Genome-wide investigation and expression analyses of the pentatricopeptide repeat protein gene family in foxtail millet

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Abstract

Background: Pentatricopeptide repeat (PPR) proteins are encoded by a large gene family of approximately 450 members in Arabidopsis and 477 in rice, which characterized by tandem repetitions of a degenerate 35 amino acid characteristic sequence motifs. A large majority of the PPR genes in the higher plants are localized in organelles. Their functions remain as yet largely unknown. The majority of characterized PPR proteins have been found to function in modulating the expression plastid and mitochondrial genes in plants.

Results: Here, a genome-wide identification and comparison of the PPR genes from 5 organisms was performed, including the moss Physcomitrella patens, the lycophyte Selaginella moellendorffii, the eudicot Arabidopsis, and the monocots rice and foxtail millet. It appears that the expansion of this gene family prior to the divergence of the euphyllophytes and the lycophytes in land plants. The duplication and divergence rates of the foxtail millet PPR genes (SiPPRs) showed that the expansion period of this gene family around 400 Mya, and indicated that genome segmental duplication was very likely the primary mechanism underlying the expansion of the PPR gene family in vascular plants. An analysis of a complete set of SiPPR genes/proteins that included classification, chromosomal location, orthologous relationships, duplication analysis, and auxiliary motifs is presented. Expression analysis of the SiPPR genes under stress conditions revealed that the expression of 24 SiPPR genes was responsive to abiotic stress. Subcellular localization analysis of 11 PPR proteins indicated that 5 proteins were localized to chloroplasts, that 4 were localized to mitochondria, and that 2 were localized to the cytoplasm.

Conclusions: Our results contribute to a more comprehensive understanding the roles of PPR proteins and will be useful in the prioritization of particular PPR proteins for subsequent functional validation studies in foxtail millet.

Keywords: Foxtail millet, Pentatricopeptide repeat (PPR) proteins, Genome segmental duplication, Phylogenesis, Responsive mechanism, Subcellular localization

Background

Foxtail millet (Setaria italica) was domesticated in neolithic China approximately 8,700 years ago. It has been regarded as an important dietary staple in China for many millennia. Although it is closely related to the major food and feed crops maize and sorghum, the required growth conditions and low productivity have limited the potential of foxtail millet [1]. Foxtail millet has a relatively small genome (515 Mb; 2n = 2x = 18); it is a C4 panicoide crop with relatively low amounts of repetitive DNA, a short life-cycle, and is self-pollinating [2, 3]. It has prolific seed production and has close phylogenetic relationships with several biofuel crops such as switch grass (Panicum virgatum), napier grass (Pennisetum purpureum), and pearl millet (Pennisetum glaucum) [1]. Unsurprisingly, considering that foxtail millet is a cereal crop with excellent drought tolerance, there is an extensive germplasm collection available for research and the plant has become a prominent model genetic system for use in the study of the evolution and
physiology of C4 photosynthesis and abiotic stress tolerance mechanisms, particularly for salinity and dehydration stress [3–5]. These features accentuate the potential of using this crop as an experimental model system to investigate the stress resistance mechanisms and of using this crop as a source from which to mine abiotic stress-responsive genes.

The pentatricopeptide repeat (PPR) proteins represent one of the largest protein families in land plants (450 members in Arabidopsis and 477 members in rice). This family was identified serendipitously over a decade ago as a result of bioinformatics analyses of the then incomplete Arabidopsis genome sequence [6]. PPR proteins are characterized by the tandem array of a PPR motif, a highly degenerate unit consisting of 35 canonical amino acids [7–9]. The vast majority of the PPR genes in each of the higher plants with sequenced genomes were found to be lacking introns, and retrotransposition has been proposed as the likely mechanism underlying the expansion of the higher plant PPR gene family [8, 10]. The majority of plant PPR proteins that have been characterized to date function in important roles in a broad range of developmental and physiological processes such as respiration, cytoplasmic male sterility (CMS), embryogenesis, and photosynthesis, and few of these proteins have been proved to play a role in post-transcriptional processes associated with RNA in plant organelles [11–14]. PPR proteins escaped identification until complete genomic sequences became available because their primary sequences are highly degenerate. While subsequent analysis revealed that the PPR proteins are found in a few animal and fungal proteins, but it is clear that the family has expanded greatly in land plants; there are with 450 PPR genes in Arabidopsis and 477 PPR genes in rice (Oryza sativa) [6–8, 15]. The vast majority of these proteins are predicted to be localized to mitochondria or chloroplasts, however, there are also a few PPR proteins that have been shown to be localized to the nucleus [16–18]. Most of the functional analyses of PPR proteins have been performed Arabidopsis, rice, or maize (Zea mays), and these studies revealed that they participate in various post-transcriptional processes related to gene expression in plant organelles [11–14, 19–22]. Functional studies of PPR proteins relating to biotic and abiotic stress response mechanisms in higher plants remain very sparse, and the characterization of these proteins remains one a major challenge in plant science.

Here, we took advantage of the existing research results [7, 8, 10] and the newly completed foxtail millet genome sequencing results to perform a genome-wide comparison of the PPR genes in 5 organisms distributed widely across plant lineages: the moss Physcomitrella patens [23, 24], the lycophyte Selaginella (Selaginella moellendorffii) [25], the dicot Arabidopsis, and the monocots rice and foxtail millet [1]. Our results allow us to draw definitive conclusions on the timing and causes of the expansion of the vascular plants PPR gene family. In addition, our complete analysis of all of the PPR proteins in foxtail millet enables a more comprehensive foundation from which to explore the functional and regulatory networks of PPR genes. Our quantitative real-time PCR (qRT-PCR)-based analysis of these gene expression patterns under specific stress treatment conditions, and our subcellular localization studies of a number of candidate SiPPRs provide empirical data that should greatly facilitate advances in understanding of the biological functions of these fascinating proteins in foxtail millet in particular, and in plants generally.

