OsCOP1 regulates embryo development and flavonoid biosynthesis in rice (Oryza sativa L.)

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Abstract

Key message Novel mutations of OsCOP1 were identified to be responsible for yellowish pericarp and embryolethal phenotype, which revealed that OsCOP1 plays a crucial role in flavonoid biosynthesis and embryogenesis in rice seed.

Abstract Successful production of viable seeds is a major component of plant life cycles, and seed development is a complex, highly regulated process that affects characteristics such as seed viability and color. In this study, three yellowish-pericarp embryo lethal (yel) mutants, yel-hc, yel-sk, and yel-cc, were produced from three different japonica cultivars of rice (Oryza sativa L). Mutant seeds had yellowish pericarps and exhibited embryonic lethality, with significantly reduced grain size and weight. Morphological aberrations were apparent by 5 days after pollination, with abnormal embryo development and increased flavonoid accumulation observed in the yel mutants. Genetic analysis and mapping revealed that the phenotype of the three yel mutants was controlled by a single recessive gene, LOC_Os02g53140, an ortholog of Arabidopsis thaliana CONSTITUTIVE PHOTOMORPHOGENIC 1 (COP1). The yel-hc, yel-sk, and yel-cc mutants carried mutations in the RING finger, coiled-coil, and WD40 repeat domains, respectively, of OsCOP1. CRISPR/Cas9-targeted mutagenesis was used to knockout OsCOP1 by targeting its functional domains, and transgenic seed displayed the yel mutant phenotype. Overexpression of OsCOP1 in a homozygous yel-hc mutant background restored pericarp color, and the aberrant flavonoid accumulation observed in yel-hc mutant was significantly reduced in the embryo and endosperm. These results demonstrate that OsCOP1 is associated with embryo development and flavonoid biosynthesis in rice grains. This study will facilitate a better understanding of the functional roles of OsCOP1 involved in early embryogenesis and flavonoid biosynthesis in rice seeds.

Introduction

Light is a crucial signaling component that controls a variety of physiological and developmental processes in morphogenesis, germination, and flowering in plants. Light signaling is also involved in metabolic alterations, including the biosynthesis of chlorophyll (Ilag et al. 1994; Pattanayak et al. 2005) and several types of pigments (Takos et al. 2006; von Lintig et al. 1997; Zoratti et al. 2014). These light signaling pathways are triggered upon perception of light signals by multiple photoreceptors, such as red/far red light-sensing phytochromes (phyA to phyE), UV-A/blue light-sensing cryptochromes (cry1 and cry2), phototropins (phot1 and phot2), Zeitlupes (ZTL), and the UV-B-sensing photoreceptor (UVR8), which transmit signals to genes that regulate developmental and metabolic processes such as anthocyanin biosynthesis in higher plants (Jiao et al. 2007; Kami et al. 2010).

The light signaling regulator CONSTITUTIVELY PHOTOMORPHOGENIC 1 (COP1) is an essential negative regulator of light-mediated plant development. In darkness, COP1 acts as an E3 ubiquitin ligase to repress light signaling by targeting numerous photomorphogenic-promoting transcription factors, including ELONGATED HYPOCOTYL5 (HY5), HY5-HOMOLOG (HYH), LONG AFTER FAR-RED1 (LAF1), and LONG HYPOCOTYL IN FAR-RED1 (HFR1), for ubiquitination and degradation in the nucleus (Holm et al. 2002; Osterlund et al. 2000; Seo et al. 2003; Yang et al. 2005). In addition to acting as a repressor of photomorphogenesis (Ang et al. 1998; Deng et al. 1991; Ma et al. 2002; Seo et al. 2003), COP1 functions as a crucial regulator in several biological processes, such as flowering

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time (Jang et al. 2015; Tanaka et al. 2011; Xu et al. 2016a), circadian regulation (Yu et al. 2008), stomatal development (Kang et al. 2009; Lee et al. 2017), UV-B signaling (Favory et al. 2009; Huang et al. 2014; Yin et al. 2015), abiotic stress responses (Kim et al. 2016), and hormone signaling (Pacin et al. 2016; Zheng et al. 2017). In particular, COP1 plays a central role in regulating anthocyanin biosynthesis and fruit coloration by complexing with other proteins (Li et al. 2012; Liang et al. 2020; Maier et al. 2013).

The COP1 protein comprises three distinct structural domains that are conserved in both higher plants and vertebrates: a zinc-binding RING finger motif, a coiled-coil domain, and a WD40 repeat domain (Stacey et al. 1999). These highly modulated domains of COP1 are involved in self-association, dimerization, and protein interactions with other proteins (Holm et al. 2001; Torii et al. 1998). The RING finger domain plays a critical role in mediating the recognition of substrates as well as the transfer of ubiquitin to substrates and to the RING finger domain itself (Joazeiro and Weissman 2000; Kosarev et al. 2002). The coiled-coil domain is involved in dimerization and interactions with other proteins, including CIP1, SPA1, and CSU2 (Hoecker and Quail 2001; Lee et al. 2018; Matsui et al. 1995; Xu et al. 2015). In COP1, the WD40 domain is found at the C-terminus and consists of seven WD40 repeats that mediate protein–protein interactions. COP1 directly interacts with several regulators, such as HY5 and HYH, and photoreceptors (cytochromes and UVR8) via its WD40 domain (Holm et al. 2001, 2002; Ponnu et al. 2019; Yin et al. 2015).

