Overexpression of OsFTL10 induces early flowering and improves drought tolerance in Oryza sativa L.

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ABSTRACT

Flowering time control is critically important for the reproductive accomplishment of higher plants as floral transition can be affected by both environmental and endogenous signals. Flowering Locus T-like (FTL) genes are major genetic determinants of flowering in plants. In rice, 13 OsFTL genes have been annotated in the genome and amongst them, Hd3a (OsFTL2) and RFT1 (OsFTL3) have been studied extensively and their functions are confirmed as central florigens that control rice flowering under short day and long day environment, respectively. In this report, a rice OsFTL gene, OsFTL10, was characterized, and its function on flowering and abiotic stress was investigated. The expression level of OsFTL10 was high in young seedlings and shown to be induced by GA3 and drought stress. Overexpression of OsFTL10 resulted in earlier flowering in rice plants by up to 2 weeks, through up-regulation of the downstream gene OsMADS15. OsFTL10 also regulated Ehd1 and OsMADS51 through a feedback mechanism. The OsFTL10 protein was also detected in both nucleus and cytoplasm. Furthermore, yeast two hybrid (Y2H) and bimolecular fluorescence complementation (BiFC) results show that OsFTL10 could interact with multiple 14-3-3s, suggesting that OsFTL10 might function in a similar way to Hd3a in promoting rice flowering by forming a FAC complex with 14-3-3, and OsFD1. Further experiments revealed that constitutive expression of OsFTL10 improved the drought tolerance of transgenic plants by stimulating the expression of drought responsive genes. These results suggest that rice FTL genes might function in flowering promotion and responses to environmental signals.

INTRODUCTION

Flowering is one of the most important physiological processes that changes the vegetative and reproductive growth of plants. Floral transition can be induced by both environmental factors and endogenous genetic networks. Upon flowering induction, multiple signals
converge in floral pathway integrators which subsequently activate downstream genes for growth phase transition and the development of flower organs (Wagner, 2017). Recently, advances in the molecular genetics of Arabidopsis and rice, have revealed the molecular nature of florigen to be a globular protein, named FLOWERING LOCUS T (FT), which is produced in the mature leaf and transported to shoot meristem to induce flowering. This process satisfies the major prerequisites of florigen as a systemic floral signal (Corbesier et al., 2007; Tamaki et al., 2007).

Hd3a, the Arabidopsis FT homolog protein in rice, belongs to the phosphatidylethanolamine binding protein (PEBP) family in animals (Kardailsky et al., 1999). Both Hd3a and FT are expressed in mature leaves when flowering-promotive day lengths are given, long days (LD) for Arabidopsis and short days (SD) for rice (Corbesier et al., 2007; Tamaki et al., 2007). Both proteins migrate from leaves to the apical meristem (SAM), where they bind to transcription factors of the bZIP family, including FD in Arabidopsis and OsFD1 in rice. The rice Hd3a and FD1 have been shown to form a florigen activation complex (FAC) with 14-3-3 (GF14 in rice) to promote floral initiation (Tamaki et al., 2007; Taoka et al., 2011).

Genome annotation results have revealed thirteen FT homologs (OsFT-LIKE genes) in the rice genome (Chardon & Damerval, 2005). Amongst the thirteen FTLs, RFT1/FT-L3 showed the highest homology with Hd3a, and was recently found to be important for rice flowering under LD conditions. FTL/FT-L1 is another member with high homology to Hd3a. Transgenic rice plants overexpressing RFT1 or FTL exhibited an early flowering phenotype, similar to Hd3a-overexpressing plants (Izawa et al., 2002; Kojima et al., 2002). Knockdown of Hd3a or RFT1 in the japonica rice cultivar Norin 8, delays rice flowering only under SDs or LDs, respectively (Komiya et al., 2008; Komiya, Yokoi & Shimamoto, 2009). In contrast, flowering in rice is almost completely blocked even under favorable SD conditions by knockdown of both Hd3a and RFT1, suggesting that these two genes have overlapping functions and are necessary for flowering (Komiya et al., 2008). Until now, studies in rice have revealed that the photoperiod is a major environmental factor influencing rice flowering time. Furthermore, photoperiodic flowering in rice is considered as two distinct pathways: one including OsGI (an orthologue of Gigantea), Hd1 (Heading date 1) and Hd3a. These proteins form a pathway of OsGI-Hd1-Hd3a, which is similar to that of Arabidopsis under SD conditions (Turck, Fornara & Coupland, 2008; Deng et al., 2017). Another one including Ghd7 (Grain Number, Plant Height and Heading Date 7), Ehd1 (Early heading date 1) and Hd3a/RFT1. These proteins form a pathway of Ghd7-Ehd1-Hd3a/RFT1 in rice under LD conditions. The major genes downstream of Hd3a and RFT1 are OsMADS14, OsMADS15, OsMADS18 and OsMADS34. Simultaneous knockdown of the above four genes completely blocked flowering in rice (Kobayashi et al., 2012; Tsuji, 2017).

