Overexpression of a methyl-CpG-binding Protein Gene OsMBD707, Leads to Larger Tiller Angles and Reduced Photoperiod Sensitivity in Rice

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Research article

Keywords: Rice, MBD, Tiller angle, Flowering time, Photoperiod sensitivity, Overexpression

DOI: https://doi.org/10.21203/rs.3.rs-95600/v1

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Abstract

Background: Methyl-CpG-binding domain (MBD) proteins play important roles in epigenetic gene regulation, and have diverse molecular, cellular, and biological functions in plants. MBD proteins have been functionally characterized in various plant species, including *Arabidopsis*, wheat, maize, and tomato. In rice, 17 sequences were bioinformatically predicted as putative MBD proteins. However, very little is known regarding the function of MBD proteins in rice.

Results: We explored the expression patterns of the rice *OsMBD* family genes and identified 13 *OsMBDs* with active expression in various rice tissues. We further characterized the function of a rice class I MBD protein OsMBD707, and demonstrated that OsMBD707 is constitutively expressed and localized in the nucleus. Transgenic rice overexpressing *OsMBD707* displayed larger tiller angles and reduced photoperiod sensitivity—delayed flowering under short day (SD) and early flowering under long day (LD). RNA-seq analysis revealed that overexpression of *OsMBD707* led to reduced photoperiod sensitivity in rice through regulating expression of key flowering regulator genes in the *Ehd1-Hd3a/RFT1* pathway.

Conclusion: The results of this study demonstrated that OsMBD707 plays important roles in rice growth and development, and should lead to further studies on the functions of OsMBD proteins in growth, development, or other molecular, cellular, and biological processes in rice.

Background

DNA methylation, a conserved epigenetic modification in plant and animal genomes, plays an important role in genome stability, genomic imprinting, and gene expression regulation, and exerts effects on various aspects of plant and animal cellular and developmental processes [1, 2]. Methyl-CpG-binding domain (MBD) proteins emerge as important interpreters of DNA methylation marks [3]. MBD proteins can bind to methylated DNA and recruit chromatin remodeling factors, histone deacetylases and methylases to regulate gene expression [4–6]. Numerous studies in mammals revealed that mutations or abnormal expression of MBD proteins occur in many neurological diseases and cancers [7]. For example, mutations in the human methyl-CpG-binding protein 2 (MeCP2) cause a postnatal neurodevelopmental disorder Rett syndrome [8], demonstrating that MBD proteins play important roles in maintaining a normal epigenetic and cellular homeostasis.

Plant MBD proteins have been identified based on amino acid sequence similarity with mammalian MBDS [9, 10]. The *Arabidopsis* genome encodes 13 MBD domain-containing proteins [11]. Studies of the *Arabidopsis* MBDs showed that plant MBDs can bind to methylated or unmethylated DNA [10, 12], and interact with chromatin-modifying or transcriptional complexes, such as the histone deacetylase AtHDA6 [10], the histone methyltransferase AtPRMT11 [13], the histone acetyltransferase IDM1 and the alpha-crystallin domain protein ROS5/IDM2 in the DNA demethylation pathway [14–16], the RNA binding proteins AtRPS2C, AtAGO4 and AtNTF2 which are involved in the RNA-mediated gene silencing pathway [17], or the chromatin remodelers CHR11/CHR17 and chromatin remodeling complex SWR1 subunit PIE1.
and ARP6 which are involved in histone H2A.Z deposition [18, 19]. The Arabidopsis MBDs play important roles in diverse molecular and biological processes. For example, knockdown of AtMBD6 or AtMBD10 leads to disturbed nucleolar dominance in *Arabidopsis suecica* [20]; mutation in AtMBD8 causes late flowering in the C24 ecotype [21]; mutation in AtMBD9 promotes early flowering and enhances shoot-branching in *Arabidopsis* [22, 23]; and knockdown of AtMBD11 results in *Arabidopsis* plants with morphological and developmental abnormalities [9].

