A Cyclophilin OsCYP20–2 Interacts with OsSYF2 to Regulate Grain Length by Pre-mRNA Splicing

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Abstract

Background: Grain size is one of the key agronomic traits that impact grain yield. Several regulatory pathways had been reported to participate in grain size determination via cell expansion or proliferation in rice. However, little is known about cyclophilin and spliceosome participation in grain shape regulation.

Results: Here, we identified OsCYP20–2, a cyclophilin that influences spliceosome assembly to determine grain length. oscyp20–2t1, a knock out mutant of OsCYP20–2 caused by T-DNA insertion, produced shorter grains with deficient cell elongation. Through yeast two-hybrid screening and pull-down assays, OsSYF2, a pre-mRNA splicing factor, was identified as an interacting protein of OsCYP20–2. The phenotypes of transgenic lines indicated that OsSYF2 positively regulates grain length via its influence on cell expansion. Transcriptomic analysis showed that OsSYF2 controls the expression and pre-mRNA alternative splicing of genes involved in sugar metabolism. In addition, these two genes have similar effects on panicle architecture.

Conclusions: Taken together, OsSYF2, an interacting protein of OsCYP20–2, controls grain length and panicle architecture by regulating the alternative splicing of pre-mRNA involved in cell elongation and sugar metabolism.

Keywords: OsCYP20–2, Grain length, OsSYF2, Spliceosome, Alternative splicing

Background

Vital agronomic traits for yield improvement in rice include inflorescence architecture and grain size. Genetic and molecular analyses have identified numerous quantitative trait loci (QTLs) and genes involved in multiple signaling pathways that regulate grain size. For example, G-protein signaling positively regulates grain length through the concerted actions of RGA1, RGB1, GS3, and DEP1 (Ashikari et al. 1999; Fan et al. 2006; Huang et al. 2009; Mao et al. 2010; Xu et al. 2019). The ubiquitin-mediated protein degradation pathway (Chen et al. 2013; Huang et al. 2017; Song et al. 2007), the mitogen-activated protein kinase signaling pathway (Wang et al. 2019), and several hormone signaling pathways (Shirley et al. 2019; Xiao et al. 2019) also affect grain size.

The protein conformational change impacts protein folding and assembling to regulate its cellular functions (Zhang et al. 2019). Proline residues can form cis and trans peptide bonds. The prolyl cis-trans isomerization functions as a molecular switch to regulate protein conformation. This prolyl cis-trans conversion can be catalyzed by peptidyl prolyl cis-trans isomerase (PPIase). PPIases can be divided into four families: protein Ser/Thr phosphatase 2A (PP2A) activator, FK506-binding proteins (FKBPs), parvulins, and cyclophilins (CYPs) (Lu et al. 2007). CYPs are evolutionarily and structurally conserved proteins, and have been identified in both prokaryotes and eukaryotes. Cyclophilins are found in all cellular compartments and are involved in multiple processes. In plants, CYPs can regulate organogenesis (Fulgosi et al. 1998), hormone signaling (such as GA,
and BR) (Li et al. 2010; Zhang et al. 2013), as well as defense responses (Berardini et al. 2001; Li et al. 2014; Trupkin et al. 2012). In Arabidopsis, the PPlase domain interacted with snRNP-specific proteins to regulate premRNA splicing (Lorkovic et al. 2004). In rice, overexpression of the Golgi-resident OsCYP21–4 can increase yield by promoting the accumulation of mannosidic glycoproteins (Park et al. 2017). OsCYP20–2 could integrate chilling tolerance and cell elongation by its dual-localization. Under low temperature, OsCYP20–2 functions with OsFSD2 to scavenge reactive oxygen species (ROS) in chloroplasts, and the nuclear-localized variant OsNuCYP20–2 targets SLENDER RICE1 to regulate GA signaling (Ge et al. 2020). However, it remains unknown if CYPs regulate grain size by other mechanisms.

Alternative splicing is crucial for plant growth and stress response. The removal of introns from pre-mRNA is subtly modulated by a variety of splicing factors. Several nuclear-localized CYPs function in spliceosome assembly in humans (Adams et al. 2015; Galat 1993; Wang and Heitman 2005). Cycophilin USA-CyP formed independent complexes with the splicing factor hPrp18 and hPrp4 to participate in pre-mRNA splicing (Horowitz et al. 2002). SYF2/NTC31/p29, is a critical splicing factor hPrp4 to participate in pre-mRNA splicing (Horowitz et al. 2002). SYF2 results in cell cycle arrest by activation of the spindle checkpoint in Saccharomyces cerevisiae (Dahan and Kupiec 2002). In humans, overexpression of SYF2 promotes cell proliferation in breast cancer (Shi et al. 2017). However, the function of SYF2 orthologs in plants is completely unknown.

