OsPPR939, a *nad5* splicing factor, is essential for plant growth and pollen development in rice

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Abstract

**Key message** P-subfamily PPR protein OsPPR939, which can be phosphorylated by OsS6K1, regulates plant growth and pollen development by involving in the splicing of mitochondrial *nad5* introns 1, 2, and 3.

Abstract In land plants, pentatricopeptide repeat (PPR) proteins play key roles in mitochondrial group II intron splicing, but how these nucleus-encoded proteins are imported into mitochondria is unknown. To date, a few PPR proteins have been characterized in rice (*Oryza sativa*). Here, we demonstrate that the mitochondrion-localized P-subfamily PPR protein OsPPR939 is required for the splicing of *nad5* introns 1, 2, and 3 in rice. Complete knockout or partial disruption of OsPPR939 function resulted in different degrees of growth retardation and pollen sterility. The dramatically reduced splicing efficiency of these introns in *osppr939-4* and *osppr939-5* led to reduced mitochondrial complex I abundance and activity and enhanced expression of alternative respiratory pathway genes. Complementation with OsPPR939 rescued the defective plant morphology of *osppr939-4* and restored its decreased splicing efficiency of *nad5* introns 1, 2, and 3. More importantly, the 12th amino acid Ser in the N-terminal targeting sequence of OsPPR939 is phosphorylated by OsS6K1, and truncated OsPPR939 with a non-phosphorylatable S12A mutation in its presequence could not be imported into mitochondria, suggesting that phosphorylation of this amino acid plays an important role in the mitochondrial import of OsPPR939. To our knowledge, the 12th residue Ser on OsPPR939 is the first experimentally proven phosphorylation site in PPR proteins. Our results provide a basis for investigating the regulatory mechanism of PPR proteins at the post-translational level.

Introduction

Cytoplasmic male sterility (CMS), a maternally inherited trait characterized by the inability of a plant to produce viable pollen grains, is caused by incompatibility between mitochondrial and nuclear genes. CMS has been detected in nearly 200 flowering plant species (Bentolila et al. 2002). The sterile phenotype of CMS lines can be restored by nuclear genes called restorer-of-fertility genes (*Rf* genes), which encode highly diverse proteins. For example, *Rf2*, the first plant restorer gene isolated in maize CMS-T, encodes an aldehyde dehydrogenase (Cui et al. 1996). *Rf17* in rice CMS-CW (Chinese wild rice) encodes a mitochondrial sorting protein containing an acyl-carrier protein synthase-like domain (Fuji and Toriyama 2009). The candidate gene for *Rf1* in sugar beet (*Beta vulgaris*) CMS-Owen, *bvORF20*, encodes a putative peptidase of the M48 family (Hagihara et al. 2005). However, most plant *Rf* genes encode pentatricopeptide repeat (PPR) proteins.
RF-related PPR proteins are commonly targeted to mitochondria and suppress male sterility by controlling the expression of CMS-associated genes at the transcriptional and translational levels. RF1A and RF1B, two mitochondrion-localized proteins in rice CMS-BT (BoroII), restore male fertility by prohibiting ORF79 production via endonucleolytic cleavage (RF1A) or degradation (RF1B) of dicistronic B-\textit{atp6/orf79} mRNA (Wang et al. 2006). In rice CMS-WA (Wild Abortive), RF4 suppresses WA352-mediated male sterility by decreasing WA352 mRNA levels (Tang et al. 2014). The PPR protein RF5 physically interacts with the glycine-rich protein GRP162 to cleave the CMS-associated \textit{atp6-orf79} transcript in rice CMS-HL (Honglian), thus restoring fertility (Hu et al. 2012). Other PPR proteins also act as fertility restorers, including RF-PPR592 in petunia (Bentolila et al. 2002), Rfo in Kosena rapeseed (Giancola et al. 2003), Rfk1 in radish (Koizuka et al. 2003), and PPR13 in sorghum (Koizuka et al. 2003).

PPR proteins are a large group of nucleus-encoded RNA-binding proteins containing a degenerate 35-amino acid motif (Small and Peeters 2000). Most PPRs are predicted to localize to plastids and/or mitochondria (Lurin et al. 2004). Structurally, PPRs are divided into the P and PLS subfamilies. P-subfamily PPR proteins contain canonical PPR motifs (P motifs, 35 amino acids long), whereas PLS-subfamily proteins possess triplets of P, L (35–36 amino acids), and S (31 amino acids) motifs, sometimes with additional C-terminal domains (the E, E+, and DYW domains) (Lurin et al. 2004; Cheng et al. 2016).

PPR proteins regulate gene expression in organelles at the post-transcriptional level, from RNA editing, intron splicing, RNA stability, and cleavage to translation (Lurin et al. 2004). The PPRs in the PLS-subfamily primarily participate in RNA editing (Choury and Araya 2006; Barkan and Small 2014) by changing cytidine to uridine at specific sites. Many PPRs, including CRR4 (Kotera et al. 2005), OTP72 (Chateignier-Boutin et al. 2013), and EMP5 (Liu et al. 2013), have been reported to involve in RNA editing in \textit{Arabidopsis thaliana} and maize. Unlike PLS-subfamily PPR proteins, P-subfamily PPRs have demonstrated roles in intron splicing (Huang et al. 2018; Sun et al. 2019), transcript stabilization (Wang et al. 2017; Zhang et al. 2017), cleavage, and translation (des Francs-Small and Small 2014; Haili et al. 2016).

Because mitochondrial RNA lost the ability to undergo self-splicing, it requires PPR proteins to act as splicing factors (Bonen 2008; Brown et al. 2014). To date, many P-subfamily PPR proteins involved in mitochondrial intron splicing have been reported. The loss-of-function of these PPR genes leads to distinct defective phenotypes, highlighting the important roles of PPR proteins in plant growth and development. For example, \textit{Arabidopsis thaliana} SLOW GROWTH 3 (SLO3) is involved in the splicing of \textit{NADH dehydrogenase subunit7} (\textit{nad7}) intron 2. \textit{slo3} mutants with defects in the splicing of this intron show delayed germination, retarded growth, and delayed development (Hsieh et al. 2015). ORGANELLE TRANSCRIPT PROCESSING 43 (OTP43) is required for the trans-splicing of \textit{nad1} intron 1. An \textit{Arabidopsis} mutant with a disrupted OTP43 gene shows curled leaves and malformed seeds (de Longevialle et al. 2007). In maize, \textit{Defective kernel 35} (\textit{DEK35}) is essential for mitochondrial \textit{nad4} intron 1 splicing. The \textit{dek35} mutant exhibits impaired mitochondrial structure and delayed seed development (Chen et al. 2017). Empty Pericarp8 (\textit{Emp8}) functions in the trans-splicing of \textit{nad1} intron 4 and the cis-splicing of \textit{nad4} intron 1 and \textit{nad2} intron 1. The \textit{emp8} mutation is embryonic lethal, resulting in severely arrested embryo and endosperm development (Sun et al. 2018). Both PPR101 and PPR231 are required for the splicing of mitochondrial \textit{nad5} introns 1 and 2, whereas PPR231 is also required for \textit{nad5} intron 3 and \textit{nad2} intron 3 splicing. Loss-of-function of PPR101 and PPR231 generates empty pericarp and small kernel phenotype, respectively (Yang et al. 2020).

Phosphorylation is a common post-translational modification that plays essential roles in many cellular signaling pathways and networks by altering the structures, localizations, and functions of proteins. The phosphorylation of the transit peptides of nucleus-encoded preproteins regulates the import of proteins into chloroplasts. For example, three plant-specific STY kinases can phosphorylate the transit peptide of the precursor of the small subunit of Rubisco (pSSU) (Lamberti et al. 2011a, b). Phosphorylated pSSU shows more import-competent than the unphosphorylated monomeric form by forming a guidance complex with heat shock protein 70 and 14-3-3 proteins (May and Soll 2000). The phosphorylation of precursors of nucleus-encoded mitochondrial preproteins might also participate in the import process (Law et al. 2015, 2018). The phosphorylation of multiple organelar RNA editing factors 3 (MORF3) is phosphorylated by cytosolic STY kinases. After phosphorylation, the import of pMORF3 into mitochondria is impeded (Law et al. 2015).

Most P-subfamily PPR proteins implicated in intron splicing in the mitochondria reported to date are from \textit{Arabidopsis} and maize. Although some PPRs have been characterized in rice, most of them are required for RNA editing of mitochondrial transcripts or RNA editing and intron splicing in chloroplasts (Toda et al. 2012; Tang et al. 2017; Xiao et al. 2018; Zhang et al. 2020). By contrast, only a few P-subfamily PPR proteins in rice were identified to function in mitochondrial intron splicing (Wu et al. 2019, 2020; Xue et al. 2019).

In this study, we characterized a P-subfamily PPR protein, OsPPR939, which is associated with the development of rice. OsPPR939 acted as a splicing factor required for the splicing of the mitochondrial \textit{nad5} introns 1, 2, and 3. The
loss-of-function of OsPPR939 seriously impaired the activities of the mitochondrial electron transport chain complexes and resulted in reduced plant height and pollen viability. We next demonstrated that the 12th residue Ser in the N-terminal sequence of OsPPR939 was phosphorylated by the kinase OsS6K1, which might be important for the import of OsPPR939 into mitochondria. Our results indicate that OsPPR939 plays crucial roles in the maintenance of mitochondrial function, plant growth, and pollen development and provide insight into the regulation of PPR proteins at the post-translational level.