In addition to the studies in the model plant *Arabidopsis*, several recent researches have shown the important roles of MBD proteins in different plant species. The wheat *TaMBD2* homoeologous genes were significantly induced by salt stress, suggesting their potential functions in stress responses [24]. *TaMBD6* was highly responsive to prolonged chilling, suggesting that TaMBD6 was potentially involved in regulating the developmental transition from vegetative to reproductive stages in wheat [25]. The maize ZmMBD101 was identified to be required for maintaining Mutator (Mu) elements chromatin in maize [26]. In tomato, SIMBD5 was identified to interact with the CUL4-DDB1-DET1 complex, which was involved in the transcriptional activation of downstream genes [27]. Overexpressing of the *SIMBD5* gene leads to dark fruit color and dwarf phenotype in transgenic plants [27]. A number of MBD genes in four different solanaceae species have been observed to be differentially induced or suppressed during fruit development or abiotic stress responses, suggesting their roles involved in these processes [28].

Rice (*Oryza sativa* L.) is one of the most important crops worldwide. In the rice genome, 17 genes were bioinformatically predicted to encode putative MBD proteins and were designated as OsMBD701 to OsMBD718 [11]. However, very little is known regarding the functions of OsMBD proteins in rice. In this study, we explored the expression patterns of the predicted OsMBD family genes and detected 13 OsMBDs with active expression in various rice tissues. We further performed functional study of a rice class I MBD protein, OsMBD707, and revealed that overexpression of OsMBD707 resulted in larger tiller angles and reduced photoperiod sensitivity in rice. Our results demonstrated that OsMBD707 plays an important role in rice growth and development.

**Results**

**Differential expression patterns of 13 OsMBD family genes in rice tissues**

A bioinformatics analysis has identified 17 putative OsMBD proteins in rice [11]. Considering the annotation updates of the rice genome over the past decades, in this study, we verified the predicted OsMBD family genes by searching the National Center for Biotechnology Information (NCBI) and MSU Rice Genome Annotation Project (RGAP) databases. Our search revealed 15 genes in both NCBI and RGAP databases matching with previously predicted OsMBD701, OsMBD703, OsMBD704, OsMBD705, OsMBD706, OsMBD707, OsMBD708, OsMBD709, OsMBD710, OsMBD711, OsMBD713, OsMBD714, OsMBD715, OsMBD717, and OsMBD718, respectively (Additional file 1: Table S1). However, no NCBI RAP locus or MSU RGAP locus has been identified that match with OsMBD712 or OsMBD716 (Additional file 1:
Table S1). In addition, a RAP locus *Os04g0192775* and a MSU RGAP locus *LOC_Os04g11510* were retrieved as genes putatively encoding MBD-containing proteins (Additional file 1: Table S1).

Quantitative RT-PCR (qRT-PCR) was performed to explore the expression patterns of the *OsMBD* family genes in the roots, stems, leaves, spikelets, seeds, and panicle axes of rice plants. Among the 17 putative OsMBD-encoding genes retrieved from NCBI and/or MSU RGAP databases, no transcripts of the predicted *OsMBD703/Os06g072100/LOC_Os06g48870*, *OsMBD713/Os04g0193900/LOC_Os04g11730*, *Os04g0192775*, or *LOC_Os04g11510* were detected in any tested tissues (Additional file 1: Table S1). While *OsMBD704* and *OsMBD714* were detected to be preferentially expressed in the seeds, *OsMBD701*, *OsMBD705*, *OsMBD706*, *OsMBD707*, *OsMBD708*, *OsMBD709*, *OsMBD710*, *OsMBD711*, *OsMBD715*, *OsMBD717*, and *OsMBD718* were observed to be differentially expressed in various tissues (Fig. 1).

