

# Genes of the RAV Family Control Heading Date and Carpel Development in Rice<sup>1[OPEN]</sup>

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In plants, correct formation of reproductive organs is critical for successful seedset and perpetuation of the species. Plants have evolved different molecular mechanisms to coordinate flower and seed development at the proper time of the year. Among the plant-specific RELATED TO ABI3 AND VP1 (RAV) family of transcription factors, only TEMPRANILLO1 (TEM1) and TEM2 have been shown to affect reproductive development in Arabidopsis (*Arabidopsis thaliana*). They negatively regulate the floral transition through direct repression of *FLOWERING LOCUS T* and *GIBBERELLIN 3-OXIDASE1/2*, encoding major components of the florigen. Here we identify RAV genes from rice (*Oryza sativa*), and unravel their regulatory roles in key steps of reproductive development. Our data strongly suggest that, like TEMs, OsRAV9/OsTEM1 has a conserved function as a repressor of photoperiodic flowering upstream of the floral activators *OsMADS14* and *Hd3a*, through a mechanism reminiscent of that one underlying floral transition in temperate cereals. Furthermore, *OsRAV11* and *OsRAV12* may have acquired a new function in the differentiation of the carpel and the control of seed size, acting downstream of floral homeotic factors. Alternatively, this function may have been lost in Arabidopsis. Our data reveal conservation of RAV gene function in the regulation of flowering time in monocotyledonous and dicotyledonous plants, but also unveil roles in the development of rice gynoecium.

In plants, the correct formation of reproductive organs is critical not only for successful seedset but also for the perpetuation of the species. Accordingly, floral evocation must take place at a favorable time of the year

to guarantee pollination and maximum survival possibilities for the offspring. Plants that are affected in their flowering time often have a lower amount of seeds resulting in yield losses. Indeed, precocious flowering is frequently associated with reduced photosynthetic capacity due to a shortened vegetative phase (Endo-Higashi and Izawa, 2011). Conversely, delayed flowering can affect seed maturation due to exposure to unfavorable conditions. A negative correlation also exists between grain size and grain number (Guo et al., 2018; Li et al., 2018), two important agronomic traits which are controlled by both genetic determinants and environmental conditions.

Plants have evolved different molecular mechanisms to coordinate flower and seed development at the proper time of the year. Actually, the switch from vegetative to reproductive growth is controlled by multiple genetic determinants that integrate the responses to environmental and physiological conditions of the plant. Ultimately, the regulatory pathways underlying the floral transition converge on floral integrators that are able to activate genes in the shoot apical meristem (SAM) that control the initiation and development of the inflorescence meristem (IM), and then of floral meristems (FM) from which floral organs differentiate. Upon fertilization, the carpel transforms into a fruit in which the seeds develop.

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M.O., M.M.K., and S.P. designed the experiments, discussed the experiments, and wrote the paper; M.O. performed and analyzed most of the experiments; L.M.-H. and A.E.A.-J. performed analyses of Arabidopsis transgenic plants expressing OsRAV genes.

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In the last decade, the molecular basis of the floral transition has been unveiled in different plant species: the florigen, a long distance signaling molecule, is first produced in leaves under favorable conditions, and then transported to the apical meristem to initiate reproductive development (Andrés and Coupland, 2012). In the model species *Arabidopsis* (*Arabidopsis thaliana*), photoperiodic flowering is triggered by FLOWERING LOCUS T (FT), a small globular protein of 21 kD (Kardailsky et al., 1999; Kobayashi et al., 1999). The expression of FT is activated under inductive long days (LD) in vascular tissues of leaves by the positive regulator CONSTANS (CO; Suárez-López et al., 2001; An et al., 2004), and precocious flowering is prevented by the counteraction of the RELATED TO ABI3 AND VP1 (RAV) transcription repressors TEMPRANILLO1 (TEM1) and TEM2 (Castillejo and Pelaz, 2008). Since prevention of precocious flowering is important for reproductive fitness other repressors such as FLOWERING LOCUS C (FLC), SCHLAFMUTZE (SMZ), SCHNARCHZAPFENZ (SNZ), TARGET OF EARLY ACTIVATION TAGGED 1 (TOE1), TOE2, FLOWERING LOCUS M (FLM) or SHORT VEGETATIVE PHASE (SVP) play key roles in the control of FT expression in response to vernalization, age, photoperiod or ambient temperature (Hartmann et al., 2000; Scortecci et al., 2001, 2003; Aukerman and Sakai, 2003; Schmid et al., 2003; Jung et al., 2007; Lee et al., 2007; Li et al., 2008; Mathieu et al., 2009; Lee et al., 2013; Posé et al., 2013). Once flowering is induced, FM identity genes, such as the MADS-box genes SUPPRESSOR OF CONSTANS1 (SOC1) and APETALA1 (AP1), are induced which in turn repress TEMs expression (Kaufmann et al., 2010; Tao et al., 2012). Under noninductive short days (SD), CO is not active and there is no CO-dependent FT induction. In this light regime, the accumulation of the plant hormones gibberellins (GA) triggers floral transition by inducing SOC1 and LEAFY (LFY; Wilson et al., 1992; Blázquez et al., 1997; Moon et al., 2003; Eriksson et al., 2006; Hisamatsu and King, 2008). Interestingly, TEMs also regulate GA accumulation by repressing GA-3-OXIDASE 1 (GA3OX1) and GA3OX2 genes (Osnato et al., 2012).

In the crop species rice (*Oryza sativa*), two closely related genes have been described as FT orthologs: *Heading date 3a* (*Hd3a*), which promotes flowering under inductive SD, and *Rice Flowering locus T 1* (*RFT1*), which does it under noninductive LD (Kojima et al., 2002; Komiya et al., 2009). An evolutionarily conserved module defined by the rice orthologs of *Arabidopsis* CO and FT controls photoperiodic flowering (Shrestha et al., 2014). The rice homolog of CO, Hd1, functions as activator of *Hd3a* in SD, and contrarily as repressor in LD (Yano et al., 2000; Nemoto et al., 2016). Nevertheless, additional rice-specific regulators have been discovered: the floral activator *Early Heading date 1* (*Ehd1*, Doi et al., 2004), and its repressor *Grain number, plant height and heading date 7* (*Ghd7*; Itoh et al., 2010). The *Ehd1-Ghd7* pathway determines the critical day-length necessary for the induction of the florigen through a double gating mechanism dependent on the circadian clock and phytochrome-mediated light perception (Itoh et al.,

2010). Additional studies on phylogenetic reconstructions revealed the presence of FLC homologs in monocots genomes (Ruelens et al., 2013). These FLC-like factors repress the floral transition in response to cold in temperate cereals (Winfield et al., 2009; Greenup et al., 2010), but appeared to have acquired an opposite function in tropical cereals. Indeed, the rice FLC homolog OsMADS51 activates the expression of the floral promoter *Ehd1* (Kim et al., 2007).

Upon FT/*Hd3a* activation, the resulting gene product moves through the phloem from the leaf to the SAM (Corbesier et al., 2007; Jaeger and Wigge, 2007; Mathieu et al., 2007; Tamaki et al., 2007; Notaguchi et al., 2008; Komiya et al., 2009) where it triggers transcriptional reprogramming able to confer the competence to form flowers (Corbesier et al., 2007; Torti et al., 2012). Intriguingly, FT-like proteins do not have DNA binding activity per se and must interact with bZIP transcription factors (TFs), i.e. FLOWERING LOCUS D (FD) in *Arabidopsis* and OsFD1 in rice (Abe et al., 2005; Wigge et al., 2005; Taoka et al., 2011; Brambilla et al., 2017; Collani et al., 2019), to activate the expression of downstream genes. The interaction of FT and FD proteins require the 14-3-3 adaptor proteins for the formation of an active Floral Activation Complex (FAC; Taoka et al., 2011; Collani et al., 2019), which in *Arabidopsis* directly activates the FM identity gene *AP1* (Wigge et al., 2005), whereas the rice FAC complex induces the expression of the *AP1-like* genes *OsMADS14/15/18* and the *SEPALLATA-like* gene *OsMADS34* that are required for the specification of IM identity (Kobayashi et al., 2012; Gómez-Ariza et al., 2019). The florigens also mediate the transcriptional repression of *PINE*, a gene encoding a Zinc Finger type TF involved in the negative regulation of stem elongation. Therefore, the FAC coordinates the formation of reproductive structures (by activating IM identity genes) and internode elongation (by repressing *PINE*), guaranteeing the emergence of the panicle from the flag leaf, known as heading, which occurs when rice inflorescence development is completed (Gómez-Ariza et al., 2019).

In the rice inflorescence, primary and secondary branches form on the flanks of the IM (rachis) and terminate in spikelet meristems that develop floret meristems from which the palea, the lemma, two lodicules, six stamens and one central carpel differentiate. The identity of different floral organs is specified by the interaction of MADS domain TFs belonging to the *SEPALLATA* (OsMADS1), *APETALA3* (OsMADS16) and *AGAMOUS* (OsMADS3–OsMADS58) subfamilies (Jeon et al., 2000; Kyojuka et al., 2000; Nagasawa et al., 2003; Dreni et al., 2011). Eventually, the FM is consumed during gynoecium development, when a single ovule primordium forms inside the carpel (Dreni et al., 2007). Recently, OsMADS1 was also shown to be essential during seed development, specifically in the regulation of grain size and shape (Liu et al., 2018).

In this study, we identified four members of the RAV family of TF in rice, which share sequence similarity with *Arabidopsis* TEMs. In silico/coexpression analyses based on available transcriptomics data revealed that *OsRAVs*

interact with genes belonging to the *MADS-box* superfamily at different developmental stages, suggesting that RAVs are unknown players in the gene regulatory network underlying reproductive development in rice. Specifically, *OsRAV8* and *OsRAV9* negatively correlate with IM identity genes of the *AP1* subfamily, whereas *OsRAV11* and *OsRAV12* act downstream of MADS-domain floral homeotic factors of the *SEP* and *AG* subfamilies. Molecular and functional studies using knock-down and knock-out mutant lines indicate a conserved function for *OsRAV9*/*OsTEM1* as repressor of photoperiodic flowering upstream of the floral activators *OsMADS14* and *Hd3a*, and reveal a role for *OsRAV11* and *OsRAV12* in the correct development of female reproductive organs, downstream of *OsMADS1* and *OsMADS13*.