Results and Discussion
Identification and classification of PPR proteins in foxtail millet

All of the 486 genes encoding PPR proteins in the genome of foxtail millet were identified in this paper. The Phytozone locus, protein length, open reading frame (ORF) length, and each of these PPR genes chromosomal location are listed in Additional file 1: Table S1. Foxtail millet PPR gene/protein sequence informations, and gene annotations in this study were downloaded from the Phytozone (http://phytozone.jgi.doe.gov/pz/portal.html).

To better understand the phylogenetic relationships of the PPR protein-endoding genes among land plants, we identified 1670 nonredundant putative PPR proteins in the Selaginella, which is a nonseed vascular plant with a dominant and complex sporophyte generation of an ancient lineage that diverged shortly after land plants evolved vascular tissues [25] (Additional file 2: Table S2). The numbers of PPR genes in foxtail millet, Arabidopsis, and rice were found to be strikingly similar. There were approximately three times as many PPR genes in Selaginella than in foxtail millet. The Selaginella plastome sequence has a large amount of RNA-edited sites, which coincides with the exceptionally large number of PPR genes in Selaginella [25].

Given the structure of the repeated motif, PPR gene family can be split into two subfamilies; these are termed the P and PLS subfamilies. On the basis of the presence of the C-terminal conserved domains, the PLS subfamily can be further split into the following three subgroups: PLS, E/E+ and DYW subgroups [8]. Figure 1a details the numbers of PPR genes split by different species and subgroups and in the context of the phylogeny of plants. A large majority of the 1403 Selaginella PPRs, and 263 SiPPRs, belong to the P subfamily; 271 Selaginella PPRs, and 223 SiPPRs, belong to the PLS subfamily, including 39 and 32 PLS subgroups, 139 and 80 DYW subgroups, 93 and 111 E/E+ subgroups PPR genes were found in Selaginella and foxtail millet, respectively.
Gene structure and chromosomal distribution of PPR genes in foxtail millet

The structures of PPR genes were determined by analyzing their exon-intron organization. The great majority of the foxtail millet PPR genes contained very few introns, as is the case with the vast majority of the Selaginella PPR genes. Figure 2 details the ratios of PPR genes in two species that contain no intron, 1 intron, 2–5 introns, and 6 or more introns. Approximately 80 % of the Arabidopsis and rice PPR genes lack introns [8, 10, 15]. Intron-rich PPR genes may represent “ancient” PPR genes previous to the occurrence of retrotransposition-mediated expansion of the PPR gene family in land plants [8, 10]. 79.94 % (1335 of 1671) and 78.60 % (382 of 486) of PPR genes of, respectively, Selaginella and foxtail millet, were predicted to lack introns. The 486 PPR genes are unevenly distributed on all 9 of the foxtail millet chromosomes. Chromosome 9 of foxtail millet possesses the highest number of SiPPRs [99 (20.49 %)], while the lowest number of SiPPRs were found on chromosome 8 [26 (5.34 %)]. For fine mapping, the exact position (in bp) of each SiPPR on the respective foxtail millet chromosome was obtained from Phytozome and is indicated diagrammatically in Fig. 3 (the exact position in bp is given in Additional file 1: Table S1). Substantial clustering of PPR genes was evident on all of the chromosomes. In total, 180 (~36.96 %) PPR genes were arranged in tandem repeats of two to seven genes, either in the same or the inverse direction (the distance between these genes being less than 100 kb), representing localized gene (tandem) duplications.

Comparing the numbers and structures of the PPR genes in all 5 species, including moss, Selaginella, Arabidopsis, rice, and foxtail millet, the numbers of Selaginella PPR genes were approximately three times as many as the other vascular plants. However, all homologous PPR genes in flowering plants also could be found in Selaginella, and the intron-rich PPR genes show similar structure in all 5 species, which suggested the PPR gene family expanded before the divergence of the euphyllophytes (ferns and seed plants) and the lycophytes (~400 Mya). The PPR genes may have been derived from a primitive
and intron-rich “ancient” PPR gene by retrotransposition [7, 8, 10, 24]. Further, we speculate that some PPR genes may be non-functional, or may even have negatively affected the transition of non-flowering to flowering in vascular plants. The expansion of the PPR gene family may be coupled with organelle evolution.

6688 (~19%) of the genes of the foxtail millet genome were segmentally duplicated [26]. Among the SiPPR genes, 190 (95 pairs; ~39.01%) were segmentally duplicated (connected by black lines in Fig. 4); note that this proportion is far higher than that of genes generally. In particular, there were several genes that were duplicated a dramatic number of times, such as the gene Si020204m (located on chromosome 1), which was duplicated eleven times, forming 66 paralogs that can be found on all 9 of the foxtail millet chromosomes (Fig. 4). These results
showed that genes segmental duplications were important in the expansion of the PPR gene family in vascular plants.