Materials and methods

Plant materials and growth conditions

The osppr939 mutants were generated using the CRISPR/Cas9 system. The CRISPR/Cas9 plasmids were constructed as described previously (Liu et al. 2017a). The 20-bp guide sequences (Table S1) within the target regions were selected, and their targeting specificity was analyzed using CRISPR-P 2.0 (Liu et al. 2017b). The osppr939 mutants with mutations in MS1, 4, and 5 were in the Nipponbare (Oryza sativa. L. Japanica) background, and the osppr939 mutants with mutations in MS2 and 3 were in the Zhonghua 11 (Oryza sativa. L. Japanica) background. Calli derived from mature seeds were used for Agrobacterium-mediated transformation. Genomic DNA was extracted from the transgenic lines by the cetyltrimethylammonium bromide (CTAB) method. The genomic region surrounding the CRISPR target sites of osppr939 was amplified by PCR using specific primers (Table S2), and the segment was subjected to Sanger sequencing to identify mutants. Rice plants used in this study were grown in the paddy field and greenhouse in Zhejiang University under proper management.

The tang2 mutant (SALK_003139) was a gift from Prof. Catherine Colas des Francs-Small (University of Western Australia). Both wild type and tang2 Arabidopsis thaliana mutants were in the Columbia-0 (Col-0) ecotype background. The plants were grown in soil at 22 °C under a 16-h light/8-h dark photoperiod.

Bioinformatic analysis

The full-length amino acid sequences of OsPPR939 and its 15 homologs were retrieved by BLASTP analysis (http://www.ncbi.nlm.nih.gov/Blast.cgi). Multiple protein sequence alignments were performed using Clustal X. A phylogenetic tree was constructed using MEGA v.6.0 (http://www.megasoftware.net/index.html) based on the neighbor-joining method with the following parameters: p-distance model, pairwise deletion, and bootstrap (1000 replicates; random seed). Bootstrap support values for each node from 1000 replicates are shown next to the branches. The 30 N-terminal amino acids of OsPPR939 and its Clustal X-aligned counterparts from the 15 homologs were used to generate a sequence logo using WebLogo (http://weblogo.berkeley.edu/logo.cgi).

RNA extraction, RT-PCR and qRT-PCR analysis

Total RNA was isolated from rice tissues using RNAiso Plus reagent (TaKaRa), including root, stem, leaf, callus, anther, pistil, palea, and lemma tissue and anther tissue at different stages (determined based on spikelet length). The quality and quantity of total RNA were analyzed using a NanoDrop 1000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA). Approximately 2 µg of RNA per sample was used to synthesize cDNA with a PrimeScript RT reagent Kit with gDNA eraser (TaKaRa). Two microliters of a tenfold dilution of the cDNA solution were used as the template for RT-PCR. qRT-PCR was performed on a LightCycler 96 (Roche Diagnostics) using TaKaRa SYBR Premix Ex Taq II with a standard two-step protocol consisting of 95 °C for 30 s, followed by 40 cycles of 95 °C for 5 s and 60 °C for 30 s. PCRs were performed in three biological replicates, each with three technical replicates. OsGAPDH was used as an internal control. All primers used for (q)RT-PCR are listed in Table S2.

Mitochondrial complex activity and immunoblotting assays

Crude mitochondria were isolated from rice calli as described previously (Cai et al. 2017). Mitochondrial proteins (130 µg) were solubilized in 1% N-dodecyl maltoside β-DM (Sigma, Santa Clara, CA, USA) and separated by 3–12.5% BN-PAGE. The gel was stained with Coomassie Brilliant Blue R-250 or subjected to an in-gel complex I activity assay as described (Meyer et al. 2009). Mitochondrial proteins and total proteins were used for immunoblot analysis. Total proteins were extracted from wild type and osppr939-4 calli as follows: the calli were homogenized in liquid nitrogen using a mortar and pestle. Each 0.1 g sample of homogenized calli was combined with 200 µl extraction buffer (50 mM Tris–HCl, pH 7.5, 150 mM NaCl, 0.5% TritonX-100, protease inhibitor [Roche]) and centrifuged at 12,000 rpm at 4 °C for 10 min. The supernatant was transferred to a new tube, and protein content was determined using an Enhanced BCA Protein Assay Kit (Beyotime, Shanghai, China). One-third volume of 4 × Laemmli buffer (250 mM Tris–HCl, pH 6.8, 8% SDS, 40% glycerol, 4% β-mercaptoethanol, 0.01% bromophenol blue) was mixed with the supernatant. The mixture was denatured at 95–100 °C for 10 min. The mitochondrial proteins (12 µg) or
total proteins (10 µg) were separated by sodium dodecyl sulfate-PAGE (SDS-PAGE), transferred onto a polyvinylidene difluoride (PVDF) membrane (Bio-Rad), and incubated with various primary antibodies against *Arabidopsis* Nad4 and Nad7 (PhytoAB; PHY0511S and PHY0513S, respectively), wheat Nad9, maize Cyt c, ATPase-α subunit, *Arabidopsis* Cox2 (Xiu et al. 2016), or rice OsGAPDH (Beijing Protein Innovation, Beijing, China). Detection was carried out using ECL Western Blotting Detection Reagents (Bio-Rad).

**Measuring ATP content and examining pollen viability**

The ATP content of calli generated from mature rice seeds was measured with a Luciferin-luciferase ATP Assay Kit (Beyotime, Shanghai, China). To analyse pollen viability, mature anthers from wild type and mutant plants were immersed into 1% (w/v) iodine and potassium iodide (I₂–KI) solution at room temperature with three biological replicates. The stained pollen grains were visualized and photographed under a Leica DMIRB fluorescence microscope.

**Scanning electron microscopy analysis**

For SEM analysis, mature anthers collected from wild type (Nipponbare), and osppr939-4 plants were fixed in 2.5% (v/v) glutaraldehyde and dehydrated through an alcohol gradient (70%, 80%, 95%, and 100%). The samples were critical point dried using a Leica EMCPD300 critical point drier (Hitachi TM-1000, Tokyo, Japan) at an accelerating voltage of 15 kV.

**In vitro kinase assay**

For the in vitro kinase assay, different HA-fused kinases were separately expressed in 4 × 10⁵ rice protoplasts and purified as follow: The protoplasts were lysed in 1 mL immunoprecipitation buffer containing 50 mM Tris–HCl pH 8.0, 100 mM NaCl, 10 mM MgCl₂, 5 mM CaCl₂, 0.1 µM ATP, and 6 µCi [γ-³²P] ATP (Perkin-Elmer) at 30 °C for 30 min. The reaction was stopped by adding 15 µL of 6 × SDS-PAGE loading buffer. After separation, the proteins on a 15% SDS-PAGE gel, protein kinase activity was detected on the dried gel using a Typhoon imaging system (GE Healthcare Life Sciences).

**Subcellular localization of OsPPR939**

To verify the subcellular localization of OsPPR939, the 840-bp OsPPR939 gene sequence beginning at the start codon was amplified by PCR from wild type Nipponbare and cloned into pENTR/D-TOPO (Invitrogen, USA). The fusion construct was introduced into the binary vector pgWB5, which contains the cauliflower mosaic virus 35S promoter, by Gateway site-specific recombination (Invitrogen). For stable expression in *Arabidopsis*, the pgWB5-OsPPR939N1−280::GFP construct was transformed into wild type *Arabidopsis* plants by the floral dip method. Protoplasts extracted from transgenic *Arabidopsis* plants expressing OsPPR939N1−280::GFP fusion protein was stained with MitoTracker Red. To further validate the contribution of the phosphorylated N-terminus to the subcellular localization of OsPPR939, we cloned gene fragments expressing OsPPR939N1−280, the truncated derivative OsPPR939N29−280, and three point mutation derivatives (OsPPR939N1−280S12A, OsPPR939N1−280S12T, and OsPPR939N1−280S12R) into the pM999-GFP vector. The primers are listed in Table S2. The plasmids were transformed into rice protoplasts isolated from 10-day-old etiolated Zhonghua 11 seedlings with the mitochondrial marker *F1-ATPase-γ:RFP* (Jin et al. 2003) by polyethylene glycol-mediated transformation. The GFP and RFP signals were detected under a Zeiss LSM780 confocal microscope.

**Complementation analysis**

For functional complementation, wild type (Nipponbare) genomic DNA was used as a template to amplify a 7191-bp genomic DNA fragment of OsPPR939 containing the 4371-bp upstream region and 2820-bp coding region. The amplified fragment was subcloned into the pCAMBIA1300 vector by In-Fusion cloning (TaKaRa Bio, Japan). The resulting plasmid containing OsPPR930pro-OsPPR939 gDNA was transformed into Agrobacterium tumefaciens EHA105 and used for floral dip transformation of tang2 plants and callus transformation of osppr939-4 plants. Transformed plants were selected on hygromycin, and independent transgenic lines were identified by PCR analysis.
RNA in situ hybridization

RNA in situ hybridization was performed as described previously (Liu et al. 2017a). Gene-specific OsPPR939 cDNA fragments generated by PCR were used to prepare antisense and sense probes under the control of the T7 and SP6 promoter with RNA polymerase using a DIG RNA Labeling Kit (Roche). PCR primers are listed in Supplemental Table S2.