## RESULTS

### The Rice Genome Contains Four RAV Genes

Genes belonging to the RAV subfamily are present in all land plant species and encode for putative TFs, which are characterized by two DNA binding domains, an APETALA2-type at the N terminus and a B3-type at the C terminus (Kagaya et al., 1999). In the model species, Arabidopsis, the subfamily of RAV genes is composed of six members (Riechmann, 2002); in addition to the *TEMs*, four other genes belong to this family: *RAV1* and *RAV1-like*, which are phylogenetically close to *TEMs*, and *RAV3* and *RAV3-like*, which are the most divergent and nothing has been reported about their function. The functional characterization of the closest four indicate their regulatory role in different stages of plant development (Hu et al., 2004; Feng et al., 2005; Castillejo and Pelaz, 2008; Osnato et al., 2012; Matías-Hernández et al., 2014, 2016; Aguilar-Jaramillo et al., 2019) and in the response to abiotic stresses (Fu et al., 2014).

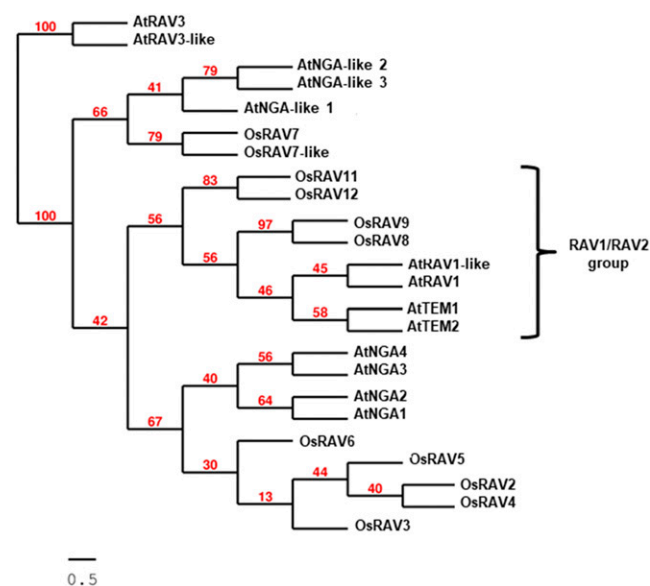
A preliminary phylogenomics analysis indicates a clear separation between AP2-B3 coding genes (*RAV*) and those encoding only the B3 domain (*ARF*, *NGA*, *VAL*), as reported in Supplemental Figure S1. Specifically in rice, although twelve genes were named *RAV* (Supplemental Table S1; Swaminathan et al., 2008), only four encode putative proteins containing both DNA binding domains as predicted by gene orthology and paralogy (*OsRAV8*, *OsRAV9*, *OsRAV11*, and *OsRAV12*; Supplemental Fig. S2A). Further analysis carried out by searching the SALAD (Surveyed conserved motif Alignment diagram and the Associating Dendrogram) database (Mihara et al., 2010) revealed the absence of RAV proteins in green and red algae, and the presence of multiple conserved motifs in addition to the AP2 and B3 domains (Supplemental Fig. S2B), including the bipartite nuclear localization signal and the B3 repression domain (Supplemental Fig. S3) together with features associated to posttranslational modifications (Supplemental Fig. S4). We also performed a phylogenetic analysis based on the deduced full-length protein sequences retrieved by a BLAST-P search against the proteomes of the two model

species (Fig. 1; Supplemental Fig. S3). Although the four *OsRAV* proteins clearly clustered together with *AtRAV1*/*AtRAV1-like* and *TEM1*/*TEM2*, *OsRAV8* and *OsRAV9* showed the highest similarity with these Arabidopsis RAV factors (Riechmann, 2002).

In the rice genome, the regions corresponding to the *OsRAV8* and *OsRAV9* loci may have originated recently, likely due to tandem duplication events after speciation. Indeed, these genes are physically linked (having a distance of about 50 Kb) at the tip of the short arm of chromosome 1 in a region that is enriched in sequences related to retro-transposons (Supplemental Fig. S5, A and B). Furthermore, when we searched the Plant Genome Duplication Database (Lee et al., 2013) using the *OsRAV8*-*OsRAV9* locus identifiers, we found intragenome syntenic relationships with a region on the long arm of chromosome 1 containing *OsRAV11* (Supplemental Fig. S5C), and a region on chromosome 5 containing *OsRAV12* (Supplemental Fig. S5D). As a result of these phylogenomics and phylogenetic analyses, *OsRAV11* and the related gene *OsRAV12* appeared as within-species paralogs of *OsRAV8*-*OsRAV9* with similar genomic structures.

### Expression Patterns of RAV Genes and Floral MADS-box Genes Are Correlated

To gain insights into the possible function of *OsRAV* genes in various biological processes and metabolic



**Figure 1.** Phylogenetic analysis of RAV proteins in model species. Tree represents the most related RAV-like proteins from Arabidopsis (At) and *Oryza sativa* (Os) retrieved using TEM1 as query in a BLAST-P search, alignment with MUSCLE, and selected with G-blocks. TEM1 and TEM2 are RAV2-like and RAV2, respectively. Phylogenetic analysis with bootstrapping procedure ( $n = 100$ ) as statistical test for branch support was carried out with PhyML, and tree visualization with TreeDyn. Scale bar indicates the number of substitutions per site. Values in red indicate percentage branch support values.

pathways, we carried out an in silico coexpression analysis by using the Rice Functionally Related gene Expression Network Database (RiceFRIEND) platform (Sato et al., 2013) and constructed a coexpressed gene list for *OsRAV9* as guide gene (Supplemental Dataset S1). Among the top 474 genes displaying a positive correlation with *OsRAV9* (Pearson's Correlation Coefficient higher than 0.3), we found enrichment for biological processes gene ontology (GO) categories related to reproductive processes, response to chemical stimuli, and response to oxidative stress (Supplemental Fig. S6, A and B). Supporting this, we also found overrepresentation for molecular function of the GO terms oxidoreductase and iron-binding activities (Supplemental Fig. S6C). Taken together, these findings suggest a possible role for RAV in reproductive development as well as abiotic stress response (similarly to Arabidopsis RAVs). In particular, peroxidases and oxidoreductases are detoxification enzymes activated upon accumulation of reactive oxygen species (Choudhury et al., 2013), and iron-binding proteins are able to sequester excess iron to avoid reaction with oxygen and the formation of damaging reactive oxygen species (Selote et al., 2015). On the other hand, GO analysis of the top 176 genes displaying a negative correlation with *OsRAV9* suggested a possible function in signal transduction pathways, due to the enrichment of the terms related to protein dephosphorylation and regulation of transcription for biological processes (Supplemental Fig. S7A). In particular, we found a significant negative Pearson's Correlation Coefficient between *OsRAV9* and genes involved in reproductive development including the florigen *Hd3a* and the IM identity genes such as *OsMADS14*, *OsMADS15*, and *OsMADS34* (Table 1). Furthermore, a recent transcriptomics analysis of apical meristems

revealed a negative correlation between the expression of the closely related gene *OsRAV8* and IM identity genes *OsMADS14*, *OsMADS15*, *OsMADS18*, and *OsMADS34* at the transition from vegetative to reproductive growth (Gómez-Ariza et al., 2019; Supplemental Fig. S7). Taken together, coexpression and available transcriptomics datasets may suggest a possible role for at least *OsRAV8* and *OsRAV9* in the negative regulation of the transition from vegetative to reproductive phase, similarly to RAV genes in Arabidopsis.

### OsRAV Genes Are Differentially Expressed during Plant Development

Because a role for *OsRAV* genes in rice plant development is yet to be elucidated, we first inferred their expression profiles by searching publicly available collections of transcriptomics data (Sato et al., 2011). *OsRAV9* and *OsRAV12* are expressed in vegetative tissues (i.e. leaves and roots), the former at early stages of plant development (Supplemental Fig. S8A), and the latter at maturity (Supplemental Fig. S8B), whereas *OsRAV11* is widely expressed in different organs with the exception of anthers (Supplemental Fig. S8C). Moreover, the expression of *OsRAV9* seems to follow a diurnal oscillation: its transcript levels are almost undetectable during the day, then increase at dusk and reach a peak in the middle of the night (Supplemental Fig. S8D). Likewise, the expression of *OsRAV11* and *OsRAV12* appeared to oscillate during the day with a peak after dusk (Supplemental Fig. S8, E and F), although with a smaller amplitude compared to *OsRAV9*.

To validate these expression data, we designed specific primers for each of the four *OsRAV* genes (Supplemental

**Table 1.** List of the top 20 genes negatively correlated with *OsRAV9*

Genes listed in bold indicate involvement in the regulation of reproductive development. ABA, Abscisic acid; PCC, Pearson's Correlation Coefficient.

| Weighted PCC | Locus Identifier | Gene name                | Description   |
|--------------|------------------|--------------------------|---|
| -0.568       | LOC_Os03g54160   | <b><i>OsMADS14</i></b>   | MADS-domain containing Transcription Factor                 |
| -0.542       | LOC_Os03g54170   | <b><i>OsMADS34</i></b>   | MADS-domain containing Transcription Factor                 |
| -0.526       | LOC_Os07g01820   | <b><i>OsMADS15</i></b>   | MADS-domain containing Transcription Factor                 |
| -0.497       | LOC_Os04g13150   |                          | Cyclin-like F-box domain containing protein                 |
| -0.491       | LOC_Os10g06560   |                          | Cyclin-dependent kinase G-1                                 |
| -0.477       | LOC_Os09g19500   |                          | Protein kinase-like domain containing protein               |
| -0.476       | LOC_Os01g11940   | <b><i>OsFT-like1</i></b> | Similar to SP3D   |
| -0.464       | LOC_Os01g19880   |                          | Conserved hypothetical protein                              |
| -0.463       | LOC_Os10g06510   |                          | Protein kinase-like domain containing protein               |
| -0.458       | LOC_Os02g55990   |                          | Longin-like domain containing protein.                      |
| -0.451       | LOC_Os07g17230   | <i>OsWRKY123</i>         | WRKY-domain containing Transcription Factor                 |
| -0.446       | LOC_Os05g43930   | <i>OMT</i>               | Similar to O-methyltransferase ZRP4 (EC 2.1.1.-)            |
| -0.434       | LOC_Os11g31770   |                          | Conserved hypothetical protein                              |
| -0.433       | LOC_Os05g38290   |                          | Similar to Protein phosphatase 2C (PP2C)                    |
| -0.4315      | LOC_Os02g52780   | <i>bZIP</i>              | Similar to ABA-responsive element binding protein 2 (AREB2) |
| -0.423       | LOC_Os03g04990   |                          | Conserved hypothetical protein                              |
| -0.424       | LOC_Os06g06320   | <b><i>Hd3a</i></b>       | HEADING DATE 3A   |
| -0.422       | LOC_Os12g13910   |                          | Conserved hypothetical protein                              |
| -0.411       | LOC_Os09g27680   |                          | Conserved hypothetical protein                              |
| -0.407       | LOC_Os08g36340   | <i>HAK4</i>              | Similar to HIGH-AFFINITY POTASSIUM TRANSPORTER 4            |

Fig. S8G). With respect to *OsRAV8*, we performed standard reverse transcription (RT)-PCR reactions, and even if a clear band was amplified using genomic DNA as control template, no amplification was observed when using cDNA obtained from different vegetative and reproductive tissues (Supplemental Fig. S8H), confirming that this gene is not transcribed at detectable levels in the samples examined. Only very recently *OsRAV8* was found to be expressed; it was absent in all previous studies likely because of its very specific expression in the apical meristem only at the time of floral transition (Gómez-Ariza et al., 2019).