**Gene ontology annotation**

Blast2GO software performed the GO slim analysis, which suggested the putative participation of SiPPRs in various biological processes, molecular functions and cellular components (Fig. 5; Additional file 3: Table S3). Out of the 486 SiPPR proteins, GO annotation of 243 sequences could not be found, and the remaining 243 SiPPRs were divided into 9 separate categories of biological processes. The results indicated that a majority of SiPPRs were likely related to primary biological process [168 (~69.13 %)], followed by post-transcriptional processes within mitochondria and chloroplasts [39 (~16.04 %)] (including RNA editing, RNA splicing, RNA cleavage and translation), and embryo development processes [17 (~7 %)]. Of note, 4 SiPPRs were predicted to participate in responses to stress, a topic that we will address later in this text. The molecular functions of 112 (~46.09 %) SiPPRs were predicted to participate in molecular function, such as basic catalysis or binding function. The second most frequently annotated molecular function was nucleic acid binding function [10 (~4.11 %)], which is in agreement with the molecular role of PPRs in binding RNA [17, 27–30]. Cellular localization prediction suggested that 203 (~83.53 %) SiPPR proteins were localized in the mitochondria, 22 (~9.05 %) were localized to chloroplasts, and 19 (7.1 %) SiPPRs were not sure localized in mitochondria or chloroplasts, 6 were localized in cytoplasm, 3 were localized in the cell nucleus, and 2 were localized in cell membranes (Fig. 5; Additional file 3: Table S3). The GO analysis results indicating that the SiPPR proteins participate in diverse biological processes, molecular functions and cellular components and will be useful for further verifications of gene functions in foxtail millet.

**Orthologous relationships**

We explored the SiPPRs orthologous relationships by using a comparative mapping-based approach in which
the physically mapped foxtail millet PPR genes were compared with those PPR genes in the chromosomes of other related grass genomes, including *B. distachyon*, rice, sorghum, and maize (Table 1; Additional file 4: Table S4 and Additional file 5: Figure S1). The specific orthologous relationships could be deduced on a mean value for ~ 65.46 % proteins. Maximal orthology of SiPPRs annotated on the *Setaria italica* chromosomes was found in *B. distachyon* (74.67 %), followed by rice (66.84 %), sorghum (66.36 %), and maize (54.01 %). The widespread synteny among *Setaria italica*, *B. distachyon*, rice, sorghum, and maize at gene level supports their close evolutionary relationships [1, 31–35]. Of note, most of the SiPPR genes had syntenic tend to special chromosomes of the four species, a finding consistent with what is known about the evolution of the *Setaria italica* genome [1, 33]. The two chromosome reshuffling events that occurred in foxtail millet, rice, and sorghum were reported [1]. Our results also suggested there may be several chromosome reshuffling events between foxtail millet and *B. distachyon* and maize, such as fusing *B. distachyon* chromosomes 1 and 4, 3 and 9 and 2 and 4 to foxtail millet chromosomes 2, 9 and 3, respectively, and maize chromosomes 4 and 5, 2 and 7 and 6 and 9 to foxtail millet chromosomes 1, 2 and 4, respectively. The comparative mapping information thus offers a helpful introduction for comprehending the evolution of PPR genes among those grasses, even the genome evolution of *Setaria italica*. In addition, this research would be helpful for the selection of candidate SiPPR genes and exploit them in genetic improvement of

**Table 1** A summary of comparative mapping of foxtail millet SiPPR genes on *Brachypodium sylvaticum*, sorghum, maize and rice

| *Setaria italica* | *Brachypodium distachyon* | *Oryza sativa* | *Sorghum bicolor* | *Zea mays* |
|-------------------|--------------------------|----------------|-------------------|------------|
| Chr1              | Chr(377.96 %)            | Chr(77.96 %)   | Chr(74.58 %)      | Chr(16.94 %),Chr(52.54 %) |
| Chr2              | Chr(61.40 %),Chr(24.56 %)| Chr(77.89 %),Chr(21.05 %) | Chr(84.21 %)      | Chr(28.07 %),Chr(49.12 %) |
| Chr3              | Chr(252.08 %),Chr(25 %)  | Chr(47.91 %),Chr(20.83 %) | Chr(941.67 %)     | Chr(33.33 %)           |
| Chr4              | Chr(76.19 %)             | Chr(59.52 %)   | Chr(1071.42 %)    | Chr(26.19 %),Chr(930.95 %) |
| Chr5              | Chr(278.2)               | Chr(73.07 %)   | Chr(70.51 %)      | Chr(39.74 %),Chr(830.76 %) |
| Chr6              | Chr(85.18 %)             | Chr(874.07 %)  | Chr(70.37 %)      | Chr(25.92 %),Chr(425.92 %) |
| Chr7              | Chr(478.2 %)             | Chr(450 %)     | Chr(650 %)        | Chr(2636.95 %)         |
| Chr8              | Chr(5769)                | Chr(142.30 %)  | Chr(6153 %)       | Chr(3461 %)            |
| Chr9              | Chr(960 %),Chr(326 %)    | Chr(353 %),Chr(1024 %) | Chr(73 %)         | Chr(155.01 %)          |

The percentages indicated among parentheses indicate the fraction of loci with conserved syntenic relationships
related grass family members. As for example, *Arabidopsis*
PpR40 as a signaling link factor for the communication of
mitochondrial electron transport and regulation of stress
and hormonal responses. Insertion mutations inactivating
PpR40 affected plant growth and enhanced mutants
sensitivity to salt, ABA, and oxidative stress [19]. There-
fore, its orthologous *Setaria italica* gene (Phytozome ID:
Si019461m) and rice gene (LOC_Os12g37100.1) may also
have the similar function.