Results

Generation of osppr939 mutants by CRISPR/Cas9-mediated genome editing

We previously identified OsPPR939 in a yeast two-hybrid screen in which five proteins including OsPPR939 were confirmed to interact with OsjBTF3 and shown to be involved in the OsjBTF3-regulated pathway controlling plant growth and pollen development in rice (Wang et al. 2012). Here, to investigate the function of OsPPR939, we selected five mutation sites (MS1–MS5) in different regions of this gene to generate osppr939 mutants using the CRISPR/Cas9 system (Fig. 1a). CRISPR/Cas9 constructs expressing guide RNA sequences targeted to the five mutation sites were independently transformed into rice calli (MS1, 4, and 5 in Nipponbare and MS2 and 3 in Zhonghua 11). Sequencing of the DNA covering the mutation sites amplified from T0 transgenic plants confirmed the existence of independent mutations at each mutation site, namely MS1 to MS5 (Table S1).

Surprisingly, all T0 plants with mutations at MS1, MS2, and MS3 were heterozygous, whereas T0 plants with mutations at MS4 and MS5 were either homozygous or heterozygous. Perhaps MS4 and MS5 are located at the C-terminus of OsPPR939 to retain part of its function and thereby ensuring homozygous mutant formation. To generate a complete loss-of-function mutant, we performed segregation analysis using osppr939-1, -2, and -3 plants heterozygous for MS1, 2, and 3, respectively. The progeny of both osppr939-1 and osppr939-2 showed 1:1 segregation of wild type (WT/WT) to heterozygous (WT/−1 bp) plants. The segregation ratio of WT/WT:WT/−1 bp in osppr939-3 was 1:2 instead of 1:1, and no homozygous mutant was isolated from the offspring of these three heterozygotes (Table 1).

We reasoned that perhaps pollen with the osppr939-1 or -2 genotype was nonviable, while pollen with the osppr939-3 genotype survived and completed double fertilization of oocytes, but the resulting homozygous osppr939 zygotes died at an early stage. To confirm our hypothesis, we performed reciprocal crosses using the osppr939-1 heterozygote (WT/−1 bp) as the female parent and the wild type (WT/WT) as the male parent or vice versa. The progeny from the forward cross included 50 WT/WT and 57 WT/−1 bp plants, corresponding to a 1:1 segregation ratio ($\chi^2 = 0.46, P > 0.05$). By contrast, all progeny from the reverse cross were WT/WT (Table 2). These results demonstrate that OsPPR939 is crucial for pollen development in rice and that its mutation results in non-viability of male gametes but does not affect female gamete development.

Phenotypic analysis of osppr939 mutants

Because osppr939-4 and osppr939-5, which were created by CRISPR/Cas9 at the MS4 and MS5 loci, respectively, were homozygous mutants with similar phenotypes, we selected these mutants for subsequent experiments. osppr939-4 contains a frame-shift due to a one base insertion, resulting in the termination of protein translation at the 17th PPR repeat (Fig. S1a, c). Similarly, osppr939-5 contains a frame-shift induced by a two base pair deletion, resulting in the termination of protein translation at the 19th PPR repeat (Fig. S1b, c). When we backcrossed osppr939-4 with the wild type, all F1 progeny displayed the wild type phenotype, and the F2 progeny showed a 3:1 segregation ratio of wild type to mutant plants (98:33, $\chi^2 = 0.0025, P > 0.05$), suggesting that this mutation is monofactorial recessive. Both osppr939-4 and osppr939-5 homozygous plants showed mildly retarded growth at the seedling stage (Fig. S2a, b), significantly reduced plant height, and shorter panicles than the wild type at the heading stage (Fig. 1b, c, Fig. S2c). However, these defects were more severe in osppr939-4 than in osppr939-5. The pollen grains of osppr939-4 and osppr939-5 were partially sterile, as revealed by iodine potassium iodide (I2-KI) staining (Fig. 1d–f, Fig. S2d), although their floral organs developed normally (insets in Fig. 1d–f).

This phenotype was further confirmed by scanning electron microscopy. Unlike the plump and regular wild type pollen grains, approximately half of the pollen grains from osppr939-4 were severely collapsed and shrunken, while the remaining pollen grains resembled those of the wild type (Fig. 1g, h). As a result, many empty grains appeared in the spikes of both mutants (Fig. S3a), and the seed setting rates were ~ 50%, i.e., significantly lower than the 82% seed setting rate of the wild type (Fig. S3b). Reverse-transcription (RT)-PCR analyses revealed that the steady-state levels of OsPPR939 in the mutants were indistinguishable from those of the wild type (Fig. S4), indicating that the deletion or insertion of a few nucleotides did not significantly affect the stability of these transcripts, which is consistent with a previous report (Hsieh et al. 2015).
OsPPR939 encodes a mitochondrion-localized P-subfamily PPR protein

Sequence analysis revealed that full-length OsPPR939 cDNA contains a single exon and encodes a protein of 939 amino acids. Database searches (TPPpred: https://toolkit.tuebingen.mpg.de/tools/tpppred) showed that OsPPR939 contains 22 PPR motifs and no E, E+, or DYW domains at
its C-terminus, indicating that OsPPR939 is a P-subfamily
PPR protein (Fig. 2a).

Most PPR proteins are targeted to mitochondria and/or
plastids (Lurin et al. 2004). Based on analysis using the
TargetP algorithm (http://www.cbs.dtu.dk/services/TargetP/)
and Predotar (https://urgi.versailles.inra.fr/predotar/predo-
tar.html), OsPPR939 was predicted to contain a mitochon-
drion-targeting peptide (MT) at its N-terminus (1–28 a.a.)
(Fig. 2a), suggesting that it localizes to mitochondria. To
experimentally determine the subcellular localization of
OsPPR939, we expressed full-length OsPPR939 fused with
green fluorescence protein (GFP) driven by the 35S pro-
moter in wild type rice. However, we did not observe any
GFP signals from seedling roots in any positive transgenic
line (as identified by PCR). Thus, we fused the 280 N-termi-
nal amino acids containing the predicted mitochondrion-tar-
getting peptide with GFP. We cotransformed rice protoplasts
with the OsPPR939:GFP construct and the mitochon-
drial marker F1-ATPase-γ:RFP (Jin et al. 2003). Both GFP

| Transgenic $T_1$ lines | Genotype | Segregation of $T_2$ plants | Segregation ratio WT/WT:WT/−1 bp |
|------------------------|----------|-----------------------------|---------------------------------|
| osppr939-1             | WT/−1 bp | 79 78 0                     | 1:1                             |
| osppr939-2             | WT/−1 bp | 68 60 0                     | 1:1                             |
| osppr939-3             | WT/−1 bp | 49 92 0                     | 1:2                             |

Table 2 Reciprocal crosses between osppr939-1 heterozygous
mutants and the wild type

| Parent | Progeny |
|--------|---------|
| Female (♀) | Male (♂) |
| osppr939-1 (WT/−1 bp) | Wild type (WT/WT) |
| Wild type (WT/WT) | osppr939-1 (WT/−1 bp) |

79 78 0 1:1 0.06 0.94
68 60 0 1:1 0.50 0.48
49 92 0 1:2 0.13 0.72

Fig. 2 Schematic of OsPPR939 and subcellular localization of
GFP-tagged OsPPR939N1−280.

a OsPPR939 protein contains 22 PPR motifs and a mito-
condrion-targeting peptide at the N-terminus.
b Subcel-
uar localization analysis of OsPPR939N1−280:GFP by
coeexpression with F1-ATPase-γ:RFP (mitochon-
drial marker) in rice protoplasts.
DIC, differential interfer-
ce contrast. Bars = 5 μm.
c Subcellular localization of OsPPR939N1−280:GFP in
the roots of stable transgenic Arabidopsis. Mitochondria were
stained with MitoTracker Red.
DIC, differential interference contrast. Bars = 20 μm.
and RFP signals were detected by confocal microscopy. The green fluorescence colocalized with the red fluorescence (Fig. 2b). We also introduced OsPPR939N1−280:GFP into wild type Arabidopsis Columbia-0 (Col-0) plants via the floral dip method and stained roots, leaves, and protoplasts from these plants with MitoTracker Red to label their mitochondria. In all tissues examined, the GFP signals of OsPPR939 N1−280:GFP overlapped with the RFP signals from MitoTracker Red (Fig. 2c, Fig. S5). These results suggest that OsPPR939 localizes to the mitochondria in rice.

**Phylogenetic analysis of OsPPR939**

To explore the evolution of OsPPR939, we performed a BLASTP search of the National Center for Biotechnology Information database (NCBI, http://www.ncbi.nlm.nih.gov/) using the full-length OsPPR939 sequence as a query. Homologs of OsPPR939 were only detected in angiosperm species, and no paralog of OsPPR939 is present in rice. Phylogenetic analysis of the 15 homologs retrieved from 15 different species, and the full-length OsPPR939 clearly divided the 16 proteins into two clusters (Fig. S6a). In the monocotyledon cluster, OsPPR939 shares sequence identities of 65.39%, 71.40%, 73.97%, 69.35%, 68.39%, and 70.82% with its orthologs in Oryza brachyantha, Aegilops tauschii, Setaria italica, Zea mays, and Sorghum bicolor, respectively, although the functions of these proteins are largely unknown. Among the dicotyledon species, Arabidopsis TANG2 (annotated as NP_173362.2) is the most closely related homolog; this protein is essential for mitochondrial intron splicing (des Francs-Small et al. 2014). Although TANG2 only shares 37.24% amino acid sequence identity with OsPPR939, the locations of most PPR repeats are highly conserved in these two proteins, except for the last two repeats, which only exist in OsPPR939 (Fig. S6b). Together, these results indicate that OsPPR939 is a typical P-subfamily PPR protein and may also be involved in mitochondrial intron splicing in rice.