**OsMBD707 is constitutively expressed and localized in the nucleus**

Rice OsMBDs could be divided into six classes [11]. In the present study, we focused on functional analysis of OsMBD707, which belongs to class [11]. Phylogenetic analysis of OsMBD707 and closely related MBPs in various plant species showed that OsMBD707 was clustered together with ObMBD11-like in *Oryza brachyantha*, hypothetical protein TVU48968.1 in *Eragrostis curvula*, SiMBD10 and SiMBD11 in *Setaria italica*, ZmMBD105 and ZmMBD106 in *Zea mays*, BdMBD11 in *Brachypodium distachyon*, and SbMBD10 and SbxP2 in *Sorghum bicolor* (Fig. 2). qRT-PCR analysis showed that the mRNA of *OsMBD707* was expressed in various tissues, although the transcript level in the spikelets was relatively low (Fig. 1). A 1931-bp promoter fragment upstream of the translational start of *OsMBD707* was cloned and fused with the GUS reporter gene. Histochemical staining of rice plants transformed with the *OsMBD707* promoter-GUS fusion construct showed that GUS was expressed throughout the tested tissues, including the roots, stems, leaves, and spikelets, although the expression level was weaker than that of 35S promoter-GUS transgenic plants (Fig. 3a). Overall, these results indicated that *OsMBD707* is constitutively expressed in various rice tissues.

*OsMBD707* was predicted to generate two alternative transcripts, *XM_015764399.1/LOC_Os12g42550.1*, and *XM_015764400.2/LOC_Os12g42550.2* (Additional file 2: Figure S1A, B). We performed RT-PCR to clone the cDNA fragment of *OsMBD707*, but obtained only *XM_015764399.1/LOC_Os12g42550.1*. An RT-PCR using primers (Additional file 2: Figure S1A) designed to distinguish the two predicted alternative transcripts was further performed. Consistently, only *XM_015764399.1/LOC_Os12g42550.1* was detected in all tested tissues, including the roots, stems, leaves, spikelets, seeds, and panicle axes (Additional file 2: Figure S1C, D), suggesting that there is only one isoform, LOC_Os12g42550.1, of OsMBD707. To explore the subcellular localization of OsMBD707, we performed transient expression of a GFP-OsMBD707 fusion construct in rice protoplasts. Microscopy revealed that the fluorescence signal of GFP-OsMBD707 was inside the nucleus region (Fig. 3b), demonstrating that OsMBD707 is a nuclear-localized protein which is consistent with a function as a methyl-CpG-binding protein.

**Overexpression of OsMBD707 causes larger tiller angles and reduced photoperiod sensitivity in rice**
To explore the function of *OsMBD707* in rice, we generated overexpression and RNAi knockdown transgenic plants. For each type, more than 40 independent transgenic T<sub>0</sub> plants were generated, and five independent plants were chosen for initial analysis. qRT-PCR analysis showed that the transcription levels of *OsMBD707* were significantly higher in plants transformed with *OsMBD707*-overexpression construct (about 12- to 43-fold) and lower in plants transformed with *OsMBD707*-RNAi construct (about 11–27%), as compared to wild-type plants (Fig. 4a, b), confirming the overexpression and knock-down of *OsMBD707*, respectively, in the transgenic plants. In addition, we generated CRISPR/Cas9 knockout plants of *OsMBD707*. Genotyping of the CRISPR/Cas9 transgenic plants identified nine independent T<sub>0</sub> plants with homozygous mutations in at least one of the single guide RNA (sgRNA) targeting sites of *OsMBD707* (Fig. 4c). Initial phenotypic observation showed that the *OsMBD707*-overexpression plants (referred to as OX707) displayed a larger tiller angle after tillering stage compared to wild-type. In contrast, no obvious morphological differences were observed among wild-type, the *OsMBD707*-RNAi plants (referred to as 707i) and the CRISPR/Cas9 knockout plants (referred to as mbd707).

*OsMBD707*-overexpression, -knockdown, and -knockout lines were generated up to T<sub>3</sub> to T<sub>4</sub> generations, and two independent overexpression lines, one knockdown line, and one knockout line were chosen for further analysis. Consistent with initial phenotypic observation, the two *OsMBD707*-overexpression lines OX707-#20 and OX707-#21 displayed larger tiller angles, compared to wild-type, the knockdown line 707i-#30, and the knockout line mbd707-#15 (Fig. 5a). In addition, we observed significant delays in flowering of the two overexpression lines grown under short day (SD) condition (Fig. 5a). We further investigated the heading dates of the *OsMBD707*-overexpression, -knockdown, and -knockout lines in growth chambers under SD and long day (LD) conditions. As showed in Fig. 5b, under SD, the flowering times of OX707-#20 and OX707-#21 were significantly delayed (about 15–17 days) compared with that of wild-type, 707i-#30 or mbd707-#15 (Fig. 5b). In contrast, under LD, the flowering times of OX707-#20 and OX707-#21 were significantly earlier (about 10.5–12.5 days) compared with that of wild-type, 707i-#30 or mbd707-#15 (Fig. 5c), indicating that overexpression of *OsMBD707* caused reduced photoperiod sensitivity in transgenic rice plants.