Mutagenesis is an important strategy to create mutations which are valuable sources for functional genomics as well as crop improvement. In particular, relatively small genome size of rice has enabled a genome-wide saturation mutagenesis and revelation of full spectrum of gene functions by generating allelic series of mutations with small population (Henikoff and Comai 2003). Over the past several decades, numerous mutagenesis methods have been developed for rice, but chemical and physical mutagens have been widely applied to induce genetic variability and to study gene function due to its simple and low cost strategy (Viana et al. 2019; Wang et al. 2013). The genome changes between wild-type and mutants caused by chemical and physical mutagen have been identified efficiently in recent years using Targeting Induced Local Lesions IN Genomes (TILLING) (Till et al. 2007) and MutMap (Abe et al. 2012), next-generation sequencing (NGS)-based gene mapping tool in rice. These researches through mutagenesis would be helpful to identify unknown pathways or genes and could be directly applied to the rice breeding.

Genome editing technology using the Clustered Regularly Interspaced Short Palindromic Repeats/Crispr/Cas9-associated Cas9 (CRISPR/Cas9) system is a powerful and promising tool for precise modification of genome sequences in crop plants. In recent years, the CRISPR/Cas9 system has been widely adopted for crop improvement due to its simplicity, specificity, efficiency, and feasibility compared with zinc finger nuclease and transcription activator-like effector nuclease editing technologies (Khan 2019). Studies employing CRISPR/Cas9 genome editing tools have been used to validate gene functions and improve useful traits in rice, such as herbicide resistance (Li et al. 2016a; Sun et al. 2016), rice blast disease resistance (Wang et al. 2016), cold tolerance (Zeng et al. 2019), salt tolerance (Zhang et al. 2019), amylose content (Sun et al. 2017), and grain size and/or yield (Li et al. 2016b; Xu et al. 2016b). Ongoing improvements in genome editing tools will accelerate the pace of crop improvement and gene validation experiments.

The function of COP1 in photomorphogenic development and anthocyanin biosynthesis has been extensively studied in Arabidopsis. However, little is known regarding the role of COP1 in flavonoid biosynthesis and embryo development in rice. In this study, novel OsCOP1 mutants that exhibited a yellowish pericarp color and seed lethality were identified, and CRISPR/Cas9 gene editing tool was used to validate the mutant phenotype. These findings provide new insights into the regulation of flavonoid biosynthesis and embryo development in rice.

Materials and methods

Plant materials and growth conditions

Three yellowish-pericarp embryo lethal (yel) mutants, yel-hc, yel-sk, and yel-cc, were derived from japonica rice cultivars Hwacheong, Samkwang, and Chucheong, respectively, by induction with N-methyl-N-nitosourea. Due to their embryonic lethality, the mutants were maintained as heterozygotes. F2 mapping populations were developed from crosses between the three heterozygous yel mutants and Milyang 23 (M.23, Korean indica-type rice). F2 populations derived from crosses between the three heterozygous yel mutants and their respective wild types, Hwacheong, Samkwang, and Chucheong, were used to calculate segregation ratios. F2 populations, wild-type varieties, and yel mutants were cultivated in a paddy field at the Experimental Farm of Seoul National University, Suwon, Korea.

Phenotypic analysis

Dimensions of dehulled rice including length, width, and thickness were measured in a total of 90 matured grains (30 seeds × 3 replicates) using digimatic calipers (Mitutoyo, Japan). Hundred-grain weights were measured (100 seeds × 3 replicates; 10% water content) using an analytical balance (CAS Corporation, NJ, USA). Phenotypic data...
collected from the three yel mutants and their respective wild types were statistically analyzed using SAS version 9.4 (SAS Institute Inc., Cary, NC, USA).

**Morphological and histological observations of developing seed**

Caryopses of Hwacheong and the yel-hc mutant were collected at several stages of seed development and imaged using HD’MEASURE software (HANA Vision Systems, Korea). For histological examination, samples were collected at 3 days after pollination (DAP), 4 DAP, 5 DAP, 7 DAP, 10 DAP, and 40 DAP (matured), and fixed in formalin-acetic acid-alcohol fixative (3.7% formaldehyde, 5% acetic acid, and 50% ethanol) at 4 °C. Seeds were dehydrated by soaking in a series of ethanol solutions of increasing concentration (50%, 70%, 80%, 90%, 95%, and 100%) for 3 h each, followed by soaking in 100% ethanol overnight at room temperature. The dehydrated samples were soaked in a series of mixtures of absolute ethanol and Histo-Clear (3:1, 1:1, 1:3) at 60 °C. Finally, samples were stored in 100% Histo-Clear overnight at room temperature. For paraffin infiltration, Para-plast® (Sigma, USA) was gradually added to Histo-Clear (3:1, 1:1, 1:3) at 60 °C. Samples were dehydrated by soaking in 100% Histo-Clear overnight at room temperature. For paraffin infiltration, Para-plast® (Sigma, USA) was gradually added to Histo-Clear (3:1, 1:1, 1:3) at 60 °C. Finally, samples were stored in 100% ethanol for 24 h at 60 °C. Paraffin-infiltrated samples were embedded in an embed block and cut into 10-μm sections using a HM 340 E Rotary Microtome (Microm Lab, Germany). The sections were stained with 1% Safranin O and observed under a CX31 Microscope (Olympus, Japan).