**Duplication and divergence rates of the SiPPR genes**

Gene duplications enable the evolution of genes with
novel functions. Functional deficiency phenotypes occur
less frequently when multiple copies of a gene are present
then when there is only a single copy of a gene. Genes
multiple copies may evolve new gene owing to evolution-
ary events such as the whole genome location (tandem)
and segmental duplications. Such gene duplication has
been described in previous reports [26, 36, 37]. Therefore,
we searched that the effect of Darwinian positive selection
in the duplication and divergence of the PPR genes. The
 synonymous (Ks) and non-synonymous (Ka) nucleotide
substitution rates per site were determined for 52 location
and 28 segmental duplication gene-pairs and between
orthologous gene-pairs of SiPPR with those genes of moss
(28 pairs), Selaginella (28 pairs), *Arabidopsis* (28 pairs),
rice (28 pairs), and maize (28 pairs) (Fig. 6; Additional
file 6: Table S5, Additional file 7: Table S6, Additional
file 8: Table S7, Additional file 9: Table S8, Additional
file 10: Table S9, Additional file 11: Table S10 and Add-
tional file 12: Table S11). The non-synonymous/syn-
onymous ratio (Ka/Ks or ω) is a measure of natural
selection acting on a protein, in which values of ω < 1
indicate negative purifying selection; ω = 1 indicates neu-
tral evolution; and ω > 1 indicates positive diversifying
selection. The mean number of synonymous substitutions
(Ks) across all tandemly duplicated gene pairs was 21.69
and segmentally duplicated gene pairs were 5.81. The
mean number of non-synonymous substitutions (Ka)
across all tandemly duplicated gene pairs was 0.9662,
and that for segmentally duplicated gene pairs was 0.29662.
Even though the ω values extensive changes
among gene pair groups, the average value was equal to
0.16631 and 0.37643 for tandemly duplicated gene pairs
and segmentally duplicated gene pairs, respectively,
suggesting that those duplicated SiPPRs were under
very strong selection pressure, which had undergone
substitution elimination and enormous selective con-
straint by natural selection during the course of evolution
since ω < 1. Our analysis indicated that the duplication
events leading to these location and segmental duplication
SiPPRs may have happened approximately 2123.51 Mya
and 447.23 Mya (Fig. 6), respectively. Presumably, the PPR
family is a quite ancient gene family and may have existed
as early as in the era of bacteria and algae (3500–1500
Mya), and PPRs were also found in ancient heterokont
species (brown algae, diatoms) and bacterial species
[7, 38]. However, there were no eukaryotes in the world at
that time, the PPR genes family may have derived from
*Bacteria* and/or *algae*, a supposition that is supported by
the hypothesis of the endosymbiosis of a green algae into
an ancestral host cell [39, 40]. Secondly, tandem duplica-
tion enabled the evolution of new functions among the
PPR genes in evolutionarily diverse taxa (2157 Mya).
The result of comparing the numbers and nature of the
PPR genes in all 5 species indicated that the PPR gene
family expanded before the divergence of the euphyllo-
phytes (ferns and seed plants) and the lycophytes (~400
Mya), and segmental duplication of SiPPR genes occurred
around 447.23 Mya, which suggests that genome segmen-
tal duplications may be responsible for the PPR gene ex-
pansion in vascular plant (Additional file 7: Tables S6).

Among the SiPPR orthologous gene-pairs with those
of other four species, moss, Selaginella, *Arabidopsis*, rice,
and maize, the value of $\omega$ was maximum between rice and foxtail millet (0.23), maize and foxtail millet (0.27), and least for moss and foxtail millet (0.032), Selaginella and foxtail millet (0.026), *Arabidopsis* and foxtail millet (0.046) (Additional file 8: Table S7, Additional file 9: Table S8, Additional file 10: Table S9, Additional file 11: Table S10 and Additional file 12: Table S11). The relatively higher $\omega$ between the SiPPR genes of maize and foxtail millet and rice and foxtail millet show their earlier divergence around 34.6 Mya and 51.9 Mya, the divergence time is in agreement with the recent time of divergence of maize-foxtail millet (~27 Mya) and rice-foxtail millet (~48 Mya, Fig. 6) [1]. However, the divergence time of moss-foxtail millet (~2947.9 Mya), Selaginella-foxtail millet (~2957.9 Mya) and *Arabidopsis*-foxtail millet (~2058.7 Mya) were far earlier than the time of divergence of monocots and dicots, which suggested the PPR gene family may share some common “ancient” PPR genes in land plants.

The estimation of tandem duplication time (~2123.51 Mya) of foxtail millet PPR genes is later than the divergence time of moss-foxtail millet (~2947.9 Mya), Selaginella-foxtail millet (~2957.9 Mya), and *Arabidopsis*-foxtail millet (~2058.7 Mya), but earlier than maize-foxtail millet (~34.6 Mya) and rice-foxtail millet (~51.9 Mya). Segmental duplication time (~447.23 Mya) of foxtail millet PPR genes is later than the divergence time of moss-foxtail millet (~2947.9 Mya), Selaginella-foxtail millet (~2957.9 Mya), and *Arabidopsis*-foxtail millet (~2058.7 Mya), but earlier than maize-foxtail millet (~34.6 Mya) and rice-foxtail millet (~51.9 Mya). Remarkably, the orthologous PPR gene-pairs between moss-foxtail millet (0.032), Selaginella-foxtail millet (0.026), and *Arabidopsis*-foxtail millet (0.046) appear to have undergone extensive intense purifying selection, as compared to average $\omega$ values of the foxtail millet-rice ($\omega = 0.23$) and foxtail millet-maize ($\omega = 0.27$) PPR gene pairs. Consequently, we can draw a conclusion that the tandemly and segmentally duplicated events, including the divergence events of the PPRs from other species, have played a critical role in the evolution process of this gene family in foxtail millet, and genome segmental duplications may be responsible for the PPR gene expansion in vascular plant.

