# MADS78 and MADS79 Are Essential Regulators of Early Seed Development in Rice<sup>1[OPEN]</sup>

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MADS box transcription factors (TFs) are subdivided into type I and II based on phylogenetic analysis. The type II TFs regulate floral organ identity and flowering time, but type I TFs are relatively less characterized. Here, we report the functional characterization of two type I MADS box TFs in rice (*Oryza sativa*), *MADS78* and *MADS79*. Transcript abundance of both these genes in developing seed peaked at 48 h after fertilization and was suppressed by 96 h after fertilization, corresponding to syncytial and cellularized stages of endosperm development, respectively. Seeds overexpressing *MADS78* and *MADS79* exhibited delayed endosperm cellularization, while CRISPR-Cas9-mediated single knockout mutants showed precocious endosperm cellularization. *MADS78* and *MADS79* were indispensable for seed development, as a double knockout mutant failed to make viable seeds. Both MADS78 and 79 interacted with MADS89, another type I MADS box, which enhances nuclear localization. The expression analysis of *Fie1*, a rice FERTILIZATION-INDEPENDENT SEED-POLYCOMB REPRESSOR COMPLEX2 component, in *MADS78* and *79* mutants and vice versa established an antithetical relation, suggesting that *Fie1* could be involved in negative regulation of *MADS78* and *MADS79*. Misregulation of *MADS78* and *MADS79* perturbed auxin homeostasis and carbon metabolism, as evident by misregulation of genes involved in auxin transport and signaling as well as starch biosynthesis genes causing structural abnormalities in starch granules at maturity. Collectively, we show that *MADS78* and *MADS79* are essential regulators of early seed developmental transition and impact both seed size and quality in rice.

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Double fertilization is a characteristic feature of angiosperm reproduction. During double fertilization, two haploid sperm cells reach the embryo sac via the pollen tube. One sperm cell fuses with a haploid egg cell to form a diploid embryo and the other fuses with a diploid central cell to form a triploid endosperm cell composed of two maternal genomes and one paternal genome (Olsen, 2004). The embryo acts as a precursor to the next generation of the plant life cycle, and the endosperm is the terminal nutritive tissue. In eudicots, the embryo represents the majority of the tissue in mature seeds and the endosperm is ephemeral (Chaudhury et al., 2001). However, in monocots such as rice (Oryza sativa), wheat (Triticum aestivum), and maize (Zea mays), the endosperm persists and becomes the dominant tissue that stores starch and proteins, thus determining the mature seed size (Chaudhury et al., 2001; Gao et al., 2013). Endosperm is the primary source of calories for human (Homo sapiens) and livestock nutrition, thereby making it a valuable biological and economic entity (Sabelli and Larkins, 2009).

Endosperm development undergoes several distinct and overlapping phases categorized as follows: (1) early, the triploid nuclei undergo multiple rounds of

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P.P. and H.W. conceived and designed the study; P.P. generated single and double knockouts of *MADS78* and 79; J.J.F. generated overexpression mutants of *MADS78* and 79 and *Fie1*; B.K.D. and J.S. generated single knockout of *Fie1*; P.P. performed experiments on *MADS78* and 79 mutants; B.K.D. analyzed *Fie1* mutants and performed RT-qPCR; Z.W. performed in situ hybridization assay; interaction assays were performed by M.M. and P.P.; T.O. and I.K. performed metabolite profiling; K.L. and C.Z. analyzed RNA-seq; H.Y. performed target gene analysis; P.S. critically reviewed and analyzed the work; P.P. analyzed the results from all the experiments and wrote the article; all authors read and approved the article.

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mitotic divisions without cytokinesis (syncytium) before initiating cell wall formation (cellularization); (2) mid, differentiation of cells, grain filling that is primarily contributed by synthesis of starch and proteins, and endoreduplication; and (3) late, maturation and dehydration (Sabelli and Larkins, 2009). The rate and duration of endosperm proliferation during the syncytial phase is an important determinant of final seed size (Garcia et al., 2005). Normal endosperm development requires a timely transition from syncytium to cellularization (Brown et al., 1996; Olsen, 2004). Genomic imbalance between the parental genomes disrupts the timely initiation of cellularization (Brown et al., 1996; Haig, 2013; Wang et al., 2018). Excessive paternal dosage delays cellularization and causes endosperm overproliferation, resulting in bigger seeds (Scott et al., 1998; Stoute et al., 2012). Conversely, increased maternal dosage leads to precocious cellularization and smaller seeds. Either case may result in seed abortion as well (Scott et al., 1998; Stoute et al., 2012).

The transition of endosperm from the syncytial to cellularized state is regulated at both the genetic and epigenetic levels and is sensitive to environmental conditions (Folsom et al., 2014). Epigenetic regulation is mediated either by DNA methylation or FERTILIZATION-INDEPENDENT SEED-POLYCOMB REPRESSIVE COMPLEX2 (FIS-PRC2; Gehring, 2013; Pires, 2014). METHYLTRANSFERASE1 (MET1) predominantly maintains CG DNA methylation, and a cross between *met1* as pollen donor and the wild type activates otherwise repressed maternal-specific genes (Finnegan et al., 1996; Genger et al., 1999; Köhler et al., 2012). The FIS-PRC2 complex regulates endosperm development by installing a repressive chromatin mark, trimethylation of histone H3 at Lys-27 (H3K27me3), on several MADS box and other seed development-related genes (Huh et al., 2007; Folsom et al., 2014; Zhang et al., 2018). MADS box transcription factors (TFs), divided into type I and type II (or MIKC type), are known for their role in floral organ identity and flowering time control. The type II genes have been extensively studied as floral homeotic genes, while only a few type I genes have been functionally characterized (Bemer et al., 2010; Masiero et al., 2011; Smaczniak et al., 2012; Chen et al., 2016). In Arabidopsis (Arabidopsis thaliana), a well-studied type I MADS box TF, AGAMOUS LIKE62 (AGL62) shows specific syncytial phase expression that declines with the progression of cellularization; the agl62 mutant undergoes precocious endosperm cellularization (Kang et al., 2008). Several type I MADS box genes in Arabidopsis are implicated in the *fis-prc2* mutants, which fail to undergo endosperm cellularization (Chaudhury et al., 1997; Grossniklaus et al., 1998; Köhler et al., 2003; Walia et al., 2009; Zhang et al., 2018). Interestingly, genes encoding the components of the FIS-PRC2 complex are also imprinted and prone to misregulation due to paternal excess (Luo et al., 2000, 2009; Jullien and Berger, 2010). Thus, normal seed development requires a multifaceted interplay between parental genome dosage, the imprinting of FIS-PRC2 components, and the role of FIS-PRC2 in the imprinting of downstream factors (Pires, 2014).

Here, we report the functional characterization of two rice type I MADS box genes, *MADS78* and *MADS79*, that are active during early seed development. We report that *MADS78* and *MADS79* are essential for normal seed development. Misregulation of either gene results in seed abnormalities due to mistimed developmental transition of the endosperm and downstream impacts on starch/grain quality. At least one of the two MADS genes is required for rice seed viability, suggesting partial functional redundancy. Overall, our work shows that *MADS78* and *MADS79* regulate agronomically important yield traits, including grain size, fertility, and grain quality, in rice.