**Map-based cloning of yel gene**

Genomic DNAs were extracted from a total of 898 matured mutant-type seeds and young heterozygous and wild-type leaves of F2 individuals derived from the yel-hc x Milyang23 population. To determine the locus of the gene underlying the yel-hc mutant, bulked segregant analysis (BSA) (Michelmore et al. 1991) was performed with the first batch of STS sequence-tagged-site (STS) markers, which were developed by designing primers based on the DNA sequence differences between indica and japonica rice cultivars (Chin et al. 2007). Next, additional STS primers were designed with Primer3 version 0.4.0 (http://frodo.wi.mit.edu/primer3) based on available rice genome sequence data (http://www.ncbi.nlm.nih.gov) and were used to refine the region of the yel-hc locus. Additionally, a total of 192 F2 individuals were used for each of the yel-sk and yel-cc mutant gene mapping. The overall mapping strategy and primers were the same as for yel-hc. Primers designed and used in this study are detailed in Table S1.

**Sequence analysis of candidate genes**

The full-length genomic DNA sequence of candidate genes in Hwacheong and yel-hc was divided into several overlapping segments and then amplified. PCR products were purified using a PCR purification kit (iNtRON Biotechnology, Korea), TA-cloned into the pGEM-T Easy Vector (Promega, USA), and transformed into E. coli strain DH5α. The obtained sequences were compared using CodonCode Aligner software (version 1.6.3; CodonCode Corporation, MA, USA).

**Vector constructs and rice transformation**

CRISPR/Cas9 vectors to knock out COP1 were constructed as described previously (Lowder et al. 2015). Briefly, two guide RNAs (gRNAs) for each target were designed using the web-based tools CRISPR RGEN Tools (http://www.rgenome.net/cas-designer/) (Park et al. 2015) and CRISPRdirect (https://crispr.dbcls.jp/) (Naito et al. 2015). The two individual gRNAs were cloned into two RNA expression vectors, pYPIQ131C (Addgene plasmid #69,284) and pYPIQ132C (Addgene plasmid #69,285), under the expression of the OsU6 promoter. The gRNA expression cassettes were then assembled into the Golden Gate recipient vector pYPIQ142 (Addgene plasmid #69,294). For the Cas9 entry vector, pYPIQ165 (Addgene #109,327), which is driven by an egg cell-specific promoter, was used. Final binary T-DNA vectors were prepared using a Gateway assembly LR reaction with the Cas9 entry vector (pYPIQ165), gRNA cassettes (pYPIQ142), and pMDC99 (binary vector). The full-length OsCOP1 cDNA was amplified from Hwacheong cDNA and used for construction of the overexpression vector. The amplified fragment was transferred into the pMDC32 Gateway binary vector containing the cauliflower mosaic virus (CaMV) 35S promoter using a pCR™ 8/GW/TOPO® TA Cloning vector containing the cauliflower mosaic virus (CaMV) 35S promoter, was denoted as 35S::OsCOP1. For the promoter-GUS (β-glucuronidase) assay, the vector construction and rice transformation

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The final constructs were transformed into seeds of the japonica cultivar Dongjin, via Agrobacterium-mediated transformation using the LBA4404 strain, as described previously (Nishimura et al. 2006), with slight modifications. Primers used for vector construction and genotyping are listed in Table S1.
Histochemical GUS assay

Histochemical GUS-staining was performed as described previously (Jefferson et al. 1987). X-Gluc buffer solution was vacuum-infiltrated into tissue samples, which were incubated overnight at 37 °C, after which the staining solution was serially replaced with 95% and 70% (w/v) ethanol to remove the chlorophyll. Staining patterns in tissue samples were examined using HD MEASURE software (HANA Vision, Korea).

Quantitative real-time PCR analysis

Total RNA was extracted from 7-day-old seeds of the three yel mutants and their respective wild types, 35S::OsCOP1, knock-out lines, and Dongjin using a TaKaRa MiniBEST Plant RNA Extraction Kit (Takara Bio, Japan). Total RNA samples were converted into first-strand cDNA using M-MLV reverse transcriptase (Promega, USA), and qRT-PCR was performed using SYBR Premix Ex Taq (Takara Bio, Japan) on a CFX96™ Real-Time PCR Detection System (Bio-Rad, USA), according to the manufacturer’s instructions. Primers used for qRT-PCR analysis are listed in Table S1. Expression levels of OsCOP1 were normalized relative to ACTIN, a housekeeping gene. Data were analyzed using the comparative Ct method. Expression levels were compared using a two-tailed Student’s t test.