Here, we demonstrated that OsCYP20–2 can interact with splicing factor OsSYF2 to impact grain length. Both the oscyp20–2 t1 mutant and OsSYF2 RNA interference (RNAi) lines exhibited shorter grain length than wild type. RNA-sequencing analysis demonstrated that OsSYF2 was involved in pre-mRNA alternative splicing and transcriptional regulation of some sugar metabolism pathway genes. Thus, our findings revealed that OsCYP20–2 and OsSYF2 interact to determine grain size in rice.

**Results**

**OsCYP20–2 Regulates Grain Size and Inflorescence Architecture**

We previously reported that CYP20–2 regulated plant height and chilling tolerance in rice (Ge et al. 2020). Unexpectedly, we found that the mutation of OsCYP20–2 also reduced grain length, while brown grain width and thickness were unchanged (Fig. 1a–c). Notably, the 100-grain weight of the knock-out mutant oscyp20–2 t1 was substantially decreased compared to that of wild type Hwayoung (HY) (Fig. 1d). Moreover, the spikelet hull and the lemma epidermal cells of oscyp20–2 t1 were shorter than that of HY (Fig. 1e). These observations suggested that OsCYP20–2 determines grain length via regulation of cell length.

Additionally, GUS staining of OsCYP20–2 promoter-driven GUS transgenic plants showed that OsCYP20–2 is highly expressed in the various stage of panicle from P2 to P6 (Jain et al. 2007) (Fig. 1f). Consistently, panicle length and branch number were decreased in oscyp20–2 t1 compared to HY (Fig. 1g).

**OsCYP20–2 Physically Interacts with OsSYF2**

To search for potential interacting proteins of OsCYP20–2, the rice yeast two-hybrid cDNA library was screened using OsCYP20–2–BD as bait. An accessory splicosome protein named Synthetic lethal with cdc forty 2 (OsSYF2) was identified. SYF2 was reported to participate in pre-mRNA splicing and gene expression regulation in yeast (Vincent et al. 2003). The interaction of OsCYP20–2 and OsSYF2 was further confirmed by yeast two-hybrid and pull-down assays (Fig. 2a, b). Furthermore, OsSYF2 shared a similar expression pattern with OsCYP20–2 (Figs. 1f and 2d). Fluorescent microscopy was used to show that OsSYF2-GFP fluorescence overlapped with the nuclear marker H2B-mCherry in rice protoplasts (Fig. 2c). These results indicated that OsSYF2 is nuclear-localized and might interact with OsCYP20–2 to regulate seed development.

**OsSYF2 Positively Regulates Grain Length**

To confirm whether OsSYF2 determines grain size or shape, the full-length CDS of OsSYF2 under the control of maize Ubiquitin 1 promoter was transformed into the japonica variety Zhonghua 11 (ZH11). Meanwhile, OsSYF2 RNA interference lines (SYF2-Ri) were obtained by transforming a construct containing a 410-bp fragment targeting the second and third exons of OsSYF2 (Fig. 3a) into ZH11. The SYF2-overexpression lines (SYF2-OE) displayed increased brown grain length and 100-grain weight compared to ZH11 (Fig. 3b, d). The RNAi lines Ri9 and Ri11 possessed similar grain phenotypes as oscyp20–2 t1, with decreased brown grain length, grain width, and 100-grain weight (Fig. 3b-d).

Subsequently, the cell size of epidermal cells in the outer glumes of OE6 and Ri9 lines were observed using scanning electron microscopy. Our results revealed that the cell length of OE6 was longer than that of ZH11, while Ri9 showed shorter cells compared to ZH11 (Fig. 3e). These results suggested that OsSYF2 positively regulates grain size by increasing cell length.

**Knock-Down of OsSYF2 Causes Similar Panicle Phenotypes as oscyp20–2 t**

Besides grain size, plant height and rachis length were also investigated in the transgenic lines. In OsSYF2-
RNAi lines, plant height showed shorter than wild type ZH11 when seedlings were at three leaf stage (Fig. 4a). Cell size and number are the major factors that contribute to plant organ size. To explore the mechanism of OsSYF2 regulation of plant height, we investigated the cell size of the second leaf sheath of wild type and RNAi lines at three leaf stage (Tang et al. 2018). The statistical results showed that the cell lengths in Ri9 and Ri11 were shorter than those of ZH11 (Fig. 4b, c).