# RESULTS

## Expression Analysis of MADS78 and 79

Based on phylogenetic analysis, type I MADS box genes are divided into three classes:  $M\alpha$ ,  $M\beta$ , and  $M\gamma$ (Parenicová et al., 2003). The rice genome has 32 type I MADS box proteins composed of 13 M $\alpha$ , nine M $\beta$ , and 10 M $\gamma$  proteins (Arora et al., 2007). Both *MADS78* and MADS79 correspond to the M $\alpha$  subclade and exhibit high sequence similarity (DNA, 83% and protein, 69%; Supplemental Fig. S1). The expression profile of these two genes in public databases was limited, as they were not represented on the microarray platforms. The RNA sequencing (RNA-seq)-based expression analysis also presents a challenge due to high sequence similarity, as one member is biased for mapping of most reads (Supplemental Table S1). Therefore, we determined the spatial and temporal expression patterns of MADS78 and MADS79 using reverse transcription quantitative PCR (RT-qPCR) assays and in situ hybridizations (Fig. 1). Both genes were preferentially expressed in pollen and ovary (Fig. 1A). Furthermore, tissue-specific expression patterns determined by in situ hybridization assays indicated that both genes were expressed in pollen grains as well as in anther walls (Fig. 1C). In developing seeds, the transcript abundance of both genes peaked at 48 h after fertilization (HAF), coinciding with the syncytium stage, where endosperm undergoes rapid nuclear division without forming cell walls (Fig. 1B). The transcript levels of both genes declined by 72 HAF and were significantly reduced at 96 HAF, which coincided with the onset and completion of endosperm cellularization, respectively (Fig. 1B). This decline in transcript levels was also confirmed via in situ hybridization assays where sense/antisense probes for both genes were used during each developmental stage (Fig. 1C). Furthermore, the preferential expression pattern of both genes during endosperm development was supported by the presence of regulatory motifs in their promoters (Supplemental Fig. S2). RY-element, Skn-1, and GCN4 motifs are known to



Figure 1. Expression analysis of MADS78 and MADS79. A and B, RT-qPCR analysis of MADS78 and MADS79 in mature pollen and unfertilized ovary (A) and early seed development (48, 72, and 96 HAF; B) relative to leaf tissue from wild-type plants. For statistical analysis, Student's t test was used, and different lowercase letters indicate significant differences (P < 0.05). Error bars indicate sp (n = 6; two biological and threetechnical replicates). C, In situ hybridization of MADS78 and MADS79 in mature anthers, unfertilized ovary, and seeds at 48 and 72 HAF. aw, Anther wall; cv, central vacuole; pg, pollen grain. Bars = 0.2 mm.

establish seed- or endosperm-specific expression (Dickinson et al., 1988; Blackwell et al., 1994; Yoshihara et al., 1996).

# Both MADS78 and 79 Heterodimerize with MADS87 and MADS89

Type I MADS box proteins in Arabidopsis have a well-characterized interactome exhibiting extensive heterodimerization (de Folter et al., 2005). We next tested the nature and extent of interactions of MADS78 and MADS79 in rice. For this, we cloned the coding sequences (CDS) of 13 rice type I MADS box proteins from all three subclades: four  $M\alpha$ (MADS70, MADS77, MADS78, and MADS79), three M $\beta$  (MADS90, MADS96, and MADS98), and six M $\gamma$ (MADS81, MADS83, MADS84, MADS87, MADS88, and MADS89). The resulting prey clones were cotransformed with MADS78 or MADS79 as bait in a bimolecular fluorescence complementation (BiFC) interaction assay. For controls, separate cotransformation of MADS78 and 79 was performed with an unrelated protein, bZIP76, as well as with an empty vector (Supplemental Fig. S3). Two Mγ member proteins, MADS87 and MADS89, showed positive interactions with both MADS78 and MADS79 (Fig. 2). Positive interactions were also observed when we cotransformed MADS87 or MADS89 with MADS78 or MADS79 by swapping the respective vectors to test the veracity of these interactions (Supplemental Fig. S4). Neither MADS78 nor MADS79 formed homodimers.

Since both MADS78 and MADS79 are TFs, they are expected to localize to the nucleus. To test this possibility, we fused GFP to the N or C terminus of the CDS for both genes. The N-terminal GFP fusion constructs of both genes localized to the nucleus as well as to the cytosol (Fig. 3). The C-terminal GFP fusion constructs for MADS78 localized exclusively to the nucleus, while MADS79 localized around chloroplasts (Supplemental Fig. S5A). In silico analysis revealed that neither gene carries a nuclear localization signal (NLS), suggesting that both genes likely require a recruiting factor or an anchor for nuclear transportation (Supplemental Table S2). Since both genes interact with the same  $M\gamma$  proteins (MADS87 and MADS89; Fig. 2), we cotransformed the nonfluorescent BiFC construct (YFN43) encoding a positive interaction partner (MADS87 or MADS89) with MADS78 and MADS79 GFP fusions. When MADS78 (fused with GFP on its N terminus) was cotransformed with MADS89, the GFP-MADS78 fusion protein specifically localized to the nucleus (Fig. 3). Similarly, MADS79 fused with GFP on its N or C terminus and cotransformed with MADS89 specifically localized to the nucleus (Fig. 3; Supplemental Fig. S5A). However, cotransformation of MADS78 or MADS79 with the other positive interactor, MADS87, failed to alter their localization (Supplemental Fig. S5, B-D).

# Overexpression of *MADS78* and 79 Leads to High Spikelet Sterility and Delayed Cellularization

To functionally characterize *MADS78* and *MADS79*, we generated ubiquitin-promoter-driven overexpression (OE) plant lines (OE78-4/OE78-8 and OE79-3/OE79-4 for *MADS78* and *MADS79*, respectively; Fig. 4). We did not detect any phenotypic differences between OE and wild-type plants during the vegetative phase (Supplemental Fig. S6). However, we observed significant spikelet sterility in most of the OE lines for both genes compared with the wild type (Fig. 4, B and C).



**Figure 2.** Both MADS78 and MADS79 heterodimerize with MADS87 and MADS89. BiFC assay shows positive interaction of MADS78 (A) and MADS79 (B) with MADS87 and MADS89. For a control, we cotransformed both constructs with an unrelated gene (*bZIP76*). Bars =  $25 \mu$ m.

Seed sterility was more severe for OE78 compared with OE79 plants, where sterility was significantly higher only for one plant line (OE79-4) relative to the wild type (Fig. 4C). To determine the basis of seed sterility, we assayed the viability of mature pollen derived from OE plants. We did not find significant differences in pollen viability between OE and wild-type plant lines (Supplemental Fig. S7). Furthermore, careful inspection of the dehusked sterile seeds at maturity revealed flattened seed mass instead of aborted ovaries, suggesting postzygotic failure as the cause of the seed

sterility phenotype (Supplemental Fig. S8). This observation prompted us to investigate postzygotic seed development during early stages (48, 72, and 96 HAF) when both MADS78 and MADS79 were preferentially expressed (Fig. 1B). We observed that developing seeds from OE lines could be categorized into two types: abnormal and normal. The abnormally developing seeds from the OE lines were distinguished by their smaller size relative to the normal-looking seeds, which were indistinguishable from the developing seeds of wild-type plants (Supplemental Fig. S9A). The percentage of normally developing seeds in the OE lines was significantly lower than in the wild type (Supplemental Fig. S9B). Consistent with the spikelet sterility phenotype, the lower percentage of normal developing seeds in the OE78 lines was much more severe than in the OE79 lines (Fig. 4C; Supplemental Fig. S9B). This suggested that the severity of sterility



**Figure 3.** MADS89 enhances the specificity of nuclear localization for MADS78 and MADS79. In the absence of MADS89, N-terminal GFP fusion constructs of MADS78 and MADS79 localize to nucleus and cytosol. Cotransformation of the respective constructs with MADS89 enhances their localization specifically to the nucleus. Empty vector was used as a control. Bar =  $25 \mu$ m.



**Figure 4.** Overexpression of *MADS78* and *MADS79* leads to partial spikelet sterility and delayed endosperm cellularization. A, RT-qPCR analysis of *MADS78* and *MADS79* expression in 11-d-old seedlings and developing seeds (48 and 72 HAF) in the wild-type (WT) and OE mutants for the respective tissue. Wild-type expression was used as a baseline for the corresponding tissue and gene. B, Representative mature panicle images of the OE mutants compared with the wild type. White arrows indicate sterile seeds. Images shown were digitally extracted and scaled for comparison. Bar = 2 cm. C, Quantification of spikelet fertility (%) in the wild type and OE mutants; n = 25 to 40 plants. D, Representative cross sections of developing seeds at 96 HAF from the wild type and OE mutants. cv, Central vacuole; ed, endosperm. Bar = 0.2  $\mu$ m. For statistical analysis in A and C, Tukey's test was used: \*\*\**P* < 0.001 and \*\**P* < 0.01. Error bars indicate sp (n = 6; two biological and three technical replicates).

observed at maturity is determined during early seed development when both MADS78 and MADS79 are transcriptionally active. We also observed that fully developed mature seeds derived from the OE plant lines had significantly higher length-to-width ratios relative to wild-type seeds at maturity (Table 1). We observed decreases in 1,000-grain weight in both OE lines; however, the decrease was only significant for OE78 lines (Table 1). To further investigate the developmental defects caused by misregulation of these two genes, we examined cross sections of developing seeds from OE and wild-type plants. Seeds from wild-type plants completed endosperm cellularization by 96 HAF (Fig. 4D). However, seeds from OE MADS78 and MADS79 lines did not completely cellularize by 96 HAF. This suggests that the higher abundance of MADS78 or MADS79 and/or their ubiquitous expression causes delayed endosperm cellularization (Fig. 4D).