Floral transition can be induced by a variety of external factors, including various environmental conditions and hormone treatments, such as the well-studied photoperiod pathway in rice and Arabidopsis, as well as the vernalization and GA3 (Gibberellic Acid) pathway in Arabidopsis (Kazan & Lyons, 2016). Drought stress influences plant growth and crop yield in many areas of the world, and has been shown to significantly influence
floral transition. During drought stress, plants may adopt the strategy of ‘drought escape’ by speeding up or delay of flowering (Bernal, Estiarte & Peñuelas, 2011; Franks, 2011), or the strategy of ‘drought avoidance’ or ‘drought tolerance’ by reducing water loss to prevent dehydration (Kooyers, 2015). In Arabidopsis, drought stress affects flowering time through transcriptional regulation of FT and TWINSISTER OF FT (TSF). Under long day conditions, induction of the expression of FT and TSF by drought stress depends on GI and abscisic acid (ABA), whilst under short days, floral repressors are induced by ABA and drought, and the transcription of TSF and FT is inhibited (Riboni et al., 2013). The Grain Number, Plant Height and Heading Date 7 (Ghd7), which delays heading and increases yield in rice under long days (Xue et al., 2008), also affects drought tolerance in rice (Weng et al., 2014; Kazan & Lyons, 2016).

In rice, the function of FTL genes on the induction of flowering has been well-characterized based on the studies of Hd3a and RFT1. The precise involvement of florigens in environmental response is largely unknown. When investigating the response of rice FTLs to hormones and abiotic stresses, we found that only OsFTL10 expression was stimulated by mannitol and ABA. In this report, we studied the expression of OsFTL10 and found that it expressed at a higher level in young leaf, and also its expression can be induced by drought stress. Overexpression of OsFTL10 can induce early flowering in rice through a similar mechanism as that of Hd3a. We also found that overexpression of OsFTL10 can enhance the drought tolerance of plants.

MATERIALS & METHODS
Vector construction and rice transformation
To obtain the OsFTL10 overexpression construct, a fragment containing the complete OsFTL10 cDNA (Os05g0518000) was amplified with the primer set Fw-OsFTL10-EcoRI/Re-OsFTL10-EcoRI (Table S1), and cloned into the EcoRI site in binary expression vector pCAMBIA1390. To make the RNAi construct, the cDNA fragment was amplified with the primer set Fw-OsFTL10-RNAi/ Re-OsFTL10-RNAi, and cloned into the pTCK303 vector (Wang et al., 2004). To examine the promoter activity in rice, the OsFTL10 promoter (pFTL10), spanning about 2,000 bp upstream of the ATG start codon of OsFTL10 was amplified using PCR. The CaMV35S promoter for β-glucuronidase (GUS) expression in pCAMBIA1301 was replaced with pFTL10. For subcellular localization studies of OsFTL10, its cDNA fragment was amplified by PCR and fused with EGFP in the binary vector pCAMBIA1390. The binary vector for expression of Hd3a was from a previous work (Li et al., 2011a; Li et al., 2011b). The above constructs were introduced into Agrobacterium tumefaciens strain EHA105 by electroporation (Li et al., 2011a; Li et al., 2011b). Agrobacterium-mediated transformation of rice (Oryza sativa L. ssp. japonica., Zhonghua 11) was performed as described (Li et al., 2011a; Li et al., 2011b).

Plant materials and growth conditions
Mature seeds of Zhonghua 11 (Oryza sativa L. ssp. japonica.) were collected from WT and homozygous T2 transgenic plants. Seeds were germinated for 2 days at 28 °C in the
dark, after which they were planted in soil and grown under greenhouse conditions to maturity (12-h light at 30 °C/12-h dark at 22 °C). Mature leaves from 30 day old plants were collected at 10 O’clock in the morning, and RNAs were extracted for qRT-PCR analysis of the expression of flowering related genes. The major agronomic traits including heading date, plant height, grain weight and number of flowers per panicle of the homozygous plants were investigated.