In OsSYF2-RNAi lines, plant height, panicle length, and branch number were all decreased compared to ZH11 at mature stage. However, no significant changes were observed in OsSYF2-overexpression lines (Fig. 4d-g). Overall, down-regulating the expression of OsSYF2 resulted in decreased plant height and panicle length.

**OsSYF2 Associates with Sugar Metabolism**

Considering OsSYF2 is a putative subunit of the spliceosome and impacts grain size, the transcription of OsSYF2 was detected in different developmental stages of the panicle. Our qRT-PCR results showed that OsSYF2 had the highest transcription levels in 6-cm panicles (Fig. 2d). Then, 6-cm panicles of ZH11 and Ri9 were harvested and used for RNA-seq analysis. There were 32,
502 and 32,228 transcripts detected in the transcriptome of ZH11 and Ri9 panicles, respectively (Fig. 5a). Two thousand seven hundred sixty genes were differentially-expressed between Ri9 and ZH11 as determined by a 2-fold cutoff (false discovery rate < 0.05). Of the 2760 differentially-expressed genes, 59.2% (1633/2760) genes were up-regulated, and 40.8% (1127/2760) were down-regulated in Ri9 related to ZH11 (Fig. 5b). The Kyoto Encyclopedia of Genes and Genomes pathway (KEGG) enrichment analysis revealed that the differentially expressed genes were enriched in the functions of RNA transport, mRNA surveillance, RNA degradation, and sugar metabolism (Fig. 5c).

Starch is the primary component of the rice endosperm and considered to be biosynthesized using ADP-glucose as a substrate (Braun et al. 2014). Sugar metabolism and transport influence starchy content and morphology (Bai et al. 2016). The contents of sucrose and maltose were decreased in Ri9 and Ri11 lines compared to ZH11 (Fig. 5d, e). Scanning electron microscopy images showed that the starch granules were smaller and more tightly arranged in the endosperm of Ri9 than ZH11 (Fig. 5f). Thus, our data suggested that OsSYF2 regulates the expression of genes involved in sugar metabolism and affects the formation of starch granules.

OsSYF2 is Significant for Alternative Splicing
To understand the effects of OsSYF2 on RNA splicing, the alternative splicing (AS) events were analyzed from our transcriptomic data. Interestingly, the ratio of skipping exon (SE) is increased by 0.6% and the ratio of intron retention (IR) is decreased by 0.4% in Ri9 compared to ZH11 (Fig. 5g). Furthermore, functional assignment of different SE event genes by KEGG analysis indicated that OsSYF2 SE events were associated with nucleotide excision repair and non-homologous end-joining (Fig. 5h). Meanwhile, the genes with differential IR events were involved in the spliceosome assembly, carbon metabolism, and the pentose phosphate pathway (Fig. 5h). These results indicated that OsSYF2 plays an important role in alternative splicing.

Discussion
Grain shape is a complex trait controlled by many factors that include transcription factors, hormones, miRNAs, and cyclophilins (Duan et al. 2015; Ishimaru et al. 2013; Park et al. 2017; Si et al. 2016). Cyclophilins participate in the photosynthetic process, plant growth and development, and stress responses (Cheong et al. 2017; Jing et al. 2015; Park et al. 2013). Here we revealed that the cyclophilin OsCYP20–2 directly interacts with
OsSYF2, a subunit of the spliceosome, to regulate grain length in rice. Both the mutation of OsCYP20–2 and knockdown of OsSYF2 resulted in short grains. Scanning electron microscopy demonstrated that both genes positively regulate cell length in the lemma (Figs. 1e and 3e). The changes in starch morphology, sucrose content, and maltose content suggested that sugar metabolism was abnormal in the OsSYF2-RNAi lines (Fig. 5e, f). In humans, eight nuclear-located cyclophilins interact with the spliceosome to regulate transcription and pre-mRNA splicing. Among the eight nuclear cyclophilins, peptidyl prolyl isomerase-like 2 (PPIL2), spliceosome-associated protein CWC27 homolog (CWC27), peptidyl prolyl isomerase G (PPIG), and peptidyl prolyl isomerase H (PPIH) had more substantial effects on splicing and spliceosome assembly than peptidyl prolyl isomerase E (PPIE), peptidyl prolyl isomerase-like 2 (PPIL2), peptidyl prolyl isomerase-like 2 (PPIL3), and peptidyl prolyl isomerase domain and WD repeat-containing protein 1 (PPWD1) (Rajiv and Davis 2018). PPIL1, a minimal signal isomerase domain cyclophilin, interacted with SYF2 assuming to regulate pre-mRNA splicing (Stegmann et al. 2010). Considering that OsSYF2 is a predicted subunit of the rice spliceosome, we examined the splicing patterns of genes related to sugar metabolism. The intron retention and skipping exon events were changed in the OsSYF2-RNAi line relative to the ZH11 control (Fig. 5g, h). Based on these results, we speculated that
the interaction between OsCYP20–2 and OsSYF2 is important for mRNA processing and splicing, thereby regulating grain size by way of cell elongation and sugar metabolism.