# Single Knockout Mutants of *MADS78* or *MADS79* Show Precocious Endosperm Cellularization

For each of the genes, we selected two homozygous single mutants (*MADS78*, *mads78*\_17 and *mads78*\_19;

MADS79, mads79 23 and mads79 24). Both mutant lines corresponding to MADS78 had deletions in the targeted regions and resulted in truncated transcripts due to premature stop codons (Supplemental Fig. S10A). MADS79 mutants, mads79\_23 and mads79\_24, carried an insertion of adenine and a 4-bp deletion, respectively (Supplemental Fig. S10A). Neither of the MADS79 mutations led to premature stop codons. RT-qPCR analysis showed a decline in transcript abundance of MADS78 and MADS79 in the mutants relative to the wild type at 48 HAF (Supplemental Fig. S10, B and C). At reproductive maturity, we did not observe severe spikelet sterility in the mutants compared with the wild type (Supplemental Fig. S10, D and E). Like the OE lines, one single knockout mutant line of both genes (mads78 19 and mads79 23) showed a significant increase in length-to-width ratio of mature seeds relative to the wild type (Table 1). We detected significant decreases in 1,000-grain weight in single knockout lines of both MADS78 and MADS79 (Table 1). Next, we performed histochemical analysis of developing seeds to check the timing of endosperm cellularization. The knockout mutants exhibited a faster rate of endosperm cellularization relative to the wild type (Fig. 5). As a result, knockout mutants underwent complete

Table 1. Physiological and morphometric measurements of OE lines and single knockout mutants relative to the	e wild type
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Spikelet fertility, 1,000-grain weight, total tiller count, and total seeds per plant were measured at the whole-plant level (n = 25-50 plants). Area, perimeter, length, width, and length-to-width ratio were measured for individual seeds (n = 700-1,000). Tukey's test was used for comparing the wild type and mutant lines: \*, P < 0.05 and \*\*, P < 0.001.

Plant Line	1,000-Grain Weight (g)	Tiller Count	Area (mm <sup>2</sup> )	Perimeter (mm)	Length (mm)	Width (mm)	Length-to-Width Ratio
Wild type	$21.60 \pm 0.8$	8.11 ± 1.0	$12.13 \pm 0.8$	$13.81 \pm 0.5$	$5.05 \pm 0.2$	$3.04 \pm 0.1$	$1.65 \pm 0.1$
OE78-4	$19.09 \pm 2.0^{*}$	$7.09 \pm 0.7$	$11.57 \pm 0.9^{**}$	$13.58 \pm 0.6^{**}$	$5.01 \pm 0.2^{*}$	$2.90 \pm 0.1^{**}$	$1.72 \pm 0.1^{**}$
OE78-8	$19.22 \pm 1.4^*$	$6.44 \pm 1.0^{*}$	$13.00 \pm 1.1^{**}$	$14.42 \pm 0.6^{**}$	$5.36 \pm 0.2^{**}$	$3.05 \pm 0.1^*$	$1.75 \pm 0.1^{**}$
OE79-3	$20.18 \pm 1.5$	$7.44 \pm 1.0$	11.34 ± 1.1**	$13.44 \pm 1.0^{**}$	$4.97 \pm 0.3^{**}$	$2.87 \pm 0.2^{**}$	$1.73 \pm 0.1^{**}$
OE79-4	$19.65 \pm 1.3$	$7.10 \pm 1.2$	$11.97 \pm 1.0$	$14.11 \pm 0.6^{**}$	$5.38 \pm 0.2^{**}$	$2.79 \pm 0.1^{**}$	$1.93 \pm 0.1^{**}$
mads78_17	19.17 ± 1.7**	$6.63 \pm 1.0$	$12.00 \pm 1.0^*$	$13.66 \pm 0.7^*$	$4.97 \pm 0.3^{*}$	$3.05 \pm 0.01$	$1.62 \pm 0.1$
mads78_19	$21.95 \pm 1.2$	$6.90 \pm 1.3$	$12.38 \pm 0.8$	$13.99 \pm 0.5^*$	$5.14 \pm 0.2^{**}$	$3.04 \pm 0.01$	$1.69 \pm 0.1^{**}$
mads79_23	$21.74 \pm 0.8$	$7.90 \pm 1.2$	$12.45 \pm 1.0^{*}$	$14.01 \pm 0.6^{*}$	$5.17 \pm 0.2^{**}$	$3.06 \pm 0.01$	$1.68 \pm 0.1^{**}$
mads79_24	$19.40 \pm 0.5^{**}$	$7.80\pm0.8$	10.92 ± 1.1**	$13.05 \pm 0.7^{**}$	$4.79 \pm 0.3^{**}$	$2.90 \pm 0.01^{**}$	$1.65 \pm 0.1$

endosperm cellularization by 84 HAF, whereas the wild type completed it by 96 HAF (Fig. 5). This suggests that seeds deficient in either of the two genes exhibit precocious endosperm cellularization.

# The Double Knockout Mutation for *MADS78* and 79 Is Lethal

Given the high percentage of sequence similarity and overlap in MADS78 and MADS79 gene activity during rice reproductive and seed development, we next examined the combined role of MADS78 and MADS79. For this, we generated CRISPR-Cas9-mediated double knockout mutants. We obtained two homozygous T0 mutants corresponding to each of the single guide RNAs (sgRNAs): sgRNA-1 (mads78-79\_5 and mads78-79\_9) and sgRNA-2 (mads78-79\_3 and mads78-79\_4; Supplemental Fig. S11A). Both mads78-79\_5 and mads78-79\_9 double mutants (sgRNA-1) had a homozygous point mutation within both genes (insertion of adenine). In addition, mads78-79\_9 possessed a substitution (guanine to thymine) in both alleles of MADS78 (Supplemental Fig. S11A). The *mads78-79\_3* (sgRNA-2) double knockout mutants had a homozygous insertion of an adenine in both genes, while mads78-79\_4 had an insertion of thymine and guanine in MADS78 and MADS79, respectively (Supplemental Fig. S11A). All double knockout mutant lines were completely sterile, as they failed to develop mature seeds (Fig. 6; Supplemental Fig. S11B). On the other hand, we obtained seed from double knockout mutants that were heterozygous for one or both genes (Supplemental Fig. S11C). These data indicated that at least one functional gene (MADS78 or MADS79) is required for seed viability in rice.

# Expression of *MADS78* and *MADS79* Is Negatively Associated with *Fie1*

Several type I MADS box genes in Arabidopsis (Walia et al., 2009; Zhang et al., 2018; Bjerkan et al., 2019) and rice (Folsom et al., 2014; Chen et al., 2018;

Wang et al., 2018) are imprinted. Therefore, we examined the DNA methylation status of MADS78 and MADS79. McrBC (an endonuclease that cleaves DNA containing methylcytosine) assay indicated that promoter regions of *MADS78* and 79 are methylated in the leaf tissue and developing seed (48 HAF; Supplemental Fig. S12, A and B). Furthermore, bisulfite sequencing confirmed methylation in the promoter region for MADS78 and the gene body for MADS79 in developing seeds (24, 48, 72, and 96 HAF; Supplemental Fig. S12C). No significant change in CG, CHG, or CHH methylation levels for either gene were detected in developing seeds (Supplemental Fig. S12C). Thus, the methylation status did not associate with changes in the gene expression profile of MADS78 and MADS79 (Fig. 1), suggesting that DNA methylation at the sites assayed is unlikely to play a direct regulatory role for these two genes.