The expression profiles of the other three *OsRAVs* were investigated by RT-quantitative PCR (qPCR) in wild-type plants at different developmental stages. During the juvenile phase, the mRNAs of *OsRAV9* and *OsRAV11* were detected in roots, basal region of the stem comprising the SAM, and young leaves (Fig. 2A), although the abundance was much higher for *OsRAV9* than *OsRAV11*. During the adult phase, the expression of *OsRAV9* became almost undetectable in vegetative tissues resembling Arabidopsis *TEM* genes expression, whereas *OsRAV11* and *OsRAV12* displayed high transcript levels in mature leaves (Fig. 2B), resembling the expression behavior of the positive regulator of leaf senescence *AtRAV1* in Arabidopsis (Woo et al., 2010). *OsRAVs* were also expressed in female reproductive organs (Fig. 2C), and their transcript levels decreased after pollination, suggesting that their activities might be restricted to the gynoecium before anthesis.

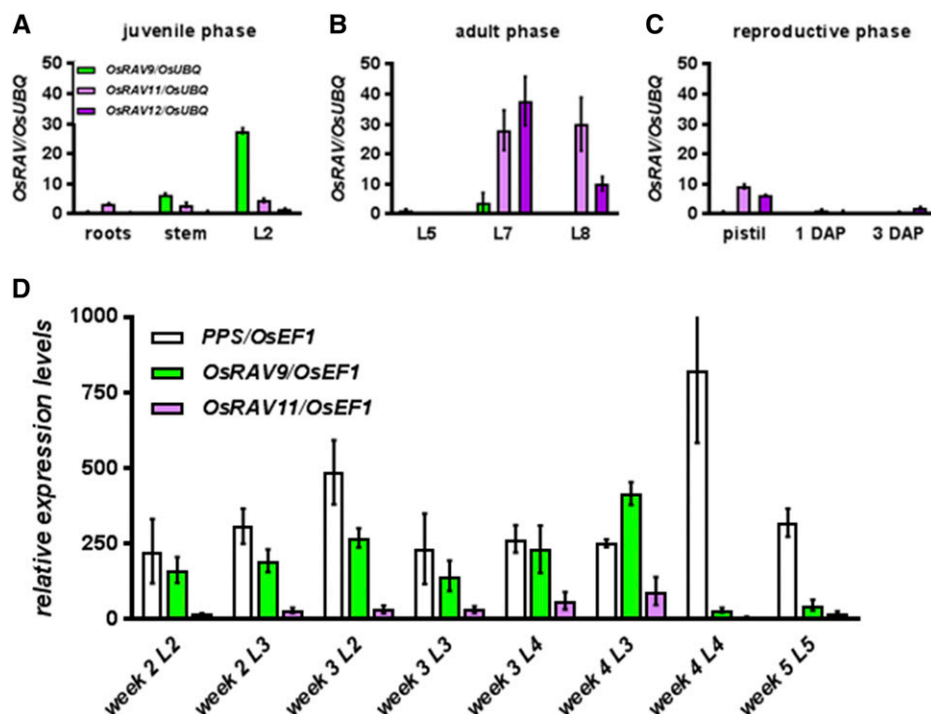
To investigate in detail a possible role of *OsRAVs* in phase changes during vegetative growth, we dissected differentiating leaves from wild-type plants to monitor their transcript levels at different developmental stages.

Precisely, in rice, the juvenile phase is limited to the second leaf (L2), because the transition to the adult phase occurs during the development of the third to fifth leaf (L3–L5) when the midrib differentiates (Itoh et al., 2005). We also analyzed the expression of the *Peter Pan Syndrome (PPS)* gene (Tanaka et al., 2011), the rice ortholog of the Arabidopsis *COP1* (Liu et al., 2008), as a marker for the transition from the juvenile to adult phase. In accordance with its function, *PPS* displayed a peak of expression in the fourth leaf four weeks after germination. *OsRAV9* exhibited the highest transcript levels in L3 and then decreased in L4 and L5 (Fig. 2D), indicating that its transcription was drastically reduced at the transition to the adult phase. Also the expression of *OsRAV11* was detected in young leaves, although at very low levels as compared with *PPS* and *OsRAV9* (Fig. 2D).

In summary, these molecular analyses suggest diversification of expression patterns for *OsRAV9* and *OsRAV11*, the former being transcribed at higher levels in the vegetative phase and specifically in juvenile leaves, and the latter at maturity, in particular in old leaves and female reproductive structures.

### *OsRAV9* Negatively Regulates Floral Transition

To investigate the functional conservation between rice and Arabidopsis *RAV* genes, we used the *Pro-35S:OsRAV9* and *Pro-35S:OsRAV11* constructs to transform Arabidopsis plants (Supplemental Fig. S9A). After selection of several independent transgenic lines (Supplemental Fig. S9B), phenotypic analyses of selected T2 generations revealed a mild late flowering



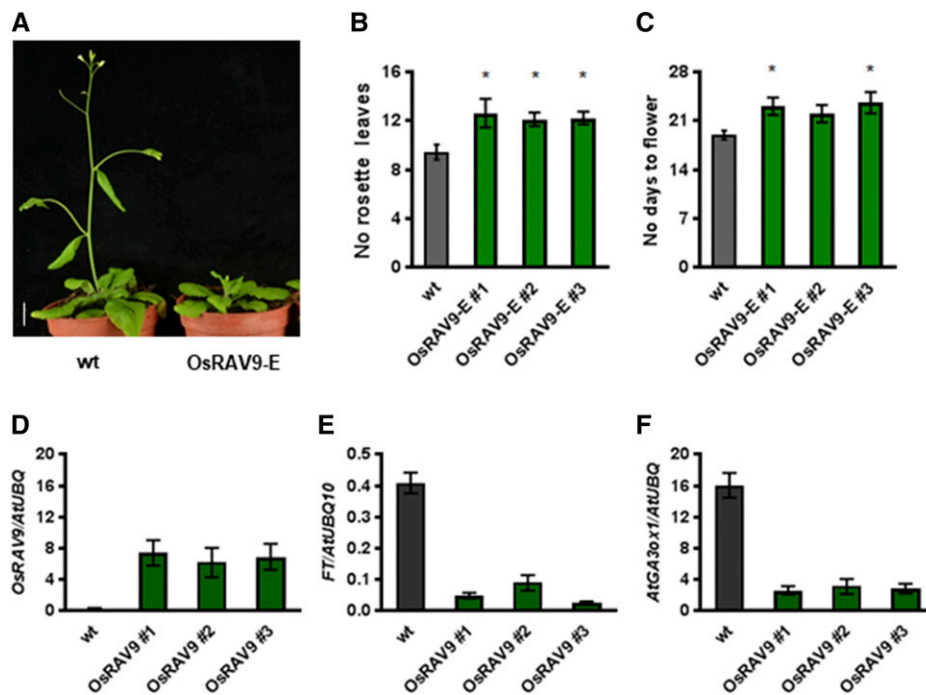
**Figure 2.** Patterns of *OsRAV* genes expression during plant development. A to C, Transcript levels of *OsRAVs*, relative to *OsUBQ*, in vegetative and reproductive tissues of wild-type plants. A, RT-qPCR in roots, stems, and L2 dissected from 1-week-old seedlings. B, RT-qPCR in mature leaves (L5, L6, L7) dissected from 10-week-old adult plants. C, RT-qPCR in mature pistils and fertilized ovaries 1 and 3 d after pollination (DAP). D, Transcript abundance of *PPS* (white), *OsRAV9* and *OsRAV11*, relative to *OsUBQ*, in juvenile (L2) and adult (L3–L4–L5) leaves. At week 2, L3 is formed and L4 is emerging. At week 3, L4 expands and L5 is emerging. At week 4, L5 is fully expanded. Expression data are mean values of three biological replicates, and error bars represent sd.

phenotype of transgenic plants expressing *OsRAV9* (*OsRAV9-E*), but no phenotypic alterations in plants expressing *OsRAV11* (Supplemental Fig. S9C). Furthermore, molecular analyses of three representative T3 lines revealed also a correlation between the late flowering phenotype of *OsRAV9-E* plants (Fig. 3, A to D) and the down-regulation of *FT* and *GA3oxidase1* (Fig. 3, E and F), two downstream targets of the TEM factors in Arabidopsis (Castillejo and Pelaz, 2008; Osnato et al., 2012). Taken together, these findings suggest that rice *OsRAV9* and the Arabidopsis *TEM* genes have an at least partial conserved function as repressor of photoperiodic flowering in Arabidopsis.