**FTL10 sub-cellular localization and BiFC analysis**

The subcellular localization of FTL10 and GF14c, as well as BiFC analysis characterizing the interaction of FTL10 and GF14c, were carried out according to (Lin et al., 2017). For subcellular localization studies of FTL10 and GF14c, the coding regions of both FTL10 and GF14c were amplified with primers (Table S1) and fused with EGFP, respectively. For BiFC analysis, GF14c and OsFTL10 were amplified by PCR and restriction-cloned (Bam HI and/or Eco RI) into the p35S::cYFP and p35S::nYFP vectors, respectively. Negative controls constructs were made by replacing GF14c or OsFTL10 with GUS (Taoka et al., 2011). The above vectors were transformed into rice protoplasts for live cell imaging. Images were captured using a LSM 710 confocal microscope (Zeiss, Jena, Germany) (Lin et al., 2017).

**Yeast two-hybrid assays**

Yeast two-hybrid (Y2H) assays were performed using the Matchmaker™ Gold Yeast Two-Hybrid Systems (Clontech, Mountain View, CA, USA) according to Lin et al. (2017). The open reading frames of OsFTL10 and 14-3-3s were inserted into pGBKTK7 and pGADT7 vectors with proper restriction enzyme sites indicated in the primers to create bait and prey, respectively.

**GUS staining of transgenic rice plants**

Homozygous transgenic lines carrying the pFTL10::GUS were randomly chosen for the study. Seeds were briefly sterilized and germinated, then planted in soil and grown under greenhouse conditions. Immature and mature leaf, roots and other tissues were collected from the plants grown at different stages, GUS staining of the plant materials was performed according to procedures described by Jefferson (1987).

**Growth regulators and abiotic stress treatments**

WT seedlings germinated for 2 weeks on half strength MS medium were collected and incubated in ddH2O containing either of the following components: 5 mmol/L IAA, 1 mmol/L 6-BA, 0.1 mmol/L MeJA, 10 mmol/L GA3, 100 µmol/L ABA, 1 mmol/L SA, 200 mmol/L Mannitol, 200 mmol/L NaCl, respectively. Treatments with IAA and BA were performed for 0, 0.5, 1 and 4 h, respectively. Treatments with ABA, SA, MeJA and GA were performed for 0, 1, 4 and 12 h, respectively. Treatments with mannitol and NaCl were performed for 0, 6, 14 and 48 h, respectively. All the treatments were started from about 10 O’clock in the morning. After treatment for the indicated times, total RNAs were extracted and reversely transcribed as the template for qRT-PCR.
qRT-PCR for gene expression analysis

Total RNAs were extracted using a TRIzol kit according to the user’s manual (Invitrogen, Carlsbad, CA, USA). For each sample, 1 µg of total RNAs was reverse-transcribed as cDNA template using a kit (TaKaRa, Kusatsu, Japan). qRT-PCR was carried out in a mixture of 20 µL consisting of cDNA, gene specific primers (Table S1) and 10 µL 2X mix (TaKaRa), water was added to the mixture to a final of 20 ul. qPCR was performed on an ABI 7900 real-time PCR machine according to the manufacturer’s instruction (Applied Biosystems). For each sample, three biological replicates were performed and cDNA were amplified in triplicate by quantitative PCR. The relative expression values were determined by using rice Ubiquitin gene as reference and the comparative Ct method (2- \Delta \Delta Ct).

Drought stress treatment

Drought stress treatment of WT and transgenic plants was performed according to Li et al. (2011a); Li et al. (2011b). Seeds of WT, overexpression transgenic lines ox2, ox5, and ox8 and RNAi lines i1,i2 and i5 (T3, homozygous seeds) were germinated, and in the same pot containing vermiculite were soaked with distilled water. Eight days later, water was withheld from the seedlings. After another 20 (for RNAi lines) or 25 days (for overexpression lines), the plants were re-watered for recovery. Three days later, plants with green, healthy leaves were regarded as viable, having survived. For study of drought related gene expression in transgenic plants, seedlings germinated for 2 weeks were incubated in a solution containing 200 mmol/L mannitol for 2, 4, 8, and 24 h, respectively. Total RNA was extracted for qRT-PCR.

RESULTS

Expression pattern of OsFTL10 in rice

OsFTL10 encodes a protein consisting of 174 amino acid residues. To investigate the temporal and spatial gene expression pattern of OsFTL10 in rice, we quantified the OsFTL10 expression level in different tissues including immature and mature leaves, stem, young panicle and floral organs from the wild-type plants grown under SD conditions. OsFTL10 expression was detected in all the above tissues. The highest expression level was observed in immature leaves with a lower level of expression found in floral organs (Fig. 1A).