**Functional conservation of PPR proteins in foxtail millet**

Most functional analyses of PPR proteins undertaken to date have only tested proteins from *Arabidopsis*, rice, or maize, and there is as yet very limited information regarding the evolutionary relationships between PPR proteins within or across species. The PPR gene family existed in the era of bacteria and green algae, and PPR proteins are mostly aimed at mitochondria or plastids in terrestrial plants [8], it has been suggested that the origin and evolution of the PPR gene family may be inextricably coupled with the endosymbiosis of mitochondria and chloroplasts in the early evolutionary history of eukaryotes and plants [7, 8].

PPR proteins can be classified based on motif structure(s) and the presence of additional C-terminal domains. Subgroupings include the P and PLS subfamilies and the P-class PPR proteins presented the presence of one or more known/unknown functional motifs or domains, such as an RNA recognition motif (RRM) [41] and LAGLIDADG domain [42]. Those PPRs with the same domain always are intron rich (1 to 26) and tended to be located in the same organelles in moss, Selaginella, *Arabidopsis*, rice and foxtail millet (Additional file 13: Table S12), respectively. Comparison of the gene structures and amino acid sequences encoded by PPR proteins containing the same domains showed that these were well conserved among the five species (Fig. 7), indicating that these attributes had been preserved during evolution. This suggested that these homologous PPRs likely share the same or similar origins and functions, and further suggest that these the intron-rich PPR genes of moss may represent “ancient” PPR genes previous to the occurrence of retrotansposition-mediated [10, 24] or/and segmental duplication-mediated the PPR family expansion in terrestrial plants. Such conservation was likely very important for the survival and development of the chloroplast and mitochondria in these 5 species, perhaps even in all land plants. For instance, the PPR-SMR motif containing proteins of *Arabidopsis* (pTAC2, SVR7, and AT5G46580) and the corresponding maize orthologs (Zm-pTAC2, ATP4, and PPR53) were found to be localized to the chloroplast and to have the same or similar functions [43–46]. The *Arabidopsis* protein (OTP51) contains two LAGLIDADG motifs that are known to specifically promote the splicing of introns, and loss of AtOTP51 affected the assembly of the PSI complex [47]. The rice protein (OsOTP51) was found to have a similar function with AtOTP51, and loss-of-function OsOTP51 affected the intron splicing of a number of plastid genes, indirectly affected the structure and functions of PSI, and resulted in severe photoinhibition and eventual plant death, even when mutant plants were grown under very low light [13]. This example of conservation suggests that such homologous PPR proteins are important for the growth and development of plants and they preserved their ancient functions.

In order to investigate the functional conservation of the PPR proteins in foxtail millet, we examined the presence of PPR-LAGLIDADG proteins in a broad range of species to explore their origins and diversification. The PPR-LAGLIDADG proteins were essentially only found in eukaryotic species and were largely limited to the Viridiplantae clade. We then investigated the function of SiOTP51. It contains a PPR motif and an LAGLIDADG_2
motif. We transformed *SiOTP51* into *Arabidopsis atotp51-1* and *atotp51-2* mutants using a plasmid containing *SiOTP51* under the control of the 35S cauliflower mosaic virus promoter. Homozygous *atotp51-1* and *atotp51-2* mutants exhibited identical phenotypes: seedlings were light straw yellow; rosettes were light yellow in standard light and pale green for young leaves in low light conditions; plants had retarded growth and greening [47] (Fig. 8). Complementation mutants of *atotp51* with the *Arabidopsis AtOTP51* gene ortholog *SiOTP51* showed that *SiOTP51* would to a certain extent reverse the mutant phenotypes. This would suggested that *SiOTP51* may also affect photosynthesis and plastid gene expression. The molecular functions of *OTP51s* in all land plants may be conserved.

**Expression of SiPPR genes under abiotic stress conditions and abscisic acid (ABA) treatment**

The exposure of plants to changes in environment conditions and the treatment of plants with phytohormones is known to elicit various physiological, biochemical, and molecular responses, causing alterations in gene expression [36]. Some PPRs have been proved to function in post-transcriptional and post-translational processes and have been classified as sequence-specific RNA-binding proteins [7, 8, 48]. However, the function of the vast majority of PPR proteins remain to be characterized, and there is very limited information about this family in higher plants. This dearth of information is especially pronounced regarding the potential function of these proteins in plant responses to abiotic stresses and phytohormone treatments. To explore the potential functions of PPR proteins in stress and/or phytohormone responses in higher plants, the expression of the foxtail millet PPR genes was analyzed in plants that had been grown under drought treatment (20 % PEG 6000). We performed an RNA-seq experiment of total RNA isolated from 4-week-old foxtail millet seedlings, including leaves and roots (Beijing Genomics Institute, BGI). 51 PPR genes were identified which were differentially expressed (≥2-fold difference, p value less than 0.05) between plants grown under drought stress conditions and control plants. A heat map was got according to the RPKM values for the 51 genes. 41 genes were up-regulated; 10 genes were down-regulated.
genes (Additional file 14: Figure S2 and Additional file 15: Table S13). Then we examined the expression patterns of 31 PPR genes, including 21 PPR genes that were up-regulated expression, 6 PPR genes were down-regulated expression, and 4 selected from GO annotation (Additional file 16: Table S14), using qRT-PCR analysis of samples from plants, including leaves and roots, subjected to drought, salt, cold, and ABA treatment. A heat map illustration of expression patterns of 31 selected SiPPR genes is exhibited in Fig. 9. The qRT-PCR results of most candidate genes under abiotic stress condition(s) were agreement with the RNA-seq experiment under drought treated (Fig. 9). Seven of these genes were up-regulated and three were down-regulated; four showed no-response under any of the treatments (Fig. 9a). Three were up-regulated under salt and cold, but down-regulated under drought and ABA; Four were up-regulated under salt, cold, and ABA, but no response under drought stress; four PPR protein-encoding genes were up-regulated under salt and cold stresses, but no response under drought stress and ABA; one showed up-regulation under drought stress and cold, but no response under salt and ABA (Fig. 9b). Three were up-regulated under salt treatment, one was down-regulated under ABA, one was up-regulated under drought stress, and one was up-regulated under cold stress (Fig. 9c).

