A genome-wide identification and analysis of the DYW-deaminase genes in the pentatricopeptide repeat gene family in cotton (Gossypium spp.)

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Abstract

The RNA editing occurring in plant organellar genomes mainly involves the change of cytidine to uridine. This process involves a deamination reaction, with cytidine deaminase as the catalyst. Pentatricopeptide repeat (PPR) proteins with a C-terminal DYW domain are reportedly associated with cytidine deamination, similar to members of the deaminase superfamily. PPR genes are involved in many cellular functions and biological processes including fertility restoration to cytoplasmic male sterility (CMS) in plants. In this study, we identified 227 and 211 DYW deaminase-coding PPR genes for the cultivated tetraploid cotton species G. hirsutum and G. barbadense (2n = 4x = 52), respectively, as well as 126 and 97 DYW deaminase-coding PPR genes in the ancestral diploid species G. raimondii and G. arboreum (2n = 26), respectively. The 227 G. hirsutum PPR genes were predicted to encode 52–2016 amino acids, 203 of which were mapped onto 26 chromosomes. Most DYW deaminase genes lacked introns, and their proteins were predicted to target the mitochondria or chloroplasts. Additionally, the DYW domain differed from the complete DYW deaminase domain, which contained part of the E domain and the entire E+ domain. The types and number of DYW tripeptides may have been influenced by evolutionary processes, with some tripeptides being lost. Furthermore, a gene ontology analysis revealed that DYW deaminase functions were mainly related to binding as well as hydrolase and transferase activities. The G. hirsutum DYW deaminase expression profiles varied among different cotton tissues and developmental stages, and no differentially expressed DYW deaminase-coding PPRs were directly associated with the male sterility and restoration in the CMS-D2 system. Our current study provides an important piece of information regarding the structural and evolutionary characteristics of Gossypium DYW-containing PPR genes coding for deaminases and will be useful for characterizing the DYW deaminase gene family in cotton biology and breeding.
Introduction

RNA editing is a post-transcriptional process affecting organellar transcripts that results in nucleotide changes from DNA to RNA. There are various types of RNA editing in different organisms. In mammals, a zinc-dependent enzyme related to cytidine deaminase is responsible for converting cytidines (C) to uridines (U) in apoB mRNA [1, 2]. In vascular plants, RNA editing occurs in chloroplasts and mitochondria, and almost exclusively involves C-to-U changes [2]. Additionally, nuclear genes encoding proteins involved in RNA editing belong to the gene subfamily associated with pentatricopeptide repeat (PPR) proteins.

The PPR protein family is one of the largest protein families in higher plants [3]. The PPR proteins usually have 2–27 repeating motifs consisting of 35 amino acids arranged in clusters [3, 4]. These motifs are divided into three classes. The canonical PPR motifs (i.e., P motifs) are common among eukaryotes, while the short PPR-like (i.e., S motifs) and long PPR-like motifs (i.e., L motifs) are present only in higher plants [4]. The PPR protein family is divided into two subfamilies (i.e., P and PLS subfamilies) based on the different motifs. The P subfamily members contain only the P motif, while the PLS subfamily members usually consist of a tandem array of degenerated P, L, and S motifs. The PLS subfamily proteins are plant-specific and usually contain an additional C-terminal extension with E, E+, and DYW domains. According to the C-terminal extension, this subfamily can be divided into the following four subgroups: PLS, E, E+, and DYW [4]. Arabidopsis thaliana carries 87 genes encoding PPR proteins with the DYW domain. The DYW domain is named for the frequent presence of an Asp–Tyr–Trp tripeptide at the C-terminal, and has been detected only in the proteins of upland plants. The E and E+ domains are usually highly degraded and difficult to identify. The DYW domain contains a highly conserved region with a-β-α secondary structure consisting of conserved amino acids (e.g., Cys and His) [5].

The DYW domain is the candidate catalytic domain for cytidine deaminase. The HxExx...CxxCH motifs in the DYW domain align with the active site of cytidine deaminase (i.e., C/HxExx...xCxxC). Additionally, the DYW domain is highly evolutionarily correlated with RNA editing. Thus, proteins containing the DYW domain have been classified as deaminases involved in RNA editing [6, 7]. As such, proteins are grouped into the nucleic acid deaminase superfamily (i.e., DYW deaminase family; Pfam: Pf14432) according to DYW domain characteristics. The DYW deaminase domain contains part of the E domain, the whole E+ domain, and most of the DYW domain [7]. Recently, Hayes et al. (2015) suggested that the conserved Glu residue of the HXE motif in the DYW domain is necessary for RNA editing. When the codon for the Glu residue in the DYW domain of OTP84 and CREF7 was mutagenized, the transgenic plants containing the mutated genes could not efficiently edit the cognate editing sites [8]. Wagoner et al. (2015) proposed that the DYW domain and the E (or E+) domain are essential for cytidine deaminase activity because truncating either domain resulted in a QED1 protein that completely lacked any editing activity. In contrast, the DYW domain was revealed to be unnecessary for four RNA editing factors (i.e., CRR22, CRR28, OTP82, and ELI1) [9]. The DYW deaminase domain is different from the DYW domain in PPR proteins, which is essential for the RNA editing activity of some enzymes but not for others. Thus, the role of the DYW domain during RNA editing has not been fully characterized. Boussardon et al. (2012) suggested that the interaction between CRR4 and DYW1 contributes to the editing of the ndhD-1 site. The DYW domain, which functioned as the catalytic domain, may have been lost in CRR4, which now requires the DYW domain from DYW1 to provide catalytic activity. The DYW1 protein carrying the E+ and DYW domains may interact with CRR4 to edit the ndhD C2 site [10]. In this case, CRR4 may use the DYW1 as the catalytic domain. In summary, as
one of the most specific subgroups of the PPR protein family, the DYW deaminase domain-containing PPR proteins are particularly striking because of their RNA editing activities.

Cotton is an economically important fiber crop as the source of the most important natural textile fiber. The genus *Gossypium* contains approximately 45 diploid species and five polyploid species. Crossing between the extant of diploid species *G. arboreum* and *G. raimondii* and chromosome doubling 1–2 million years ago resulted in the emergence of allotetraploid cotton species, including cultivated *G. hirsutum* (accounting for ca. 95% world cotton production) and *G. barbadense* [11]. The genomes of the two ancestral diploids (i.e., *G. arboreum* and *G. raimondii*) and the two tetraploids (i.e., *G. hirsutum* and *G. barbadense*) have recently been sequenced, providing an opportunity to study cotton gene families. PPR genes including those responsible for fertility restoration to cytoplasmic male sterility (CMS) have been cloned with functions analyzed in several plant species. For instance, *Rf4* encoded a PPR protein for wild abortive-type CMS of rice, was confirmed to reduce the *orf352*-containing transcripts to restore pollen fertility [12], and a PPR protein *Rf6* function with hexokinase 6 (*OsHXK6*) to promote the processing of the aberrant CMS-associated transcript *atp6-orfH79* to rescue HL-CMS of rice [13]. In sorghum, two PPR genes were identified for fertility restoration of A1 cytoplasm [14, 15]. However, no such information is currently available in cotton. In this study, we performed an *in silico* analysis of the four *Gossypium* genome sequences to identify PPR genes encoding for the DYW-containing deaminases. We focused on gene structure and expression with a specific focus on CMS and fertility restoration in *G. hirsutum*. The results of this study will contribute to the understanding of the distribution and functions of DYW deaminase genes in cotton.