**Expression analysis of OsPPR939**

We examined the expression pattern of OsPPR939 by quantitative real-time (qRT)-PCR using total RNA extracted from vegetative and reproductive organs of wild type plants. OsPPR939 was ubiquitously expressed in all tissues examined, with peak expression in calli and a relatively high expression level in spikelets at stage 6 (S6), which gradually decreased during later stages of development. OsPPR939 was also highly expressed in anther and pistil tissue and relatively weakly expressed in root, stem, leaf, palea, and lemma tissue (Fig. 1i).

To investigate the spatial and temporal expression patterns of OsPPR939 during pollen development, we performed RNA in situ hybridization using wild type anther sections. Strong hybridization signals were detected in both the tapetum and male meiocytes at stages 6, 8b, and 10, while only background levels of signal were detected with the sense probe (Fig. 1j). Since the tapetum plays a crucial role in regulating pollen formation, these data suggest that OsPPR939 is involved in pollen development.

**OsPPR939 functions in mitochondrial nad5 intron splicing**

PPR proteins are RNA-binding proteins that function in RNA metabolism, including RNA splicing (Barkan and Small 2014). Many PPR proteins, especially members of the P-subfamily, participate in organellar RNA intron splicing. To verify whether this was the case for OsPPR939, we measured the levels of 45 mitochondrial-encoded transcripts in wild type and osppr939 plants by RT-PCR analysis. We extracted total RNA from wild type and mutant leaves and performed RT-PCR using specific primers designed to amplify the transcript of each mitochondrial gene (Table S2). *nad5* transcript levels were dramatically reduced in the mutants, whereas the levels of most transcripts were indistinguishable between the wild type and osppr939 (Fig. 3a). These results suggest that OsPPR939 might be essential for the splicing or stability of *nad5* transcript.

The mitochondrial gene *nad5* encodes a subunit of the mitochondrial respiratory complex I (NADH:ubiquinone oxidoreductase) (Meyer 2012). Rice *nad5* consists of five coding segments, which are located in a wide region of the mitochondrial genome (Notsu et al. 2002). The formation of the mature *nad5* transcript requires two *cis*-splicing events of intron 1 and 4 as well as two *trans*-splicing events of introns 2 and 3 (Fig. 3b) (Knoop et al. 1991). To examine whether the weak accumulation of mature *nad5* transcript in the *osppr939* mutants was caused by a deficiency in intron splicing, we examined the splicing of *nad5* precursor transcript. Four specific primer pairs were designed to amplify fragments containing each of the four introns in the *nad5* transcript (Fig. 3b). In both osppr939-4 and osppr939-5, the levels of the spliced fragments joining exon 1 and exon 2 were much lower, and the levels of the unspliced fragments containing *nad5* intron 1 were higher than those in the wild type when we used the primer pair F1 + R1 for amplification (Fig. 3c). The levels of intron 2 and intron 3 spliced fragments, which were amplified using primer pairs F2 + R2 and F3 + R3, respectively, were also lower in the mutants than the wild type. However, intron 2 and intron 3 unspliced transcripts were not detected in wild type or mutant plants due to the failure to perform PCR amplification of such long fragments containing both intron 2 and intron 3 (Fig. 3c). However, there was no difference in the abundance of intron 4-spliced and intron 4-unsplliced fragments between wild
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The osppr939 mutants show reduced mature mitochondrial nad5 transcript levels and deficient splicing of nad5 introns 1, 2, and 3. a RT-PCR analysis of the transcript levels of 45 mitochondria-encoded genes in the wild type (WT; left lane), osppr939-4 (middle lane), and osppr939-5 (right lane). Total RNA was isolated from the leaves of plants at the flowering stage. OsGAPDH was used as an internal control. b Structure of the rice mitochondrial nad5 gene. Introns 1 and 4 are cis-spliced introns and introns 2 and 3 are trans-spliced introns. The expected amplification products of different primer pairs are indicated. c RT-PCR analysis of the intron splicing efficiency of nad5 in wild type and osppr939 plants. Solid arrows correspond to unspliced fragments. d qRT-PCR analysis of the splicing efficiency of all 23 group II introns in osppr939-4, osppr939-5, and wild type plants. Error bars indicate SEs based on three biological replicates.

Fig. 3 The osppr939 mutants show reduced mature mitochondrial nad5 transcript levels and deficient splicing of nad5 introns 1, 2, and 3. a RT-PCR analysis of the transcript levels of 45 mitochondria-encoded genes in the wild type (WT; left lane), osppr939-4 (middle lane), and osppr939-5 (right lane). Total RNA was isolated from the leaves of plants at the flowering stage. OsGAPDH was used as an internal control. b Structure of the rice mitochondrial nad5 gene. Introns 1 and 4 are cis-spliced introns and introns 2 and 3 are trans-spliced introns. The expected amplification products of different primer pairs are indicated. c RT-PCR analysis of the intron splicing efficiency of nad5 in wild type and osppr939 plants. Solid arrows correspond to unspliced fragments. d qRT-PCR analysis of the splicing efficiency of all 23 group II introns in osppr939-4, osppr939-5, and wild type plants. Error bars indicate SEs based on three biological replicates.

To further investigate the changes in splicing in the osppr939 mutants, we designed specific primers for qRT-PCR to detect the splicing efficiency of 23 mitochondrial introns in both osppr939-4 and osppr939-5. Primers across adjacent exons and across adjacent exons and introns were used to amplify the intron-spliced fragments and unspliced fragments, respectively. Splicing efficiency was calculated as the ratio of spliced to unspliced forms of each transcript in the mutants normalized to the same ratio in the wild type. The splicing efficiency of nad5 introns 1, 2 and 3 was dramatically reduced in both mutants (Fig. 3d). Importantly, the significantly reduced splicing efficiency resulted from a reduction in nad5 spliced forms (exon-exon) and an increase in nad5 unspliced forms (exon-intron) (Fig. S9). However, other mitochondrial introns were spliced normally in osppr939 (Fig. 3a, d). These results demonstrate that OsPPR939 participates in the cis-splicing of nad5 intron 1 and the trans-splicing of nad5 introns 2 and 3.

Although most PPR proteins involved in RNA editing belong to the PLS-subfamily, P-subfamily PPR proteins also play roles in RNA editing (Doniwa et al. 2010), suggesting that OsPPR939 might also function in the editing of mitochondrial transcripts. To investigate this possibility, we used RT-PCR and bulk sequencing of the amplified cDNAs to compare the editing efficiencies of mitochondrial transcripts in osppr939-4 versus the wild type. Of the 491 mitochondrial RNA editing sites examined (Notsu et al. 2002), no significant difference in editing efficiency was observed.
between osppr939-4 and the wild type. Hence, OsPPR939 is not essential for the editing of these RNAs in rice.

**Complementation of the osppr939-4 mutant**

To confirm that the mutant phenotypes and defective intron splicing were due to the lack of functional OsPPR939, we performed a complementation assay of osppr939-4. We amplified the native promoter along with the full-length coding sequence of OsPPR939 from genomic DNA, inserted it into the pCAMBIA1300 vector, and introduced the construct into the osppr939-4 mutant. In all seven independent positive transgenic lines, the osppr939-4 phenotypes of reduced plant height and low pollen viability were successfully rescued (Fig. 4a, images 1 and 2, Fig. S8a, b). The reduced splicing efficiency of introns 1, 2, and 3 resulted in the weak accumulation of nad5 transcript (Fig. 3a, c). To investigate whether the splicing of these introns was recovered in the transgenic complementation lines, we performed RT-PCR to detect the mature nad5 transcript using primer pair F1 and R4 (Fig. 3b). Complementation with OsPPR939pro:OsPPR939 completely restored the intron splicing efficiency of the mutant to wild type levels (Fig. 4a, image 3). These results confirm that the defective intron splicing in the osppr939 mutants is due to impaired OsPPR939 function, and that the abnormal growth is indeed caused by defects in intron splicing.

Tang2, the Arabidopsis ortholog of OsPPR939, is involved in the trans-splicing of nad5 intron 3. Mutants of Tang2 display retarded growth compared to the wild type. To investigate whether OsPPR939 could rescue the defective plant morphology of tang2, we transformed this mutant with the OsPPR939pro:OsPPR939 construct for functional complementation. We performed RT-PCR using the primer pair Aranad5-F and Aranad5-R, located in exon 2 and 4, respectively (Fig. 4b, image 1), to examine the splicing efficiency of nad5 intron 3. The results showed that intron 3-spliced transcripts were restored to wild type levels in the complemented lines (Fig. 4b, image 2), and the mutant phenotype of tang2 was fully rescued (Fig. 4b, image 3). These data suggest that the functions of OsPPR939 and TANG2 are partially conserved.

**Mitochondrial complex I abundance and activity are reduced in osppr939-4**

The substantially reduced nad5 gene expression in osppr939 mitochondria may result in the weak accumulation of Nad5
protein. Nad5 is a central subunit of mitochondrial complex I, one of the five complexes in the OXPHOS electron transport chain. To investigate the effect of the intron splicing deficiency of nad5 on mitochondrial complex I, we examined complex I abundance by Blue Native Polyacrylamide Gel Electrophoresis (BN-PAGE) and complex I activity with an in-gel NADH dehydrogenase activity assay. We isolated crude mitochondria from osppr939-4 and wild type calli and examined mitochondrial respiratory complex abundance by BN-PAGE. As indicated by Coomassie Brilliant Blue (CBB) staining, the abundance of complex I was reduced in osppr939-4 compared to the wild type. The levels of both complex III and complex V were slightly higher in osppr939-4 than in the wild type (Fig. 5a), perhaps due to responsible regulation, as described previously (Xiu et al. 2016). An in-gel NADH dehydrogenase activity assay confirmed that complex I activity was indeed dramatically reduced in osppr939-4 (Fig. 5b).