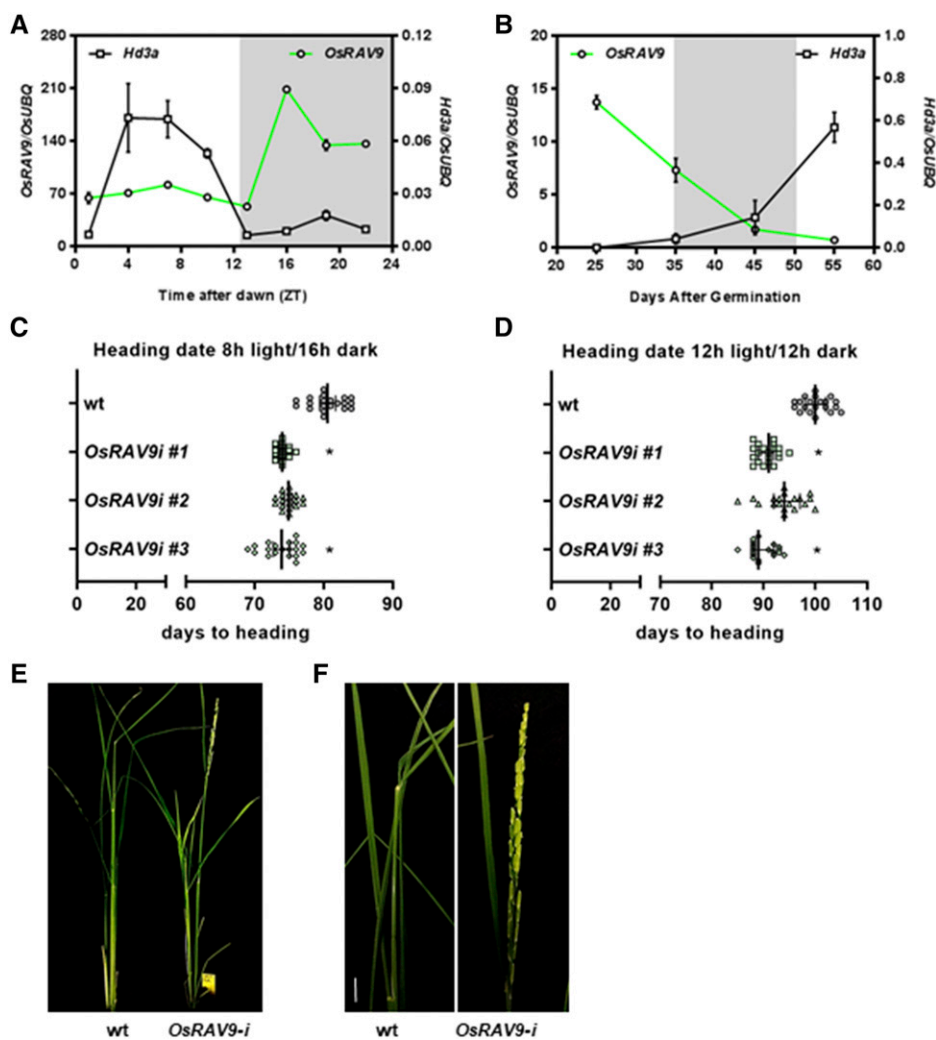
Besides the high sequence identity between *OsRAV9* and *AtTEMs*, their expression patterns were similar as both displayed high transcripts levels in the juvenile phase that decreased as the plant aged shortly before the transition from vegetative to reproductive growth (Fig. 2; Castillejo and Pelaz, 2008). Consistently, expression analyses in wild-type plants showed a mutual exclusive pattern for *OsRAV9* and the florigen *Hd3a*, not only during the day (Fig. 4A) but also throughout plant development (Fig. 4B), similarly to the opposite expression patterns of *TEM* and *FT* in Arabidopsis

(Castillejo and Pelaz, 2008). Actually, the expression of *OsRAV9* was high at early stages and dropped at around the transition to the adult phase. Conversely, the expression of *Hd3a* is almost undetectable at early stages of vegetative growth, and increases in the adult phase (Fig. 4B; Kojima et al., 2002; Komiya et al., 2009). The florigen begins to be produced in adult leaves 4 weeks after germination under inductive conditions, and triggers floral transition in adult plants when it reaches its maximum accumulation around 6 weeks.

Because the dynamics of *OsRAV9* expression strikingly resembled those of *TEM* genes, we decided to investigate if the function was also conserved in rice. We used an RNA interference (RNAi) silencing strategy due to the absence of insertion mutants in public collections for *OsRAV9*. Wild-type rice calli were transformed with an RNAi construct carrying a Gene Sequence Tag specific for the 3' end of the gene under the control of a constitutive promoter (Supplemental Fig. S10A). We obtained eleven independent transgenic lines for the silencing construct, and selected T1 *OsRAV9*-RNAi transformants by monitoring its transcript levels at the peak of expression (Supplemental Fig. S10B). T2 generation lines were



**Figure 3.** Late flowering phenotype of Arabidopsis plants expressing *OsRAV9*. A, Representative images of wild-type (wt; left) and *OsRAV9-E* transgenic (right) plants grown for 4 weeks under LD. Scale bar = 1 cm. B and C, Flowering time scored as number of rosette leaves and number of days to flower of wild-type (gray) and *OsRAV9-E* lines (green) grown under LD. D to F, Relative expression levels of *OsRAV9* and TEM1 downstream targets in wild-type and representative T3 *OsRAV9-E* lines grown for 1 week under LD. D, Ectopic expression of *OsRAV9* in transgenic Arabidopsis lines. E and F, Down-regulation of *FT* and *AtGA3ox1* in *OsRAV9-E* lines compared with wild-type. Flowering time data are the average of 25 plants each genotype, with SEM. Three biological replicates gave similar results, and one was chosen as representative. Data were analyzed by one-way ANOVA followed by Dunnett's multiple comparisons test between wild-type and transgenic lines. Asterisks indicate statistical significance ( $*P < 0.05$ ). Expression data are reported as mean values of three biological replicates; error bar represents the SEM. *AtUBQ10* was used for normalization.



**Figure 4.** Role of *OsRAV9* in heading date. A, Diurnal oscillation of *Hd3a* and *OsRAV9* expression in leaves of 4-week-old wild-type (wt) plants. Gray block indicates night. B, Mutually exclusive expression patterns of *Hd3a* and *OsRAV9* in leaves throughout wild-type (wt) plant development. Gray block indicates floral transition. C and D, Scatter plots representing heading date as number of days to flower of wild-type (in gray) and *OsRAV9-i* plants (in light green) grown under inductive photoperiods (8-h light/16-h dark, 12-h light/12-h dark). Lines represent the median with 95% of confidence interval. E, Early flowering phenotype of a selected *OsRAV9* silencing line (*RAV9-i 3*, right) compared with wild-type (left) 100 d after germination (DAG). F, Close-up view showing wild-type panicle at the booting stage and *OsRAV9-i* panicle at anthesis. For molecular analyses, plants were grown under 12-h light/12-h dark at 28°C, whereas for heading date plants were grown under different daylengths. Bars = 10 cm. Expression data are reported as mean values of three biological replicates; error bar represents the SEM. Flowering time data are the average of 18 to 20 plants each genotype. Three biological replicates gave similar results, and one was chosen as representative. Data were analyzed by one-way ANOVA followed by Dunnett's multiple comparisons test between wild-type and transgenic lines. Asterisks indicate statistical significance (\* $P < 0.05$  and \*\* $P < 0.033$ ).

obtained by self-pollination, and three lines with the highest silencing of *OsRAV9* were characterized. The RNAi lines (reported as *OsRAV9-i*) flowered slightly earlier than the wild-type under inductive photoperiods, more clearly under SD than under 12 h of light (Fig. 4, C and D; Supplemental Fig. S10C), but not under noninductive LD (Supplemental Fig. S10C). Under SD, the down-regulation of *OsRAV9* resulted in significantly early flowering plants (Fig. 4C). Under 12-h light/12-h dark regime, transgenic lines flowered on average 90 DAG and underwent anthesis 1 week later, whereas wild-type

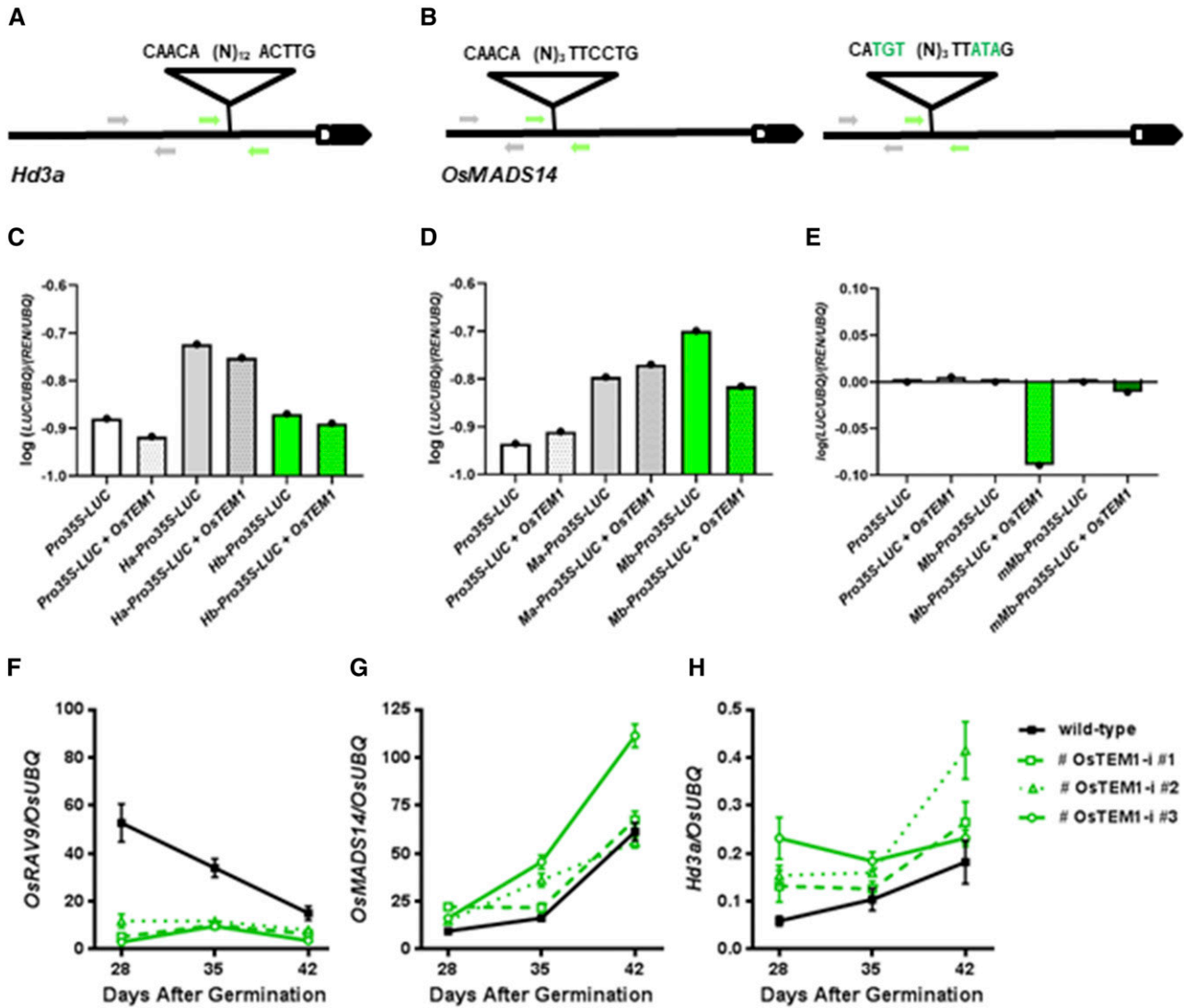
plants were still at the booting stage (Fig. 4, E and F). These findings suggested that *OsRAV9* functions as floral repressor in rice like TEMs do in Arabidopsis, likely upstream of the florigen *Hd3a* under inductive conditions. Therefore, hereafter we refer to *OsRAV9* as *OsTEM1*.