Quantitative analysis of major flavonoids using HPLC

HPLC sample preparation and quantitative analysis were performed as described previously (Kim et al. 2018). Whole grain, embryo, and endosperm tissues from yel-hc, Hwacheong, and transgenic overexpression seeds were powdered, and three technical replicates were prepared for quantitative analysis of six representative flavonoid compounds (isorientin, isoorientin 2″-O-glucoside, vitexin 2″-O-glucoside, isoscoparin 2″-O-glucoside, isoscoparin, and isovitexin). HPLC was performed on a XBridge C18 column (250 × 4.6 mm, 5 μm; Waters, Milford, MA, USA) using an Ultimate 3000 UHPLC system (Thermo Fisher Scientific, Inc., MA, USA) equipped with an autosampler, a gradient system, and a diode array detector (DAD). Chromatograms were acquired at 365 nm, and photodiode array spectra were recorded from 190 to 800 nm. The injection volume of each sample was 5 μL.

Results

Morphological characterization of yel mutants

Three yellowish-pericarp embryo lethal (yel) mutants, yel-hc, yel-sk, and yel-cc, were derived from japonica rice cultivars Hwacheong, Samkwang, and Chucheong, respectively. The most prominent characteristic of the mutants was a bicolor grain phenotype. Embryos were black in all three yel mutants, and the pericarp was yellowish in yel-hc and yel-sk grains and mixed yellow-purple in yel-cc grains (Fig. 1a). The black pigmentation in yel mutant embryos was first observed at approximately 5 DAP and the yellowish pericarp color was clearly visible at 35 DAP, the late-maturing stage (Fig. 1b). All three yel mutants exhibited significantly reduced grain length and thickness compared with wild type, and grain width was also significantly lower in the yel-sk and yel-cc mutants. Correspondingly, hundred-grain-weight was also significantly lower in yel mutants than in wild type. However, there was no significant difference between yel mutants and wild type in the length-to-width ratio of de-hulled rice grains (Table 1). These results indicated that the yel mutations influenced embryo and pericarp color, grain size, and grain weight in rice.

Embryo development of yel mutants

Homozygous yel mutant seeds failed to germinate under both standard germination testing and on MS medium (data not shown). Longitudinal sections of yel-hc seeds were examined to determine when defects occurred during yel mutant embryo development. Normal embryonic development was observed in both yel-hc and wild-type embryos during early embryogenesis, including at the globular stage (Fig. 2a and f). The coleoptile primordium was differentiated in both the wild-type and yel-hc mutant, and no morphological differences were observed at 4 DAP (Fig. 2b and g). In wild-type embryos, the coleoptile, epiblast, and scutellum had formed, and the shoot and root meristems had begun to differentiate, at 5 DAP. However, the overall morphology of yel-hc embryos differed from that of wild type at 5 DAP. It was not possible to determine organ differentiation in the yel-hc embryo as the shape and structure of organs were unclear (Fig. 2c and h). At more advanced developmental stages, all organs were normally differentiated and developed in wild-type embryos, whereas abnormally differentiated organs were observed in yel-hc embryos (Fig. 2d, e and i, j). In the mature embryo, all of the organs were evident in wild type, but, despite completion of embryogenesis, irregular and unstructured organs were observed in the yel-hc mutant (Fig. S1). These results demonstrated that the mutation of YEL caused defects in embryonic differentiation and development, resulting in seed lethality and failure to germinate in yel mutants.

Genetic analysis and map-based cloning of the YEL gene

As homozygous yel mutants were lethal, heterozygotes of each yel mutant and their respective wild type were used in
The segregation ratios of wild-type and heterozygous plants for yel-hc, yel-sk, and yel-cc in F1 progenies were 12:14, 11:10, and 14:11, respectively. These ratios were close to a 1:1 ratio, and the segregation ratios of wild-type and yel mutant grains produced by F1 plants fitted a 3:1 ratio (WT:yel-hc = 479:145, \( \chi^2 = 1.034 < \chi^2_{0.05(1)} = 3.841, p = 0.31 \); WT:yel-sk = 496:154, \( \chi^2 = 0.593 < \chi^2_{0.05(1)} = 3.841, p = 0.44 \); WT:yel-cc = 418:128, \( \chi^2 = 0.706 < \chi^2_{0.05(1)} = 3.841, p = 0.40 \)). These results indicated that the yel phenotype was controlled by a single recessive gene.