We then measured the expression levels of several mitochondrial complex subunits by immunoblot analysis using total proteins extracted from calli. No difference was detected between wild type and osppr939-4 proteins using antibodies against Nad4 and Nad9 (subunits of complex I), whereas the level of Nad7 (a subunit of complex I) was slightly reduced in the mutant. The levels of Cytc1 (a subunit of complex III), Cox2 (a subunit of complex IV), and ATPase-α (a subunit of complex V) were higher in osppr939-4 than in the wild type (Fig. 5c). Similar results were obtained using purified mitochondria for immunoblot analysis (Fig. S10).

Since ATP production in the mitochondria is coupled to electron transfer through complex I, we measured ATP levels in the calli. The ATP levels in osppr939-4 and osppr939-5 were only 55.9 and 65.6% that of the wild type, respectively (Fig. 6a). These results suggest that the lower level of mature nad5 in the osppr939 mutants due to the reduced splicing efficiency of nad5 introns 1–3 indeed leads to reduced levels of complex I, which might stimulate the accumulation of other mitochondrial complexes via unknown retrograde signaling.

Fig. 5 Complex I abundance and activity are reduced in the osppr939-4 mutant. a BN gels stained with CBB. Mitochondrial complexes were isolated from rice calli and subjected to a 3% to 12% BN-PAGE. WT, wild type. b In-gel NADH dehydrogenase activity assay of mitochondrial complex I performed using dihydrolipoamide dehydrogenase. DLDH was used as a loading control. The positions of mitochondrial complexes are indicated. C I, complex I. c Total proteins extracted from wild type and osppr939-4 calli were used for immunoblot analysis with antibodies against Nad4, Nad7, and Nad9 of complex I, Cytc1 of complex III, Cox2 of complex IV, and ATPase-α of complex V. All blots were loaded in the same order. OsGAPDH and CBB staining are shown as a sample loading control. The molecular weights of the protein markers are indicated on the left.

Accumulation of alternative respiratory pathway transcripts in osppr939

A previously described splicing defect of nad transcripts led to partial or complete loss of complex I activity and
thus increased levels of alternative oxidases (AOX) (de Longevialle et al. 2007; Xiu et al. 2016). To investigate whether the nad5 splicing defect had similar effects in the osppr939 mutants, we performed RT-PCR to measure the expression levels of AOX genes, including AOX1a, AOX1b, and AOX1c, in wild type (WT), osppr939-4, and osppr939-5 plants. The numbers of PCR cycles are indicated on the left. The expression levels were normalized against OsGAPDH.

![Fig. 6 ATP contents and OsAOX1a, OsAOX1b, and OsAOX1c transcript levels in the osppr939 mutants. a ATP contents of calli generated from mature seeds. Error bars indicate the SE based on three biological replicates. b RT-PCR analysis of OsAOX1a (AB004864.1), OsAOX1b (AB004865.1), and OsAOX1c (AB074005.1) transcripts in wild type (WT), osppr939-4, and osppr939-5 plants. The numbers of PCR cycles are indicated on the left. The expression levels were normalized against OsGAPDH. c qRT-PCR analysis of OsAOX1a, OsAOX1b, and OsAOX1c transcripts. Error bars indicate SEs based on three biological replicates.](image)

OsS6K1 phosphorylates the N-terminal sequence of OsPPR939

The phosphorylation of the N-terminal targeting peptide of nucleus-encoded mitochondrial proteins regulates the import of these proteins (Law et al. 2018). To examine whether the presequence of OsPPR939 contains a phosphorylation site, we used the web server PPSP (http://ppsp.biocuckoo.org/) to examine the first 28 amino acid residues of this protein. Three residues in this region were predicted to be phosphorylation sites: S12, S22, and S23. To determine whether these potential phosphorylation sites are conserved among the 15 homologous proteins of OsPPR939, we analyzed the N-terminal sequences of the homologs using the WebLogo program (Crooks et al. 2004) to create a sequence logo. As shown in Fig. 7a, the predicted phosphorylation site S12 was highly conserved among the homologous proteins in the plant species examined.

To experimentally validate the predicted phosphorylation sites, we expressed the N-terminal targeting peptide (OsPPR939N1−28) and the mutated peptide with the 12th residue changed from Ser-to-Ala (OsPPR939N1−28S12A) in E. coli, followed by purification (Fig. 7b). We then examined the abilities of several kinases to phosphorylate the S12 site. Among the kinases tested, only the kinase OsS6K1 phosphorylated the targeting peptide (Fig. S12), and OsS6K1 only phosphorylated the peptide OsPPR939N1−28, and not the mutated form (Fig. 7c).

To evaluate the importance of the first 28 residues and the phosphorylation of S12 for the mitochondrial localization of OsPPR939, we expressed two forms of GFP fusion protein, OsPPR939N1−28GFP and OsPPR939N1−28S12A:GFP, in rice protoplasts with the mitochondrial reporter F1-ATPase-γ:RFP. As shown in Fig. 7d and e, most GFP signals did not overlap with RFP signals in the mitochondria, indicating that deleting the first 28 amino acid residues or the lack of phosphorylation at S12 seriously impeded the import of OsPPR939 into mitochondria.

To confirm that the impaired mitochondrial localization of OsPPR939N1−28S12A was not due to the Ser-to-Ala substitution, we examined the phosphorylation of the mutated peptides OsPPR939N1−28S12T and OsPPR939N1−28S12R, with the 12th residue Ser replaced by Thr or Arg, respectively (Fig. S12a). OsPPR939N1−28S12T but not OsPPR939N1−28S12R was phosphorylated by OsS6K1 (Fig. S13a). Finally, we detected the subcellular localization of the corresponding GFP fusion proteins in rice protoplasts. The GFP signals from both OsPPR939N1−28S12T:GFP and OsPPR939N1−28S12R:GFP colocalized with RFP signals from mitochondrial markers (Fig. S13C and D), indicating that the presence of a phosphorylated or positively charged residue at the N-terminus is essential for the mitochondrial localization of OsPPR939. Together, these results suggest
that phosphorylation of the S12 in N-terminal targeting sequence of OsPPR939 is important for its mitochondrial localization.

**Discussion**

Introns are rather common in the mitochondrial genomes of land plants, with 22 in the maize mitochondrial genome and 23 in *Arabidopsis* and rice (Bonén 2008). Because plant mitochondrial introns cannot undergo self-splicing, they require the help of nucleus-encoded splicing cofactors, which interact with intron RNA to catalyze the formation of its active conformation (Brown et al. 2014). So far, most PPR proteins acted as splicing factors in mitochondria have been identified in *Arabidopsis* and maize. In this study, we demonstrated that the P-subfamily PPR protein OsPPR939 is required for cis-splicing of nad5 intron 1 and trans-splicing of nad5 introns 2 and 3 in rice mitochondria. Two *osppr939* mutants (*osppr939-4* and *osppr939-5*) showed identical defects in the splicing of *nad5* introns, whereas other mitochondrial introns were correctly spliced in the mutants (Fig. 3), suggesting that a single *osppr939* mutation impaired the splicing of three *nad5* introns. By contrast, TANG2, the homolog of OsPPR939 in *Arabidopsis*, is specifically involved in *trans*-splicing of *nad5* intron 3 (des Francs-Small et al. 2014). Our findings indicate that the molecular functions of these two proteins are only partially conserved.

In *Arabidopsis*, the splicing of *nad5* intron 3 is impaired in *tang2* mutants, six-week-old mutant plants exhibit dark curled foliage when grown under a 16-h light/8-h dark photoperiod (des Francs-Small et al. 2014). In rice *osppr939* mutants, the splicing of *nad5* intron 1, 2, and 3 are defective, which results in far more serious phenotypes (Fig. 1b–h, Tables 1 and 2). The finding that mutations at different sites of a single gene cause mutant phenotypes of different severities has been widely documented (Cushing et al. 2005; Huang et al. 2018). One possible explanation is that mutations at different sites of a coding sequence impair gene function to different extents, with some mutations totally abolishing gene expression and others resulting in only a partial reduction in gene function, leading to very different morphological consequences. In this study, to explore the biological function of *OsPPR939* in “ideal” homozygous mutants (with severe but viable phenotypes), we targeted five sites for CRISPR/Cas9-mediated mutation. A mutation at the 5′-terminus (MS1) or middle regions (MS2 and MS3) of *OsPPR939* yielded no homozygous mutants, but homozygous mutants were obtained when a mutation occurred at the 3′-terminus (MS4 or MS5) of *OsPPR939* (Table 1). Genetic experiments demonstrated that pollen grains with the *osppr939* genotype from *osppr939-1* and *osppr939-2* heterozygous mutants were nonviable (Table 2). However, pollen grains with a mutated *OsPPR939* gene from *osppr939-3* could complete the process of fertilization, but the resulting zygotes failed to develop. By contrast, homozygous mutants *osppr939-4* and *osppr939-5* showed weaker compromised phenotypes, with only partially sterile pollen and reduced plant height (Fig. 1b–h, Fig. S2). These results imply that it is possible to obtain completely male sterile rice plants by knocking out *OsPPR939* to some extent.