**Materials and methods**

**Materials**

The *Gossypium hirsutum* variety Zhong10 was used for different tissue expression analysis. Materials were planted under greenhouse. Roots, stems and leaves were collected in one-month-old seedlings, flowers were collected from plants grown two-month-old on the day of flowering, all tissue samples were obtained from 3 individual plants. Cytoplasm male sterility (CMS-D2) A S(*rf1rf1*) with cytoplasm from *Gossypium harknessii* and its isogenic restorer line R S(*Rf1rf1*) were planted under normal conditions in the farm. Floral buds about 3 mm in length (nearly represent the stage of male meiosis) were collected from the near isogenic line with three biological replicates as previous studies, respectively [16, 17]. All collected samples were immediately frozen in liquid nitrogen and stored at -76˚C before use. Above the materials were provided by Cotton Research Institute (CRI), Chinese Academy of Agricultural Science (CAAS), Anyang, Henan, China.

**Local databases of sequence data and RNA-seq**

Local databases including genome sequence and annotation information files of *Gossypium* species (*G. raimondii*, *G. arboreum*, *G. hirsutum* and *G. barbadense* (xinhai)) and other organisms (*Cyanidioschyzon merolae*, *Physcomitrella patens*, *Selaginella moellendorffii*, *Sorghum bicolor*, *Oryza sativa*; Japonica Group) were established. The genome sequence and annotation information of *Gossypium* species was downloaded from the Cottongen (https://www.cottongen.org/). Other organism genome information was downloaded from the NCBI. Arabidopsis thaliana genome sequence were downloaded from TAIR (http://www.arabidopsis.org/). The extraction of total RNA from flower buds about 3 mm in length (at roughly the stage of male meiosis) of CMS cotton (CMS-D2) A and its isogenic restorer line R was performed by using Spectrum™ Plant Total RNA Kit according to the manufacturer’s guide. RNA concentration was
measured using NANODROP 2000. Six transcriptome libraries (A1-3, R1-3) were constructed by using equal amounts of RNA from the three biological replicates. The libraries preparation and sequencing were performed following standard Illumina methods on the Illumina sequencing platform HiSeq™ 2500 system (2×150bp read length). All of the raw sequence data from the Illumina sequencing platform can be found in the Short Read Archive (SRA) database of the National Center for Biotechnology Information (NCBI) under accession number SRX2578795. The FPKMs (mean value of three biological replicates) of gene was further used to perform a differential expression analysis. The criterion was set as: fold change = |log₂ (sample1/sample2) | ≥ 1 and P ≤ 0.05. The differentially expressed genes between CMS cotton (CMS-D2) A and its isogenic restorer line R were shown in S1 Table. RNA-seq data of gene expression during fiber development stages at 5, 10, 20, 25 dpa and in root, stem, leaf, petal, stamen, and pistil in G. hirsutum were downloaded from ccNET database (http://structuralbiology.cau.edu.cn/gossypium/) [18].

DYW-deaminase proteins sequence identification and domain analysis

Download hidden Markov model information (PF14432) of the unique DYW deaminase domain from Pfam29.0 database, then use Pfam file PF14432 to retrieve annotated protein sequences file of Gossypium species and other organisms with Hmmssearch program (E value as the default) in the HMMER 3.0 software, obtained sequences above the threshold from totally protein sequences. Domain analysis manually one by one on candidate protein sequences were performed by using online software SMART (http://smart.embl-heidelberg.de/), removed these sequences without DYW deaminase domain.

Genome wide synteny analysis of DYW deaminase genes

A comparative syntenic map of the DYW deaminase genes from tetraploid cotton species and two diploid cotton species was constructed by using circos-0.69–3 software package, with parameters set as the default.

Conserved domain alignment and analysis of DYW deaminase proteins in Gossypium

By using MEME online software analysis, DYW deaminase protein sequences was used to obtain consensus sequence and Logo of DYW domain and DYW deaminase domain, and compared consensus sequence with tomato and Arabidopsis thaliana.

Phylogenetic tree construction, gene structure analysis, chromosomal and subcellular location of DYW-deaminase proteins

The multiple Sequence alignment of DYW deaminase domain sequence in four cotton species was accomplished by using ClustalX2 software with parameters set as default; the nonroot phylogenetic tree was constructed by the neighbour joining tree (NJ), the bootstrap set for 1000 replications in MEGA 6 software.

According to GFF information, the gene structure display system of Peking University (http://gsds.cbi.pku.edu.cn/) was used to draw the gene structure and DYW deaminase gene were mapped on chromosomes by using Mapchart software.

Using the signal peptide prediction program Target P to predict the subcellular location of DYW-deaminase proteins.
The GO and expression analysis of DYW-deaminase genes

GO analysis of DYW-deaminase genes completed by blast2go software, and the picture was drawn by online software WEGO (http://wego.genomics.org.cn/cgi-bin/wego/index.pl).

The extraction of RNA from root, stem, leaf and flower was performed using RNAprep Pure Plant kit (TIANGEN, China). Equal amounts of RNA from three biological replicates were equally pooled together and constructed the total RNA of each sample. Reverse transcription was conducted by using 0.5 μg total RNA with PrimeScript™ RT reagent kit (Takara Bio Inc., Dalian, China) to 10 μl reaction volume following the manufacturer’s guides and diluted to 100 μl before use. For real-time PCR, 1 μl diluted cDNA was mixed with 10 μl 2 × SYBR Green Mix (Takara) and 1.2 μl primers in a total volume of 20 μl. The PCR was performed using SYBR Green as fluorescence dye and run on Eppendorf Real plex system with melt curve program. The PCR amplification reaction was performed as follows: 94˚C for 3 min, 40 cycles of 94˚C for 5 s, 56˚C for 20 s and 72˚C for 25 s, then ending with the melting curve. Ghhis3 was used as the reference gene for normalization. All reactions were done in three biological replicates, the melting curve was used for each qRT-PCR to determine PCR performance, Ct value in three biological replicates of each gene in four tissues was calculated by 2^-ΔΔCt method [19]. Primers of 16 selected DYW deaminase genes were designed through Primer Premier 5.0 (Table 1).