The FIS-PRC2 complex regulates imprinting by installing histone methylation marks on the target genes (Folsom et al., 2014). In this framework, we asked if Fie1, an essential component of the FIS-PRC2 complex, is a potential regulator of MADS78 and MADS79. The transcript abundance of Fie1 was negligible at 48 HAF but accumulated to significant levels by 72 HAF (Fig. 7A). Conversely, the transcript levels of *MADS78* and MADS79 were higher at 48 HAF and declined considerably by 72 HAF in normally developing seeds (Fig. 7A). We next examined the expression of Fie1 in MADS78 and MADS79 mutant lines. Fie1 exhibited lower transcript abundance at 72 HAF in the OE lines compared with the wild type (Fig. 7B). The single knockout mutants of MADS78 and MADS79 accumulated more Fiel transcripts at 48 HAF relative to the wild type (Fig. 7B). Moreover, in OE *Fie1* mutants, the transcripts of both MADS78 and MADS79 were not detected at 48 HAF, when otherwise they were abundantly expressed in wild-type seeds (Fig. 7C). Also, expression of MADS78 and MADS79 was up-regulated at 72 HAF in knockout mutants of Fie1 (Fig. 7D). The expression analysis of *Fie1* in *MADS78* and *MADS79* mutants and expression of MADS78 and MADS79 in *Fie1* mutants establishes an inverse transcriptional association between MADS78 and MADS79 and Fie1.



**Figure 5.** Single knockout mutants of *MADS78* and *MADS79* show precocious cellularization. Representative cross sections of developing seeds are shown at 72, 84, and 96 HAF in single knockout mutants of *MADS78* and *MADS79* compared with the wild type (WT). The red arrowhead points to the central vacuole region (cv). ed, Endosperm. Bar =  $0.2 \mu m$ .

Collectively, our data suggest that *MADS78* and *MADS79* regulation is likely linked to the abundance of *Fie1* and potentially mediated by the FIS-PRC2 complex.

# Misregulation of *MADS78* and/or 79 Disturbs Auxin Homeostasis

To further explore the basis of the observed phenotypes with respect to misregulated endosperm cellularization, we performed transcriptome analysis of developing seed from mutants (48 and 72 HAF; Supplemental Table S3). We searched the transcriptome data set for genes that were (1) preferentially activated during the syncytial stage (48 HAF) and down-regulated with the onset of cellularization (72 HAF; syncytial-associated genes) and (2) suppressed during the syncytial stage but activated with the progression of endosperm cellularization, referred to here as cellularization-associated genes (Folsom et al., 2014; Chen et al., 2016). As expected, OE mutants exhibiting delayed cellularization had higher transcript abundance for syncytiumassociated genes at 48 and/or 72 HAF (Supplemental Table S3). Conversely, single knockout mutants with precocious cellularization exhibited increased transcript abundance of cellularization-associated genes at 72 HAF relative to the wild type (Supplemental Table S4).

Auxin (indole-3-acetic acid [IAA]) plays an important role in the initiation of endosperm development and its maintenance during the proliferative phase in Arabidopsis (Figueiredo et al., 2015; Batista, 2019). Auxin synthesized in endosperm is exported to integuments via the ABCB-type transporter, P-GLYCOPROTEIN10 (PGP10; Figueiredo et al., 2016). Auxin blocks the PRC2 complex and thus facilitates cell elongation and differentiation. Seeds deficient in AGL62 have reduced auxin export due to suppressed PGP10, which releases PRC2 suppression. The activated PRC2 in turn inhibits cell elongation and differentiation leading to precocious cellularization (Figueiredo et al., 2016; Batista et al., 2019a). Since MADS79 is one of the orthologs of AGL62 and knocking out MADS78 or MADS79 accelerates cellularization, we explored the link between auxin homeostasis and MADS78 and MADS79 in our transcriptome data. For this, we combined the transcriptome analysis with a search for potential target genes of MADS78 and/or MADS79 (Supplemental Table S5). The CArG box motif is a signature binding motif for MADS box TFs (both type I and II TFs; Batista et al., 2019b). The rice ABCB-type transporter, ABCB4, which carries a CArG box-like DNA-binding motif in its promoter, showed lower transcript abundance in the absence of MADS78 or MADS79 compared



**Figure 6.** Double knockout mutation is lethal. A, Representative mature panicle images of T0 homozygous double knockout mutants compared with the wild type (WT). *mads78-79\_5* and *mads78-79\_9* are derived from sgRNA-1, while *mads78-79\_3* and *mads78-79\_4* correspond to sgRNA-2 (Supplemental Fig. S11). Bar = 2 cm. B, Representative mature seed images of the T0 homozygous double knockout mutants with husk. Bar = 1 cm. Images shown in A and B were digitally extracted and scaled for comparison.

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**Figure 7.** FIS-PRC2 potentially regulates *MADS78* and *MADS79*. A, RT-qPCR analysis of *MADS78*, *MADS79*, and *Fie1* at 48 and 72 HAF in the wild type (WT). Expression levels at 48 HAF were considered as baseline for each gene. B, RT-qPCR analysis of *Fie1* in OE and single knockout mutants for *MADS78* and *MADS79* at 72 and 48 HAF, respectively. C and D, RT-qPCR analysis of *MADS78* and *MADS79* in OE and single knockout mutants for *Fie1* at 48 and 72 HAF, respectively. For B to D, wild type expression was used as a baseline. For statistical analysis, the Holm-Sidak method was used: \*\*\*, P < 0.001 and \*\*, P < 0.01. Error bars indicate sD (n = 6; two biological and three technical replicates).



with the wild type at 72 HAF (Fig. 8A). Likewise, other ABCB-type transporters, such as *ABCB11*, *ABCB14*, and ABCB22 (orthologs of Arabidopsis PGP10), also had lower transcript levels in single knockout mutants, while expression in OE lines was unaltered (Fig. 8A; Supplemental Table S5). Genes responsible for maintaining auxin levels either by oxidation (DIOXYGEN-ASE FOR AUXIN OXIDATION [DAO]; Zhao et al., 2013) or amino acid conjugation (GRETCHEN HAGEN3 [GH3]; Jain and Khurana, 2009) also showed lower transcript abundance in single knockouts at 72 HAF (Fig. 8A; Supplemental Table S5). Furthermore, auxin signaling genes (IAA10, IAA19, and IAA24) exhibited lower transcript abundance in single knockout mutants at 72 HAF (Fig. 8A; Supplemental Table S5). Previous work has shown that ABCB4/11/14/22, DAO, GH3s, and IAAs are up-regulated in developing rice seeds at 24 HAF (Jain and Khurana, 2009; Uchiumi and Okamoto, 2010; French et al., 2014; Basunia and Nonhebel, 2019). TRYPTOPHAN AMINOTRANSFER-ASE-RELATED PROTEIN2 (TAR2), the auxin biosynthesis gene that is preferentially expressed prior to cellularization (Abu-Zaitoon et al., 2012), exhibited lower transcript levels in MADS78 and MADS79 mutants, while YUCCA7 was unaltered in the MADS78 mutant and showed higher transcript abundance in the MADS79 mutant (Fig. 8A). On the other hand, auxin biosynthesis genes (YUCCA12 and TAR1) preferentially expressed in postcellularization seeds (Abu-Zaitoon et al., 2012; Basunia and Nonhebel, 2019) exhibited higher transcript abundance in single knockouts at 72 HAF (Fig. 8A; Supplemental Table S5). YUCCA12 and TAR1 promoters have CArG boxbinding motifs (Supplemental Table S5). These changes in transcript abundance and the presence of CArG box-binding motifs in promoters of auxin biosynthesis, transporters, and signaling-related genes (Supplemental Table S5) suggest a regulatory role for *MADS78* and *MADS79* in maintaining auxin levels in developing rice seeds (Fig. 8).