The expression pattern of OsFTL10 was also investigated by GUS staining assay in stable transgenic plants. The promoter region of OsFTL10 was cloned and a construct was made by placing the GUS reporter under the OsFTL10 promoter. The construct was transformed into rice cell and stable transgenic plants were obtained. GUS expression was found in immature leaf, mature leaf, flower, young panicle, pollen and shoot tip (Figs. 1B–1H). Strong GUS staining was found in immature leaf tissue and very faint blue staining was found in the pollen (Fig. 1G).

Expression of OsFTL10 under growth regulators and abiotic stress treatments

To determine whether OsFTL10 expression is influenced by environmental factors, qRT-PCR was performed to study the expression of OsFTL10 under different growth regulators
**Figure 1** Temporal and spatial expression *OsFTL10* in rice. (A) The expression of *OsFTL10* in rice tissues by qRT-PCR analysis. (B) GUS assay of *OsFTL10* promoter activity in immature leaf. (C) Mature leaf. (D) Flower. (E, F) Young panicle. (G) Anther. (H) Germinated seedling (with amplified leaf tip).

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and abiotic stress treatments. Seedlings germinated for 2 weeks on half strength MS medium were collected and treated with different growth regulators and abiotic stresses for different time intervals, and total RNAs were extracted for qRT-PCR. Amongst the treatments of different growth regulators, ABA at 100 µM stimulated the expression of OsFTL10 to nearly two-fold after 4 hrs (Fig. 2C). The addition of GA₃ at 10 mM greatly increased the expression of OsFTL10 to more than ten-fold its basal level (Fig. 2F). Treatments with SA and MeJA also showed slightly increased expression of OsFTL10 (Fig. 2D, 2E), whilst treatment with IAA or BA reduced the expression level of OsFTL10 (Figs. 2A, 2B). Abiotic stress treatment was performed using 200 mM mannitol or 200 mM NaCl. Treatment with 200 mM mannitol resulted in increased OsFTL10 expression to four-fold basal levels at 6 hrs, and the expression dropped after 24 h (Fig. 2G). 200 mM NaCl slightly increased the expression of OsFTL10 which dropped after 24 h (Fig. 2H). In accordance with the above results, some cis elements involved in drought stress and ABA response were found in OsFTL10 promoter (Table S1). As ABA and mannitol are related to drought stress, we further studied the drought tolerance of OsFTL10 transgenic plants in addition to its function in flowering.

**Overexpression of FTL10 induced earlier flowering in rice**

As OsFTL10 is one of the homologs of Hd3a, we expected that overexpressing this gene would also promote rice flowering. We generated OsFTL10 overexpression transgenic plants and examined the transgene expression by RT-PCR (Fig. 3A). At the same time, we obtained RNAi transgenic plants for OsFTL10, and in some lines OsFTL10 expression was greatly reduced (Fig. 3B). Representative pictures of the overexpressing line ox8 (Fig. 3A) and the RNAi line (Fig. 3B) are shown. To investigate the agronomic traits of the transgenic plants, the homozygous T3 generation of the three independent overexpressing lines ox2, ox5, ox8 and RNAi lines i1, i2, i5 were grown under short day conditions, and their phenotype data compared with WT plants (Fig. 3). The average heading date of the overexpressing lines was 53 days, which was about 2 weeks less than of the WT plants (about 68 days). The RNAi lines did not show significant difference in heading time compared with the WT (Fig. 3C). The plant height for the overexpressing lines was on average 65 cm (Fig. 3D), which was about 19 cm shorter than that for the WT (84 cm). The number of grains per panicle and grain weight were also reduced significantly in the overexpressing lines, compared to that in WT (Figs. 3E, 3F). In the RNAi lines, the above traits were not significantly changed (Figs. 3C–3F). These results showed that overexpression of OsFTL10 could induce early flowering, which influenced the agronomic traits of the plants. The down-regulation of endogenous OsFTL10 exerted little effect on plant growth. When compared with Hd3a overexpression plants, we found that heading time of OsFTL10 overexpression plants were about 3 weeks longer (Table S1).

