blot hybridization demonstrating partial rescue of *petL* transcripts in *pgr3* mutants expressing dPPR^{*petL*}. 2 μ g of leaf RNA was loaded in each lane. The methylene blue stained blot is shown below to demonstrate equal sample loading. 25S and 18S are cytosolic rRNAs. (E) Primer extension assays demonstrating stabilization of the expected novel *petL* 5' terminus by dPPR^{*petL*}. The two panels show analysis of four independent pools of *pgr3* mutants expressing dPPR^{*petL*}. The positions of size markers are shown to the left. The last lane in each gel (WT + ddGTP) shows a primer extension reaction that used wild-type RNA and a trace amount of ddGTP to induce termination at C residues in the template (see panel A). The gel on the left used 4 μ g of RNA from the WT (Ws-0) and *pgr3* samples, and 14 μ g from *pgr3*:dPPR^{*petL*}. The gel on the right used 2.5 μ g of RNA from the WT (Ws-0) and *pgr3* samples, and 7.5 μ g from *pgr3*:dPPR^{*petL*}.

to be insoluble when expressed in heterologous systems (7), which limits the ability to confirm sequence specificities of variants prior to *in vivo* applications. Synthetic consensus PPR repeats (dPPR) were established to overcome these issues (16–18) but their *in vivo* functionalities have just started to be investigated (21).

MRL1 and PGR3 are 'pure' (P-type) chloroplastic PPR proteins that lack any accessory domains. MRL1 and PGR3 consist of 10 and 27 PPR repeats, respectively (22,42). Whereas MRL1 specifically stabilizes processed *rbcl* mRNAs in Arabidopsis, PGR3 binds three RNA targets (*petL* 5' UTR, *ndhG* 5' UTR and *rpl14-rps8* inter-

genic) with effects on RNA stability, translation, or both (23,42,45). However, we expected dPPR^{*petL*} to bind only the *petL* site because it did not include repeats that mimic the C-terminal region of PGR3, where specificity to the *ndhG* and *rpl14-rps8* sites is encoded (23,45) (Figure 6). The footprints of MRL1 and PGR3 in the *rbcl* and *petL* 5'-UTR, respectively, are represented by abundant sRNA of ~20–30 nucleotides that accumulate *in vivo* (Figure 6). However, like many other P-type PPR proteins, some PPR motifs in MRL1 and PGR3 lack canonical amino acids at the specificity-determining positions and the amino acid codes of some motifs do not match the presumed aligned

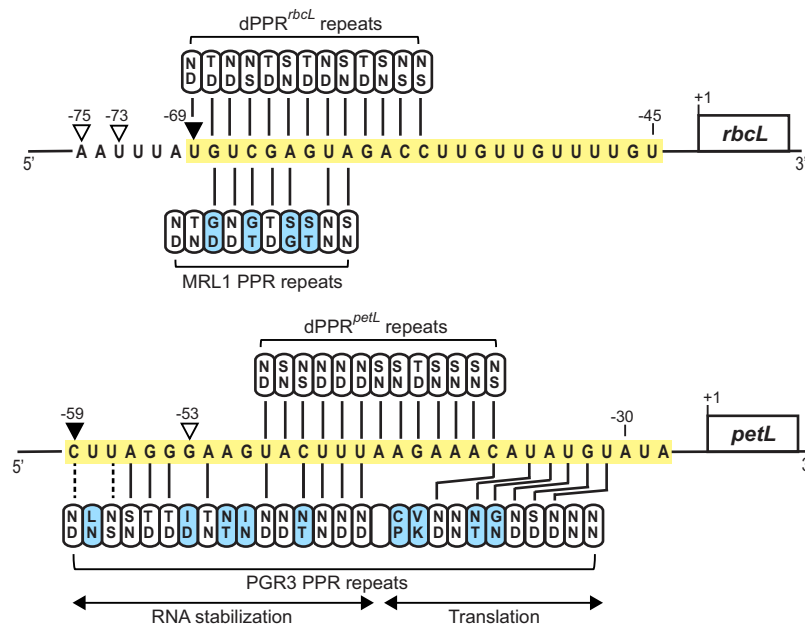


Figure 6. Alignments between dPPR^{rbcL}, dPPR^{petL} and MRL1, PGR3 and their respective designated or predicted *rbcL* and *petL* binding sites. The *in vivo* footprints of MRL1 and PGR3 are highlighted in yellow. The MRL1- and PGR3- or dPPR^{rbcL}- and dPPR^{petL}-dependent *in vivo* 5' ends are indicated by black or white arrowheads, respectively. The specificity-determining amino acids in each PPR motif (positions 5 and 35) are indicated. The PPR motifs that exhibit canonical amino acid combinations according to the original PPR code (9) are colored in white while the others are in blue. PPR repeats whose amino acid combinations match the nucleotide binding prediction according to a recently extended PPR code (11) are indicated with black lines. Dotted lines indicate compatible PPR nucleotide binding but with low affinity. PGR3 PPR repeats promoting RNA stabilization or translation are underlined (44,45).