Amino acids at specific positions in each PPR motif ensure that a given PPR protein can bind to single-stranded RNA in a sequence-specific fashion (Yin et al. 2013). Perhaps the severely defective phenotypes of the mutants could be explained by the following example. The *osppr939-1* mutant produces a truncated OsPPR939 protein with only three intact PPR motifs, which is not sufficient to allow the protein to recognize and splice the RNA sequence of *nad5*. As a result, no mature *nad5* RNA accumulates, leading to a complete loss of complex I activity, thereby resulting in pollen death. By contrast, *osppr939-4* and *osppr939-5* produce truncated OsPPR939 proteins with 16 and 18 motifs, respectively, which have retained partial functions for *nad5* intron splicing, resulting in reduced complex I abundance and partial pollen sterility. Overall, these findings indicate that *OsPPR939* plays crucial roles in plant growth and pollen development in rice, and that *osppr939-4* and *osppr939-5* are two weak alleles among the mutants created in this study.

*Nad5* is one of nine Nad subunits of respiratory complex I (NADH:ubiquinone oxidoreductase) encoded by the mitochondrial genome (Meyer 2012). Before these *nad* mRNAs are translated into functional proteins, several post-transcriptional processes are required, including RNA editing, intron splicing, and RNA stability. Defects in any of these processes impair the assembly and activity of complex I. Several PPR proteins that participate in the intron splicing of *nad* mRNAs have been reported in *Arabidopsis* (de Longevialle et al. 2007; Liu et al. 2010; des Francs-Small et al. 2014; Hsieh et al. 2015; Haili et al. 2016) and maize (Xiu et al. 2016; Cai et al. 2017; Chen et al. 2017; Sun et al. 2018). The splicing defect of certain *nad* mRNAs leads to the partial or complete loss of complex I assembly and activity. In the current study, we determined that *OsPPR939* encodes a P-subfamily PPR protein that is required for the splicing of *nad5* introns 1, 2, and 3. The deficient splicing of these three introns in *osppr939* led to a complete or partial reduction in the levels of processed *nad5* transcript (Fig. 3a, c, d). As a result, mitochondrial complex I could not be assembled effectively due to lack or deficiency of the Nad5 subunit, and its activity was greatly weakened (Fig. 5a, b). This abnormality eventually resulted in growth retardation in the *osppr939* mutants, along with pollen death or partial sterility (Fig. 1b–f and Fig. S2, Tables 1 and 2). Likewise, the splicing defect of *nad5* intron 3 in *tang2* significantly reduces
the amount of fully processed \textit{nad5} transcript, resulting in severely reduced levels of complex I (des Francs-Small et al. 2014).

In mitochondria, complex I connects ATP synthesis with electron transfer (Millar et al. 2011). Defects in complex I block electron transfer, leading to a deficiency in oxidized NAD$^+$, ultimately resulting in a loss in ATP production and changes in cellular metabolism. However, these abnormal signals can be also perceived by the alternative respiratory pathway, which is characterized by the expression of
OsPPR939 is crucial for intron splicing of \textit{nad5}, which encodes a core subunit of electron transport chain complex I. In mitochondria, a majority of ATP production is coupled to normal electron transfer. Therefore, the correct translocation of OsPPR939 to mitochondria is important for energy homeostasis. In the current study, we demonstrated that abolishing the phosphorylation of the 12th residue on the N-terminal sequence of OsPPR939 by inducing a Ser-to-Ala conversion largely impeded the localization of OsPPR939 to mitochondria (Fig. 7d). The deletion of the first 28 amino acids of OsPPR939 (Fig. 7e) also impaired the mitochondrial localization of OsPPR939, whereas the Ser-to-Thr-mutation did not (Fig. S13c). These results suggest that the phosphorylation of the N-terminal sequence of OsPPR939 is important for its mitochondrial localization (Fig. 7f).

Surprisingly, the truncated OsPPR939 protein with the 12th residue Ser mutated to a non-phosphorylatable Arg localized to the mitochondria (Fig. S13D), suggesting that the mitochondria import process of OsPPR939 is associated with a different mechanism. Indeed, the phosphorylation of transit peptides of some nucleus-encoded mitochondrial or chloroplast preproteins by specific kinases affects the efficiency of their import into intracellular destinations (Law et al. 2015; Lamberti et al. 2011a, b). In addition, mitochondrial targeting peptides are enriched in positively charged residues (Arg in particular). These residues are conducive to the formation of the positively charged face of a protein and provide net positive charges, which are important for the correct translocation of mitochondrial proteins (Emanuelsson and von Heijne 2001; Neupert and Herrmann 2007). Thus, perhaps the phosphorylation of the targeting peptide of OsPPR939 promotes its interaction with auxiliary factors to form a guidance complex for mitochondrial localization. Indeed, Arg residues provide preproteins with a more positive charge, which is important for the mitochondrial localization of many nucleus-encoded proteins. However, more precise assays, such as an in vitro mitochondria import assay, are needed to test this hypothesis.

Much is known about the role of PPR proteins in regulating the expression of organelle genes at the post-transcriptional level, but little is known about the post-translational regulation of PPR proteins. To our knowledge, the 12th residue Ser on OsPPR939 is the first experimentally proven phosphorylation site in PPR proteins. This finding provides not only a new insight into investigating how PPR proteins are regulated at post-translational level but also a theoretical basis for chemical emasculation and heterosis utilization. Torin-1 is a specific inhibitor of TOR that can modulate the activity of S6K1 (Ma and Blenis 2009; Thoreen et al. 2009; Schepetilnikov et al. 2011). Therefore, torin-1 can be used to inhibit the phosphorylation activity of TOR, which consequently impedes the phosphorylation of OsPPR939 by blocking the activity of OsS6K1 and makes OsPPR939

alternative NADH dehydrogenases and alternative oxidases (AOX) (Kühn et al. 2015). Several mutants with dysfunctional complex I that accumulate alternative NADH dehydrogenase and AOX transcripts have been reported (Keren et al. 2012; Xiu et al. 2016; Cai et al. 2017; Qi et al. 2017; Ren et al. 2017; Zhang et al. 2017; Sun et al. 2018; Sun et al. 2019). Consistent with these findings, we detected a significant reduction in ATP content in the \textit{osppr939} mutants (Fig. 6a). Moreover, AOX1a, NDA2, and NDB3 transcript levels were slightly higher in the \textit{osppr939} mutants than the wild type (Fig. 6b, c and Fig. S10), indicating that the alternative respiratory pathway was activated in the mutants. Taken together, these data indicate that the defective phenotypes observed in \textit{osppr939} plants were due to the reduced abundance of complex I.

Chloroplasts and mitochondria supply energy to living cells in plants. The biogenesis and functioning of these two organelles requires the involvement of many nucleus-encoded proteins. The N-terminal sequences of most nucleus-encoded preproteins determine their correct intracellular destinations (Chotewutmontri et al. 2012; Schleiff and Becker 2011). The phosphorylation of transit peptides of chloroplast preproteins and the presequences of mitochondrial proteins functions in protein import between the cytosol and organelles (Law et al. 2015; Nickel et al. 2015). In the current study, we demonstrated that the 12th amino acid residue (Ser) on the N-terminal targeting peptide of OsPPR939 is phosphorylated by OsS6K1 (Fig. 7c), a member of the AGC family (PKA, PKG, and PKC) of Ser/Thr kinases (Jacinto and Lorberg 2008). The conserved kinase S6K1 is a major substrate and crucial effector of TOR (target of rapamycin) kinases (Wullschleger et al. 2006). In animals, “mTORC-S6K1” signaling controls fundamental cellular process by integrating multiple signals such as nutrient, stress, and energy signals (Magnuson et al. 2012). However, the regulatory mechanism of S6K1 in plants is poorly understood.
unable to correctly locate in mitochondria, thus leading to rice pollen sterility. In this way, we can easily obtain chemical induced sterile lines for hybrid production. In practical application, the advantage of chemical emasculation is that the preferred parental combination is not restricted by the relationship between male sterile line, restorer line and maintainer line, and male sterile line does not need to be selected by special program. But at present, this is only a theoretical assumption, and further research needs to be carried out on a large scale.

In conclusion, our results suggest that nucleus-encoded OsPPR939 controls the assembly of mitochondrial complex I and energy generation by participating in splicing of mitochondrial nad5 introns 1, 2, and 3, thus playing an important regulatory role in rice plant growth and pollen development. In addition, the 12th amino acid in the N-terminal targeting sequence of OsPPR939 protein can be specifically phosphorylated by rice kinase OsS6K1, and phosphorylation of this site may plays an important role in the mitochondrial localization of OsPPR939. These results provide a theoretical basis for the creation of chemically induced male sterile line, and also point out a direction for investigation the regulatory mechanism of PPR proteins, especially those Rf-related PPRs, at the post-translational level.