To elucidate the function of OsFTL10 on flowering induction, we first examined the subcellular localization of the OsFTL10 and GF14c protein. Constructs were made to fuse the OsFTL10 or GF14c with EGFP protein, under the CaMV35S promoter (CaMV35S::OsFTL10-EGFP). The constructs were transformed into rice leaf mesophyll protoplasts, and the EGFP signal was analyzed. The EGFP signal of OsFTL10 was observed
Figure 2  Expression of OsFTL10 in rice seedlings treated with different growth regulators and stresses. Expression of OsFTL10 in rice seedlings treated with IAA (A), BA (B), ABA (C), SA (D), MeJA (E), GA (F), mannitol (G), and NaCl (H). All data represent the mean of three biological replicates, with error bars indicating SD. (* indicates a significant difference in comparison with the control (0 h) at $P < 0.05$).
Figure 3  Phenotype of FTL10 overexpression and RNAi transgenic plants. (A) Representative picture of WT and overexpression line ox8 after heading. (B) Representative picture of WT and RNAi line i5 after heading. Performance of the transgenic lines in greenhouse on heading date (C). Grain weight (D). Number of flowers (E). Plant height (F). Data represent the mean of 15 to 24 plants, with error bars indicating SD. (*) indicates a significant difference in comparison to the WT at $P < 0.05$.

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in both the nucleus and cytoplasm, whilst the EGFP signal of GF14c was observed mainly in the cytoplasm (Fig. 4A). These results indicated that FTL10 had a similar subcellular localization to Hd3a.

Hd3a was found to interact with 14-3-3, and form a complex with OsFD1 in the nucleus for activation of the downstream flowering genes (Taoka et al., 2011). The interaction of OsFTL10 with GF14c in the rice protoplast was examined by BiFC. A strong signal was detected in the nucleus, which demonstrated the interaction of the two proteins. At the same time, the negative controls showed no obvious signal (Fig. 4B). The interactions of OsFT10 with multiple 14-3-3 s were also investigated by the yeast two-hybrid system. The results showed that OsFTL10 interacted with different 14-3-3 homologs in rice, including GF14a, GF14b, GF14c, GF14d, GF14e (Fig. 4C). These results suggested that OsFTL10 functioned similarly to Hd3a to bind with 14-3-3 s, which was a key step for entering the nucleus in the formation of the FAC complex with OsFD1 (Taoka et al., 2011).

Flowering in rice is mainly influenced by the photoperiodic pathway (Tsuji, Taoka & Shimamoto, 2011). OsGI (an orthologue of Gigantea) is an integrator of photoperiod pathway in rice (Kim et al., 2008). Under short day (SD) conditions, OsGI up-regulates the expression of Hd1 (a CONSTANS-like gene), and Hd1 will induce the expression Ghd8 (grain yield, heading date and plant height 8). Ghd8 encodes the OsHAP3 subunit of a CCAAT-box binding protein and can activate the expression of Hd3a (Yan et al., 2011). Ehd1 (Early heading date 1) is a B type response regulator, which up-regulates the expression of FT-like genes (Matsubara et al., 2011). OsGI can also up-regulate Ehd1
Figure 5 Expression of major flowering related genes in transgenic plants. Expression of OsGI (A), Hd1 (B), Ehd1 (C), MADS51 (D), RFT1 (E), Hd3a (F), FTL1 (G), MADS14 (H), MADS15 (I) in transgenic plants. All data represent the mean of three biological replicates, with error bars indicating SD. (* indicates a significant difference in comparison to the WT at $P < 0.05$).

Through activating MADS51 under SD conditions (Kim et al., 2008). To dissect the genetic networks of OsFTL10 on flowering, we performed qRT-PCR analysis of the genes that played important roles in rice flowering. The cDNA template was obtained from RNA extracted from mature leaves of WT, overexpressing lines ox2 and ox8, and the RNAi lines i2 and i5 before heading. For genes upstream of Hd3a and RFT1, GI, Hd1, Ehd1 and MADS51 were selected. qRT-PCR results showed that overexpression of FTL10 had no significant influence on the expression of GI and Hd1 in both ox2 and ox8, as their expression levels were similar to those in WT (Figs. 5A and 5B). Ehd1 was significantly up-regulated in both overexpressing lines at 1.5 and 2 fold (Fig. 5C). However, MADS51 expression was largely suppressed in the overexpressing lines (Fig. 5D). Amongst the FTL genes, Hd3a, and FTL1 were found to be up-regulated in both ox2 and ox8 (Figs. 5F, 5G), whilst the expression of RFT1 slightly decreased (Fig. 5E). For the major genes downstream of Hd3a (Taoka et al., 2011), we found that MADS15 expression was significantly increased (4 to 6 fold) in ox2 and ox8 (Fig. 5I), and MADS14 was slightly changed in these two lines (Fig. 5H). The results from the RNAi lines i2 and i5, showed similar results to those in the WT line (Fig. 5). These results suggested that overexpression of OsFTL10 induced early
flowering, through direct enhancement of downstream MADS15 expression and feedback regulation of its upstream genes.