An F2 population derived from a cross between a yel-hc heterozygous mutant and Milyang23 (M.23) was used to map the locus responsible for the yel-hc phenotype. For the first step of genetic mapping, BSA was performed using 84 polymorphic STS markers, and the yel-hc locus...
was localized between two flanking markers, S02135 and S02140, on chromosome 2 (Fig. 3a). To refine the flanking region, BSA was again conducted using an F2 population of 840 individuals with newly designed STS markers between the two flanking markers (Table S1). The yel-hc locus was finally mapped to a region of approximately 55 kb between the S02K-1 and S02K markers (Fig. 3a). Seven predicted open reading frames (ORFs) were located within this candidate region, and analysis of wild-type and yel-hc mutant sequences revealed a 706 bp deletion in the yel-hc mutant at LOC_Os02g53140 (Os02g0771100), an ortholog of AtCOP1 (Fig. 3b). This 706 bp deletion included part of the 5′ untranslated region (−19 to −1), the entire coding sequence (CDS) of exon 1 (+1 to +342), and part of intron 1 (+343 to +687), resulting in the loss of the conserved RING finger motif of LOC_Os02g53140 (Fig. 3b). A similar mapping strategy and markers confirmed that yel-sk and yel-cc harbored mutations in the same gene as yel-hc. In the yel-sk mutant, a single base change (G to A) was discovered at the splicing donor site of the 5′ end of intron 2, within the coding region for the coiled-coil motif (Fig. 3b). In the yel-cc mutant, a single point mutation (G to T) was identified in exon 7 that generated a premature stop codon (GAG to TAG; Glu409 to stop) in the WD40 repeat motif (Fig. 3b). Three transcripts were identified in homozygous yel-sk seed. The first, most abundant, transcript contained the entire 473 bp of intron 2; the second included 64 bp of intron 2; and the third transcript type was the wild-type form. The insertion of the intron 2 sequence was predicted to introduce a premature termination codon at residue 145, resulting in the loss of the coiled-coil and WD40 functional motifs (Figure S2). These results indicated that the mutations identified in LOC_Os02g53140 were responsible for the yellowish pericarp and embryo lethal phenotypes in yel mutants.

Validation of mutations causing the yel phenotype

To assess the functions of the individual OsCOP1 domains and the effects of the yel mutations, sequences for each domain were disrupted using the CRISPR/Cas9 system. Two gRNAs for each of the three constructs were designed to target crucial mutation points or domains, including the RING finger, coiled-coil, and WD40 repeat motifs (Fig. 3b). An egg cell-specific Cas9 promoter was used to overcome lethality during tissue culture and increase the frequency of mutated homozygous T1 seeds generated from positive T0 transgenic plants. T1 seeds exhibiting the yel mutant phenotype were observed when the RING finger and WD40 repeat motifs of the OsCOP1 gene were targeted (Fig. 4a), but no T1 seeds exhibiting the yel phenotype were generated upon targeting of the coiled-coil motif, even though several positive T0 transgenic events occurred. The mutations were confirmed by Sanger sequencing the target region of each domain in T1 seeds displaying the yel mutant phenotype. The most frequently identified mutations were single base pair

Fig. 2 Embryo development in wild-type and yel mutant. Upper and lower panels represent wild-type (Hwacheong) and yel-hc embryo development, respectively. a and f Embryo at 3 DAP (bar = 20 μm), b and g 4 DAP (bar = 100 μm), c and h 5 DAP (bar = 200 μm), d and i 7 DAP (bar = 300 μm), and e and j 10 DAP (Bar = 300 μm). Cp, coleoptile; Ep, epiblast; Rd, radicle; Sc, scutellum; Sm, shoot meristem; Rm, root meristem; Sa, shoot apex
insertion and deletion mutations. A large fragment deletion between two gRNAs induced in transgenic seeds targeting the WD40 repeat domain was also observed (Fig. 4b). The insertions and deletions introduced into the OsCOP1 gene in seeds with yel phenotypes were all predicted to lead to frameshifts and premature stops, and thereby to produce aberrant proteins. These results demonstrated that the yel mutant phenotype was caused by loss-of-function of crucial domains in OsCOP1.

To complement the yel phenotype, the full-length OsCOP1 cDNA was introduced into wild-type Dongjin rice under the control of the 35S promoter. Eight positive transgenic lines were produced and crossed with a heterozygous yel-hc mutant to introduce the overexpression construct (35S::OsCOP1) into the homozygous yel-hc genetic background. F1 plants harboring 35S::OsCOP1 were selected, and the phenotypes of F2 progenies from the self-pollinated F1 plant were observed. As shown in Fig. 4c, three different types of seed were segregated from the F2 progenies: normal-type (normal pericarp and embryo color), yel mutant-type (yellowish pericarp and black colored embryo), and partial recovery-type (normal pericarp color but black colored embryo). A total of 93 F2 seeds were genotyped to find the fully recovered seeds containing 35S::OsCOP1 transgene in homozygous yel-hc mutant background. All 20 yel mutant-type seeds showed yel-hc homozygous genotype (yel/yel), and all the partial recovery-type seeds contained 35S::OsCOP1 transgene in homozygous yel-hc mutant background (yel/yel + 35S_OsCOP1). However, among the normal-type seeds, none of the fully recovered transgenic yel seeds carrying 35S::OsCOP1 transgene were detected. (Table S2).