# Misregulation of *MADS78* and/or 79 Alters Carbon Metabolism

We observed a chalky grain appearance for mature seeds of OE lines and knockout mutants (Fig. 9A). Further investigation via scanning electron microscopy (SEM) revealed structural abnormalities in the starch granules of the mutants. In contrast to densely packed polygonal starch granules in the wild type, mutants possessed loosely packed spherical starch granules resulting in larger openings between granules (Fig. 9A). Quantification of total starch content from mature seed as well as developing seed (10 DAF) did not show a significant difference between the mutants and the wild type (Supplemental Fig. S13A). However, we observed higher total protein content for developing seed in OE and single knockout mutants relative to wild-type seeds (Supplemental Fig. S13B). Abnormal starch granule packaging and higher total protein content are likely related to altered carbon and nitrogen metabolism. To investigate the metabolic status in the developing seeds (10 DAF), we performed metabolite profiling by gas chromatography (GC) coupled to mass spectrometry. Alterations in sugar metabolism were observed as evidenced by significant elevations in Glc, Fru, and myoinositol in OE lines and knockout mutants relative to the wild type (Supplemental Table S6). Taken together, our results suggest impaired metabolism of carbohydrates in the developing seeds of the mutant lines.

Genes involved in starch biosynthesis exhibited differential transcript abundance between the genotypes (Supplemental Table S7). We validated the expression of a selected subset of these genes by RT-qPCR (Fig. 9B). In this context, the seed-specific isoform of Suc synthase (*Susy3*), the two subunits of ADP-Glc pyrophosphorylase



YUCCA12

TAR1

YUCCA7

IAAs/GH3s

(?)MADS78/79

TARI

В

Endospern

YUCCAT

IAAs/GH3s

?]*MADS*78/79

Figure 8. Misregulation of MADS78 and/or MADS79 disturbs auxin homeostasis. A, Heat map based on normalized read counts for genes involved in auxin homeostasis in the wild type (WT) and single knockout mutants (mads78 and mads79) in developing seeds (72 HAF). The color key represents a range of normalized values. B, MADS78 and MADS79, auxin, and FIS-PRC2 collectively regulate the timely initiation of endosperm cellularization. MADS78 and MADS79 regulate some of the auxin biosynthesis (YUCCAs), signaling (IAAs and GH3s), and transport (ABCB-type transporters) related genes in developing endosperm. In normally developing syncytial stage seeds, auxin is exported from developing endosperm to the seed coat and blocks FIS-PRC2, thus ensuring normal endosperm proliferation and seed coat cell elongation. Seeds deficient in MADS78 and MADS79 have reduced auxin export, thus releasing auxin-mediated FIS-PRC2 suppression. The activated FIS-PRC2 blocks cell elongation and differentiation, which causes cellularization. The accelerated cellularization triggers the activation of auxin biosynthesis genes YUCCA12 and TAR1 that may promote endoreduplication, differentiation, and starch deposition during later stages of seed development. ? indicates additional, uncharacterized factors that might participate in this model.

(AGPS2b and AGPL2), and its transporter BRITTLE1 showed higher transcript abundance in knockout mutants at 72 HAF, while the transcript levels were unaltered in OE lines (Fig. 9B; Supplemental Table S7). The granule-bound starch synthase (GBSSI), which is involved in amylose biosynthesis (Waxy), and its upstream regulators, NF-YB1, exhibited higher transcript levels in knockout mutants at 72 HAF (Fig. 9B; Supplemental Table S7). One of the amylopectin biosynthesis genes (starch synthase [SSIIa]), cytosolic pyruvate

orthophosphate dikinase (PPDK), and the known regulators for seed storage proteins, rice basic Leu zipper (RISBZ1/bZIP58) and rice prolamin box binding factor (RPBF), were also altered. These genes showed higher transcript abundance in the developing seeds (72 HAF) of OE lines and knockout mutants (Fig. 9B; Supplemental Table S7). Collectively, our results indicate that misregulation of MADS78 and MADS79 during early seed development results in compromised seed quality.

## DISCUSSION

Functional characterization of MADS78 and MADS79 revealed six key findings: (1) misregulation of the genes by either OE or knockout leads to delayed or precocious endosperm cellularization, respectively; (2) the double knockout mutant is not viable, thereby suggesting an indispensable role for MADS78 and MADS79 in rice seed development; (3) MADS78 and MADS79 physically interact with MADS89, which enhances their specificity to localize in the nucleus; (4) MADS78 and MADS79 are potentially regulated by the FIS-PRC2 complex; (5) MADS78 and MADS79, auxin, and FIS-PRC2 are important regulators of the timely transition of endosperm from syncytium to cellularization; and (6) misregulation of MADS78 and/or MADS79 alters grain quality and metabolism.

Functional characterization of MADS78 and MADS79 indicates that both genes are involved in regulating the rate of endosperm cellularization. Seeds deficient in either of the two genes exhibit slightly precocious cellularization (Fig. 5), and overabundance of either gene results in delayed cellularization (Fig. 4D). Knockout and OE lines of both genes do not arrest the initiation of cellularization but likely determine the duration required to complete cellularization. Seeds that carry mutations in both MADS78 and MADS79 are nonviable, indicating that at least one of these two genes is required for producing viable rice seeds (Fig. 6). This also suggests that MADS78 and MADS79 have partial functional redundancy. OE of MADS78 has a greater penalty on spikelet fertility compared with MADS79 (Fig. 4C). Although MADS78 and MADS79 do not dimerize with each other, their nuclear localization is enhanced when they heterodimerize with MADS89 (Fig. 3). We found gene expression of Fie1, a rice FIS-PRC2 member, to be negatively associated with the abundance of MADS78 and MADS79 transcripts in knockout and OE lines, suggesting a possible link between MADS78, MADS79, and Fiel in regulating the rate of endosperm cellularization (Fig. 7).

### MADS89 Enhances Nuclear Localization of MADS78 and MADS79

Both MADS78 and MADS79 belong to the M $\alpha$  subclade and heterodimerized with two  $M\gamma$  proteins, MADS87 and MADS89. Furthermore, RNA-seq analysis

Figure 9. Misregulation of MADS78 and MADS79 causes impairment of starch granules. A, Representative images of mature seeds from MADS78 and MADS79 mutants (left column; bar = 1 cm) and their cross sections (right column; bar = 20  $\mu$ m) using SEM. Images shown were digitally extracted and scaled for comparison. B, RT-qPCR analysis of selected genes involved in carbon metabolism at 72 HAF in OE and knockout mutants relative to the wild type (WT). The genes were selected based on RNA-seq analysis. For statistical analysis, Student's t test was used: \*\*\*, *P* < 0.001; \*\*, *P* < 0.01; and \*, P < 0.05. Error bars indicate sp (n = 6; two biological and three technical replicates).



revealed that both MADS87 and MADS89 coexpressed with MADS78 and MADS79 (Supplemental Table S2). No homodimers were observed for either MADS78 or MADS79. In Arabidopsis, MADS box partnerships are limited to subclass interactions, as interclass interactions (e.g. between type I and type II MADS) have not been reported (de Folter et al., 2005). Furthermore, interactions within a specific subclade of type I proteins  $(M\alpha, M\beta, and M\gamma)$  are rare, and members of a subclade mainly interact with a protein corresponding to another subclade. M $\alpha$  proteins dimerize preferentially with members of  $M\beta$  and  $M\gamma$ , and only a few interactions exist between M $\beta$  and M $\gamma$ . Thus, it is proposed that  $M\alpha$  proteins are central components stabilizing higher order protein complexes (Immink et al., 2009). The necessity of at least one of MADS78 and MADS79 being functional is indicated by the complete sterility of double knockout mutants. Physical interactions of both MADS78 and MADS79 with MADS89 suggest partial functional redundancy. One possible role for this interaction could be to enhance the nuclear localization of MADS78 and MADS79 in the presence of MADS89. Both MADS78 and MADS79 lack a clear NLS (Supplemental Table S2). The retargeting experiment suggests that MADS78 and MADS79 require MADS89 and not MADS87 to increase the specificity of their localization to the nucleus (Fig. 6; Supplemental Fig. S4). The in silico analysis indicates that MADS89 carries an NLS on its N terminus, whereas MADS87 lacks an NLS (Supplemental Table S2). Likewise, in Arabidopsis, AGL62 (an ortholog of MADS79) heterodimerizes with AGL80 and AGL61, which are orthologs of MADS89 and MADS78, respectively (de Folter et al., 2005). Both AGL80 and AGL61 play crucial roles in the initiation of endosperm development (de Folter et al., 2005; Portereiko et al., 2006; Steffen et al., 2008).