**Overexpression of FTL10 improved drought tolerance in rice**

As OsFTL10 expression can be influenced by different growth regulators and abiotic stresses such as ABA and mannitol treatments, the drought stress tolerance in transgenic plants was investigated. Transgenic and WT seedlings grown under normal conditions with similar vitality were used for treatment (Figs. 6A, 6D). When water was withheld for 25 days from the seedlings of the OsFTL10 overexpressing plants (ox2, ox5, and ox8) and the WT plants, obvious drought stress phenotypes were observed (Fig. 6B). Compared with the WT plants, leaf rolling and browning were delayed in the OsFTL10 overexpressing plants. Three days after re-watering, most of the OsFTL10 overexpressing plants survived, whereas the WT plants withered and died (Fig. 6C). The average survival rate for WT, ox2, ox5 and ox8 plants were 6%, 84%, 88% and 91%, respectively (Fig. 6G). Similar treatment was performed on OsFTL10 RNAi lines (i1, i2, and i5) and the WT plants, except that water withholding time was reduced to 20 days (Fig. 6E). The OsFTL10 RNAi plants were found to be more sensitive to drought stress, and after re-watering, most of the WT remained viable, whereas the RNAi plants died (Fig. 6F). The survival rate for WT, i1, i2 and i5 plants were 92%, 14%, 6% and 8%, respectively (Fig. 6H). These results suggested correlation of drought tolerance with OsFTL10 expression. In transgenic rice plants overexpressing OsFTL10 acquired significantly drought tolerance, whilst the suppression of OsFTL10 resulted in drought sensitivity.

**Influence of OsFTL10 on the expression of drought stress-related genes**

Environmental stresses and other external signals can induce complex responses in plants including changes in gene transcription, protein synthesis and metabolism. Studies in different plant species have revealed key genes and proteins involved in stress signaling and transcriptional regulation (Ito et al., 2006; Shinozaki & Yamaguchi-Shinozaki, 2007; Molina et al., 2008; Nakashima et al., 2011; Rai & Penna, 2013; Janiak, Kwaśniewski & Szarejko, 2016). To elucidate the mechanisms of OsFTL10 on drought tolerance, the expression of stress-related transcription factors in stable transgenic plants under drought stress was revealed. Seedlings from WT, i5 and ox8 were collected for drought stress treatment, and RNAs were extracted for qRT-PCR analysis.

OsDREB2A is a drought inducible transcription factor found in rice, which shows increased expression upon drought stress treatment and constitutive expression of this gene results in drought tolerance of transgenic plants (Ito et al., 2006). The expression of this gene was rapidly induced in WT plants and ox8 upon drought stress, a response which continued for 24 h. However, the expression level increased significantly after 8 h of treatment in ox8 compared to WT plants. In i5, the expression of OsDREB2A decreased after 8 h of treatment (Fig. 6I).

bZIP23, a transcription factor that confers ABA-dependent drought resistance in rice (Xiang et al., 2008), has also been shown to be induced by drought stress. Overexpression
of this gene resulted in drought tolerance in transgenic plants. The expression of bZIP23 was rapidly induced in WT, ox8 and i5 plants under drought stress, whilst its expression decreased after 8 h in both WT and i5 plants. In ox8, the high expression level of this gene was maintained for 24 h (Fig. 6).
SNAC1 gene belongs to the stress-related NAC superfamily of transcription factors. SNAC1 expression was induced rapidly in WT, i5 and ox8 plants. In ox8, the expression of this gene showed higher expression levels after induction which were maintained after 24 h. Together, these results suggested that overexpression of OsFTL10 improved drought stress tolerance in rice by modulating the expression of stress responsive genes (Fig. 6K).

DISCUSSION

In this report, we studied the function of OsFTL10 and found that overexpression of this gene induced earlier flowering in rice. These data are similar to the well-studied rice florigen Hd3a and RFT1, which promoted rice flowering under short and long day conditions, respectively. We also found that overexpression of OsFTL10 enhanced the tolerance of rice plants to drought stress.