**Global transcriptome analysis reveals transcriptional changes in key flowering regulator genes induced by overexpression of MBD707**

RNA-seq-based transcriptome analysis was performed to investigate global transcript changes in the *OsMBD707*-overexpression transgenic line OX707-#21 under both SD and LD conditions. Under SD, about 1,026 genes were identified that were differentially expressed between OX707-#21 and wild-type, and of these, 616 genes were up-regulated, whereas 410 genes were down-regulated in OX707-#21 (Additional file 3: Figure S2A, Additional file 4: Table S2). Under LD, about 1,653 differentially expressed genes (DEGs) were identified between OX707-#21 and wild-type, including 997 up-regulated genes and 656 down-regulated genes in OX707-#21 (Additional file 3: Figure S2A, Additional file 5: Table S3). In total, about 2,353 DEGs were identified under SD and/or LD (Additional file 3: Figure S2B).
Gene Ontology (GO) analysis of these 2,353 DEGs showed that the most common biological process categories were associated with translation, peptide biosynthetic, peptide metabolic, amide biosynthetic, and cellular amide metabolic processes; the most common cellular component categories were ribosome, ribonucleoprotein complex, non-membrane-bounded organelle, intracellular non-membrane-bounded organelle, and cytoplasmic part; and the top three common molecular function categories were structural molecule activity, structural constituent of ribosome, and transferase activity/transferring glycosyl groups (Additional file 6: Figure S3, Additional file 7: Table S4). Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis revealed that the identified DEGs were significantly enriched in ribosome and phenylpropanoid biosynthesis pathways (Additional file 8: Figure S4, Additional file 9: Table S5).

We further surveyed the DEGs with known or putative functions involved in tiller angle or flowering time regulation. The phytochromeinteracting factorlike protein gene OsPIL15 (Os01g0286100) that negatively regulates tiller angle [29] was significantly down-regulated in OX707-#21 under both SD and LD (Additional file 4: Table S2, Additional file 5: Table S3). Notably, a number of genes with functions in controlling flowering time were identified among the DEGs, including FLAVIN-BINDING, KELCH REPEAT, F-BOX 1 (OsFKF1) [30], Early heading date 1 (Ehd1) [31, 32], Days to heading on chromosome 2 (DTH2) [33], Heading date 3a (Hd3a) and RICE FLOWERING LOCUS T1 (RFT1) [34, 35], OsMADS14 and OsMADS15 [36], and Flowering Locus T gene homologs FT-L7, FT-L8 and FT-L12 that promote flowering, and Grain number, plant height, and heading date 2 (Ghd2) [37] that inhibits flowering. Under SD, Hd3a, RFT1, FT-L7, OsMADS14, and OsMADS15 were down-regulated in OX707-#21. In contrast, OsFKF1, Ehd1, Hd3a, RFT1, FT-L8, FT-L12, and OsMADS14 were up-regulated, whereas Ghd2 was down-regulated in OX707-#21 under LD (Fig. 6a, b). The transcriptional changes of these flowering regulator genes in OX707-#21 were consistent with the delayed flowering and early flowering phenotypes of the MBD707-overexpression line under SD and LD, respectively, except that the minor-effect heading promoting gene DTH2 was paradoxically down-regulated under LD (Fig. 6a, b). The transcriptional profiles of five key flowering regulator genes, Ehd1, Hd3a, RFT1, OsMADS14, and OsMADS15 were verified by qRT-PCR, and the results were consistent with the RNA-seq data (Fig. 6c).

**Discussion**

MBD family proteins have been functionally characterized in various plant species, including Arabidopsis [9–23], wheat [24, 25], maize [26], and tomato [27, 28]. MBD proteins play pivotal roles in plant growth, development, and stress responses. In the present study, we characterized OsMBD707 in rice and demonstrated its roles in regulating tiller architecture and flowering time.