#### **OsTEM1 Regulates Floral Activators *OsMADS14* and *Hd3a***

Based on the results obtained, it was tempting to speculate a role for *OsTEM1* in the direct repression of

*Hd3a* as a noncanonical RAV binding site is present in its regulatory region (Fig. 5A). However, we could not exclude an indirect effect on *Hd3a* via additional transcriptional regulators. Actually, a negative correlation also exists between *OsTEM1* and additional floral activators including *OsFT-like1* and IM identity genes (Table 1), previously shown to act in a regulatory loop with

the florigen, upstream of *Hd3a* in the leaf and downstream of the *Hd3a*/14-3-3/*OsFD1* complex in reproductive meristems (Kobayashi et al., 2012). *OsMADS14* is also expressed in adult leaves, and a perfect RAV binding site was found in its promoter (Fig. 5B). Therefore, we used a transient expression system based on a dual Renilla-Luciferase assay to investigate the



**Figure 5.** Interaction between *OsRAV9/OsTEM1* and floral activators. A, Noncanonical RAV binding site in the promoter of *Hd3a*, 785 bp upstream of the transcription starting site. B, Perfect RAV binding site in the promoter of *OsMADS14*, 2250 bp upstream of the transcription starting site, and mutated version (right). Arrows represent oligonucleotides used to amplify fragments of *ProHd3a* (H) and *ProOsMADS14* (M) without RAV binding sites (Ha and Ma, in gray) and with RAV binding sites (Hb and Mb, in green). C to E, Transactivation activity of *OsTEM1* in transiently transformed protoplasts, reported as ratio of transcript levels of *LUC* and *REN* reporter genes relative to *UBQ*. C, RT-qPCR of protoplasts cotransformed with *Pro-35S:OsTEM1* and reporter vectors containing sequences of *ProHd3a* (Ha in gray, Hb in green). D, RT-qPCR of protoplasts cotransformed with *Pro-35S:OsTEM1* and reporter vectors containing sequences of *ProOsMADS14* (Ma in gray, Mb in green). E, Analysis of protoplasts cotransformed with *Pro-35S-OsTEM1* and reporter vectors containing sequences of *ProOsMADS14* (Mb in green, mutated Mb in dark green). Values are the mean of three independent replicates. F to H, Expression analysis of genes involved in heading date in independent T2 lines (green) compared with wild-type plants grown for 28, 35, and 42 d under inductive conditions. F, Down-regulation of *OsRAV9/OsUBQ* in silencing lines (green). G and H, Up-regulation of the floral activators *OsMADS14* and *Hd3a* in silencing lines (green lines) compared with wild-type (black line). Expression data are mean value of three biological replicates with three technical replicates each, and error bars represent SD.



direct interaction between *OsTEM1* and potential target genes. The effector vector *Pro-35S:OsTEM1* was transiently coexpressed with reporter vectors containing different regulatory regions of *Hd3a* and *OsMADS14* (Fig. 5, C and D; Supplemental Fig. S11). We evaluated transactivation ability of the floral repressor on target promoters by measuring the relative expression of the *LUCIFERASE* reporter genes from firefly (*LUC*) and from renilla (*REN*) in cell lysates. Despite the biological variability between independent replicates, a clear reduction of *LUC/REN* relative transcript levels was always observed when *OsTEM1* was cotransformed with a reporter vector carrying promoter sequences of *OsMADS14* containing the RAV binding site but not *Hd3a* (Fig. 5, C and D; Supplemental Fig. S11, B and C). However, this reduction was abolished when we cotransfected the effector vector with a mutated version of the reporter vector carrying the RAV binding site of the *OsMADS14* promoter. For statistical analyses purposes, the values of these three replicates including intact and mutated RAV binding sites are shown as logarithmic values (Fig. 5E; Supplemental Fig. S11D). Therefore, the transient cotransformation assays soundly suggest that transcription repression mediated by *OsTEM1* is stronger on DNA regulatory sequences of *OsMADS14* and is likely mediated by the RAV binding site.

Finally, we monitored the expression levels of *OsTEM1* and floral activators shortly before and around the floral transition in wild-type and *OsTEM1-i* lines grown under inductive conditions. As expected, transcripts levels of *OsTEM1* were confirmed to be higher at week 4 in wild-type plants, and strong down-regulation was observed in the transgenic lines (Fig. 5F). Conversely, the expression of *OsMADS14* increased from week 4 to 6 in wild-type plants, and a clear up-regulation was detected in *OsTEM1-i* lines 2 and 3 at 35 DAG (Fig. 5G). The related IM genes *OsMADS15* and *OsMADS34* were expressed at extremely low levels in leaves at floral transition (Supplemental Fig. S12A); however, *OsMADS18* was transcribed at higher levels in vegetative tissues, and alteration of its mRNA abundance was found in the transgenic line with highest *OsRAV9* silencing (Supplemental Fig. S12B). Furthermore, considerable increase in *Hd3a* mRNA levels was also detected in silencing lines around floral transition (Fig. 5H). Taken together, transient cotransformation of protoplasts and comparative expression analysis of wild-type and transgenic RNAi lines suggest that *OsTEM1* controls heading date via direct repression of *OsMADS14*, although it also modulates *Hd3a* expression.

We also tested the effect of the down-regulation of *OsTEM1* on other genetic pathways involved in the control of heading date (Supplemental Fig. S12C), and, although variable, we found up-regulation of the flowering inductors *OsMADS50*, *OsMADS51*, and *Ehd1* in silencing lines (Supplemental Fig. S12, D and E), perhaps suggesting additional roles in parallel pathways. As a consequence, down-regulation of the *OsTEM1*

repressor and up-regulation of different floral activators in transgenic lines resulted in early flowering phenotype.

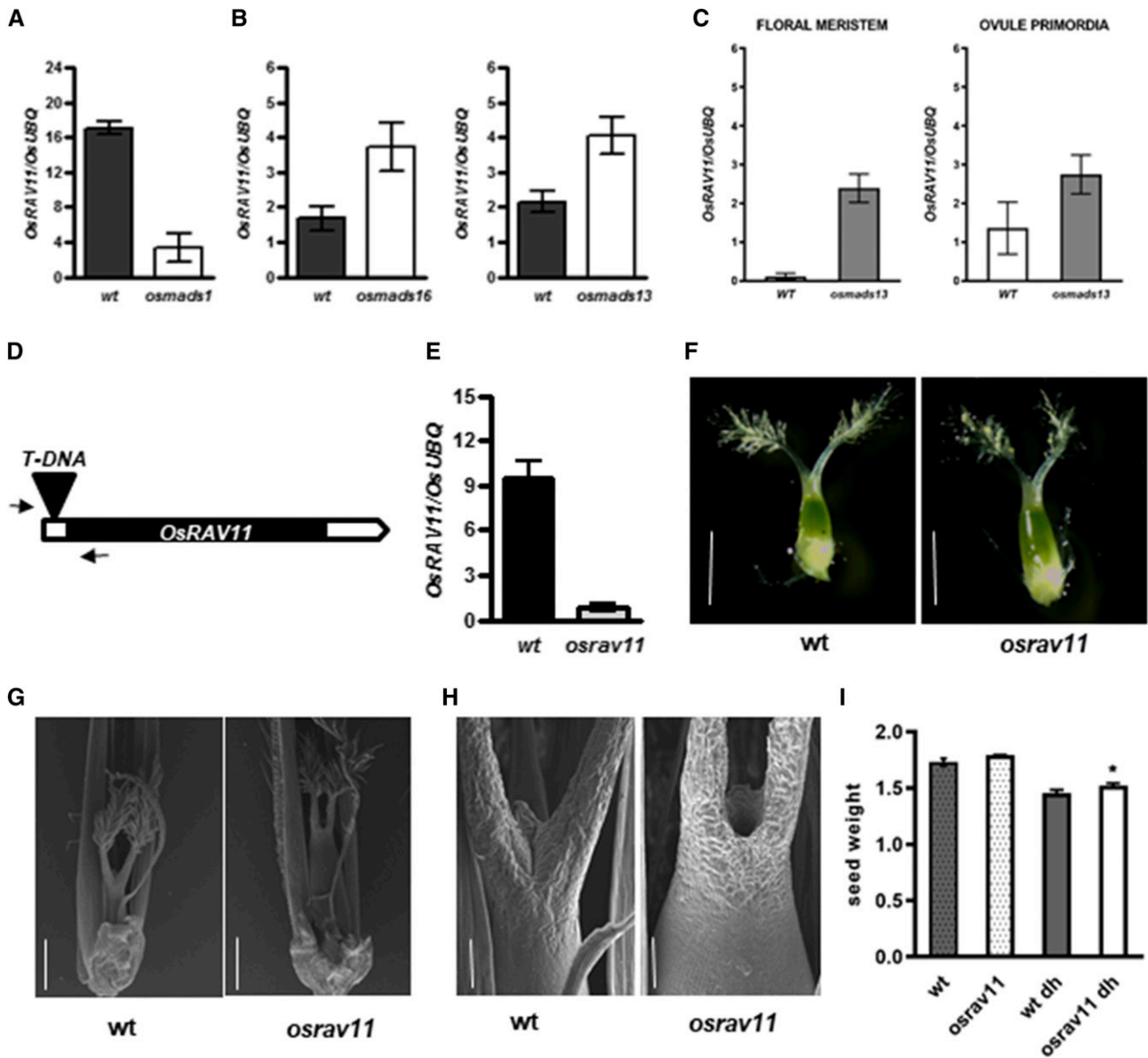
### *OsRAV11* and *OsRAV12* Regulate Carpel Development

The fact that *OsRAV11/OsRAV12* have diversified from *OsTEM1* in their expression patterns and coding sequences prompted us to hypothesize that these genes might have acquired different roles in plant development, likely after floral induction based on their expression pattern in the reproductive phase (Fig. 2C). A preliminary spatio-temporal expression analysis indicated that *OsRAV11* was expressed at early stages of flower development before organ primordia differentiation because its mRNA was detected in the spikelet meristem (Supplemental Fig. S13A). Later on, transcripts became first restricted to the carpel primordia similar to carpel identity genes, and afterward specifically limited to the apical part of the developing gynoecium (Supplemental Fig. S13B).