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Author contributions statement JMT and YJL designed the experiments; PZ, HD, and YJL performed most of the experiments and analyzed the data; XJL and YQH provided assistance in western blot and subcellular localization analyses; WYW, HC, MJ, YY, and CCZ performed the field experiments; FS supervised and helped to perform the BN gel analysis; PZ and YJL wrote the manuscript; BCT, HD, and JMT revised the manuscript. All authors read and approved the final manuscript.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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References

Barkan A, Small I (2014) Pentatricopeptide repeat proteins in plants. Annu Rev Plant Biol 65:415–442
Bentolila S, Alfonso AA, Hanson MR (2002) A pentatricopeptide repeat-containing gene restores fertility to cytoplasmic male-sterile plants. Proc Natl Acad Sci USA 99:10887–10892
Bonen L (2008) cis- and trans-splicing of group II introns in plant mitochondria. Mitochondrion 8:26–34
Brown GG, des Francs-Small CC, Ostersetzer-Biran O (2014) Group II intron splicing factors in plant mitochondria. Front Plant Sci 5:35
Cai MJ, Li SZ, Sun F, Sun Q, Zhao HL, Ren XM, Zhao YX, Tan BC, Zhang ZX, Qiu FZ (2017) Emp10 encodes a mitochondrial PPR protein that affects the cis-splicing of nad2 intron 1 and seed development in maize. Plant J 91:132–144
Chateignier-Boutin A, des Francs-Small CC, Fujii S, Okuda K, Tanz S, Small I (2013) The E domains of pentatricopeptide repeat proteins from different organelles are not functionally equivalent for RNA editing. Plant J 74:935–945
Chen XZ, Feng F, Qi WW, Xu LM, Yao DS, Wang Q, Song RT (2017) Dek35 Encodes a PPR Protein that affects cis-splicing of mitochondrial nad4 intron 1 and seed development in maize. Mol Plant 10:427–441
Cheng SF, Gutmann B, Zhong X, Ye YT, Fisher MF, Bai FQ, Castleden I, Song Y, Song B, Huang JY, Liu X, Xu X, Lim BL, Bond CS, Yiu SM, Small I (2016) Redefining the structural motifs that determine RNA binding and RNA editing by pentatricopeptide repeat proteins in land plants. Plant J 85:532–547
Chotewutmontri P, Reddick LE, McWilliams DR, Campbell IM, Bruce BD (2012) Differential transit peptide recognition during preprotein binding and translocation into flowering plant plastids. Plant Cell 24:3040–3059
Choury D, Araya A (2006) RNA editing site recognition in heterologous plant mitochondria. Curr Genet 50:405–416
Crooks GE, Hon G, Chandonia JM, Brenner SE (2004) WebLogo: a sequence logo generator. Genome Res 14:1188–1190
Cui X, Wise RP, Schnable PS (1996) The r2 nuclear restorer gene of male-sterile T-cytoplasm maize. Science 272:1334–1336
Cushing DA, Forsthoefel NR, Gestaut DR, Vernon DM (2005) Arabidopsis emb175 and other ppr knockout mutants reveal essential roles for pentatricopeptide repeat (PPR) proteins in plant embryogenesis. Planta 221:424–436
de Longevialle AF, Meyer EH, Andres C, Taylor NL, Linur C, Millar AH, Small ID (2007) The pentatricopeptide repeat gene OTP43 is required for trans-splicing of the mitochondrial nad1 intron 1 in Arabidopsis thaliana. Plant Cell 19:3256–3265
des Francs-Small CC, Small I (2014) Surrogate mutants for studying mitochondrially encoded functions. Biochimie 100:234–242
des Francs-Small CC, de Longevialle AF, Li YH, Lowe E, Tanz SK, Smith C, Bevan MW, Small I (2014) The pentatricopeptide repeat proteins TANG2 and organellar transcript processsiNG439 are involved in the splicing of the multipartite nad5 transcript encoding a subunit of mitochondrial complex I. Plant Physiol 165:1409–1416
Doniwa Y, Ueda M, Ueta M, Wada A, Kadowaki K, Tsutsumi N (2010) The involvement of a PPR protein of the P subfamily in partial RNA editing of an Arabidopsis mitochondrial transcript. Gene 454:39–46
Emanuelsson O, von Heijne G (2001) Prediction of organellar targeting signals. Biochim Biophys Acta 1541:114–119
Fujii S, Toriyama K (2009) Suppressed expression of retrograde-regulated male sterility restores pollen fertility in cytoplasmic male sterile rice plants. Proc Natl Acad Sci USA 106:9513–9518

Giancola S, Marhadrour S, Desloire S, Clouet V, Falentin-Guyomarc’h H, Lalou W, Falentin C, Pelletier G, Renard M, Bendahmane A, Delourme R, Budar F (2003) Characterization of a radish introgression carrying the Oguira fertility restorer gene Rfo in rapeseed, using the Arabidopsis genome sequence and radish genetic mapping. Theor Appl Genet 107:1442–1451

Hagihara E, Ichoda N, Habu Y, Iida S, Miikami T, Kubo T (2005) Molecular mapping of a fertility restorer gene for Owen cytoplasmic male sterility in sugar beet. Theor Appl Genet 111:250–255

Haili N, Planchard N, Arnal N, Quadrado M, Vrielynck N, Dahan J, des Fujii S, Toriyama K (2009) Suppressed expression of retrograde-regulated male sterility restores pollen fertility in cytoplasmic male sterile rice plants. Proc Natl Acad Sci USA 106:9513–9518

Huang W, Zhu Y, Wu W, Li X, Zhang D, Yin P, Huang J (2018) The directional evolution of mitochondrial gene order in Arabidopsis. Plant Physiol 170:354–366

Hsieh WY, Liao JC, Chang CY, Harrison T, Boucher C, Hsieh MH (2018) Suppressed expression of retrograde-regulated male sterility restores pollen fertility in cytoplasmic male sterile rice plants. Proc Natl Acad Sci USA 106:9513–9518

Hu J, Wang K, Huang WC, Liu G, Gao Y, Wang JM, Huang Q, Ji YX, Qin XJ, Wan L, Zuo RS, Li SQ, Yang DC, Zhu YG (2012) The rice pentatricopeptide repeat protein RFS5 Restores fertility in Hong-Lian cytoplasmic male-sterile lines via a complex with the glyoxine-rich protein GRP162. Plant Cell 24:109–122

Huang W, Zhu Y, Wu W, Li X, Zhang D, Yin P, Huang J (2018) The pentatricopeptide repeat protein SOTS/EMB2279 is required for plastid rpl2 and trnK intron splicing. Plant Physiol 177:684–697

Jin JB, Bae H, Kim SJ, Jin YH, Goh CH, Kim DH, Jin YH, Tse YC, Jiang L, Hwang I (2003) The Arabidopsis dynamin-like proteins ADL1C and ADL1E play a critical role in mitochondrial morphogenesis. Plant Cell 15:2357–2369

Keren I, Tal L, des Frances-Smith CC, Mireau H (2016) The MTLL1 pentatricopeptide repeat protein is required for both translation and splicing of the mitochondrial NADH dehydrogenase subunit7 mRNA in Arabidopsis. Plant Physiol 170:354–366

Khoop V, Schuster W, Wissinger B, Brennicke A (1991) Trans splicing integrates an exon of 22 nucleotides into the nad5 mRNA in higher plant mitochondria. EMBO J 10:3483–3493

Koizuka N, Imai R, Fujimoto H, Hayakawa T, Kimura Y, Kohno-Murase J, Sakai T, Kurose A, Inamura J (2003) Genetic characterization of a pentatricopeptide repeat protein gene, orf687, that restores fertility in the cytoplasmic male-sterile Kosaen radish. Plant J 34:407–415

Kotera E, Tasaka M, Shikanai T (2005) A pentatricopeptide repeat protein is essential for RNA editing in chloroplasts. Nature 433:326–330

Kuhn K, Yin GK, Duncan O, Law SR, Kubiszewski-Jakubiak S, Kaur P, Meyer E, Wang Y, Small CCD, Giraud E, Narasi R, Whelan J (2015) Decreasing electron flux through the cytochrome and/or alternative respiratory pathways triggers common and distinct cellular responses dependent on growth conditions. Plant Physiol 167:228–250

Lamberti G, Drurey C, Soll J, Schwenkert S (2011) The phosphorylation state of chloroplast transit peptides regulates preprotein import. Plant Signal Behav 6:1918–1920

Lamberti G, Gugel IL, Meurer J, Soll J, Schwenkert S (2011) The cytosolic kinases STY8, STY17, and STY46 are involved in chloroplast differentiation in Arabidopsis. Plant Physiol 157:70–85

Law YS, Zhang RS, Guan XQ, Cheng SF, Sun F, Duncan O, Mur- cha MW, Whelan J, Lim BL (2015) Phosphorylation and dephosphorylation of the presequence of precursor multiple organelar RNA editing factor3 during import into mitochondria from Arabidopsis. Plant Physiol 169:1344–1355

Law YS, Ngn L, Yan JR, Kwok LY, Sun YZ, Cheng SF, Schwenkert S, Lim BL (2018) Multiple kinases can phosphorylate the N-terminal sequences of mitochondrial proteins in Arabidopsis. Plant J 63:749–765

Liu Y, He JN, Chen ZZ, Ren XZ, Hong XH, Gong ZZ (2010) ABA is also sensitive to 5 (ABO5), encoding a pentatricopeptide repeat protein required for cis-splicing of mitochondrial nad2 intron 3, is involved in the abscisic acid response in Arabidopsis. Plant J 63:479–475

Liu YJ, Xiu ZH, Meeley R, Tan BC (2013) Empty pericarp5 encodes a pentatricopeptide repeat protein that is required for mitochondrial RNA editing and seed development in maize. Plant Cell 25:868–883

Liu H, Ding YD, Zhou YQ, Jin WQ, Xie KB, Chen LL (2017) CRISPR-P 2.0: an improved CRISPR/Cas9 tool for genome editing in plants. Mol Plant 10:530–532