In our previous study, we demonstrated that high levels of flavonoids accumulated in the embryo and endosperm of yel-hc mutant grains (Kim et al. 2018). The contents of six major flavonoids were estimated in the embryo, endosperm, and whole grain of wild-type, yel-hc and partial recovery-type seeds. The total amounts of all six flavonoids were significantly reduced by approximately 46%, 23%, and 98% in the whole grain, embryo, and endosperm of partial recovery-type (35S::OsCOP1) seeds, respectively, compared with the yel-hc mutant seeds (Fig. 4d–f). The lack of accumulation of flavonoids was observed in the endosperm of partial recovery-type seeds overexpressing the 35S::OsCOP1 transgene, resulting in reduced flavonoid levels and restoration of wild type pericarp color. However, although the total amounts of the six main flavonoids were significantly reduced in the partial recovery-type seeds, high levels of flavonoids still accumulated in the embryo. In addition, partial recovery-type seeds still failed to germinate on both MS media and under standard germination conditions (data not shown). These results suggested that overexpression of the OsCOP1 gene in
**yel-hc** seeds partially restored the yel phenotype by reducing flavonoid accumulation in the embryo and pericarp.

**Relative expression of YEL**

Quantitative real-time PCR (qRT-PCR) analysis using developing seeds (7 DAP) showed that expression of OsCOP1 was significantly higher in the yel-sk mutant than in wild type, but no significant differences were observed between wild-type and yel-hc and yel-cc mutants (Fig. 5a).

No statistically significant differences in relative OsCOP1 expression level were observed between wild-type and CRISPR/Cas9 knockout seeds, which was consistent with the expression pattern of the yel-hc and yel-cc mutants. By contrast, OsCOP1 transcript levels were significantly...
higher in 35S::COP1 transgenic T2 seeds than in wild-type and yeI mutant seeds or knockout lines (Fig. 5b).

**OsCOP1 promoter–GUS expression pattern**

Transgenic plants were generated containing the GUS reporter gene under the control of the OsCOP1 promoter region, and GUS expression was observed in several organs. GUS signals were detected primarily in pollen, node, embryo, and pericarp tissues, but no expression was detected in leaf, spikelet, stigma, or endosperm tissues (Fig. 5c–j). Interestingly, GUS staining in seed was concentrated in the embryo and pericarp, consistent with the flavonoid accumulation pattern in yeI mutant seed.

**Discussion**

COP1 has been extensively studied as a repressor of plant photomorphogenesis, alongside DET (DE-ETIOLATED) and FUS (FUSCA), which also exhibit light-grown phenotypes in darkness. Many cop/det/fus mutants have been screened and identified in Arabidopsis (McNellis et al. 1994; Wei and Deng 1996), with the majority shown to cause...
constitutive photomorphogenesis. However, some cop/det/fus mutant alleles, such as cop1-5, cop1-8, det1-6, and fus7-1, had high levels of anthocyanin accumulation in cotyledons and leaves, and exhibited seedling lethal phenotypes under dark conditions (Castle and Meinke 1994; Ma et al. 2003; Stacey et al. 2000). Many of these phenotypically similar mutants exhibiting the ‘fusca’ phenotype were revealed to be allelic to each mutant group (Castle and Meinke 1994; Chory et al. 1989; Misera et al. 1994). However, unlike Arabidopsis, few cop1 mutants have been reported in rice. A loss-of-function mutation in PETER PAN SYNDROME (PPS), an ortholog of Arabidopsis COP1, produced dwarfed dark-green plants that showed a prolonged juvenile phase and early flowering in both short-day and long-day conditions (Tanaka et al. 2011). In this study, three allelic cop1 mutants were isolated that exhibited embryonic lethality and accumulated flavonoids in the pericarp and embryo (Figs. 1, 3). The three cop1 mutants are likely to be complete loss-of-function mutants and, based on the severity of the phenotype, the yel phenotype likely represents the cop1 null phenotype in rice. To the best of our knowledge, cop1 null mutants exhibiting a fusca phenotype similar to those of Arabidopsis mutants have not yet been reported in rice.

Two distinctive differences were apparent between rice yel mutants and Arabidopsis cop1 null mutants with the fusca phenotype, such as cop1-5, namely, differences in embryonic development and lethality and differences in pigmentation. First, despite germination delays, Arabidopsis cop1 null mutant seeds completed embryogenesis and germinated successfully, although seedling growth was severely retarded and viable plants did not develop (Castle and Meinke 1994; McNellis et al. 1994; Stacey et al. 2000). In Arabidopsis cop1 null mutants, morphological abnormalities in embryogenesis were not observed for all the lethal alleles (McNellis et al. 1994). By contrast, in rice, all the homozygous recessive yel mutant seeds failed to germinate, and all showed defective embryo morphologies during embryogenesis (Fig. 2 and Fig. S1). Due to this embryonic lethality, it was impossible to observe the photomorphogenic phenotype in homozygous yel mutant plants. Studies on embryogenesis in plants indicate that there are crucial developmental events in early embryogenesis that are important for establishing seedling architecture after germination, such as embryo pattern formation, polarity, positioning, and cell fate determination. Although these embryogenic processes are similar among plant species, there are some differences in the patterns of cell division and organ differentiation positioning (Radoeva et al. 2019; Zhao et al. 2017). Embryogenesis in Arabidopsis and rice differ in their cell division patterns, position of shoot apical meristem with respect to embryo polarity, position of radicle differentiation, and the effects of embryo-endosperm interactions on embryo development (Itoh et al. 2005; Nagasawa et al. 2013). Rice and Arabidopsis also differ in their embryogenesis-related gene expression patterns and regulatory networks (Itoh et al. 2016). Taken together, this suggests that the differences in morphological defects between rice and Arabidopsis embryos in cop1 null mutants may be due to differences in developmental processes between monocots and dicots during early embryogenesis.