Results

Identification and structural analysis of Gossypium genes encoding DYW deaminases

A total of 227 G. hirsutum genes were identified as encoding DYW deaminases belonging to the PPR family. Most of these proteins contained multiple PPR domains, and 190 proteins contained intact DYW deaminase domain structures. An analysis of the 227 predicted G. hirsutum DYW deaminases indicated that the number of amino acids in the protein family ranged from 52 to 2016. Furthermore, we identified 126, 97, and 211 DYW deaminases in G. raimondii, G. arboreum, and G. barbadense, respectively. Information about the DYW deaminase genes in

Table 1. The primers of 16 selected PPR DYW deaminase genes for quantitative RT-PCR.

| Gene ID | Forward primer (5’- 3’) | Reverse primer (5’- 3’) |
|---------|------------------------|------------------------|
| Gh_D04G0152 | CTCACCCTTGGATTTCTCGTT | ACCAAGTAACCCAACCTCCTGC |
| Gh_D09G0761 | AAATGCCCTGACCAGACTTTTACCA | AACCCTGCTCATCACTCCTAACCC |
| Gh_A02G0419 | TCCCCACCGCTCTCTC | TGCTGGGGAAGTCTAGGTC |
| Gh_D08G2381 | TTATACATGGCAGAGGCTCT | TGCAATGCTCAACTCCAGGCT |
| Gh_D05G0911 | AACCAAGCTTACCTGAAAGAACCC | CGTATATTTCGAGATCGGGGTG |
| Gh_A03G1665 | TTGTCGGAACTGCCCTCATT | TGACCCGGCTTCAACTAACCC |
| Gh_A01G1752 | TCTGTTCAAAAATTTGCC | TGCTTTCTGCTTGACCAT |
| Gh_A07G1662 | TGGATCTAGTGCTTGCAGGGG | CCAATTTTACTGCCCCGCT |
| Gh_A12G0594 | AAAAAACCCGAAACAGCCC | CATCAGTCTGCAACAGGAAC |
| Gh_D05G0556 | TGGACTTTAGATAGTTACTCCACCAGG | CCTGGAATGAGTAGAAGGAGAG |
| Gh_D02G1741 | GTATGGTGAGATGCTGGAGGCA | GGTGTGCTTAGATCTCCGTT |
| Gh_D01G15053 | TGGGACCCGATGATGACTACG | TCTGCTGACCAAATCGCTC |
| Gh_A02G0212 | TGGGCTGGGTATTGGTTGAGGCAG | GCTCCCAGGATGTTAGGCC |
| Gh_A12G0486 | GCATACGAGTGTAGAGATTGGTGGC | AGCCGACACTGGCTGTAT |
| Gh_A13G1386 | ACGAGAATGAGGGCCGTA | CCGATGCCCAAAATGTC |
| Gh_A01G1103 | GGGAATCTTCTTGATGCTTACACC | TGCTGCTTTACCTGGAGCCT |

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**Gossypium** is summarized in S2 Table. All of the DYW deaminases in the four sequenced cotton species had similar protein structures and contained the DYW deaminase domain. Additionally, most of the DYW deaminases also contained PPR domain tandem repeats. Most of the PPR genes had no introns, which is also an important structural feature of the sequences in DYW deaminase genes. In *G. hirsutum*, 81% of the genes lacked an intron, while 12% contained only one intron and 7% consisted of two or more introns (S1 and S2 Figs). Similar proportions were observed in *G. raimondii* (i.e., 75, 16, and 9%, respectively) and *G. arboreum* (i.e., 81, 12 and 7%, respectively). However, in *G. barbadense*, only 47% of the DYW deaminase genes lacked introns, while the proportion of genes with introns was relatively high (i.e., 24% with one intron and 27% with two or more introns).

**Conserved sequences in the DYW and DYW deaminase domains**

All of the 227 analyzed Upland cotton protein sequences contained the DYW deaminase domain. Additionally, 190 proteins contained a complete and conserved DYW domain, while the remaining 37 proteins contained only a degenerated E+ domain or part of the DYW domain. Using Upland cotton as an example, we obtained the consensus sequences and motif logos for the DYW domains of 68 randomly selected *G. hirsutum* protein sequences using the MEME online program. We also compared the sequences of these proteins with the DYW domain consensus sequences from tomato and *Arabidopsis thaliana* (Fig 1). The obtained consensus sequence of the *G. hirsutum* DYW domain was as follows: AGYyPDTSVLHveEeKEXMLXYHSEkLaIAFGlisTPgTPIReifKNlRvCGDtAhAIK1sIK1tGReiVRDrNRFHFFkDgScSCGdYW.

A comparison of the domain sequences revealed that the DYW domains of the above three different species were similar (Fig 1). The sequences contained conserved HxEx...CxxCH motifs, which corresponded to the cytidine deaminase active site (i.e., C/HxEx...CxxC). Cytidine deaminase is a Zn-dependent enzyme in which the CxxCH motif functions as a binding site [20]. The conserved His and Cys residues in the DYW domain combine with Zn ions, while the conserved Glu is required for deamination [8]. Thus, the DYW domain may be involved in nucleotide deaminations. Additionally, a previous study revealed that if CxxCH in the DYW domain mutated into GxxGH, the RNA cleavage activity decreases significantly [5], suggesting that the CxxCH motif in the DYW domain is necessary for endonuclease activity. Intriguingly, we detected a CxCx motif near the conserved DYW tripeptide or its variants, which may be crucial for protein functions. These results indicate that the DYW domain may have multiple functions.

![Fig 1](https://doi.org/10.1371/journal.pone.0174201.g001)

**Fig 1.** a: The Logo of DYW domain in *G. hirsutum*. The higher the letter, the higher the conservation; b: The comparison of DYW domain consensus sequence in upland cotton, tomato and *Arabidopsis*. The capital letters represent highly conserved, lowercase letters indicate lower conservative.

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We also obtained the complete consensus sequence of the *G. hirsutum* DYW deaminase domain containing part of the E domain, the whole E+ domain, and the DYW domain (Fig 2). We compared this sequence with the DYW deaminase domain sequences from other three cotton species. Furthermore, we obtained a conserved PG box sequence of the E and E+ domains as described by Hayes et al. (2013). The PG box may bind to different factors to generate editing components, although the role of the PG box in editing events has not been clarified [21]. The intact DYW deaminase domain sequences were highly similar among the different analyzed cotton species, and contained the PG box, HxEx ... CxxCH, and CxCxDYW motifs. These results indicate the DYW deaminase and DYW domains may exhibit similar molecular characteristics and functions.

Fig 2. a: The alignment of the DYW deaminase domain in *G. hirsutum*; b: The Logo of DYW deaminase domain in *G. hirsutum*; c: The consensus sequence in *G. hirsutum* and comparison of DYW deaminase domain in *Gossypium*. The capital letters represent highly conserved, lowercase letters indicate lower conservative.