nucleotides in the binding sites (Figure 6). Similarly to what has been reported for the well-characterized PPR protein PPR10 (15), some of these non-canonical interactions might still be critical for the high-affinity binding of these proteins to their RNA binding sites.

Our functional complementation assays of Arabidopsis mutants lacking MRL1 and PGR3 by dPPRs built from 13 synthetic PPR repeats add to the evidence that dPPRs bind their designated RNA targets with specificity *in vivo*, and additionally revealed that they can protect RNA downstream from their binding site against the action of exoribonucleases. Similarly to the action of native PPR proteins, this dPPR-dependent RNase blockade mechanism allowed the definition of novel 5'-end processed mRNAs in chloroplasts and control of RNA stability. Our RIP-seq experiment showed that dPPR^{rbcL} binds weakly to several off target RNAs *in vivo*, but this had no apparent effects on plant growth, pigmentation, or the accumulation of photosynthetic complexes harboring plastid-encoded subunits. Weak off-target binding without phenotypic impact was also observed in the sole prior study of *in vivo* dPPR protein expression (21). Therefore, some degree of off-target binding can be tolerated and does not preclude the use of this protein design to achieve a variety of practical purposes.

Nevertheless, the dPPR^{rbcL} and dPPR^{petL} data revealed interesting differences in their effects in comparison with their natural PPR counterparts. Whereas dPPR^{rbcL} fully complemented the *rbcL* mRNA stabilization function of MRL1, dPPR^{petL} only weakly stabilized its cognate *petL*

mRNA ligand as compared to PGR3. This difference might result simply from different expression levels. However, we suspect that the design of the dPPR tracts is a major factor underlying the different degree of complementation we observed. dPPR^{rbcL} harbors more PPR repeats than its counterpart MRL1 (Figure 6). By contrast, dPPR^{petL} contains fewer than half of the PPR motifs as PGR3 (Figure 6). Thus, dPPR^{petL} provides many fewer protein-RNA contacts than does PGR3, which presumably results in much lower RNA binding affinity. Furthermore, roughly 15 of the nucleotides that are masked upon PGR3 binding at *petL* will be left exposed upon dPPR^{petL} binding; should these exposed sequences be susceptible to endonucleolytic cleavage, this would compromise the ability of dPPR^{petL} to stabilize the *petL* 5'-end. PGR3 functions have been shown to partition between its N- and C-terminal PPR tracts: the *petL* RNA stabilization function requires only the N-terminal 16 PPR motifs, whereas the *ndhG* and *petL* translation activation functions also require its C-terminal ~11 PPR motifs (44) (Figure 6). dPPR^{petL} is not expected to mimic this latter function because it was not designed to interact with the relevant nucleotides at *petL* (Figure 6).

In conclusion, our results demonstrate that synthetic PPR proteins can be used for the targeted stabilization of specific chloroplastic mRNAs and therefore, the manipulation of organellar gene expression. Finally, this study provides knowledge upon which to develop strategies for the regulation of RNA turn-over in plant organelles by the use of designer PPRs.

DATA AVAILABILITY

The NGS data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus (46) and are accessible through GEO Series accession number GSE146249 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE146249>).

Sequence data from this article can be found in the EMBL/GenBank libraries under the following accession numbers: *MRL1* (At4g34830) and *PGR3* (At4g31850) from *A. thaliana*, *A. thaliana* chloroplast genome (NC_000932.1).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

We are grateful to Olivier Vallon (CNRS UMR7141) for the gift of *mr11* seeds and Toshiharu Shikanai (Kyoto University) for the gift of *pgr3* seeds and NdhL antibody.

FUNDING

Agence National de la Recherche [ANR-18-CE20-0013 to K.H.]; IdEx Unistra from the Investments for the future program of the French Government [to KH]; European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie action [794377 to N.M.]; U.S.-Israel Binational Agricultural Research and Development Grant [IS-5205-19 to A.B.]. Funding for open access charge: Centre National de la Recherche Scientifique. *Conflict of interest statement.* None declared.

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