OsCYP20–2 and OsSYF2 had variable expression in different developmental stages of the panicle (Figs. 1f and 2d). Moreover, OsCYP20–2 and OsSYF2 are necessary for panicle length regulation in rice (Figs. 1g and 4a). Our transcriptome data showed that OsSYF2 affects the expression of genes related to RNA surveillance, RNA degradation, RNA transport, and sugar metabolism (Fig. 5c). Comparing the AS events in ZH11 and Ri9, we found that OsSYF2 majorly impacts the frequency of skipping exon (SE) and intron retention (IR) events (Fig. 5g). The genes with differential SE and IR were enriched for functions in carbon metabolism, spliceosome assembly, and nucleotide excision repair (Fig. 5). The genes with differential SE and IR were enriched for functions in carbon metabolism, spliceosome assembly, and nucleotide excision repair (Fig. 5). Among the differential IR events between ZH11 and Ri9, LOC_Os03g15050 encodes a phosphoenol/pyruvate carboxykinase (PEPCK), which facilitates the conversion of lipids to sugars (Penfield et al. 2004; Malone et al. 2007). LOC_Os03g15050 encodes a transaldolase, which plays an important role in glycolysis and the pentose phosphate pathway (PPP) (Caillau and Paul Quick 2005). In addition, LOC_Os02g44550 (an NADP-dependent malic enzyme) and LOC_Os07g42440 (a glycolate oxidase) have important functions in photosynthesis (Cui et al. 2016; Tao et al. 2016). It was recently found that splicing factors participated in flower and embryo development in Arabidopsis (Xiong et al. 2019a, b), and seed development in maize (Fouquet et al. 2011). Taken together, OsSYF2 may regulate the frequency of IR and ES events participating in panicle development.

Conclusions

Overall, we speculated that OsCYP20–2 interacted with OsSYF2 to regulate gene expression and the frequency of AS events to affect grain shape and panicle length. However, how OsSYF2 affects AS events is still unknown. This study provides further insight into the genetic mechanisms that impact grain yield in rice and provides new targets for molecule breeding to improve rice yield.

Methods

Plant Materials and Growth Conditions

The line oscyp20–2t1 is a T-DNA insertion mutant in the Hwayoung background (a japonica variety of Oryza sativa). Two independent RNAi lines (Ri9 and Ri11) and three overexpression lines (OE3, OE6, and OE9) of OsSYF2 were generated in the Zhonghua 11 (ZH11, a japonica variety of Oryza sativa) background, separately.
The 410-bp fragment from 398 to 807 in the coding sequence of OsSYF2 and the full-length OsSYF2 coding sequence were inserted into pTCK303 and pUN1301, respectively, to generate transgenic materials. The 2018 bp sequence upstream of the OsCYP20–2 coding sequence and the 2067 bp sequence upstream of the OsSYF2 coding sequence fragment were inserted into pCAMBIA1391z vectors to generate GUS constructs. The GUS constructs were transformed into ZH11 to generate transgenic plants. The young panicles from P2 to P6 stage (Jain et al. 2007) were selected for GUS staining.

Rice plants were cultured in the field in the summer in Beijing (40°06'N), China, under normal conditions,
including irrigation and fertilization (Che et al. 2015). Within a row, the distance between the plants was 15 cm, which the distance was 20 cm between the rows. After all measured materials fertilization on October, statistical analysis was performed on agriculture traits of transgenic plants.