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Chromosomal localization and evolutionary relationships among *Gossypium* DYW deaminase genes

Of the identified 227 *G. hirsutum* DYW deaminase genes, 203 genes were mapped to chromosomes, with 97 genes localized in the A subgenome and 106 genes localized in the D subgenome (Fig 3). Each chromosome contained an average of 7.8 DYW deaminase genes. In the A subgenome, chromosomes 4 and 10 carried the fewest DYW deaminase genes (i.e., only four each), while chromosome 12 contained the most (i.e., 14). In the D subgenome, chromosome 3 and 10 had the fewest DYW deaminase genes (i.e., only 3 and 4, respectively), and chromosome 12 carried the most DYW deaminase genes (i.e., 12). Furthermore, some genes clustered together on chromosomes. For example, four DYW deaminase genes localized within a 3.2-Mb region on chromosome 5 in the D subgenome (i.e., *Gh_D05G0556, Gh_D05G0696, Gh_D05G0877*, and *Gh_D05G0911*), while five DYW deaminase genes were mapped to a 2.9-Mb region on chromosome 12 of the same subgenome. Additionally, in *G. hirsutum*, the location of genes on chromosomes 9, 10, and 13 in the A subgenome exhibited a good collinearity with that on homoeologous chromosomes 9, 10, and 13 in the D subgenome. In contrast, the chromosomal localization of DYW deaminase genes in *G. raimondii* and *G. arboreum* did not exhibit a good collinearity with that in *G. hirsutum* because of differences in gene evolution (S2 Fig).

Due to the evolutionary relationships between diploid and tetraploid cotton species, the DYW deaminases in the A and D subgenomes of the tetraploid species are homologous to those in diploid species *G. arboreum* and *G. raimondii*, respectively. We analyzed the evolutionary relationships between the *G. hirsutum* and *G. barbadense* DYW deaminases and their corresponding proteins in *G. raimondii* and *G. arboreum*. An analysis of the DYW deaminases in the two tetraploid cotton species revealed that the *G. hirsutum* proteins had more tandem duplications. A comparative syntenic map of the DYW deaminase genes from tetraploid cotton species and the two diploid cotton species was constructed for a further analysis of genetic origins and evolution (Fig 4). During the evolution of *Gossypium* DYW deaminase genes, one *G. hirsutum* gene and seven *G. barbadense* genes did not have homologous genes in the two diploid cotton species (S2 Table). These genes may be new, with distinct functions in tetraploid cotton species. Additionally, 19 and 36 *G. raimondii* genes and 11 and 25 *G. arboreum* genes are not present in the *G. hirsutum* and *G. arboreum*.
barbadense genomes, respectively. Furthermore, 32 and 30 G. raimondii genes and 15 G. arboreum genes had more than one homologous gene in G. hirsutum and G. barbadense, respectively. We also observed that 75 and 60 G. raimondii genes and 71 and 57 G. arboreum genes each had one homologous gene in G. hirsutum and G. barbadense, respectively. Further studies are necessary to clarify the functions of the unique genes in the two tetraploid cotton species.

Conserved DYW tripeptides and disordered amino acids in the C-terminal of DYW deaminases

In this study, we observed that the DYW and DFW tripeptides were the most frequently observed tripeptides in the C-terminal regions of DYW deaminases (Table 2). For example, there were 131 and 27 proteins that contained the DYW and DFW tripeptides in the C-terminal of G. hirsutum DYW deaminases, respectively. There were 113 and 28 G. barbadense DYW deaminases with DYW and DFW tripeptides, respectively. Of the diploid Gossypium species, 74, 19 and 58, 8 proteins contained the DYW and DFW tripeptide in G. raimondii and G. arboreum, respectively. Though the number of DYW tripeptides differed among Gossypium species, its proportions in the whole gene family were similar, ranging from 53.55% to 59.79%.

The types and number of tripeptides present in the C-terminal of DYW deaminases vary depending on organisms, including Cyanidoschyzon merolae, Physcomitrella patens, Selaginella moellendorffii, A. thaliana, Oryza sativa, and Sorghum bicolor (Table 2). Interestingly, some sequences contained other types of C-terminal tripeptides (with disordered amino acid residues), accounting for different proportions of the tripeptides in diverse organisms. For example, there are no DYW deaminases in C. merolae, while P. patens contained only DYW deaminases with C-terminal DYW and DFW tripeptides. However, 15 DYW deaminases in S. moellendorffii (i.e., 8.67%) contained a disordered amino acid residue in the C-terminal region.
Additionally, 17.7% and 5.38% of the *Gossypium* species and *A. thaliana* DYW deaminases contained a disordered amino acid residue, respectively.

**Subcellular localization and gene ontology analysis of DYW deaminases**

To predict the subcellular locations of DYW deaminases, the Target P online program was used to examine protein transport. The DYW deaminases were detected in similar subcellular locations in all four analyzed cotton species. In *G. hirsutum*, 120 DYW deaminases (i.e., 53%)
were localized to the mitochondria and chloroplasts, while 16 proteins (i.e., 7%) containing an N-terminal signal peptide were transported to other locations. However, we were unable to predict the subcellular locations of 40% of the predicted DYW deaminases, suggesting the need for further investigations. In other cotton species, the distribution of DYW deaminases in specific subcellular locations was similar to that of G. hirsutum (S3 Fig).

Gene ontology (GO) analysis revealed that G. hirsutum DYW deaminases may be important for developmental processes (Fig 5). The cellular component proteins were mainly associated with intracellular or membrane-bound organelles, which is consistent with the subcellular localization results for DYW deaminases. The primary molecular functions included binding as well as hydrolase and transferase activities. The main biological process terms were related to cellular metabolic processes, macromolecule metabolic processes, nitrogen compound metabolic processes, and primary metabolic processes.

**Phylogenetic of *Gossypium* DYW deaminases**

To clarify the evolutionary relationships among *Gossypium* DYW deaminases, we constructed a phylogenetic tree using the MEGA 6 program (Fig 6). The DYW deaminases were divided into five groups (i.e., A–E) based on protein sequence similarities, with groups C and E containing the most and fewest proteins, respectively.

**Gossypium DYW deaminase gene expression patterns at different developmental stages and tissues**

To understand the temporal and spatial expression patterns of these PPR DYW deaminase genes in Upland cotton, a publicly deposited RNA-seq data were used to assess the expression across tissues and developmental stages. Among the 227 PPR DYW deaminase genes, there
were 219 genes with an FPKM > 0 in at least one of the 10 investigated tissues and developmental stages (Fig 7). The expression for the remaining 8 genes was not detected and will not be discussed in this study. We observed that the expression of DYW deaminase domain-containing PPR genes were different in specific tissues and developmental stages. For example, most of the genes were upregulated in root, stem, leaf and pistil, and fiber tissues, while they were downregulated in petal and stamen tissues except for Gh_A12G0429, Gh_A13G155, Gh_D08G0677 and Gh_D12G2546. Some genes were specifically expressed in different tissues. For example, Gh_A13G1722 gene was petal-specific, while Gh_D05G3688 was root-specific, because they were highly expressed in petal and root, respectively.