Of the 450 predicted PPR proteins in Arabidopsis, only five mitochondrion-localized proteins and one cytosol-nucleus dual-localized PPR protein were proved to function in response to abiotic stresses and/or ABA. Arabidopsis PPR40-mediated ubiquinol-cytochrome c oxidoreductase activity in mitochondrion complex III and it was related to oxidative respiration that also conducive to abiotic stress tolerance in Arabidopsis [19]. ABO5 was required for cis-splicing of the mitochondrial nad2 intron 3 (nad2 is one subunit in mitochondrion complex I) and altered the expression of several stress-responsive and nuclear-encoded genes to affect the ABA signaling pathway [11]. The PGN can regulate the ROS homeostasis in plants mitochondria during abiotic and biotic stress responses may occur through the regulation of mitochondria-nucleus retrograde signaling. AHG11 regulates the nad4 (mitochondrion complex I) transcriptional level and thus led to changes in oxidative levels by controlling RNA editing events in plants mitochondria and affecting plant responses to ABA [21]. SLG1 regulated the nad3 (mitochondrion complex I) transcript by regulating RNA editing events in mitochondria and affecting the expression of genes which were involved in the alternative respiratory pathway [22]. SOAR1 (cytosol-nucleus dual-localized) was identified recently as a crucial regulator in the CHLH/ABAR (Mg-chelatase H subunit/putative ABA receptor)-mediated signaling pathway that acts downstream of CHLH/ABAR and upstream of a nuclear ABA-responsive bZIP transcription factor, ABI5 [18]. qRT-PCR results showed that 11 SiPPRs were up-regulated under any three abiotic stress treatment (Fig. 9). These SiPPRs may have a similar stress response mechanism with Arabidopsis PPRs [11, 18–22]. In addition, the class of PPR proteins that can edit RNA are of the PLS subfamily, and the E+ motif is required for the conversion of C to U RNA editing [49]. Consistent with their functions, six stress-related PPR proteins, including PGN, AHG1, and SLG1, belonged to the E subgroup, and PPR40, ABO5, and SOAR1 belonged to the P

![Fig. 8](Image)

The phenotype of Arabidopsis seeding at two weeks old under low light conditions, which transformed SiOTP51 into atotp51-1 and atotp51-2 mutants. (a) Schematic representation of ATOTP51 and SiOTP51 showed the gene or protein structure. Boxes and lines indicate exons and introns, respectively. (b) Positions of the two T-DNA insertions in the atotp51 gene. Salk_112013 was named atotp51-1 and Sail_713D08 was named atotp51-2. (c) Phenotype of two atotp51 mutant seedling and reverse mutation of atotp51 mutants with SiOTP51 gene in low light conditions at two weeks old.
It has been suggested that the mitochondrial localization of P subgroup or E subgroup PPR proteins may participate in plant responses to biotic and abiotic stress.

Subcellular localization of foxtail millet PPR proteins

Based on the expression of SiPPR genes in plants grown under abiotic stress conditions, 11 candidates PPR proteins were selected for verification of the subcellular localization, including 7 up-regulated under any of the treatments, and 4 up-regulated under salt, cold, and ABA. Among these 11, 4 proteins were predicted by TargetP to be localized to chloroplasts, 6 proteins were predicted to be targeted to mitochondria, and 1 protein was predicted to be localized to cytoplasm [50]. Those foxtail millet PPR genes were cloned and inserted into a subcellular localization vector that included a GFP protein-encoding gene under the control of the 35S promoter; these vectors were transformed into foxtail millet protoplasts.

To confirm that the putative mitochondrion-targeted PPR proteins were expressed in the mitochondria, we stained transformed foxtail millet protoplasts with a mitochondria-specific dye [51], and then observed the samples with 488 and 543 nm illumination. 5 of these proteins were localized to chloroplasts and 4 were localized to mitochondria and 2 located to the cytoplasm (Fig. 10). Among 11 SiPPR genes, 6 belong to the P subfamily and 5 belong to the PLS subfamily. Most of PPRs located to chloroplasts or mitochondria were consistent with the results predicted by TargetP. However, Si032871m and Si006059m, which were predicted to be localized to mitochondria, were localized to the cytoplasm. PPR proteins involved in abiotic stresses and ABA were localized to the cytoplasm and/or mitochondria [11, 18–22].

These results suggest that the 4 mitochondrion-localized genes and the 2 cytoplasm-localized SiPPR proteins may be viewed as candidates implicated in plant responses to adverse environmental conditions and exogenous phytohormones.

Chloroplasts and mitochondria are the two main plant cellular organelles with independent genomes. Under unfavorable conditions, the accumulation of ROS in both chloroplasts and mitochondria increases oxidative stress and this acts as a signal to help the plant respond to unfavorable environmental conditions. It is known that PPR proteins are mostly aimed at plastids and mitochondria in plants, and the mitochondrial/chloroplast PPR proteins play various and crucial roles in plant developmental processes and responses to environmental stresses [6, 8, 16, 52–61]. However, there is not much information about the chloroplast localization of PPR proteins in response to abiotic stresses. To date, the only reported chloroplast-localized PPR protein, GUN1, was found to be a central regulator of plastid to nucleus retrograde signaling, and its impairment led to ABI4-mediated repression of nuclear-encoded genes and enhanced sensitivity to ABA [57, 62]. The subcellular localization assays of 11 foxtail millet PPR proteins in our study revealed that 5 members were...
located to chloroplasts, and these candidate proteins seem to be responsive to diverse abiotic stresses (Figs. 9 and 10).