# Misregulation of *MADS78* and *MADS79* Affects Early Seed Development

*MADS78* and *MADS79* are preferentially expressed during the syncytial stage of endosperm development, and transcript levels decline as the endosperm cellularizes. Overexpressing either gene prevents the decline in their transcript pools at 72 HAF (Fig. 4A). Higher abundance of MADS78 or MADS79 in OE lines is concomitant with delayed cellularization in seeds that reach maturity. Seeds from wild-type plants are cellularized by 96 HAF. It is noteworthy that MADS78 and MADS79 OE seeds have a range of phenotypes. While some seeds reach maturity, others can be scored as abnormal as early as 48 HAF (before cellularization; Supplemental Fig. S9). These abnormal seeds are randomly distributed on the panicle. The development of individual seeds is partially independent of other seeds. Therefore, we hypothesize that certain developing seeds experience relatively higher syncytial activity than others and thus fail to cope with this abnormal developmental pace, eventually resulting in seed abortion. Seeds that abort during the early developmental stage contribute to the increased sterility of mature panicles. On the other hand, some seeds endure the abnormal developmental progression but get penalized with respect to delayed endosperm cellularization. Delayed cellularization in a subset of OE seeds could be due to (1) delayed transition from syncytium to cellularization of the endosperm and/or (2) reduced rate of cellularization because of overabundance of MADS78 and MADS79. Knockout mutants of MADS78 or MADS79 show precocious cellularization but do not impact seed viability (Fig. 5; Supplemental Fig. S10, D and E). These observations are consistent with the reported role of AGL62 (an ortholog of MADS79 in Arabidopsis) in endosperm cellularization (Kang et al., 2008; Bemer et al., 2010). agl62 mutant seeds show precocious cellularization, thus establishing its role in suppressing endosperm cellularization (Kang et al., 2008). Plants with mutations in both genes do not produce viable seed, indicating that at least one of these two genes should have a functional allele to produce viable seeds in rice (Fig. 6; Supplemental Fig. S11).

# FIS-PRC2 Potentially Regulates MADS78 and MADS79

Normal endosperm development is tightly controlled by genome dosage, as paternal excess leads to endosperm overproliferation, while increase in maternal dosage results in precocious endosperm cellularization (Scott et al., 1998; Stoute et al., 2012). FIS-PRC2 is one of the key components that aids in maintaining genome dosage by regulating seed-specific type I MADS box genes (Folsom et al., 2014). In this context, *Fie1*, which itself is regulated by repressive histone methylation (H3K9me2), regulates type I MADS box genes (MADS82, MADS87, and AGL36) by H3K27me3 histone methylation in rice (Folsom et al., 2014). The expression of type I MADS box genes MADS78 and MADS79 is negatively associated with Fie1 (Fig. 7). Overexpressing either MADS78 or MADS79 prolongs the syncytium stage, thereby delaying the induction of Fiel expression (Fig. 7B). On the other hand, in single knockout mutants of MADS78 and MADS79, *Fie1* transcripts accumulate at 48 HAF while remaining absent in the wild type (Fig. 7B). Furthermore, expression of *MADS78* and *MADS79* is repressed at 48 HAF in the mutant lines overexpressing *Fie1*, whereas their expression is temporally extended in *Fie1* knockout mutants (Fig. 7, C and D). Our results are consistent with previous work that reported an up-regulation of *AGL62* (an ortholog of *MADS79*) in PcG mutants of Arabidopsis (Kang et al., 2008).

# Interplay between Auxin, Type I MADS Box TFs, and FIS-PRC2 Ensures Timely Endosperm Cellularization

We detected CArG box-binding motifs in the promoter of an auxin transporter, ABCB4 (Supplemental Table S5), which exhibits lower transcript abundance in knockout mutants (Fig. 8A). Moreover, auxin biosynthesis (TAR2) and signaling (IAAs and GH3s) genes, which are preferentially expressed before cellularization (Abu-Zaitoon et al., 2012), also showed lower transcript abundance in knockout mutants, resulting in altered auxin homeostasis (Fig. 8). This could lead to decreased auxin export, which relieves FIS-PRC2 suppression in the seed coat (Figueiredo et al., 2016), as evidenced by increased transcript abundance of *Fie1* in knockout mutants (Fig. 7B). The activated FIS-PRC2 likely blocks cell elongation and differentiation, which induces precocious cellularization (Batista et al., 2019a). Early cellularization in the absence of MADS78 or MADS79 triggers activation of the CArG box-containing auxin biosynthesis genes YUCCA12 and TAR1 (Fig. 8A) that may have a role in subsequent endoreduplication, differentiation, and starch deposition during later stages of seed development (Abu-Zaitoon et al., 2012; Basunia and Nonhebel, 2019). Detection of CArG box-binding motifs in promoters of auxin biosynthesis enzyme-, transporter-, and signaling-related genes further supports the regulatory role of type I MADS box TFs in maintaining auxin levels and/or distribution (Supplemental Table S5). Collectively, our findings present evidence that type I MADS box TFs (MADS78 and MADS79 and potentially other type I genes), in conjunction with auxin homeostasis and *Fie1*, regulate the timely transition of endosperm from syncytial stage to cellularization, which is necessary for normal seed development (Fig. 8B).

## Misregulation of Early Seed Development Alters Grain Quality Traits

Timely transition of endosperm from syncytial to cellularization ensures optimal nutritional quality of the seed, which is also dependent upon successful import of photoassimilates from source tissues (Aguirre et al., 2018). Misregulation of *MADS78* and/or *MADS79*, associated with mistimed endosperm cellularization, also causes structural abnormalities in kernel starch (Fig. 9A). The starch biosynthesis genes were either

unchanged or down-regulated in OE lines and upregulated in single knockout mutants (Fig. 9B; Supplemental Table S7). This alteration implied that OE mutants exhibiting delayed cellularization postpone the transition of developing seed from the prestorage phase to the storage phase, while single knockout mutants with precocious cellularization reach the storage phase earlier. The transcript level changes for starch biosynthesis genes did not corroborate with starch quantity; however, abnormal starch granule structures were observed (Fig. 9A). This is not entirely unexpected, because the starch granule structure is determined not only by starch content but also by protein content, protein-starch interactions, amylose-lipid complexes, and short and long chains of amylopectin, which collectively define cooking and milling qualities of rice (Singh et al., 2007; Chen et al., 2012). The discrepancy between starch quantity and related gene expression is linked to the cytosolic (cy) isoform of *PPDK*, which is preferentially expressed in endosperm, especially during syncytial/endosperm cellularization, and down-regulated during subsequent stages by posttranslational mechanisms (Kang et al., 2005; Chastain et al., 2006). cyPPDK catalyzes a reversible reaction between pyruvate and phosphoenolpyruvate, thereby modulating carbon metabolism in favor of amino acid or lipid biosynthesis (Aoyagi and Bassham, 1984; Kang et al., 2005). In our study, cyPPDK showed higher transcript abundance in single knockout mutants (Fig. 9B; Supplemental Table S7). These results indicate that carbon influx is possibly directed toward either amino acid or lipid biosynthesis. Higher transcript abundance for SSIIa, RISBZ1/bZIP58, and RPBF was detected for both OE lines and knockout mutants (Fig. 9B; Supplemental Table S7). Since normal starch deposition and endosperm development are impaired in both OE lines and knockout mutants, it is plausible that SSIIa and potentially other genes are misregulated (at the transcript level) as a consequence of abnormal seed development triggered by overabundance or deficiency of MADS78 and MADS79. It is important to note that in normally developing seeds, MADS78 and MADS79 transcript abundance declines as endosperm initiates starch biosynthesis in nuclei that have cellularized.