**Quantitative real-time PCR (qRT-PCR) analysis**

A quantitative real-time PCR was used to investigate the expression levels of 16 randomly selected PPR DYW deaminase genes in roots, stem, leaves, and flowers of Upland cotton (Fig 8). All of
the 16 selected genes were expressed in the four analyzed tissues, with different tissue-specific expression profiles. Among these genes, \textit{Gh\textunderscore D09G0761}, \textit{Gh\textunderscore D05G0556}, and \textit{Gh\textunderscore A03G1665} were the most highly expressed genes in flowers. \textit{Gh\textunderscore D09G0761} and \textit{Gh\textunderscore A03G1665} produced similar expression profiles (i.e., gradual increase in the four examined tissues). The results imply that these genes may have a special role during flower development. \textit{Gh\textunderscore A01G1752}, \textit{Gh\textunderscore D08G2381}, \textit{Gh\textunderscore D04G0152}, \textit{Gh\textunderscore A12G0594}, \textit{Gh\textunderscore A02G0419}, \textit{Gh\textunderscore A07G1662}, and \textit{Gh\textunderscore A13G1386} were the most highly expressed genes in leaves. Additionally, highest expression levels for \textit{Gh\textunderscore D02G1741}, \textit{Gh\textunderscore D01G0153}, \textit{Gh\textunderscore A12G0486}, \textit{Gh\textunderscore A01G1103}, and \textit{Gh\textunderscore A02G0212} were observed in stems. Finally, \textit{Gh\textunderscore D05G0911} was highly expressed in the roots. These results indicate that DYW deaminases may be important for cotton development.

Differentially expressed PPR genes between CMS-D2 and restorer

In a genomewide transcriptome analysis of floral buds between near-isogenic CMS-D2 (i.e., A line) and its restorer (R line) differing in the \textit{Rf1} locus, 6 differentially expressed (DE) PPR genes were identified (S1 Table). However, no DYW deaminase domain in C-terminal was found in these 6 PPR genes. Among these 6 PPR genes, \textit{Gh\textunderscore A02G1102} (PLS subgroup), \textit{Gh\textunderscore A11G0734} and \textit{Gh\textunderscore D11G0852} (E subgroup) belong to the PLS subfamily, while \textit{Gh\textunderscore A06G0542}, \textit{Gh\textunderscore D05G3392} and \textit{Gh\textunderscore D06G0610} belong to the P subfamily. It is interesting to note that, as compared with the R line, 3 PLS subfamily genes (\textit{Gh\textunderscore A02G1102}, \textit{Gh\textunderscore A11G0734} and \textit{Gh\textunderscore D11G0852}) were up regulated and 3 P subfamily genes (\textit{Gh\textunderscore A06G0542}, \textit{Gh\textunderscore D05G3392} and \textit{Gh\textunderscore D06G0610}) were down regulated in the A line. Furthermore, the PPR gene \textit{Gh\textunderscore D05G3392} is located on the chromosome where the restorer gene \textit{Rf1} resides. The \textit{Gh\textunderscore D05G3392} protein belongs to P subfamily and contain 14 repeats of PPR motifs. According to the previous studies [22–26], we
adopted the three number PPR code as proposed by Yin et al [24] and predicted the amino acids in 3 particular positions (1, 4 and ii; residues 2, 5 and 35, respectively, as used in this study) within Gh_D05G3392 protein motifs. In Fig 9, we aligned the PPR motif and highlighted the residues at 2, 5 and 35 positions. Additionally, we also predicted the RNA sequence (UUUUUUUUUU CUU) recognized by the Gh_D05G3392 according to the rule proposed by Yagi et al [23]. Interestingly, one of the blast results in NCBI showed that this sequence (UUUUUUUUUCUU) is a partial sequence of a pseudogene in *G. hirsutum* and SSR marker GNCOT-C1-F (GenBank: KX090570.1). However, it is outside of the Rf1 locus [27], while other five genes were even not located on the Rf1-carrying chromosome (Gh_D05). Thus, with the introduction of the dominant restorer gene Rf1 to the A line containing the CMS-inducing cytoplasm, the expression of the

![Fig 8. Expression analysis of 16 selected DYW deaminase genes in different tissues through qRT-PCR. a: High expression in the flower; b: Expression peaks in the leaf; c: High expression in the stem; d: High expression in the root.](https://doi.org/10.1371/journal.pone.0174201.g008)
three PLS subfamily genes (PLS and E subgroup) was suppressed, while other three genes in the P subfamily were activated for male fertility restoration in the R line. These results indicated that DYW deaminase domain-containing PPR genes may not directly function in the CMS occurrence or fertility restoration. This is consistent with previous results in cloned restorer genes from other plant species.

Discussion

Relationship between DYW deaminase and DYW domains

In this study, we identified 227 DYW deaminase genes in the \textit{G. hirsutum} genome. We also obtained the consensus sequence of the \textit{Gossypium} DYW deaminase domain. Compared with the DYW domain, the complete DYW deaminase domain contained the E and E+ domains, which was consistent with the findings of a previous study [7]. Additionally, a PG-box was detected in the DYW deaminase domain, spanning the E and E+ domains. The functions of this PG box have not been characterized.

Molecular evolution of DYW deaminases

Allotetraploid \textit{Gossypium} species evolved from two diploid cotton species through hybridizations and genome doubling that occurred 1–2 million years ago [11]. Therefore, diploid and allotetraploid cotton species are useful models for investigating the evolution of plants. The whole genome sequences of \textit{Gossypium} species have enabled researchers to study the molecular evolutionary characteristics of \textit{Gossypium} DYW deaminases. In this study, we analyzed the types and number of DYW tripeptides or their variants in four \textit{Gossypium} species (Table 2).

By comparing the types and number of tripeptides present in the C-terminal of DYW deaminases from red algae (\textit{C. merolae}), bryophytes (\textit{P. patens}), lycophyta (\textit{S. moellendorffii}), monocots (\textit{Sorghum bicolor} and \textit{Oryza sativa}), and eudicots (\textit{A. thaliana} and \textit{Gossypium} species), we determined that diverse plant species evolved differently in terms of the types and number of tripeptides. For example, \textit{C. merolae} lacks DYW deaminases, while \textit{P. patens} produces 10 DYW deaminases with a conserved C-terminal tripeptide (i.e., either DYW or DFW). \textit{Selaginella moellendorffii} rapidly evolved 10 types of tripeptides, and the proportion of
conserved DYW tripeptides decreased. Additionally, a disordered amino acid residue is present in the C-terminal of some *S. moellendorffii* DYW deaminases, similar to corresponding DYW deaminases in monocots and eudicots. This indicates that the types of conserved DYW tripeptide increased during the evolution from algae to angiosperms. Selection pressures may have induced the adaptation of DYW tripeptide types to evolutionary processes. Previous studies indicated that the phylogenetic distribution of DYW domains is highly correlated with RNA editing, with the number of RNA editing events increasing during evolution [6, 28]. Thus, the types and number of conserved tripeptides in DYW domains may be highly correlated with RNA editing events.