Conclusions
Pentatricopeptide repeat (PPR) proteins are extensive in terrestrial plants, and the genomes of terrestrial plants typically include several hundred genes encoding these proteins. Most of PPRs are targeted to plastids and mitochondria, where they play crucial roles in mechanisms controlling post-transcriptional processes. However, functional studies of PPRs in terrestrial plants remain scarce. This study provides evidence that the period of the expansion of the PPR gene family occurred prior to the divergence of the euphyllophytes (ferns and seed plants) and the lycophytes (~400 Mya). Our results also suggest that genome segmental duplications were responsible for the expansion of the PPR gene family in vascular plants. In addition, we conducted a comprehensive analysis of the SiPPR genes/proteins that included classification, chromosomal location, orthologous relationships, duplication analysis, and auxiliary motifs. We conducted expression pattern analysis of 31 candidate SiPPR genes in plants grown under stress conditions, and performed subcellular localization analysis of 11 PPR proteins. Our study establishes an empirical foundation for further investigations seeking to elucidate the precise role of individual PPRs in foxtail millet.

Methods
Database searching
The Hidden Markov Model (HMM) profile of PPR motif (PF01535) was acquired from the Pfam v28.0 database (http://Pfam.sanger.ac.uk/), and was queried against the HMMER (http://www.ebi.ac.uk/Tools/hmmer/) of foxtail millet, with a threshold E-value ≤ 10. Ultimately, 486 and 1670 nonredundant PPR proteins in foxtail millet and lycophyte Selaginella, respectively, displayed the presence of PPR motif with confidence (E-value less than 1.0) in Pfam searches, and SMART (http://smart.embl-heidelberg.de/). The sequence data of the PPR genes and the gene annotations
used in this study were for moss, *Arabidopsis*, and rice as obtained according to the previous description [8, 10]. The hmmssearch program from the HMMER package (Eddy 1998) was applied to the translated sequence data to identify clusters of all of the PPR motifs (P, L, S, L2, E/E+, and DYW).

Physical locations, gene structure, and genomic distribution
All the sequenced contigs of foxtail millet have been physically constructed as pseudo molecules at Phytozome v10 (http://phytozome.jgi.doe.gov/pz/portal.html). Subsequently, each of the PPRs were positioned on the nine foxtail millet chromosome pseudo molecules based on their ascending order of physical position (bp) according to BLASTN searching against the Phytozome database adopting default settings. The analytical tools available at the Plant Genome Duplication Database were used to identify segmentally duplicated PPR gene in foxtail millet [63], and detailed methods according to the previous description [64, 65]. Segmental duplications were drew using Circos 0.67 (http://circos.ca) [66]. Tandem duplications were characterized according to the previous description [65]. Manual curation and assessment of the numbers of introns and exon-intron positioning of the genes was based on comparison of the full-length cDNA or the predicted coding sequence (CDS) of SiPPR genes with their corresponding genomic sequence.

Gene ontology (GO) annotation
Blast2GO (http://www.blast2go.com) software was used to perform the functional annotation of the SiPPR gene sequences and the subsequent analysis of annotation results, and detailed methods according to the previous description [67].

Comparative physical mapping of PPRs between foxtail millet and other grass species
The comparative orthologous relationships of the PPR proteins between foxtail millet and other grass species were performed according to the previous description by Mishra et al. (2014) and Lata et al. (2014) [26, 37], and visualized by using Circos 0.67 [66].

Estimation of duplication and divergence rates
The corresponding amino acids and their cDNA sequences of paralogous and orthologous gene-pairs of the SiPPR proteins in moss, *Selaginella, Arabidopsis*, rice, and maize were aligned using a ClustalW-based multiple sequence alignment tool and analyzed using the tool of PAL2NAL (http://www.bork.embl.de/pal2nal/) [68]. Time (million years ago, Mya) of duplication and divergence of each SiPPR gene was calculated according to the previous description [69, 70].

Functional conservation of PPR proteins in foxtail millet
To create the complementation construct pSiOTP51, the full-length opening reading frames of *SiOTP51* (*Si016353m*) was obtained from Yugu 1 cDNA with the primers 5′-ATCCAAAGGTATCTCCCTCCAGC-3′ and 5′-GTGAAACATACTGTCTTTACCT-3′, and the resulting fragments were inserted into the plant binary vector pCAMBIA1302, in which the *SiOTP51* gene was under the control of the CaMV 35S promoter. The constructs were introduced into the *atotp51-1* and *atotp51-2* mutant by *Agrobacterium tumefaciens*-mediated transformation as described previously [71]. The *Arabidopsis* Columbia-0 (Col) T-DNA mutants and wild-type plants were germinated on ½ MS medium, with 2 % sucrose, at 22 °C, with a 16-h photoperiod and a light intensity of 50 μmol photons/m². Seedlings were transferred to Nutrition Soil (Pindstrup Mosebrug A/S) with vermiculite (nutrition soil:vermiculite, 2:1) and grown with the same conditions described above.

RNA-seq experiment of drought-treated foxtail millet
Isolation of foxtail millet seedlings total RNA using TRIzol reagent (Invitrogen) and treated with RNase free DNase I (Takara) according to the manufacturer’s recommendations. Subsequently, RNA-seq experiment of drought-treated foxtail millet was completed by Beijing Genomics Institute, and detailed description according to the previous description [72].