Seventeen sequences were predicted as putative MBD proteins from the rice genome (Additional file 1: Table S1). In the present study, 13 predicted OsMBD genes were detected to be actively expressed in the roots, stems, leaves, spikeletes, seeds, or panicle axes (Fig. 1). However, no transcripts were detected in any tested tissues for the rest four predicted sequences (Additional file 1: Table S1). Whether these four sequences are pseudogenes, or their expression is restricted to other specific tissues or developmental stages, remains to be further elucidated. The plant MBDs were grouped into eight classes, but among
them, class IV and class VI were present only in dicots [11]. OsMBD707 belongs to the class I MBD proteins [11]. Previous studies have demonstrated that the Arabidopsis class I MBD proteins AtMBD10 and AtMBD11 had important functions in regulating nucleolar dominance [20], and morphological development [9]. RNAi-mediated knockdown of AtMBD10 revealed that AtMBD10 was required for rRNA gene silencing in nucleolar dominance [20], and knockdown of AtMBD11 led to aerial rosettes, serrated leaves, abnormal flower position, late flowering, and reduced fertility in Arabidopsis [9]. In the present study, we observed that overexpression of OsMBD707 caused larger tiller angles and reduced photoperiod sensitivity in rice. On the contrary, both knockdown and knockout of OsMBD707 did not result in observable changes in the morphology of rice plants.

Both tiller architecture and flowering time are important traits for rice cultivation. Over the past decades, several key/major regulators of the tiller angle have been identified, such as LAZY1 [38], Tiller Angle Control 1 (TAC1) [39], PROSTRATE GROWTH 1 (PROG1) [40, 41], Loose Plant Architecture 1 (LPA1) [42], PLANT ARCHITECTURE AND YIELD 1 (PAY1) [43], TAC3 and D2 [44], HEAT STRESS TRANSCRIPTION FACTOR 2D (HSFA2D) [45], TILLER INCLINED GROWTH 1 (TIG1) [46], Tiller Angle Control 4 (TAC4) [47], etc. However, RNA-seq analysis did not identify any of these tiller angle regulating genes as DEGs between the OsMBD707-overexpression line OX707-#21 and wild-type, except that an OsPIL15 gene negatively regulating tiller angle [29] was found to be down-regulated in OX707-#21. Since RNA-seq was conducted using RNAs extracted from fully expanded leaves, the reason for rare detection of the key/major tiller angle regulating genes among the DEGs could be that these genes were expressed abundantly in tissues involved in tillering (e.g., stems, tiller base, and tiller node), but not expressed or expressed at very low levels in mature leaves [38, 39]. Nevertheless, the mechanism by which OsMBD707 regulates tiller angle remains to be further explored.

Rice is an SD plant whose flowering is promoted under SD, but postponed under LD. Rice flowering time is largely determined by photoperiod sensitivity, which is controlled by an intricate genetic network of an antagonistic regulation of flowering promotion under SD, and repression under LD [48]. In the present study, a number of flowering time genes were identified as DEGs between OX707-#21 and wild-type (Fig. 6a, b). Among these differentially expressed flowering time genes, Hd3a, RFT1, and Ehd1 are primary integrators of photoperiodic signals. Hd3a and RFT1 are two florigens of rice and Ehd1 promotes the expression of Hd3a and RFT1 to induce heading [31, 34, 35]. Via RNA-seq, Hd3a and RFT1 were found to be down-regulated in OX707-#21 under SD, but up-regulated under LD (Fig. 6). Ehd1 was identified to be up-regulated in OX707-#21 under LD (Fig. 6a), and although RNA-seq did not detect Ehd1 as a DEG under SD, qRT-PCR verified that Ehd1 was down-regulated in OX707-#21 under SD (Fig. 6b). Besides, RNA-seq revealed that three flowering-promoting genes/homologs involved in the Ehd1-Hd3a/RFT1 pathway, OsMADs14, OsMADS15, and FTL7, were down-regulated in OX707-#21 under SD; and four genes/homologs OsFKF1, OsMADs14, FTL8, and FTL12 were up-regulated in OX707-#21 under LD (Fig. 6a, b). Although a DTH2 gene having a function in promoting heading by inducing Hd3a and RFT1 was paradoxically down-regulated in OX707-#21 under LD (Fig. 6a, b), its effect did not seem to change the up-regulation of Hd3a and RFT1. Overall, RNA-seq results indicated that overexpression of OsMBD707
led to transcriptional changes in key flowering regulator genes in the *Ehd1-Hd3a/RFT1* pathway, resulting in reduced photoperiod sensitivity in transgenic rice.