RNA Isolation and qRT-PCR
For identifying the transgenic plants, three-leaf stage seedlings of ZH11 and OsSYF2 transgenic lines were selected for RNA isolation via an RNA extraction kit (Qiagen, Germany). Various tissues, including root, young stem, young leaf, mature leaf, flower, panicles of 3–4 cm, 4–5 cm, 6 cm, 8 cm, 10 cm, and 12 cm, were harvested until RNA isolation. Subsequently, 2 μg RNA was used to produce cDNA with SuperScript reverse transcriptase (Invitrogen) with random primers. For qRT-PCR, the UBIQUITIN gene was used as an internal control. Each biological sample consisted of three technical replicates. The primers are listed in Table S1.

Subcellular Localization
To study the subcellular localization of OsSYF2, GFP was fused to the C terminus of OsSYF2 in the pBI221 vector. The plasmids were co-transformed into rice protoplasts, with the nuclear marker construct AtH2B-mCherry via the polyethylene glycol (PEG 4000) method (Bart et al. 2006). After culturing overnight, the transformed protoplasts were observed with a Leica TCS SP5 fluorescence microscope.

Yeast Two-Hybrid Assays
The full-length of OsCYP20–2 and OsSYF2 ORFs were separately cloned into pGBK7T7 and pGADT7, respectively. Then, the constructs were transformed into the yeast strain AH109 and cultured on SD/−Trp-Leu-His-Ade medium. After 3 days, positive clones were selected to culture; YX, KC, and QG analyzed the data; YX, KC, JZ and QG wrote the paper. The author(s) read and approved the final manuscript.

Scanning Electron Microscopy
Fresh panicles were fixed in FAA at 4 °C overnight and dehydrated in an ethanol series ranging from 75% ~ 100% by every for 30 min at each step. Then, the samples were critical point dried by CO2 and imaged on a scanning electron microscope (S-4800 FESEM, Japan).

GST-Pull Down
For GST pull-down assays, 50 μL of glutathione sepharose beads (GE Health, Glutathione Sepharose 4B) was washed with 1 × PBS three times. Fifty micrograms of purified pCold-His-SYF2 protein was dissolved in reaction buffer (20 mM Tris, 150 mM NaCl, 2 mM DTT, 5% glycerol, and PMSF) and mixed with Sepharose beads conjugated to GST or GST-CYP20–2 for 2 h at 4 °C. After washing three times, the target protein was centrifuged and analyzed by 10% SDS-PAGE and western blotting with anti-GST or anti-His antibodies.

**Determination of Glucose, Sucrose, and Maltose Content**
The mature seeds of ZH11 and OsSYF2-RNAi were dried and peeled. Seeds were ground into a powder and 2 mL ddH2O was added to 0.2 g sample, then subjected to ultrasonic extraction for 60 min. Samples were centrifuged at 15,000 rpm for 10 min at room temperature. The supernatant was filtered through the water system and the samples were measured via HPLC (Agilent 1260 Infinity). The velocity of flow was 1 mL/min in Agilent Zorbax NH2 (250 mm×4.6 mm, 5 μm) at room temperature.

**RNA-Seq and Data Analysis**
RNA-Seq libraries were constructed according to the manufacturer’s protocol. An Illumina HiSeq 2000 platform was selected for RNA-seq by the Beijing Genomics Institute. After discarding the low-quality reads, Bowtie was used to rebuild the genome, which used the MSU Rice Genome Annotation Project Release 7 (http://rice.plantbiology.msu.edu/) as the reference genome.

MeV v4.9 (Saeed et al. 2003) was used to generate heat maps showing gene expression levels. Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways analysis was performed using KOBAS v.2.0 (Xie et al. 2011). Significant enrichment was detected with a P value < 0.05.

**Supplementary information**
Supplementary information accompanies this paper at https://doi.org/10.1186/s12284-020-00425-0.

**Additional file 1: Supplemental Table S1.** Primers used in this study.

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**Authors’ contributions**
YX, KC, and QG designed the project; QG performed the physiological, biochemical and genetic experiments; YT and WL performed the material culture; YX, KC, and QG analyzed the data; YX, KC, JZ and QG wrote the paper. The author(s) read and approved the final manuscript.

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Availability of Data and Materials
The datasets in this study are available from the corresponding author on reasonable request.

Ethics Approval and Consent to Participate
Not applicable.

Consent for Publication
Not applicable.

Competing Interests
The authors declare that they have no competing interests.

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