Second, the seed coat and cotyledons in Arabidopsis cop1 null mutants were dark purple as a consequence of anthocyanin accumulation (Castle and Meinke 1994; McNellis et al. 1994; Stacey et al. 2000). In rice, yel mutants displayed yellowish pericarps and black embryos in dried seeds. Our previous study showed that the accumulated pigments in the embryo and pericarp were mainly C-glycosyl flavones, such as isoorientin, isovitexin, and isoscoparin, and were not anthocyanins or their derivatives (Kim et al. 2018). Flavonoid biosynthesis is controlled by structural and regulatory genes, and the genes involved in the flavonoid pathway are differentially regulated in monocot and dicot species, as indicated by the regulation patterns of regulatory genes (Petroni and Tonelli 2011), functional similarities of homologous genes (Spelt et al. 2000), and species-specific enzymes (Tohge et al. 2017). Taken together, this suggests that the biosynthetic pathways of anthocyanins and C-glycosyl flavones are regulated by transcription factors that interact with COP1, but that these transcription factors and/or their interactions with COP1 may differ among plant species. This also suggests that the transcription factors and genes downstream of COP1 that are involved in the flavonoid biosynthesis pathway may be differentially regulated in monocots and dicots.

In Arabidopsis, cop1 mutants have been separated into three classes, weak, strong, and lethal, according to their light reactivity (McNellis et al. 1994). Homozygous cop1 mutant seedlings in both the weak and strong categories exhibited light-grown characteristics when grown under dark conditions, albeit with a strongly reduced seed set. Homozygous lethal cop1 mutant seedlings exhibited a similar light-grown phenotype in darkness, but also had dark purple seeds and were seedling lethal. Mutations in the C-terminal region of the COP1 protein, containing the WD40 repeat regions, produced a much more severe phenotype, similar to that of complete loss-of-function mutants, than mutations in the N-terminal region. Many cop mutants have mutations within the WD40 repeat region, including cop1-5, cop1-7, and cop1-8, all of which display a lethal phenotype (Ma et al. 2003; McNellis et al. 1994; Zhou et al. 1998). In rice, the pps-1 mutation is located between the coiled-coil and WD40 domains, similar to weak cop1 alleles in Arabidopsis such as cop1-4 and cop1-6, which contain mutations in the N-terminal (Stacey et al. 2000). The rice pps-1 mutation was dark-green in color and had a dwarf phenotype, contrasting with pps-2,
which had a premature stop codon in the WD40 domain and exhibited ~5% germination (Tanaka et al. 2011). Of the two pps mutants, it is likely that the pps-2 mutant allele, which has a mutation in the C-terminal of COP1, represents a more severe phenotype than pps-1. In this study, three different mutations in the RING finger, coiled-coil, and WD40 repeat domains of YEL caused similar phenotypic changes: embryonic lethality and accumulation of flavonoids in the pericarp and embryo. Interestingly, although the pps-2 mutation was located between the mutations in the CPR_WD-23 and CPR_WD-13 knockout lines (Fig. 4a and b), which generated premature stop codons three amino acids before and eight amino acids after the pps-2 mutation point, respectively, pps-2 did not exhibit the same phenotype as the yel mutants, although the pps-2 germination rate was extremely low and plants died within a week. These results suggest that the yel phenotype and other severe phenotypes can be induced by knocking out the crucial domain of COP1 in rice, irrespective of whether the mutation is in the N-terminal or C-terminal.