**Conclusion**

In this study, the bioinformatically predicted *OsMBD* family genes were verified and 13 *OsMBDs* were identified to be actively expressed in various rice tissues. We further performed functional study of *OsMBD707*, and demonstrated that *OsMBD707* is constitutively expressed and localized in the nucleus. Overexpression of *OsMBD707* causes larger tiller angles, delayed flowering under SD and early flowering under LD in rice. RNA-seq analysis revealed that overexpression of *OsMBD707* led to reduced photoperiod sensitivity in rice by down-regulating flowering-promoting genes under SD and up-regulating flowering-promoting genes under LD. Our results demonstrated the important roles of *OsMBD707* in rice growth and development, and lay the foundation for future studies on the function of *OsMBD* proteins in molecular, cellular, and biological processes in rice.

**Methods**

**Plant materials and growth conditions**

*Oryza sativa* L. ssp. *japonica* (cv. Nipponbare) (maintained in Biotechnology Research Institute, Fujian Academy of Agricultural Sciences) was used in this study. Rice plants were cultured in a greenhouse under partially regulated conditions of 26–32 °C with a 14-h light/10-h dark cycle. For flowering time investigation, rice plants were grown in environmentally controlled growth chambers under SD (9.5-h light, 28 °C/14.5-h dark, 26 °C) and LD (14.5-h light, 28 °C/9.5-h dark, 26 °C) conditions, respectively.

**Gene expression analysis by qRT-PCR**

Total RNAs were extracted from rice tissues using TRIzol reagent (Invitrogen, USA). The RNA samples were treated with DNase I (Takara, Dalian, China). The first-strand complementary DNA was generated from 0.5 ug RNA using a ReverTra Ace qPCR RT Kit (TOYOBO, Japan). qRT-PCR was performed on an ABI QuantStudio 6 Flex System (Applied Biosystems, USA) using a FastStart Universal SYBRgreen Master (ROX) (Roche, Germany). Three replications were conducted for each sample. The primers used for analysis are listed in Additional file 10: Table S6.

**Phylogenetic analysis**

The full-length amino acid of *OsMBD707* was used as a query to search against the PLAZA database (https://bioinformatics.psb.ugent.be/plaza/). The retrieved sequences were verified against the NCBI non-redundant (NR) protein database (http://blast.ncbi.nlm.nih.gov/). The homologs were aligned using Clustal X program [49], and the phylogenetic tree was constructed using the neighbor-joining algorithm with 1,000 bootstrap replicates in MEGA X [50].

**Plasmids construction**
A 1935-bp DNA sequence upstream of the ATG start codon of OsMBD707 was amplified by PCR from Nipponbare genomic DNA. The amplified fragment was cloned into pCXGUS-P [51], resulting in a binary vector containing a fusion of the OsMBD707 promoter and a GUS reporter gene. The open reading frame (ORF) of OsMBD707 was amplified by using specific primers and was cloned into pCXUN [51], resulting in a binary vector pCXUN-OsMBD707 in which OsMBD707 was driven by the maize ubiquitin-1 promoter. A 213-bp DNA fragment of OsMBD707 and a 388-bp stuffer DNA fragment from the GUS gene were amplified by using specific primers. Overlapping PCR was performed using the two amplified fragments as templates. The resultant hairpin RNAi fragment was cloned into pCXUN [51], resulting in a binary RNAi vector pCXUN-OsMBD707-RNAi. Two sgRNA sequences targeting the coding region of OsMBD707 were designed according to Shan's program [52]. Construction of the CRISPR/Cas9 vector was carried out as previously described [53]. Two DNA oligos corresponding to the designed sgRNAs were synthesized and the dimer was cloned into pYLgRNA-OsU6a and pYLgRNA-OsU6b, respectively. The resultant sgRNA expression cassettes were thus cloned into pYLCRISPR/Cas9Pubi-H [53]. To generate subcellular localization construct for OsMBD707, the ORF of a GFP gene was amplified and cloned into pCXSN [51], resulting in a binary vector pCS-NGFP. The OsMBD707 ORF digested from pCXUN-OsMBD707 by BamHI was then cloned into the BamHI-digested pCS-NGFP to fused in-frame with the GFP gene. All primers or oligos used for plasmid construction are listed in Additional file 10: Table S6.