In rice, 13 FT like genes have been found based on sequence similarity, but only three genes, OsFTL1, Hd3a and RFT1, have been studied. Amongst these genes, Hd3a and RFT1 have been extensively studied and shown to be responsive for rice flowering under short and long day conditions, respectively (Komiya et al., 2008). Studies have revealed that Hd3a proteins are produced in mature leaves, and then transported to the shoot meristem. Hd3a protein binds 14-3-3 in the cytoplasm, and is imported to nucleus to form a complex (FAC), which binds to C box of MADS15 promoter region, to stimulate the expression of MADS15 and initiate flower transition (Taoka et al., 2011). To determine whether OsFTL10 also formed a FAC complex to stimulate the floral transition, BiFC was performed and showed that OsFTL10 interacted with GF4c in the nucleus. Yeast two-hybrid results also showed that OsFTL10 interacted with major 14-3-3 homologs in rice, including GIF14b and GIF14c, two 14-3-3 proteins interacting with Hd3a (Fig. 4). We also found that OsFTL10 up-regulated the downstream flowering gene OsMADS15 (Fig. 5). These data suggested that OsFTL10 had a function similar to Hd3a.

As a homolog of Hd3a, overexpression of OsFTL10 induced early flowering in rice, but its flowering promotion effect was weaker than Hd3a. The number of days to heading in CaMV35S::Hd3a plants was less than 30 days (Supplemental File 1) compared to 53 days in CaMV35S::OsFTL10 plants. As overexpression of both genes under the same promoter resulted in a different flowering time, this effect is probably caused by the amino acids composition of the two proteins. In addition, we found some differences between the two FTLs: (1) OsFTL10 was found to have a high expression level in young leaves (Fig. 1A), whilst Hd3a was normally highly expressed in mature leaves; (2) although the localization of both OsFTL10 and Hd3a proteins was found in both the cytoplasm and nucleus (Fig. 4), our BiFC results showed the interaction of OsFTL10 and GF14c mainly occurred in the nucleus, whilst Hd3a and GF14b interactions took place in the cytoplasm (Taoka et al., 2011); (3) overexpression of OsFTL10 enhanced the expression of Ehd1 (Fig. 5), Ehd1 could activate the expression of FTLs, which explains why the expressions of other FTLs were influenced. We also found that the expression of OsMADS51 was suppressed in OsFTL10 transgenic plants (Fig. 5). Thus, the function of OsFTL10 may be slightly different from the major florigens Hd3a and RFT1 in promotion of flowering.
Ehd1 can promote rice flowering under both SD and LD. Under SD, Ehd1 is involved in two modules OsGI-OsCO-Ehd1-Hd3a and OsGI-OsMADS15-Ehd1-Hd3a, in which Ehd1 functions upstream of Hd3a and promotes flowering by directly up-regulating Hd3a expression (Yan et al., 2011). The expression of OsFTL10, together with the increased expression of Ehd1 and Hd3a in overexpression lines may lead to earlier flowering phenotype in the transgenic plants. As Ehd1 is a central player which integrates flowering signaling from both SDs and LDs, and is controlled by multiple genes, the feedback regulation of Ehd1 by OsFTL10 needs further investigation. OsMADS51 is a flowering promoter under SDs, which is downstream of OsGI (Kim et al. 2008). The decreased expression level of this gene in OsFTL10 overexpression lines is somehow in contradictory with their early flowering phenotype. However, the effects of OsMADS51 may be reduced by the increased expression of Ehd1 downstream of it. We also found that RFT1, the florigen responsible for flowering under LDs, are slightly down-regulated. We suspect that the expression of this gene was influenced by OsFTL10 or other flowering related genes, as similar effects had been observed between Hd3a and RFT1. When Hd3a expression was knocked down by RNAi, RFT1 was activated for promoting flowering under SDs (Komiya, Yokoi & Shimamoto, 2009).