Knockout of essential genes generally leads to lethality or sterility, and knockout approaches are therefore limited in their capacity to yield informative results regarding gene function when the gene plays a crucial role in the organism (Zimmer et al. 2019). However, in this study, complete gene knockout using the CRISPR/Cas9 system was preferable to knockdown with RNA interference (RNAi), as difficulties were experienced in obtaining transgenic RNAi plants with the yel phenotype by reducing COP1 expression. Although several transgenic plants carrying RNAi constructs were generated, the yel phenotype was not observed in T1 or T2 seeds (data not shown). Attempts were also made to generate COP1 loss-of-function plants through anther culture. However, no homozygous recessive plants were obtained from a total of 293 double haploid plants derived from anther culture (data not shown). These experiences indicated that the RNAi approach did not downregulate COP1 expression to a sufficiently low level to induce the yel phenotype, and that recessive deleterious or lethal genes, such as cop1, might cause callus cell death, eliminating haploid plants with homozygous recessive cop1 alleles during anther culture. However, CRISPR/Cas9 editing produced several heterozygous T0 transgenic plants that produced segregating yel phenotype seeds in the T1 generation. Moreover, an egg cell-specific promoter rather than the 3SS promoter was used to drive Cas9 expression (Lee et al. 2019; Wang et al. 2015) to overcome potential lethality during tissue culture regeneration of T0 plants and to increase the frequency of the yel phenotype in T1 seeds. Subsequently, homozygous mutant seeds displaying the yel phenotype were observed in the T1 generation (Fig. 4a and b). In this study, targeted gene editing was successfully used to show that COP1 was essential for flavonoid biosynthesis and embryo development, demonstrating the utility of CRISPR/Cas9 editing for examining previously uninducible phenotypes masked by lethality.

Previous studies have shown that the cop1-5 null mutant exhibited seedling lethal and high level of anthocyanin accumulation was successfully restored by full-length AtCOP1 and OsCOP1 (Holm et al. 2001; Ranjan et al. 2014; Stacey et al. 2000) in Arabidopsis. In addition, overexpression of the COP1 (1–282) fragment restored a wild-type phenotype in the cop1-5 mutant seedling in darkness (Stacey et al. 2000). However, normal pericarp color but black colored embryo seeds (partial recovery-type) were produced when the full-length OsCOP1 cDNA was overexpressed in a homozygous recessive yel-hc background seed (Fig. 4a and c). Even though the contents of flavonoids were significantly reduced in grain, embryo, and endosperm, the level of flavonoids in the embryo was still higher than that of wild-type (Fig. 4d–f). Moreover, the embryonic lethality was not rescued in the partially restored seeds. It is known that COP1 interacts with SUPPRESSOR OF PHYA-105 (SPA1) through their coiled-coil regions (Hoeger and Quail 2001), and the COP1/SPA complex targets transcription factors, such as ELONGATED HYPOCOTYL5 (HY5), HY5 HOMOLOG (HYH), LONG HYPOCOTYL IN FAR-RED 1 (HFR1), and PRODUCTION OF ANTHOCYANIN PIGMENT 1 and 2 (PAP1 and PAP2) involved in photomorphogenesis and anthocyanin biosynthesis for ubiquitination and degradation in Arabidopsis (Holm et al. 2002; Jang et al. 2005; Maier et al. 2013; Saijo et al. 2003). In particular, HY5, a bZIP transcription factor, positively regulates anthocyanin biosynthesis related structural genes through direct binding to their promoters (Shin et al. 2007). In the process of restoration, the competition between the mutated form of YEL protein (yel-hc) and normal OsCOP1/YEL to interact with SPA1 may reduce E3 ligase activity of COP1, and it may prevent ubiquitination and degradation of HY5 through the 26S proteasome pathway. Thus, remained HY5 enables to accumulate the flavonoids by inducing the expression of transcription factor in the seed. Therefore, it is assumed that the weakened E3 ligase activity of COP1 caused by the mutated COP1 that interfered the interaction between normal COP1 and SPA1 led to the incomplete COP1-mediated ubiquitination and degradation of HY5, resulting in partial restoration of yel phenotype in rice.

It was previously reported that expression of OsCOP1 was observed in almost all of rice tissues, such as calli, roots, and leaves (Tsuge et al. 2001). In addition, PPS, an ortholog of Arabidopsis COP1 was primarily expressed in leaves than shoot apices and young panicles during the juvenile-adult transition stage in rice; the lowest expression of PPS was shown in second leaves, and the expression was reached a peak in fourth and fifth leaves, but declined in higher leaves (Tanaka et al. 2011). However, unlike previous studies,
negligible GUS activity of OsCOP1 was detected in leaf in this study (Fig. 5f). It is probably because the leaf was sampled at the flowering stage, which was after finishing the juvenile-adult transition, or might be sampled at the lower leaf of plant, which had the low expression level of OsCOP1 for GUS staining.

In this study, we identified three novel mutations responsible for the yellowish-pericarp and embryo lethal (yel) phenotype, and showed that COP1, a key repressor of light signaling, played a role in flavonoid biosynthesis and embryogenesis in rice seed. Further examination of the YEL locus will facilitate a better understanding of the molecular mechanisms involved in embryo development, and will provide novel insights into the roles of flavonoid biosynthesis-related genes and their interactions in rice.

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**Author contribution statement** BK designed and performed research, analyzed data, and wrote the manuscript. RP, GL, EK, YL, SW, and R performed research and analyzed data. WJ developed the plant materials. HJK, ES, and MT designed and supervised the experiment and improved the manuscript. All authors read and approved the final manuscript.

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**Declarations**

**Conflict of interest** On behalf of all authors, the corresponding author states that there is no conflict of interest.

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