The frequency of residue changes differs among conserved tripeptides. In the DYW tripeptide, Trp is the most conserved residue, followed by Asp. We observed that most DYW deaminases contain a CxCx motif near the conserved tripeptide. However, some protein sequences lacking the CxCx motif may also be missing the HxExx...CxxCH motif. This may be because during early plant evolutionary processes, the conserved DYW tripeptide sequence became flexible and distinct tripeptides due to residue changes increased the number of tripeptide types. Some flexible tripeptides were gradually lost during evolution, which resulted in the loss of the CxCx motif and other sequences in the DYW domain following the tripeptide. Based on the relationship between the conserved DYW tripeptide and CxCx motif, we hypothesized that the conserved DYW tripeptide functions like a protective cap to prevent the loss of the CxCx motif. As distinct parts of DYW deaminases, the roles of the CxCx motif and conserved DYW tripeptide should be more thoroughly characterized in future studies.

PPR gene family and male fertility restoration

The PPR gene family constitutes a large family of RNA binding in plants, it involves in many cellular functions and biological processes in organelles including gene expression, RNA stabilization, RNA cleavage and RNA editing [25, 29]. Previous studies indicated the PPR proteins involved in RNA editing events mostly present a characteristic motif, which contain a PPR tract followed by C-terminal motifs (E, E+ and DYW), although the E+ and DYW domain are often missing [30]. Additionally, a conserved motif HxE(x)nCxxCH exist in DYW domain is required for C to U conversion, rather than for recognition the editing site [31]. In this study, we found the complete DYW deaminase domain contained the part of E, E+ and DYW domains, and most DYW deaminases contain a CxCx motif near the conserved tripeptide. However, some protein sequences lacking the CxCx motif may also be missing the whole or part of HxExx...CxxCH motif. The DYW deaminase domain as the catalytic domain maybe promote C to U conversion. As the tandem arrays of PPR motifs, Yagi et al. (2013) found amino acid residues at 3 particular position (1, 4, and ii) within the PPR motif recognizing the upstream nucleotide sequence for the editing sites [23]. Furthermore, Takenaka et al. (2013) found that combining the P, L and S motifs will improve the prediction of RNA editing target sites [26]. In this study, we predicted the PPR code in 3 particular positions (1, 4 and ii; residues 2, 5 and 35, respectively, as used in this study) within Gh_D05G3392 motifs and predicted the RNA sequence (UUUUUUUUUU CUU) recognized by Gh_D05G3392, which are part of a pseudogene in *Gossypium hirsutum* and SSR marker GNCOT-C1-F (GenBank: KX090570.1).

CMS in flowering plants is characterized by a maternally inherited trait that lacks functional pollen grains [32], and this trait can be restored by nuclear genes known as restorer-of-fertility (*Rf*) genes [33]. Previous studies indicated that RNA editing events may have certain relationship with CMS occurrence in many crops. For example, Wei et al. (2008) found that the editing of atp9 change an arginine codon into a termination codon, shorting the protein to the standard in Ying xiang B while in Ying xiang A, which has no termination codon, cannot be a
normal protein [34]. Wang et al. (2009) found that RNA editing occurrence rates in atp6 and
cox2 generally increase in sterile lines [35]; Chakraborty et al. (2015) indicated that an over-
expression of the unedited mitochondrial orfB gene generated male sterility in fertile rice lines
in a dose-dependent manner, in which the unedited orfB gene expression increased, resulting
in a low level of ATP synthase activity of F1F0-ATP synthase (complex V) [36]. In cotton,
Suzuki et al. (2013) identified RNA editing sites in eight mitochondrial genes among CMS-D8
three line systems, and found that the restorer gene may change RNA editing efficiency [37].
However, little is known about the RNA editing events of mitochondrial genes in CMS-D2.
Wu et al. (2011) found a C-U and a C-G editing site at position 1178 and 1214 bp
downstream of the ATG codon in atpA and a A-U and a C-U editing site at position 18 and 35 bp
upstream of ATG codon in nad6 between maintainer and CMS-D2 lines, respectively [16]. Thus, we
speculate that RNA editing events in D2 mitochondrial genes may regulate ATP synthase
activity, and the impaired ATP synthesis results in CMS occurrence.

Fertility restoration generally requires the reduction of CMS-associated RNAs or proteins
by the action of the Rf gene(s). Most of the restorer genes cloned so far are reported to encode
a PPR protein. For instance, the radish Rfo [38], and two PPR genes in sorghum [14, 15], and
Rf4 and Rf6 gene in rice [12, 13] each encodes a different PPR protein. Additionally, previous
studies indicated that most of them belong to P subfamily, except for that fact that the Rf1 gene
in sorghum encoded a PPR protein which is classified as member of E subgroup of PLS sub-
family [14, 39][14, 33]. In this study, 6 differentially expressed (DE) PPR genes were identified
through a genomewide transcriptome analysis between A and R lines. Interestingly, 3 PLS sub-
family genes (Gh_A02G1102, Gh_A11G0734 and Gh_D11G0852) were up regulated and 3 P
subfamily genes (Gh_A06G0542, Gh_D05G3392 and Gh_D06G0610) were down regulated in
the A line. These results indicated that DYW deaminase domain-containing PPR genes may
not directly function in the CMS occurrence or fertility restoration, while P subfamily genes
might have critical role in the process of fertility restoration. And this is consistent with the
previous studies that most of Rf genes belong to P subfamily [39]. Previous studies indicated a
small subgroup of PPR genes existed in plant genomes that share high similarity with the iden-
tified Rf-PPRs, called as Rf-like PPRs (RFL) genes [38, 40]. For example, in radish Rfo
loci, three similar PPRs were identified, which may arise through gene duplication events [38]. In
this study, five genes were not located on the Rf1-carrying chromosome (Gh_D05), while the
PPR gene Gh_D05G3392 is located on the chromosome where the restorer gene Rf1 resides
[27], though it is outside of the Rf1 locus and its protein was predicted to target in the chloro-
plasts, and the physical distance is short. Therefore, we speculate that the Gh_D05G3392 gene
may share a high similarity with Rf1 in cotton but it is not the candidate gene for Rf1. As for
the mechanism of CMS restoration in cotton, due to the limited study, we speculate that the
Rf1 may be a PPR gene belonging to the P subfamily. Takenaka et al. (2013) proposed that the
P subfamily PPR proteins often bind tightly to their RNA targets and cannot be removed [26].
And Suzuki et al. (2013) indicated that the restorer gene may alter RNA editing efficiency
among the CMS-D8 three line systems [37]. Thus, Rf1-PPR protein maybe binds tightly to
their CMS gene or product, resulting in that the DYW deaminase gene cannot edit the CMS
gene product, so the male fertility is restored. However, the RNA editing events in CMS-D2
cotton and the role of Rf1 gene in RNA editing still need further studies.