Photoperiod and vernalization are two environmental factors which influence flowering time in many plant species (Khan, Ai & Zhang, 2014). In the model plant Arabidopsis, several photoperiod and circadian clock genes have been found to affect the expression of drought stress related genes. Mutations in clock components, such as the triple mutant prr5 (PSEUDO-RESPONSE REGULATOR 5), 7, 9 are more tolerant to high salinity and drought stresses. TOC1 (timing of CAB expression 1) RNAi plants showed improved drought tolerance. GI overexpression plants show enhanced salt sensitivity, whereas gi mutants show enhanced salt tolerance and enhanced survival under drought stress and oxidative stress (Grundy, Stoker & Carre, 2015). Phytochrome B (PhyB) is also a negative regulators of dehydration response (Legnaioli, Cuevas & Mas, 2009). In barley, variation at the photoperiod response and clock genes Ppd-H1 and HvELF3 can affect the expression level of osmotic stress genes (Habte et al., 2014). Stresses such as drought and salt, can also regulate flowering times (Kazan & Lyons, 2016). Early flowering can be induced by drought stress under LD conditions, and ABA can up-regulate the expression of major flower integrators FT/TSF and SOC1 in a photoperiod-dependent manner (Riboni et al., 2013). Interestingly, most of the above flowering related genes were found to be negative regulators of stress responses. OsGI was recently found to be a negative regulator of osmotic stress response in rice (Li et al., 2016). As OsGI is a positive regulator of OsMADS51, mutation of OsGI may lead to reduced expression of OsMADS51. In our OsFTL10 overexpression lines, the expression of OsMADS51 was decreased. This result indicates that overexpression of OsFTL10 might have similar effects with OsGI mutants on drought tolerance. Ghd7 functions in rice as a negative regulator flowering time under long days (Xue et al., 2008), was also found to be a negative regulator of drought tolerance in rice (Weng et al., 2014). As Ghd7 acts as a suppressor of Ehd1 under LDs, the increased expression of Ehd1 in OsFTL10 transgenic plants, suggests that drought tolerance conferred by OsFTL10 may be similar to Ghd7 mutant. The link between flowering and drought
tolerance was also demonstrated by a recent work. Overexpression of WOX13 under a drought inducible promoter Rab21 resulted in drought resistant and early flowering in rice. In these plants, drought stress and flowering related genes such as OsDREB1A, OsDREB1F, Hd3a and MADS14 were up-regulated (Minh-Thu et al., 2018). Although the above analysis provide clues that drought tolerance by OsFTL10 might be related to OsGI and (or) Ghd7, through controlling the expression of MADS51 and (or) Ehd1, much work including the analysis of the mutants involved in the flowering signaling pathways are needed to clarify their regulation networks. The OsFTL10 transgenic plants displayed enhanced drought stress tolerance, as revealed by the higher survival rate and better growth performance under stress conditions (Fig. 6). Cis elements related to drought stress and ABA response were found in the OsFTL10 promoter (Supplemental Information 1), and qRT-PCR revealed that OsFTL10 expression was induced by both ABA and mannitol treatments (Fig. 2). Investigation of the drought related gene expression in the OsFTL10 transgenic lines further confirmed the drought tolerance of the plants (Fig. 6). Many transcription factors can be induced by drought stress in rice, and constitutive expression of these transcription factors have been found to improve plant tolerance to drought stress (Shinozaki & Yamaguchi-Shinozaki, 2007; Nakashima et al., 2011; Rai & Penna, 2013; Janiak, Kwaśniewski & Szarejko, 2016). We studied the expression of three transcription factors including OsDREB2A, bZIP23 and SNAC1, and found they were induced to a higher level of expression in the OsFTL10 overexpressing line ox8. These results correlated well with the performance of the drought stress test results. DREBs/CFBs are transcription factors that can be induced by stresses. They can activate the expression of stress-inducible genes by binding to the dehydration-responsive element/C-repeat (DRE/CRT) cis-acting elements in their promoter region. OsDREB2A is a rice DREB homolog which has been found to be induced by salt and drought. The overexpression of OsDREB2A in rice plants showed both drought and salt stress tolerance (Ito et al., 2006). Abscisic acid is regarded as a drought stress hormone in plants. The transcription factor bZIP23 plays an important role in ABA signaling by binding to the promoter of PP2C49 to improve its expression. As a co-receptor, PP2C can increase ABA binding affinity toward the formation of the Receptor-ABA-PP2C complex (Moreno-Alvero et al., 2017). bZIP23 has been also found to improve drought tolerance in rice (Xiang et al., 2008). The SNAC1 gene belongs to the stress-related NAC superfamily of transcription factors, which has been expressed in plant for improving drought tolerance (Nakashima et al., 2011). Although the induced expression of the above three transcription factors can be considered as evidence of OsFTL10 tolerance to drought stress, mechanistic links between OsFTL10 and transcription factors were not fully determined.

CONCLUSIONS

In summary, results presented in this study reveal that OsFTL10 functioned similarly to Hd3a in rice by promoting floral transition, and overexpression of OsFTL10 improved the drought stress tolerance of the plants. Further investigations on the identification of OsFTL10 regulated target genes will be helpful for fully elucidating the linkage of flowering and drought stress in the monocot model rice.
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ADDITIONAL INFORMATION AND DECLARATIONS

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Author Contributions
- Maichun Fang and Meiru Li conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Zejiao Zhou performed the experiments, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Xusheng Zhou performed the experiments, approved the final draft.
- Huiyong Yang performed the experiments, contributed reagents/materials/analysis tools, approved the final draft.
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