## CONCLUSION

*MADS78* and *MADS79* play an important role in regulating rice seed size and shape. Their misregulation is associated with temporal defects in early seed development and abnormal starch deposition. Since rice yield and milling quality are significantly impacted by unfavorable environments, such as drought and temperature stress, it will be useful to explore if *MADS78* and *MADS79* could be mediating the stress sensitivity of rice yield and quality traits in future research.

### MATERIALS AND METHODS

#### Protein Sequence Alignment and Promoter Analysis

Protein sequences for *MADS78* (LOC\_Os09g02830) and *MADS79* (LOC\_Os01g74440) were downloaded from RGAP7. The SMART database (Letunic et al., 2012) was used to predict domains. The T-COFFEE program was used for aligning protein sequences of *MADS78* and *MADS79* (Di Tommaso et al., 2011). For promoter analysis, the regions 2 kb upstream of the transcription start sites of both *MADS78* and *MADS79* were downloaded from RGAP7 and analyzed using the PlantCARE database (Lescot et al., 2002).

#### **Generation of Transgenic Plants**

For generating overexpression constructs, full-length CDS corresponding to MADS78 and MADS79 were amplified. OE mutants of Fie1 (LOC\_Os08g04290) were considered from Folsom et al. (2014). The amplified products were cloned into pENTR/D-TOPO entry vector (Invitrogen). PCR products were moved from entry to destination vector (Ubi-NC1300) using LR clonase (Invitrogen). The final constructs were transformed into Agrobacterium tumefaciens strain EHA105. Calli from rice (Oryza sativa 'Kitaake') were transformed as described by Cheng et al. (1998). Hygromycin was used as a selective agent to screen the transformed calli. To generate CRISPR-Cas9 mutants, sgRNAs were designed using CRISPR-P 1.0 (http://crispr.hzau.edu.cn/CRISPR/) to specifically target either MADS78 or MADS79 and Fie1 (Lei et al., 2014). Considering the high percentage of sequence similarity between MADS78 and MADS79, sgRNAs were designed to target nonconserved genic regions. For targeting MADS78, an sgRNA that targeted a region downstream of the coiled-coil domain was selected. An sgRNA targeting a region between the MADS and coiled-coil domains was designed for MADS79 (Supplemental Fig. S10A). A list of potential off-targets corresponding to genic regions (exonic or intronic) for each sgRNA is provided in Supplemental Table S8. To create double knockout constructs, we designed two sgRNAs (sgRNA-1 and -2) targeting both MADS78 and MADS79 in their conserved MADS and coiled-coil domains (Supplemental Fig. S11A). With the aim of introducing point mutations in both genes, separate transformations for both sgRNAs were performed. The primer sequences used in the study are listed in Supplemental Table S9

CRISPR/Cas9 constructs were developed following the protocols described by Lowder et al. (2015). pYPQ141C was used to clone all sgRNAs, which were then transferred to a destination vector (pANIC6B) along with a vector containing Cas9 (pYPQ167) using LR clonase. The destination vector was transformed into *A. tumefaciens*, which was further used to infect calli. For single knockout mutants, T1 segregates carrying mutations and lacking Cas9 (confirmed by GUS screening assay) were considered for downstream analysis (Lowder et al., 2015). T0, T1, and T2 plants were screened for mutations by Sanger sequencing. T2 or later generations of mutant plant lines were used in the study. For experiments, mature rice ('Kitaake') seeds were manually dehusked and sterilized with bleach (40%, v/v) and water. The seeds were germinated in the dark on one-half-strength Murashige and Skoog medium. The germinated seedlings were transplanted in soil amid controlled greenhouse conditions: 16 h of light and 8 h of dark at  $28^{\circ}$ C  $\pm 1^{\circ}$ C and  $23^{\circ}$ C  $\pm 1^{\circ}$ C, respectively, and a relative humidity of 55% to 60% (Sandhu et al., 2019).

#### Genomic DNA and RNA Extraction, RT-qPCR

To screen for mutations in the single and double knockout mutants, genomic DNA was isolated from leaves using the Suc method (Berendzen et al., 2005). The region of interest was amplified using Kapa3G Plant PCR Kits (Kapa Biosystems) according to the manufacturer's protocol. The amplicon was then sequenced for genotyping. Total RNA was extracted from the following tissues: seedling, leaves, mature pollen, unfertilized ovary, and developing seed (24, 48, 72, and 96 HAF), using the RNeasy Plant mini kit (Qiagen). RNA was subsequently DNase treated. Early seed developmental stages corresponding to 24, 48, 72, and 96 HAF (normally developing seed tissues: category B; Supplemental Fig. S9), as well as unfertilized ovaries, were collected as described (Folsom et al., 2014). Spikelets were marked at the time of fertilization, which is evident by the opening of a floret, to track the precise developmental stage. Mature pollen grains were collected as described (Paul et al., 2016, 2017) with minor adjustments. After anthesis, the protruded anthers were collected in 500 μL of germination solution (2 mM boric acid, 2 mM calcium nitrate, 2 mM magnesium sulfate, and 1 mm potassium nitrate). The collected anthers were vortexed, filtered through muslin cloth, and centrifuged for 1 min at 1,000g.

The pelleted pollen grains were washed multiple times in 200  $\mu$ L of germination solution to remove unwanted material. The collected pollen grains were immersed in liquid nitrogen and stored at  $-80^{\circ}$ C.

One microgram of total RNA was used for cDNA synthesis using the SuperScript VILO cDNA synthesis kit (Invitrogen). The RT-qPCR (10  $\mu$ L) comprising gene-specific primers, SYBR Green Master Mix (Bio-Rad), and template was conducted using the Lightcycler 480 Real-Time PCR System (Roche). Thermal cycling conditions were used as described by Fragkostefanakis et al. (2015). As an endogenous control, ubiquitin (*UBQ5*) genes were used as reference genes (Jain et al., 2006). Data were analyzed using standard methods (Livak and Schmittgen, 2001). For RT-qPCR assays, a minimum of two independent biological replicates and three technical replicates were conducted.

## Pollen Viability Assay, Agronomic and Morphometric Measurements

For pollen viability assays, mature pollen was collected and subjected to staining with Alexander dye (Alexander, 1980) and Lugol's solution ( $I_2/KI$ ). Pollen number was determined using a Fuchs-Rosenthal cell counter (Fragkostefanakis et al., 2016). Three independent biological replicates were performed for the pollen viability assay. For determining grain morphology, seeds were harvested at maturity, and only fully developed seeds were scored to calculate percentage of spikelet fertility. The fully developed seeds were scanned using an Epson Expression 12000 XL scanner and were analyzed by SmartGrain software to obtain seed size measurements (Tanabata et al., 2012). For developing seed analysis, florets marked at the time of fertilization were tracked for 4 d after fertilization (24, 48, 72, and 96 HAF). Each marked flore was opened and aborted seeds were representative across all three replicates.

### **DNA Methylation Assay**

Genomic DNA was extracted from two biological replicates of seeds at 24, 48, 72, and 96 HAF using the Qiagen DNeasy Plant Mini Kit. Genomic DNA was then treated with sodium bisulfite (EZ DNA Methylation Kit; Zymo Research). Whole-genome bisulfite sequencing was performed using an Illumina high-throughput sequencing platform to generate 126-bp paired end reads. After filtering the low-quality regions using Trimmomatic, v 0.32 (Bolger et al., 2014), reads were mapped to the reference genome (rice ssp. *japonica* MSU v7.0), and methylated sites were called using the Bismark v0.16.3 pipeline (Krueger and Andrews, 2011). Methylation levels of sites corresponding to the genomic regions of *MADS78* and *MADS79* were plotted using R-studio. McrBC assay was performed as mentioned by Folsom et al. (2014).