**Rice protoplast transfection and stable transformation**

Rice protoplasts were prepared from the sheath and stem tissues of 2-week-old etiolated seedlings. The GFP alone and GFP-OsMBD707 fusion constructs were transfected, respectively, into rice protoplasts via a PEG-mediated method as previously described [54]. Rice stable transformation was conducted as previously described [55]. The binary vectors were introduced into Agrobacterium tumefaciens EHA105, and the transformant strains were used to transform rice calli of cv. Nipponbare. Homozygous transgenic plants were screened in T1 generation derived from self-pollination of T0 plants, and were maintained up to T3 to T4 generations.

**GUS staining and GFP detection**

GUS histochemical staining was performed following the procedure described by [56]. Rice tissues were immersed in X-Gluc (Thermo Fisher Scientific, USA) staining solution at 37 °C overnight and were subsequently rinsed in 70% ethanol at room temperature for 1 or 3 days. Pictures were taken with an Olympus SZX12 stereo microscope. The transfected rice protoplasts were incubated at room temperature for 16–20 h. Fluorescence microscopy was carried out on a Leica DMi8 Laser Scanning Confocal microscope (Leica, Germany) with Excitation/emission wavelengths 488/535 nm for green fluorescence.

**RNA-seq analysis**

Fully expanded leaves of the overexpression line OX707-#21 and wild-type were collected at about 50 days after sowing. Total RNAs extracted from leaf tissues were subjected to RNA-seq analysis at Novogene (Beijing, China). Three to four biological replicates of each sample were used for RNA-seq analysis. Briefly, sequencing libraries were generated using NEBNext UltraTM RNA Library Prep Kit for
Illumina (NEB, USA) following manufacturer’s instructions. The libraries were sequenced on an Illumina Novaseq platform. Raw reads were processed through in-house perl scripts, and the filtered clean reads were mapped to the reference genome using Hisat2 v2.0.5. FPKM (expected Fragments Per Kilobase of transcript per Million fragments sequenced) of each gene was calculated for estimating gene expression levels [57]. Gene expression difference between OX707-#21 and wild-type was performed using the DESeq2 [58]. Genes with padj < 0.05 and log2FoldChange > 1 were assigned as differentially expressed. GO and KEGG enrichment analysis of DEGs were implemented by the clusterProfiler [59].

Abbreviations

MBD: methyl-CpG-binding domain; NCBI: National Center for Biotechnology Information; RGAP: Rice Genome Annotation Project; qRT-PCR: quantitative RT-PCR; SD: short day; LD: long day; DEGs: differentially expressed genes; GO: Gene Ontology; KEGG: Kyoto Encyclopedia of Genes and Genomes; ORF: open reading frame; sgRNA: single guide RNA; FPKM: expected Fragments Per Kilobase of transcript per Million fragments sequenced.

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and material

The datasets supporting the conclusions of this article are included within the article and its additional files.

Competing interests

The authors declare that they have no competing interests.

Funding

National Key Research and Development Program of China (2016YFD0102103) to SZ (conceived and designed the experiments) and Technological Innovation Projects of Fujian Academy of Agricultural Sciences (ZYTS2019014) to MW (analyzed the data and contributed reagents and tools).

Authors’ contributions
SZ and SC conceived and designed the experiments. MQ, ZZ, TL, PN, and ZQC performed the experiments. MQ, MW, WC, and SC analyzed the data. MW and ZJC contributed reagents and tools. MQ and SC wrote the manuscript. All authors have read and approved the manuscript.

Acknowledgements

Not applicable.

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