The analysis of available transcriptomics data sets performed on floral homeotic mutants that revealed *OsRAV11* down-regulation in *OsMADS1-RNAi* inflorescences (Supplemental Table S2; Khanday et al., 2013), and conversely up-regulation in *osmads13* mutant (M. Osnato and M.M. Kater, unpublished data), suggested an interaction between *RAV* and *MADS-box* genes controlling floral organ development. This could be a direct effect since we found CARG-boxes, consensus sequences recognized and bound by *MADS*-domain TFs, in *OsRAV11* and also in *OsRAV12* regulatory sequences (Supplemental Fig. S13D). The regulation by *OsMADS1* and *OsMADS13* is further supported by the fact that both appear as regulators of *OsRAV* genes in the Environmental Gene Regulatory Influence Networks (Wilkins et al., 2016). Therefore, *OsRAV11* might have a role in the development of the pistil, likely downstream of class-D and class-E *MADS*-domain floral homeotic factors.

Consistently, we found down-regulation of *OsRAV11* in developing panicles of the *osmads1* mutant (Fig. 6A), characterized by the conversion of floral organs into glume-like structures (Agrawal et al., 2005; Khanday et al., 2013), and up-regulation in young panicles of the floral homeotic mutants *osmads16* and *osmads13* (Fig. 6B), in which stamens and ovules are homeotically converted into extra carpels respectively (Nagasawa et al., 2003; Dreni et al., 2007) perhaps because the ectopic activation of *OsRAV11*. Accordingly, we observed strong up-regulation of *OsRAV11* specifically in *osmads13* ovule primordia (Fig. 6C; M. Osnato and M.M. Kater, unpublished data), which later develop as carpels.

Therefore, we explored the function of *OsRAV11* using a knock-out mutant characterized by the insertion of T-DNA in the 5' untranslated region of the gene (Fig. 6, D and E). The loss of *OsRAV11* function did not cause alteration of heading date, but elongated carpels (Fig. 6F). A detailed phenotypic analysis by SEM upon fertilization indicated alteration in size and shape of the



**Figure 6.** Molecular and functional characterization of *OsRAV11*. A to C, Misregulation of *OsRAV11* in floral homeotic mutants. A, Down-regulation of *OsRAV11* in *osmads1* developing inflorescences. B, Up-regulation of *OsRAV11* in *osmads16* and *osmads13* developing inflorescences. C, Strong activation of *OsRAV11* in specific cell types (floral meristem and ovule primordia) isolated from *osmads13* mutant flowers. D, Schematic representation of the T-DNA insertion in the 5' untranslated region of *OsRAV11*. Arrows indicate primers used for genotyping. E, Down-regulation of *OsRAV11* in pistils dissected from *osrav11* mutant flowers at maturity. F to H, Morphological analyses of female reproductive structures at maturity. F, Representative images of wild-type and *osrav11* carpels dissected from mature flowers at anthesis obtained by optical microscopy. G and H, Representative images of reproductive structures obtained by scanning electron microscopy (SEM). G, Wild-type and *osrav11* carpels upon fertilization. Glumes were partly removed to show female reproductive organs. H, Apical tissues of wild-type and *osrav11* gynoecia after pollination. Bars = 1 mm (F), 500  $\mu$ m (G), and 100  $\mu$ m (H). I, Bar plots representing the weight of seed produced by wild-type (in gray) and *osrav11* (in white) plants grown in the greenhouse under inductive conditions. Expression data are reported as mean value of three biological replicates; error bar represents the SEM. Phenotypic data are the average of three biological replicates of the weight of 100 seeds each genotype, with SEM. Statistical significance was examined by Student's two-tailed unpaired *t* test (\**P* < 0.05).

pistil, with an enlarged ovary compared to wild-type pistils (Fig. 6G). Specifically, differentiation of apical tissues of the gynoecium were altered in the *osrav11* mutant as carpel tissues did not fuse (Fig. 6H). Likewise, we observed alteration of seed morphology (Supplemental Fig. S13): mutant plants produced seeds with increased seed length and decreased seed width (Supplemental Fig. S3G), resulting in statistically significant alterations of length-to-width ratio and circularity (Supplemental Fig. S3H). Interestingly, we observed a slight increase in seed weight after dehusking (Fig. 6I).

Ultimately, to investigate the possible redundancy between *OsRAV11* and the closely related *OsRAV12* gene in the formation of female reproductive organs, we generated transgenic rice plants in which the expression levels of both genes were reduced by RNA interference (Supplemental Fig. S14). Vegetative growth of knock-down lines (from now on *OsRAV11/12-i*) was normal, whereas flowers showed alterations caused by abnormal morphology of female reproductive organs (Fig. 7A). Indeed, elongated cylindrical pistils with highly reduced stigmas and enlarged ovaries (Fig. 7B) were observed in transgenic plants with reduced levels of both *OsRAV11* and *OsRAV12* in the gynoecium

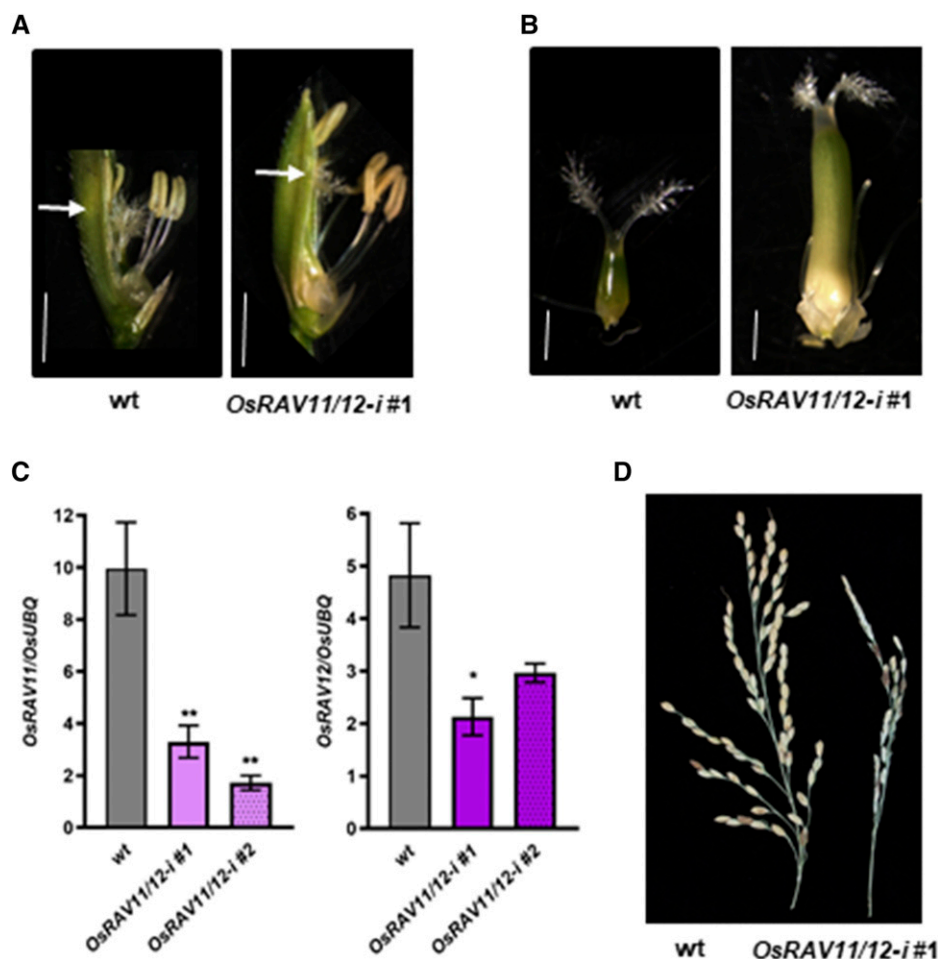
(Fig. 7C), indicating that these two genes redundantly regulate the basal-apical patterning of the gynoecium. Interestingly, flowers of these transgenic lines produced viable pollen, but most of the ovules were not fertilized, resulting in very poor seedset. Consequently, most of the transgenic lines displayed severe fertility defects (Fig. 7D).

To conclude, molecular and functional characterization of *OsRAV11* and *OsRAV12* suggest that these two genes might redundantly regulate the differentiation of the female reproductive structures. Intriguingly, a decreased activity of *OsRAV11* correlated with an increase in seed weight, whereas the loss of *OsRAV11* and *OsRAV12* activity associates with sterility problems.

## DISCUSSION

### Arabidopsis and Rice RAV Genes Are Closely Related

RAV proteins belong to the plant-specific B3 superfamily of TFs (Swaminathan et al., 2008), and are characterized by an additional AP2 DNA binding domain at the N terminus. AP2-B3 type proteins were not found in the green algae *Chlamydomonas reinhardtii*, but



**Figure 7.** Morphogenetic effects of *OsRAV11/OsRAV12* silencing in reproductive phase. A, Representative images of mature flowers dissected from wild-type (wt) and *OsRAV11/12-i* inflorescences. Arrows indicate the position of the stigmas. B, Representative images of mature carpels dissected from wild-type and one T3 *OsRAV11/12-i* flowers obtained by optical microscopy. Bars = 1 mm. C, Down-regulation of *OsRAV11* and *OsRAV12* in mature pistils of two representative T3 *OsRAV11/12-i* lines as compared with wild-type. Expression data are reported as mean values of three biological replicates with SEM. Data were analyzed by ordinary one-way ANOVA followed by Dunnett’s multiple comparisons test between wild-type and transgenic lines. Asterisks indicate statistical significance (\* $P < 0.05$  and \*\* $P < 0.033$ ). D, Fertility defects of mature *OsRAV11/12-i* panicles compared with wild-type.

they appeared early in the evolution of land plant species. The presence of two DNA binding domains suggests that these TFs achieve high affinity by specifically binding bipartite sequences in regulatory regions of downstream targets (Kagaya et al., 1999; Castillejo and Pelaz, 2008; Osnato et al., 2012; Matías-Hernández et al., 2016).