## Histochemical Analysis and RNA in Situ Hybridization Assay

Normally developing seed tissue (category B; Supplemental Fig. S9) corresponding to 72, 84, and 96 HAF was harvested and fixed in 1 mL of formaldehyde (2%, v/v), acetic acid (5%, v/v), and ethanol (60% [v/v]; FAE solution) for 16 h, followed by a 70% (v/v) ethanol wash and stored overnight at 4°C. The dehydration procedure was performed with an ethanol series (85%, 95%, and 100% [v/v]) for 1 h each at room temperature. Samples were incubated in ethanol:xylene (1:1) for 1 h, followed by xylene (100%) for 2 h, and then transferred to 500 µL of xylene and Paraplast tablets at 60°C. Samples were then transferred and embedded in Paraplast. Ten-micrometer sections were made using a rotary microtome (Leica RM2125 RTS). RNA in situ hybridization was performed following the protocol described by Chen et al. (2016). Gene-specific fragments were amplified for MADS78 and MADS79. The respective amplicons were cloned into pGEM-T Easy vector (Promega). RNA probes were generated by in vitro transcription using the DIG RNA labeling kit (Roche). The RNA transcripts by T7 and SP6 polymerase were applied to slides as sense and antisense probes, respectively. Images from cross sections and in situ hybridization were observed using a bright-field microscope (Leica DM-2500) and represent six independent biological replicates.

### Localization and BiFC

For localization, the CDS regions of *MADS78* and *MADS79* were cloned into pGWB5 and pGWB6 vectors (obtained from TAIR), generating protein fusions

between GFP and their C and N termini, respectively. Empty vector was used as a control. For BiFC, full-length CDS of 11 rice type I MADS box genes (MADS70, MADS77, MADS81, MADS83, MADS84, MADS87, MADS88, MADS89, MADS90, MADS96, and MADS98) were amplified and cloned into either YFC43 or YFN43 containing the C- or N-terminal half of yellow fluorescent protein (YFP), respectively. Gene selection for the interaction study was based on their spatiotemporal coexpression with MADS78 and MADS79 (Chen et al., 2016). To rule out any false positives, we swapped the vectors for MADS78 and MADS79 with positive interactors (MADS87 and MADS89). Empty vector and bZIP76, an unrelated TF that is expressed during early seed development, were used for controls. Also, we cotransformed the MADS78 and MADS79 GFP fusions (cloned in pGWB5 and pGWB6) with a nonfluorescent BiFC construct (YFN43), encoding a positive interaction partner (MADS87 or MADS89), to test the translocation efficacy of each gene into the nucleus. For both localization and BiFC experiments, Nicotiana benthamiana plants were hand infiltrated using needleless syringes containing suspensions of A. tumefaciens (EHA105 strain) transformed with each vector corresponding to a gene of interest. Two days later, fluorescence signals were observed using a confocal laser scanning microscope (Nikon A1). The following conditions were used for imaging: GFP, excitation at 488 nm and emission at 500 to 550 nm; YFP, excitation at 514.5 nm and emission at 525 to 555 nm (pseudocolored green); chlorophyll, excitation at 640.9 nm and emission at 663 to 738 nm. The images shown for localization and BiFC experiments are representative of four independent experiments with three to four biological replicates in each experiment. For in silico NLS prediction, we used cNLS mapper (http://nls-mapper.iab.keio.ac.jp/cgi-bin/ NLS\_Mapper\_form.cgi; Kosugi et al., 2009).

#### SEM

For SEM, mature rice grains were transversely cut, placed on conductive tape attached to aluminum SEM stubs, and vacuum dried in a sample drying oven at 40°C for 1 week. Afterward, samples were sputter coated with chromium using a Denton Vacuum Desk V sputter coater and imaged with the Hitachi-S4700 Field-Emission SEM device at  $50 \times$ ,  $200 \times$ , and  $2,000 \times$  (Dhatt et al., 2019). Three independent biological replicates were used for the observations.

### Metabolite Profiling and Quantification of Total Starch and Protein Content

Polar metabolites were extracted from 25 mg of developing seeds (10 DAF), and 50 µL of extract was dried and derivatized as described (Lisec et al., 2006). Five biological replicates were considered for metabolite profiling, which was performed using a 7200 GC-QTOF system (Agilent) with an HP-5MS UI column (30 m, 0.25 mm, and 0.25  $\mu$ m; Agilent). GC and ionization parameters followed those described by Lisec et al. (2006). Mass spectrometry detection was performed in TOF mode with a 50-Hz scan rate. Peaks in the reference sample containing equal volumes of all samples were annotated with the metabolites in the Fiehn GC/MS Metabolomics RTL Library (Agilent) by MassHunter Unknown Analysis (Agilent) with manual curation. Peak areas of each representative ion were quantified by MassHunter Quantitative Analysis (Agilent). Following background subtraction, peak area was normalized to that of the internal standard (ribitol) and sample fresh weight to calculate relative metabolite levels with orbital unit. The parameters used for peak annotation and peak area values are listed in Supplemental Table S10. Starch and total protein contents were determined in the pellet of metabolite extraction as described (Gibon et al., 2009).

#### **RNA-Seq Analysis**

Isolation of total RNA from developing seed (48 and 72 HAF) of the wild type, OE lines, and single knockout mutants was performed as described above. The NuGen Universal Plus mRNA-Seq kit was used for preparing RNA-seq libraries. The libraries were sequenced on a single-read, 75-bp high-output flow cell on the NextSeq 550 device. For the analysis, each RNA-seq read was trimmed using Trimmomatic to ensure that it had a minimum length of 70 bp and an average quality score greater than 30 (Bolger et al., 2014). All trimmed short reads were mapped to the rice genome (RGAP v7.0) using TopHat, allowing up to two base mismatches per read (Trapnell et al., 2009). Reads mapped to multiple locations were not considered. Numbers of reads in genes were counted by the software tool HTSeq-count using corresponding rice gene

annotations and the union resolution mode (Supplemental Table S4; Anders et al., 2015).

### **Target Gene Analysis**

FIMO (http://meme-suite.org/tools/fimo) was used to search for MADS box-binding sites (Grant et al., 2011). Promoter sequences (-2,000 bp upstream) downloaded from the MSU7 annotation (Kawahara et al., 2013) were scanned for matches with the CArG box-binding sequence (Batista et al., 2019b) obtained from JASPAR 2018 (http://jaspar.genereg.net/; Khan et al., 2018).

### Accession Numbers

Gene accession numbers are displayed in Supplemental Table S2.

### Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Protein sequence alignment for *MADS78* and *MADS79*.

Supplemental Figure S2. In silico promoter analysis.

- Supplemental Figure S3. Empty vector used as an additional control for BiFC assays.
- Supplemental Figure S4. Confirmation of positive protein-protein interactions.
- Supplemental Figure S5. Localization of MADS78 and 79 in the absence and presence of the positive interactors, MADS89 and 87.
- Supplemental Figure S6. Representative pictures for overexpression mutants at the vegetative stage.
- Supplemental Figure S7. Pollen viability assay.
- Supplemental Figure S8. Mature sterile seeds from overexpression mutants of *MADS78* and *MADS79*.
- Supplemental Figure S9. Seed abortion analysis during early seed development.
- Supplemental Figure S10. Single knockout mutants for *MADS78* and *MADS79*.
- Supplemental Figure S11. Double knockout mutations are lethal.
- Supplemental Figure S12. DNA methylation analysis.
- **Supplemental Figure S13.** Quantification of total starch and total protein content from developing and mature seeds.
- Supplemental Table S1. RNA-seq-based normalized read counts for *MADS78* and *MADS79*.
- **Supplemental Table S2.** Normalized read counts for type I MADS box genes in developing rice seeds (48 and 72 HAF).
- Supplemental Table S3. Summary of RNA-seq libraries.
- Supplemental Table S4. Normalized read count for syncytial- and cellularization-associated genes.
- Supplemental Table S5. Potential target genes for MADS78 and MADS79.
- Supplemental Table S6. Levels of metabolites in the developing grains (10 DAF) of *MADS78* and *MADS79* mutants.
- Supplemental Table S7. Normalized read count for carbon metabolismrelated genes in mutants and the wild type at 72 HAF.
- Supplemental Table S8. List of potential off-targets (as per CRISPR-P 1.0) corresponding to genic region (exonic or intronic) for sgRNAs and their normalized read counts in single knockout mutants.
- Supplemental Table S9. List of primers used in the study.
- **Supplemental Table S10.** Parameters used for peak annotation and row peak area data for metabolite profiling.

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