Conclusion
We identified DYW deaminases based on four sequenced Gossypium species genomes. We
conducted chromosomal and subcellular localization experiments as well as gene structure,
expression, and GO analyses for the identified genes and proteins. Most of the Gossypium
DYW deaminases contained the conserved Asp-Tyr-Trp tripeptide (or variants) in the C terminal. The DYW deaminases lacking the conserved tripeptide may have been gradually lost during evolution. The PPR proteins containing the DYW domain are ancient RNA-editing proteins. With increasing numbers of editing sites, some amino acids in the DYW domain were gradually lost during evolution, resulting in an increase in the abundance of proteins from other subgroups (e.g., when the DYW domain was converted to an E domain). Therefore, the DYW deaminase domain might influence nucleotide deamination, which is consistent with the predicted molecular function based on GO analysis. Furthermore, regarding subcellular localizations, DYW deaminases were predicted to mainly localize in mitochondria or chloroplasts, which was consistent with GO analysis results related to cell components. In addition, the transcriptome analysis indicated no differentially expressed DYW deaminase-coding PPRs were directly associated with male sterility and restoration in the CMS-D2 system. In summary, our study provides basic information regarding the structural and evolutionary characteristics of DYW deaminases in cotton. Our findings may be useful for improving cotton production in future studies.

Supporting information
S1 Fig. The number of introns in DYW deaminase genes in *Gossypium*.
(TIF)

S2 Fig. The gene structure of DYW deaminase in *Gossypium*.
(EPS)

S3 Fig. Mapping of the DYW deaminase genes in the chromosomes of *G. arboreum, G. raimondii* and *G. barbadense*.
(EPS)

S4 Fig. Subcellular localization of DYW deaminases in *Gossypium*.
(TIF)

S1 Table. Differentially expressed genes between CMS-D2 and restorer.
(XLSX)

S2 Table. Information about the DYW deaminase genes in *Gossypium*.
(XLSX)

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References
1. Wedekind JE, Dance GSC, Sowden MP, Smith HC. Messenger RNA editing in mammals: new members of the APOBEC family seeking roles in the family business. Trends in Genetics. 2003; 19(4):207–16. https://doi.org/10.1016/S0168-9525(03)00054-4 PMID: 12683974
2. Shikanai T. RNA editing in plants: Machinery and flexibility of site recognition. Biochimica et Biophysica Acta (BBA)—Bioenergetics. 2015; 1847(9):779–85.
3. Small ID, Peeters N. The PPR motif—a TPR-related motif prevalent in plant organellar proteins. Trends in Biochemical Sciences. 2000; 25(2):45–7.
4. Lurin C, Andres C, Aubourg S, Bellaloui M, Bitton F, Bruyere C, et al. Genome-wide analysis of Arabidopsis pentatricopeptide repeat proteins reveals their essential role in organelle biogenesis. Plant Cell. 2004; 16(8):2089–103. Epub 2004/07/23. PubMed Central PMCID: PMCPmc519200. https://doi.org/10.1105/tpc.104.022236 PMID: 15269332
5. Nakamura T, Sugita M. A conserved DYW domain of the pentatricopeptide repeat protein possesses a novel endoribonuclease activity. FEBS Lett. 2008; 582(30):4163–8. Epub 2008/12/02. https://doi.org/10.1016/j.febslet.2008.11.075 PMID: 19041647
6. Salone V, Rudinger M, Polsakiewicz M, Hoffmann B, Groth-Malonek M, Szurek B, et al. A hypothesis on the identification of the editing enzyme in plant organelles. FEBS Lett. 2007; 581(22):4132–8. Epub 2007/08/21. https://doi.org/10.1016/j.febslet.2007.07.075 PMID: 17707818
7. Iyer LM, Zhang D, Rogozin IB, Aravind L. Evolution of the deaminase fold and multiple origins of eukaryotic editing and mutagenic nucleic acid deaminases from bacterial toxin systems. Nucleic Acids Res. 2011; 39(22):9473–97. Epub 2011/09/06. PubMed Central PMCID: PMCPmc3239186. https://doi.org/10.1093/nar/gkr691 PMID: 21890906
8. Hayes ML, Dang KN, Diaz MF, Mulligan RM. A conserved glutamate residue in the C-terminal deaminase domain of pentatricopeptide repeat proteins is required for RNA editing activity. J Biol Chem. 2015; 290(16):10136–42. Epub 2015/03/06. PubMed Central PMCID: PMCPmc440329. https://doi.org/10.1074/jbc.M114.631630 PMID: 25739442
9. Wagoner JA, Sun T, Lin L, Hanson MR. Cytidine deaminase motifs within the DYW domain of two pentatricopeptide repeat-containing proteins are required for site-specific chloroplast RNA editing. J Biol Chem. 2015; 290(5):2957–68. Epub 2014/12/17. PubMed Central PMCID: PMCPmc4317000. https://doi.org/10.1074/jbc.M114.622084 PMID: 25512379
10. Boussardon C, Salone V, Avon A, Berthomé R, Hammami K, Okuda K, et al. Two interacting proteins are necessary for the deaminase fold and multiple origins of eukaryotic editing and mutagenic nucleic acid deaminases from bacterial toxin systems. Nucleic Acids Res. 2011; 39(22):9473–97. Epub 2011/09/06. PubMed Central PMCID: PMCPmc3239186. https://doi.org/10.1093/nar/gkr691 PMID: 21890906
11. Hu G, Hawkins JS, Grover CE, Wendel JF. The history and disposition of transposable elements in polyploid Gossypium. 2010; 53(8):599–607. https://doi.org/10.1199/gt-01-038 PMID: 20725147
12. Kazama T, Tomyama K. A fertility restorer gene, Rf4, widely used for hybrid rice breeding encodes a pentatricopeptide repeat protein. Rice. 2014; 7(1):1–5.
13. Huang W, Yu C, Hu J, Wang L, Dan Z, Zhou W, et al. Pentatricopeptide-repeat family protein RF6 functions with hexokinase 6 to rescue rice cytoplasmic male sterility. Proc Natl Acad Sci U S A. 2015; 112 (48):14984–9. https://doi.org/10.1073/pnas.1511748112 PMID: 26578614
14. Klein RR, Klein PE, Mullet JE, Minx P, Rooney WL, Schertz PF. Fertility restorer loci Rf1 of sorghum (Sorghum bicolor L.) encodes a pentatricopeptide repeat protein not present in the colinear region of rice chromosome 12. Theoretical and Applied Genetics. 2005; 111(6):994–1012. https://doi.org/10.1007/s00122-005-0311-8 PMID: 16078015
15. Jordan DR, Mace ES, Henzell RG, Klein PE, Klein RR. Molecular mapping and candidate gene identification of the Rf2 gene for pollen fertility restoration in sorghum [Sorghum bicolor (L.) Moench].
16. Wu J, Gong Y, Cui M, Qi T, Guo L, Zhang J, et al. Molecular characterization of cytoplasmic male sterility conditioned by Gossypium harknessii cytoplasm (CMS-D2) in upland cotton. Euphytica. 2011; 181 (1):17–29.