In this study, we focused on the four rice *RAV* genes, which display striking similarities with four *RAV* genes of Arabidopsis (Supplemental Fig. S15) that were already described as regulators of different aspects of plant development and stress responses. Specifically, *AtRAV1* and *AtRAV1*-like redundantly control leaf senescence (Woo et al., 2010), *AtRAV1* and *TEM2* modulate sensitivity to drought and salinity (Fu et al., 2014), *TEM1* and *TEM2* repress trichome formation (Matías-Hernández et al., 2016), and floral transition under inductive and noninductive photoperiods (Castillejo and Pelaz, 2008; Osnato et al., 2012) as well as in response to low temperatures (Marín-González et al., 2015) and to plant age (Aguilar-Jaramillo et al., 2019). Interestingly, preliminary coexpression analyses suggested that *OsRAV9* and *OsRAV11* could act in signal transduction pathways activated in response to abiotic stresses. Nevertheless, *OsRAV9* and the closely related *OsRAV8* could also act in Gene Regulatory Networks controlling the transition from vegetative to reproductive growth in the leaf and in the apical meristem, respectively.

### OsRAV9/OsTEM1 Is a New Player in Flowering

Although phylogenomics and phylogenetic analyses revealed that *OsRAV* genes might have originated from duplication events from a common ancestor after speciation and separation of monocots and dicots, it is difficult to draw conclusions from orthology with *AtRAVs*. Furthermore, the expression domains of the four *OsRAV* paralogous genes appear to have diversified likely due to polymorphisms in their regulatory regions. Regardless of the remarkable similarities in the gene structures and coding sequences, *OsRAV8* and *OsRAV9* are expressed in different tissues. Indeed, *OsRAV9* mRNA is abundant in juvenile leaves, and its reduction marks the transition to the adult phase when plants acquire the competence to flower, whereas the transcripts of *OsRAV8* are detected in the apical meristem and its levels also decrease at floral transition when IM identity genes are activated (Gómez-Ariza et al., 2019). Therefore, at least *OsRAV9* displayed an expression pattern that resembles that of *TEM* genes in Arabidopsis. Supporting the functional conservation, the ectopic expression of *OsRAV9* in Arabidopsis plants correlated with the repression of the *TEM* targets *FT* and *AtGA30X1* and delayed flowering time. In addition, silencing of *OsRAV9/OsTEM1* in transgenic rice lines resulted in early flowering due to up-regulation of the floral activators *OsMADS14* and *Hd3a*.

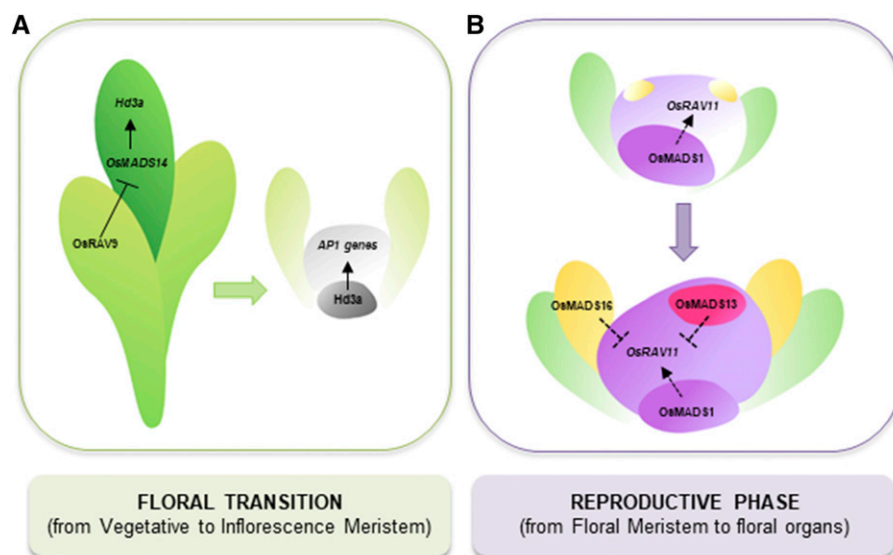
The mechanism of action seems to be different in the two species. To avoid precocious flowering, Arabidopsis *TEMs* directly target the florigens *FT* (Castillejo and Pelaz, 2008) and *AtGA3ox1/2* (Osnato et al., 2012), whereas *OsTEM1* might regulate *Hd3a* indirectly via repression of *OsMADS14* as proposed in Figure 8A. Although this *AP1/FUL-like* gene is expressed at high levels in reproductive tissues, its mRNA accumulates also during the vegetative phase. Accordingly, an additional role for *OsMADS14* as activator of the florigen in the leaf has been previously proposed, likely acting via a positive regulatory loop with *Hd3a* (Kobayashi et al., 2012). Actually, knock-down lines silencing *OsMADS14*, *OsMADS15*, *OsMADS18*, and *OsMADS34* display delayed flowering (Kobayashi et al., 2012), and conversely transgenic rice ectopically expressing *OsMADS14* and *OsMADS18* are early flowering (Jeon et al., 2000; Fornara et al., 2004). A very recent study also indicates that *AP1-like* genes could regulate drought-escape; indeed, the early flowering phenotype under drought conditions correlates with increased expression of *OsMADS18* (Groen et al., 2020).

Upon floral transition, the FAC activates the expression of *OsMADS14*, together with *OsMADS15*, *OsMADS18*, and *OsMADS34*, which specify IM identity and trigger the development of reproductive structures (Kobayashi et al., 2010; Kobayashi et al., 2012). Based on transcriptomics analysis, we speculate that *OsRAV8* could play a role in the maintenance of the vegetative state of the apical meristem, thus preventing the formation of the rachis under noninductive conditions.

### Mechanisms Controlling Floral Transition in Cereals

It has long been known that *AP1/FUL-like* proteins control seasonal flowering in cereals growing in temperate regions (Fjellheim et al., 2014). Precisely, the floral transition is triggered by the activation of *VERNALIZATION 1-like (VRN1-like)* and *FT-like* genes in wheat leaves in response to increasing day-length and prolonged exposure to low temperatures (Danyluk et al., 2003; Shimada et al., 2009; Winfield et al., 2009). At least in wheat, *VRN1* regulates flowering by directly binding the promoter of the downstream target *FT-like1* (Deng et al., 2015; Tanaka et al., 2018). Surprisingly, the function of *AP1/FUL-like* proteins as floral activators acting in leaves seems to be well conserved in rice (Jeon et al., 2000; Kobayashi et al., 2012), despite the fact that it is a cereal crop of tropical origin that does not require vernalization. In this study, we propose a new mechanism that is largely independent from previously described molecular networks determining heading date in rice (Tsuji et al., 2013). Interestingly, the presence of *TEM* orthologs in the Poacea family (Supplemental Fig. S1) opens up the intriguing possibility that the *RAV-AP1-FT* regulatory module could be conserved among tropical (*Oryza* spp. tribe) and temperate (Triticeae tribe) cereals.

Moreover, as mentioned above, *AtRAV1* and *TEM2* seem to modulate sensitivity to drought and salinity



**Figure 8.** Model for interaction between RAV and MADS factors during reproductive growth. A, *OsRAV9* represses the transcription of the florigen *Hd3a*, in the leaf. Upon floral transition, the complex *Hd3a*-*OsFD* activates the expression of IM identity genes *OsMADS14-15-18-34* in the apical meristem. B, The expression of *OsMADS1* in developing flowers marks the formation of lemma/palea (in green) and central carpel (in pink) from the FM, whereas the presence of *OsMADS16* and *OsMADS13* prevents the expression of genes involved in carpel development in stamen primordia (in yellow) and in ovule primordium (in violet).

(Fu et al., 2014). *OsTEM1* has been recently shown to play a role in response to abiotic stresses (*OsRAV2*; Duan et al., 2016), and *OsTEM1* expression was reduced in plants growing under drought conditions (Plessis et al., 2015) where the transcription of *OsMADS18* was increased (Groen et al., 2020). Therefore, RAV genes could be involved in adaptive growth by modulating heading date in response to environmental limitations and fluctuations, by integrating external and internal physiological conditions.

### Mechanisms Controlling Floral Organ Development in Rice

In the last decade, genetic and functional genomics analyses in different plant species revealed that flower development is governed by a complex framework based on MADS-domain TFs. Rice members of the grass-specific LOF-SEP clade sequentially regulate different steps of flower development upon the vegetative to reproductive phase change. First, *OsMADS34* forms tetrameric complex with the AP1-like factors *OsMADS14* and *OsMADS15* in the IM to initiate inflorescence branch meristem primordia development from which secondary branches and spikelets differentiate (Kobayashi et al., 2012). After the formation of rudimentary glumes and sterile lemmas, the spikelet meristem is converted into floret meristem, which produces different floral organs. Another grass-specific LOF-SEP factor, *OsMADS1*, plays a central role in the determination of floral organ identity, as it interacts physically and genetically with AP3-like and AG-like factors, which are involved in the development of male and female reproductive organs (Prasad et al., 2001; Li et al., 2011; Khanday et al., 2013, 2016). Recently, transcriptomics analysis performed on *OsMADS1* knock-down panicles at very early stages of flower development (Khanday et al., 2013) unveiled misregulation of

floral homeotic *MADS-box* genes as well as down-regulation of genes encoding B3-type TFs, including *OsARFs* and *OsRAV11* (Supplemental Table S2). Precisely, the ARF-type TFs *OsETTIN1* and - *OsETTIN2* control the differentiation of apical tissues of the carpel, and at least *OsETTIN2* is directly regulated by *OsMADS1* (Khanday et al., 2013). Intriguingly, the aberrant carpel morphology of loss of *OsETTINs* (Khanday et al., 2013) is similar to that of transgenic lines with reduced level of *OsRAV11* and *OsRAV12* (Fig. 7B).

### *OsRAV11* and *OsRAV12* Regulate the Development of Gynoecium

In Arabidopsis, the early flowering *TEM* loss of function mutants do not display evident alterations of flower development. Nonetheless, the late flowering *TEM* overexpressing lines show fertility defects and produce shorter siliques containing fewer seeds. This phenotype could be related to decreased content of GA, or alternatively to misregulation of genes involved in the regulation of later organ development. In rice, further molecular analyses suggest that RAV genes might have acquired additional functions at later stages of vegetative and reproductive growth. Indeed, *OsRAV11* and *OsRAV12* are highly expressed not only in mature leaves at ripening, indicating a possible regulatory role in leaf senescence similarly to *AtRAV1*, but also in the gynoecium before fertilization. Actually, functional characterization of knock-down and knock-out mutants points at a new function for *OsRAV11* and *OsRAV12* in the correct formation of female reproductive organs. Indeed, the whole basal-apical pattern was distorted; the ovary was enlarged in *osrav11* and misshapen in *OsRAV11/12-i* plants. Down-regulation of both genes resulted in reduced stigmas and larger carpels than in single *osrav11* mutants, which may suggest

a redundant function of these genes in carpel development and differentiation. Therefore, we hypothesize a role for *OsRAV11* and *OsRAV12* in the determination of the basal-apical pattern of the pistil, perhaps in parallel with ARFs. Furthermore, we propose that at least *OsRAV11* may control the differentiation of the gynoecium downstream of MADS-domain TFs (Fig. 8B), due to its mis-regulation in the floral homeotic mutants *osmads1*, *osmads13* and *osmads16* (Fig. 6, A and C).