Plant materials and stress treatments
Seeds of foxtail millet Yugu 1 were grown in a growth chamber (MGC-350HP-2; Blue Leopard) at 28 ± 1 °C day/23 ± 1 °C night with 70 ± 5 % relative humidity and a photoperiod of 14 h. For stress treatments, 21-day-old seedlings, including leaves and roots, were exposed to drought (20 % PEG 6000), salt (250 mM NaCl), cold (4 °C), or 100 μM ABA treatment for 0, 6 and 24 h. Unstressed plants were maintained as controls. After the treatments, the seedlings were immediately frozen in liquid nitrogen and stored at −80 °C until RNA isolation. These experiments were repeated thrice.

RNA extraction and qRT-PCR analysis
Total RNA from Yugu 1 was extracted with an RNA extraction kit (Takara) according to the manufacturer’s instructions. The cDNA synthesis was conducted as previous description [73]. qRT-PCR for examination were carried out with TransScript® II Probe One-Step qRT-PCR Super-Mix (TransGen Biotech) followed by amplification on an ABI 7500 System. The qRT-PCR primers used in this study are listed in Additional file 17: Table S15.
Additional files

Additional file 1: Table S1. PPR genes in foxtail millet. Detailed genomic information, including domain/class present, ORF length, predicted protein length, genomic locus (chromosomal location), number of introns within the ORF, and the subcellular localization of the PPR proteins for each PPR gene. (DOCX 62 kb)

Additional file 2: Table S2. PPR genes in Selaginella. The domain/class, open reading frame length, predicted protein length, chromosomal location, number of introns within the ORF, and the subcellular localization of the PPR proteins of each gene are listed. (DOCX 194 kb)

Additional file 3: Table S3. Blast2Go annotation details of SiPPR protein sequences. (XLSX 28 kb)

Additional file 4: Table S4. The physically mapped PPR genes of foxtail millet were compared with those in the chromosomes of other related grass genomes, including B. distachyon, rice, sorghum, and maize. (XLSX 133 kb)

Additional file 5: Figure S1. Localization analysis of the PPR proteins in foxtail millet. (DOCX 14 kb)

Additional file 6: Table S5. The Ka/Ks ratios and the estimated divergence times for putatively tandemly duplicated SiPPR genes. (DOCX 20 kb)

Additional file 7: Table S6. The Ka/Ks ratios and the estimated divergence times for putatively segmentally duplicated SiPPR genes. (DOCX 30 kb)

Additional file 8: Table S7. The Ka/Ks ratios and the estimated divergence times for orthologous SiPPR proteins between foxtail millet and moss. (XLSX 29 kb)

Additional file 9: Table S8. The Ka/Ks ratios and the estimated divergence times for orthologous SiPPR proteins between foxtail millet and Arabidopsis. (XLSX 36 kb)

Additional file 10: Table S9. The Ka/Ks ratios and the estimated divergence times for orthologous SiPPR proteins between foxtail millet and A. thaliana (setaria). (XLSX 36 kb)

Additional file 11: Table S10. The Ka/Ks ratios and the estimated divergence times for orthologous SiPPR proteins between foxtail millet and rice. (XLSX 38 kb)

Additional file 12: Table S11. The Ka/Ks ratios and the estimated divergence times for orthologous SiPPR proteins between foxtail millet and maize. (XLSX 34 kb)

Additional file 13: Table S12. The PPRs with putative auxiliary motifs or domains in moss, Selaginella, Arabidopsis, foxtail millet, and rice. The domain/class, number of introns within the ORF, and the subcellular localization of the PPR proteins are provided for each PPR gene. (DOCX 29 kb)

Subcellular localization of foxtail millet PPR proteins

11 candidate SiPPR genes were amplified from the cDNA of Yugu 1 with the primers listed in Additional file 18: Table S16. These genes were inserted into the subcellular localization vector p16318, which contains the CaMV 35S promoter and a C-terminal green fluorescent protein (GFP) domain [73]. The transient expression assays were performed as previously described [73, 74]. Transfected foxtail millet protoplasts were stained with a mitochondria-specific dye (MitoTracker Orange [Invitrogen catalogue no. M7510]) [51] and then observed with 488 and 543 nm illumination using a Zeiss LSM700 microscope.

Additional file 14: Figure S2. The heat map of the expression of 51 PPR genes in foxtail millet under drought treatment (20 % PEG 6000) based on RPKM values. (TIF 163 kb)

Additional file 15: Table S13. RNA-seq analysis of PPR gene expression under drought stress treatment (20 % PEG 6000) in foxtail millet. (XLSX 15 kb)

Additional file 16: Table S14. PPR genes differentially expressed under drought treatment (20 % PEG 6000), salt treatment (250 mM NaCl), cold treatment (4 °C), and with 100 μM ABA, at the 0, 6, and 24 h time points. (DOCX 19 kb)

Additional file 17: Table S15. Primer sequences used for real-time PCR analysis. (DOCX 15 kb)

Additional file 18: Table S16. Primer sequences used for the subcellular localization analysis of the PPR proteins in foxtail millet. (DOCX 14 kb)

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Availability of data and material

All data generated or analysed during this study are included in this published article and its supplementary information files.

Authors’ contributions

ZSX coordinated the project, conceived and designed experiments, and edited the manuscript; JML performed experiments; analyzed data and wrote the first draft of the manuscript; PPL and WWL performed experiments; MC provided analytical tools and managed reagents; CHG contributed with valuable discussions; YZM coordinated the project and contributed with valuable discussions. All authors have read and approved the final version of the manuscript.

Competing interests

The authors declare they have no competing interests.

Consent for publication

Not applicable.

Ethics approval and consent to participate

Not applicable.

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