Liu YJ, Liu XJ, Chen H, Zheng P, Wang WY, Wang LC, Zhang JH, Tu JM (2017) A plastid-localized pentatricopeptide repeat protein is required for both pollen development and plant growth in rice. Sci Rep 7:11484

Lurin C, Andres C, Aubourg S, Bellaoui M, Bitton F, Brugere C, Caboche M, Debast C, Gualberto J, Hoffmann B, Lecharchy A, Le Ret M, Martin-Magniette ML, Mireau H, Peeters N, Renou JP, Szurek B, Taconnat L, Small I (2004) Genome-wide analysis of Arabidopsis pentatricopeptide repeat proteins reveals their essential role in organelle biogenesis. Plant Cell 16:2089–2103

Ma XM, Blenis J (2009) Molecular mechanisms of mTOR-mediated translational control. Nat Rev Mol Cell Biol 10:307–318

Magnuson B, Ekim B, Finger DC (2012) Regulation and function of ribosomal protein S6 kinase (S6K) within mTOR signalling networks. Biochem J 441:1–21

May T, Soll J (2000) 14-3-3 proteins form a guidance complex with chloroplast precursor proteins in plants. Plant Cell 12:53–63

Meyer EH (2012) Proteomic investigations of complex I composition: how to define a subunit? Front Plant Sci 3:106. https://doi.org/10.3389/fpls.2012.00106

Meyer EH, Tomaz T, Carroll AJ, Estavillo G, Delannoy E, Tanz SK, Small ID, Pogson BJ, Millar AH (2009) Remodeled respiration in ndufs4 with low phosphorylation efficiency suppresses Arabidopsis germination and growth and alters control of metabolism at night. Plant Physiol 151:603–619

Millar AH, Whelan J, Soole KL, Day DA (2011) Organization and regulation of mitochondrial respiration in plants. Annu Rev Plant Biol 62:79–104

Neupert W, Herrmann JM (2007) Translocation of proteins into mitochondria. Annu Rev Biochem 76:723–749

Nickel C, Soll J, Schwenkert S (2015) Phosphomimicking within the transit peptide of pHCFl36 leads to reduced photosystem II accumulation in vivo. FEBS Lett 589:1301–1307

Notsu Y, Masood S, Nishikawa T, Kubo N, Akiduki G, Nakazono M, Hirai A, Kadowaki K (2002) The complete sequence of the rice (Oryza sativa L.) mitochondrial genome: frequent DNA sequence acquisition and loss during the evolution of flowering plants. Mol Genet Genom 268:434–445

Qi W, Tian Z, Lu L, Chen X, Chen X, Zhang W, Song R (2017) Editing of mitochondrial transcripts nad3 and cox2 by Deyk10 is essential for mitochondrial function and maize plant development. Genetics 205:1489–1501

Ren X, Pan Z, Zhao H, Zhao J, Cai M, Li J, Zhang Z, Qiu F, Leubner-Gerns (2017) Empty pericarp11 serves as a factor for splicing of mitochondrial nad1 intron and is required to ensure proper seed development in maize. J Exp Bot 68:4571–4581
Schepetilnikov M, Kobayashi K, Geldreich A, Caranta C, Robaglia C, Keller M, Ryabova LA (2011) Viral factor TAV recruits TOR/S6K1 signalling to activate reinitiation after long ORF translation. EMBO J 30:1343–1356

Schleiff E, Becker T (2011) Common ground for protein translocation: access control for mitochondria and chloroplasts. Nat Rev Mol Cell Bio 12:48–59

Small ID, Peeters N (2000) The PPR motif—a TPR-related motif prevalent in plant organellar proteins. Trends Biochem Sci 25:46–47

Sun F, Zhang X, Shen Y, Wang H, Liu R, Wang X, Gao D, Yang Y, Liu Y, Tan BC (2018) The pentatricopeptide repeat protein EMPTY PERICARP8 is required for the splicing of three mitochondrial introns and seed development in maize. Plant J 95:919–932

Sun F, Xia Z, Jiang R, Liu Y, Zhang X, Yang YZ, Li X, Zhang X, Wang Y, Tan BC (2019) The mitochondrial pentatricopeptide repeat protein EMP12 is involved in the splicing of three nad2 introns and seed development in maize. J Exp Bot 70:963–972

Tang HW, Luo DP, Zhou DG, Zhang QY, Tian DS, Zheng XM, Chen LT, Liu YG (2014) The rice restorer Rf4 for wild-aborptive cytoplasmic male sterility encodes a mitochondrial-localized PPR protein that functions in reduction of WA352 transcripts. Mol Plant 7:1497–1500

Tang J, Zhang W, Wen K, Chen G, Sun J, Tian Y, Tang W, Yu J, An H, Wu T, Kong F, Terzagli W, Wang C, Wan J (2017) OsPPR6, a pentatricopeptide repeat protein involved in editing and splicing chloroplast RNA, is required for chloroplast biogenesis in rice. Plant Mol Biol 95:345–357

Thoreen CC, Kang SA, Chang JW, Liu Q, Zhang J, Gao Y, Reichling LJ, Sim T, Sabatini DM, Gray NS (2009) An ATP competitive mammalian target of rapamycin inhibitor reveals rapamycin-resistant functions of mTORC1. J Biol Chem 284:8023–8032

Toda T, Fujii S, Noguchi K, Kazama T, Toriyama K (2012) Rice MPB25 encodes a pentatricopeptide repeat protein and is essential for RNA editing of nad5 transcripts in mitochondria. Plant J 72:450–460

Wang ZH, Zou YJ, Li XY, Zhang QY, Chen L, Wu H, Su DH, Chen YL, Guo JX, Luo D, Long YM, Zhong Y, Liu YG (2006) Cytoplasmic male sterility of rice with boro II cytoplasm is caused by a cytotoxic peptide and is restored by two related PPR motif genes via distinct modes of mRNA silencing. Plant Cell 18:676–687

Wang Y, Zhang XB, Lu SJ, Wang MG, Wang LC, Wang WY, Cao FL, Chen H, Wang J, Zhang JW, Tu JM (2012) Inhibition of a basal transcription factor 3-like gene OsHIBgBTF3 in rice results in significant plant miniaturization and typical pollen abortion. Plant Cell Physiol 53:2073–2089

Wang C, Aube F, Planchard N, Quadrado M, Dargel-Graffin C, Nogueira F, Mireau H (2017) The pentatricopeptide repeat protein MTSF2 stabilizes a nad1 precursor transcript and defines the 3′ end of its 5′-half intron. Nucleic Acids Res 45:6119–6134

Wu MM, Ren YL, Cai MH, Wang YL, Zhu SS, Zhu JP, Hao YY, Teng X, Zhu XP, Jing RN, Zhang H, Zhong MS, Wang YF, Lei CL, Zhang X, Guo XP, Cheng ZJ, Lin QB, Wang J, Jiang L, Bao YQ, Wang YH, Wan JM (2019) Rice FLOWERY ENDOSPERM10 encodes a pentatricopeptide repeat protein that is essential for the trans-splicing of mitochondrial nad1 intron 1 and endosperm development. New Phytol 223:736–750

Wu MW, Zhao H, Zhang JD, Guo L, Liu CM (2020) RADICLELESS 1 (RL1)-mediated nad4 intron 1 splicing is crucial for embryo and endosperm development in rice (Oryza sativa L.). Biochem Bioph Res Co 523:220–225

Wullschleger S, Loewith R, Hall MN (2006) TOR signaling in growth and metabolism. Cell 124:471–484

Xiao H, Zhang Q, Qin X, Xu Y, Ni C, Huang J, Zhu L, Zhong F, Liu W, Yao G, Zhu Y, Hu J (2018) Rice PPS1 encodes a DYW motif-containing pentatricopeptide repeat protein required for five consecutive RNA-editing sites of nad3 in mitochondria. New Phytol 220:878–892

Xiu ZH, Sun F, Shen Y, Zhang XY, Jiang RC, Bonnard G, Zhang JH, Tan BC (2016) EMPTY PERICARP16 is required for mitochondrial nad2 intron 4 cis-splicing, complex I assembly and seed development in maize. Plant J 85:507–519

Xue MY, Liu LL, Yu YF, Zhu JP, Gao H, Wang YH, Wan JM (2019) Loss-of-function of a rice nucleo-localized pentatricopeptide repeat protein is responsible for the floory endosperm14 mutant phenotypes. Rice 12:100

Yang H, Xiu Z, Wang L, Cao SK, Li X, Sun F, Tan BC (2020) Two pentatricopeptide repeat proteins are required for the splicing of nad5 introns in maize. Front Plant Sci 11:732

Yin P, Li QX, Yan CY, Liu Y, Liu JJ, Yu F, Wang Z, Long JF, He JH, Wang HW, Wang JW, Zhuk JK, Shi YG, Yan NE (2013) Structural basis for the modular recognition of single-stranded RNA by PPR proteins. Nature 504:168–171

Zhang YF, Suzuki M, Sun F, Tan BC (2017) The mitochondri昂-targeted PENTATRICOPEPTIDE PEPEAT78 protein is required for nad5 mature mRNA stability and seed development in maize. Mol Plant 10:1321–1333

Zhang QN, Xu YH, Huang JH, Zhang K, Xiao HJ, Qin XJ, Zhu LL, Yu YH, Gu J (2020) The rice pentatricopeptide repeat protein PPR756 is involved in pollen development by affecting multiple RNA editing in mitochondria. Front Plant Sci 11:749

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