17. Suzuki H, Rodriguez-Uribe L, Xu J, Zhang J. Transcriptome analysis of cytoplasmic male sterility and restoration in CMS-D8 cotton. Plant Cell Reports. 2013; 32(10):1531–42. https://doi.org/10.1007/s00299-013-1465-7 PMID: 23743655

18. You Q, Xu W, Zhang K, Zhang L, Yi X, Yao D, et al. ccNET: Database of co-expression networks with functional modules for diploid and polyploidGossypium. Nucleic Acids Research. 2017;45(Database issue):D1090–D9.

19. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods. 2001; 25(4):402–8. https://doi.org/10.1006/meth.2001.1262 PMID: 11846609

20. Sugita M, Ichinose M, Ide M, Sugita C. Architecture of the PPR gene family in the moss Physcomitrella patens. RNA Biol. 2013; 10(9):1439–45. Epub 2013/05/07. PubMed Central PMCID: PMCPmc3858427. https://doi.org/10.4161/rna.24772 PMID: 23645116

21. Hayes ML, Giang K, Berhané B, Mulligan RM. Identification of two pentatricopeptide repeat genes required for RNA editing and zinc binding by C-terminal cytidine deaminase-like domains. Journal of Biological Chemistry. 2013; 288(51):36519–29. https://doi.org/10.1074/jbc.M113.485755 PMID: 24194514

22. Shen C, Zhang D, Guan Z, Liu Y, Yang Z, Yang Y, et al. Structural basis for specific single-stranded RNA recognition by designer pentatricopeptide repeat proteins. Nature Communications. 2016; 7:11285. https://doi.org/10.1038/ncomms11285 PMID: 27088764

23. Yagi Y, Hayashi S, Kobayashi K, Hirayama T, Nakamura T. Elucidation of the RNA recognition code for pentatricopeptide repeat proteins involved in organelle RNA editing in plants. PLoS One. 2013; 8(3):e57286. Epub 2013/03/09. PubMed Central PMCID: PMCPmc3589468. https://doi.org/10.1371/journal.pone.0057286 PMID: 23472078

24. Yin P. Structural basis for the modular recognition of single-stranded RNA by PPR proteins. Nature. 2013; 504(7478):168. https://doi.org/10.1038/nature12651 PMID: 24162847

25. Takenakaa M, Zehrmann A, Brennicke A, Graichen K. Improved computational target site prediction for pentatricopeptide repeat RNA editing factors. Plos One. 2013; 8(6):e65343. https://doi.org/10.1371/journal.pone.0065343 PMID: 23762347

26. Knoop V, Rudinger M. DYW-type PPR proteins in a heterolobosean protist: plant RNA editing factors involved in an ancient horizontal gene transfer? FEBS Lett. 2010; 584(20):4287–91. Epub 2010/10/01. PMID: 20888816

27. Schmitzinneweber C, Small I. Pentatricopeptide repeat proteins: a socket set for organelle gene expression. Trends in Plant Science. 2008; 13(12):663–70. https://doi.org/10.1016/j.tplants.2008.10.001 PMID: 19004664

28. Okuda K, Myougaf, Motoharsish, Shinokaziskih, Shikanait. Conserved domain structure of pentatricopeptide repeat proteins involved in chloroplast RNA editing. Proc Natl Acad Sci U S A. 2007; 104 (19):8178–83. Epub 2007/05/08. PubMed Central PMCID: PMCPmc1876591. https://doi.org/10.1073/pnas.0700865104 PMID: 17483454

29. BoussardonC, Avon A, KindgrenPBondCS, ChallengerMLurinc, et al. The cytidine deaminase signature HxEx(n)CxxC of DYW1 binds zinc and is necessary for RNA editing of ndhD-1. New Phytologist. 2014; 203(4):1090–5. https://doi.org/10.1111/nph.12928 PMID: 25041347

30. HansonMR, BentolliaS. Interactions of Mitochondrial and Nuclear Genes That Affect Male Gametophyte Development. The Plant Cell. 2004; 16(suppl 1):S154–S69.

31. WangF, StewartJM, ZhangJ. Molecular markers linked to the RI2 fertility restorer gene in cotton. Genome / National Research Council Canada = Genome / Conseil national de recherches Canada. 2007; 50(9):818–24.

32. WeiL, DingYY. Mitochondrial RNA editing of F0-ATPase subunit 9 gene (atp9) transcripts of Yunnan purple rice cytoplasmic male sterile line and its maintainer line. Acta Physiologicae Plantarum. 2008; 30 (5):657–62.
35. Wang J, Cao MJ, Pan GT, Lu YL, Rong TZ. RNA editing of mitochondrial functional genes atp6 and cox2 in maize (Zea mays L.). Mitochondrion. 2009; 9(5):364–9. https://doi.org/10.1016/j.mito.2009.07.005 PMID: 19666144

36. Chakraborty A, Mitra J, Bhattacharyya J, Pradhan S, Sikdar N, Das S, et al. Transgenic expression of an unedited mitochondrial orfB gene product from wild abortive (WA) cytoplasm of rice (Oryza sativa L.) generates male sterility in fertile rice lines. Planta. 2015; 241(6):1463–79. https://doi.org/10.1007/s00425-015-2269-5 PMID: 25754232

37. Suzuki H, Yu J, Ness SA, O’Connell MA, Zhang J. RNA editing events in mitochondrial genes by ultra-deep sequencing methods: a comparison of cytoplasmic male sterile, fertile and restored genotypes in cotton. Molecular Genetics and Genomics. 2013; 288(9):445–57. https://doi.org/10.1007/s00438-013-0764-6 PMID: 23812672

38. Brown GG, Formanová N, Jin H, Wargachuk R, Dendy C, Patil P, et al. The radish Rfo restorer gene of Ogura cytoplasmic male sterility encodes a protein with multiple pentatricopeptide repeats. Plant Journal. 2003; 35(2):262–72. PMID: 12848830

39. Dahan J, Mireau H. The Rf and Rf-like PPR in higher plants, a fast-evolving subclass of PPR genes. Rna Biology. 2012; 10(9):1469–76.

40. Fujii S, Bond CS, Small ID. Selection patterns on restorer-like genes reveal a conflict between nuclear and mitochondrial genomes throughout angiosperm evolution. Proc Natl Acad Sci U S A. 2011; 108 (4):1723–8. https://doi.org/10.1073/pnas.1007667108 PMID: 21220331