Besides the specification of the identity of different floral organs, *OsMADS1* has been proposed as a key trait of agronomical interest. During spikelet development, *OsMADS1* also interacts with  $G\gamma$  subunits of *GS3* and *DEP1* (Fan et al., 2006; Huang et al., 2009; Liu et al., 2018), which regulate its transcriptional activity on a common set of target genes involved in the determination of seed size and shape. The dominant negative mutation *osmads1(lgy3)* causes an alternatively spliced protein variant and correlates with more slender grain, as the mutated protein promotes cell proliferation in longitudinal direction (Liu et al., 2018). Pyramiding of *lgy3* and *dep1-1* alleles in a japonica cultivar resulted not only in a 10% increase in grain yield, but also improved grain length-to-width ratio and grain chalkiness (Liu et al., 2018). We can hypothesize that the elongated seed phenotype associated to down-regulation of *OsMADS1* (Liu et al., 2018) could be mediated through the down-regulation of B3 genes including *OsRAV11* since a similar slender phenotype is observed in *osrav11* and *OsRAV11/12-i* carpels. Further analyses are required to understand the interactions with putative upstream regulators, interacting proteins and downstream targets constituting the molecular network that regulates the development of the gynoecium in rice.

## MATERIALS AND METHODS

### RAV Sequence Analyses

*OsRAV8* and *AtTEM1* were used as queries in phylogenomics analyses in sequenced land plants species (eudicotyledons, monocotyledons, Amborellales) using gene tree tool of *Pan-taxonomic Compara* (<http://www.gramene.org/>). *TEM1* protein sequence was used as query in a BLAST-P ([https://blast.ncbi.nlm.nih.gov/Blast.cgi?LINK\\_-LOC=blasthome&PAGE\\_TYPE=BlastSearch&PROGRAM=blastp](https://blast.ncbi.nlm.nih.gov/Blast.cgi?LINK_-LOC=blasthome&PAGE_TYPE=BlastSearch&PROGRAM=blastp)) search against the proteomes of *Arabidopsis thaliana* and rice (*Oryza sativa*).

TF binding sites (CArG box for MADS-domain proteins, consensus sequence composed of CAACA and CCTG elements at a distance of 3-9 nucleotides for RAV proteins) were searched in the regulatory regions of genes of interest by using the Promoter Analysis tool of Plant PAN3.0 (<http://plantpan.itsp.ncku.edu.tw/index.html>).

Gene and protein sequences were retrieved from The Arabidopsis Information Resource ([www.arabidopsis.org](http://www.arabidopsis.org)) and GRAMENE ([www.gramene.org](http://www.gramene.org)).

### Phylogenetic Analyses

Analysis of phylogenetic relationships between 24 RAV-related full-length protein sequences and construction of phylogenetic tree were performed using tools available at <http://www.phylogeny.fr/> (Dereeper et al., 2008). MUSCLE was used for protein sequences alignment, and G-blocks for a more stringent selection (<https://www.ebi.ac.uk/Tools/msa/muscle>). PhyML was used for phylogenetic analysis with bootstrapping procedure ( $n = 100$ ) as statistical

test for branch support ([http://phylogeny.lirmm.fr/phylo.cgi/one\\_task.cgi?task\\_type=phym1](http://phylogeny.lirmm.fr/phylo.cgi/one_task.cgi?task_type=phym1)), and TreeDyn for tree visualization ([http://www.phylogeny.fr/one\\_task.cgi?task\\_type=treedyn](http://www.phylogeny.fr/one_task.cgi?task_type=treedyn)). Sequence Diversity Diagram (SeDD; <http://vda-lab.github.io/sedd.html>) was used to compare two sets of RAV protein sequences from *Arabidopsis* and *Oryza sativa* and visualize conserved versus diversified positions in the two species. Motif clustering analysis was carried out using SALAD version 3 (<https://salad.dna.affrc.go.jp/salad/en/>) with *OsRAV9* sequence, and the prediction of functional and structural motifs in RAV protein sequences via web-based tools of ExPASy (<https://prosite.expasy.org/scanprosite>; de Castro et al., 2006).

### Expression Profiles

Expression profiles of rice RAV genes were inferred from a large collection of microarray data derived from different tissues at different developmental stages under natural field conditions (<http://ricexpro.dna.affrc.go.jp/>). Co-expression analyses were carried out by using *OsRAV9* as single guide genes and searching RiceFRIEND (<http://ricefriend.dna.affrc.go.jp/>), the Plant Co-expression Database (PLANEX; Yim et al., 2013), and the Rice Oligonucleotide Array Database (Cao et al., 2012).

### Plant Material and Growth Conditions

Wild-type and transgenic *Arabidopsis* (Col-0 background) seeds were sown on soil pots, and plants were grown under LD (16-h light/8-h dark at 22°C) until maturity. Wild-type and transgenic rice (*O. sativa* ssp. *japonica*) seeds were surface sterilized and sown on Murashige Skoog medium with 30 g L<sup>-1</sup> Suc. After germination, seedlings were transferred to soil pots and grown under controlled conditions until maturity (SD: 8-h light at 28°C, 16-h dark at 24°C; 12/12: 12-h light at 28°C, 12-h dark at 24°C or LD; LD:16-h light at 28°C, 8-h dark at 24°C). For functional characterization of *OsRAV* genes, transgenic lines (cv Nipponbare) and insertion mutant (PFG\_2A-10680, cv Hwayoung) were used. For expression analysis, segregating progenies of floral homeotic mutants (cv Dongjin) were genotyped (Supplemental Table S3), grown for 10 weeks in LD and then transferred to inductive conditions. Pools of developing inflorescences were harvested 3 weeks after floral transition from homozygous mutants (*osmads1*, *osmads13*, *osmads16*) and wild-type plants.

### Cloning and Generation of Arabidopsis and Rice Transgenic Plants

The coding sequence of *OsRAV9/OsTEM1* was amplified using primer sets SPp538-SPp541, subcloned in pCRII and pENTR-3C by restriction/ligation, and introduced in pALLIGATOR2 vector downstream of Pro-35S by Gateway technology (Supplemental Table S4). *Arabidopsis* plants (Col-0, *tem1-tem2*) were transformed with the *Pro-35S:OsTEM1* construct by floral dip, and GFP-positive seeds were selected by fluorescence microscopy. For the generation of the RNAi constructs, the Gene Sequence Tags specific for the 3' ends of *OsRAV9/OsTEM1* and *OsRAV11* were amplified using primer sets SPp516-SPp539 and SPp527-SPp537, respectively, then subcloned in pCRII, and afterward cloned in pENTR-3C by restriction/ligation (as *KpnI-EcoRV* fragments). The resulting constructs were digested with *PvuI* before LR recombination to pBios-378 plant expression vector. Scutellum-derived rice calli ('Nipponbare') were transformed by *Agrobacterium* spp. cocultivation. Independent transformation events were selected, 11 for *OsRAV9* and 7 for *OsRAV11* constructs, regenerated and propagated. Plants that underwent regeneration but did not contain the transgenic cassette were used as transformation control. The primers used for genotyping and cloning are listed in Supplemental Tables S3 and S4.

### Direct Binding of OsTEM1 to Downstream Genes

To generate the set of reporter vectors, different promoter regions of *OsMADS14* and *Hd3a* were cloned as *Sall-PstI* fragments in a modified pGreenII 0800-LUC carrying *Pro-35S:LUC* and *Pro-35S:REN* (as internal control to estimate the proportion of transformed protoplasts). Primers SPp1764- SPp1765 were designed to introduce mutations at the RAV binding site contained in *ProOsMADS14* by PCR-based method. The corresponding reporter vector was used as template for site-directed mutagenesis (15 cycles: 10' at 98°C, 20' at 66°C, 20' at 66°C), and the resulting vector used in transactivation assays. Protoplasts were isolated from calli by digesting the cell wall with Macerozyme R-10 and

Cellulase (Yakult Pharmaceuticals), and transfected with different combinations of reporter *Pro-35S:OsTEM1* and effector constructs using polyethylene glycol. After 18 h incubation in darkness at 24°C, transformed protoplasts were pelleted and resuspended in homogenization buffer for RNA extraction. Transactivation activity of *OsTEM1*, based on the relative ratio of mRNA abundance of *LUC* and *REN* reporter genes, was assessed by RT-qPCR. The primers used for cloning and expression analyses are listed in Supplemental Tables S3 and S4.

## RNA Extraction and Expression Analyses

For expression analyses in *Arabidopsis*, pools of 20 seedlings grown for 1 week under LD were collected at ZT12. For expression analyses in rice, pools of 10 to 15 samples from different tissues and/or developmental stages were collected at ZT13 unless otherwise stated. RNA was extracted with PureLink RNA mini kit (Ambion) and treated with DNaseI RNase free (Ambion). For large-scale experiments, RNA was extracted with Maxwell RSC Plant RNA kit (Promega), and DNase treatment was performed on-column. Then 1  $\mu$ g of DNase-treated RNA was retro-transcribed with SuperScript III (Invitrogen), and cDNA was used for RT-qPCR with Light Cycler 480 SYBR Green I master on Light Cycler 480 II (Roche). Three biological replicates and three technical replicates were performed. In situ hybridization was performed as previously reported (Dreni et al., 2007). The primers used for expression analyses are listed in Supplemental Table S5.

## Phenotypic analyses

Morphological analysis of reproductive structures was performed by Optical Microscopy (Olympus DP71) and SEM. For SEM, flowers at anthesis were fixed in 2.5% (v/v) glutaraldehyde in 0.1 M P-buffer (pH 7.4) overnight at 4°C, washed 4 times for 10 min in 0.1 M P-buffer, postfixed in 1% osmium tetroxide with 0.7% ferrocyanide in P-buffer, washed in water, dehydrated in an ascending ethanol series (50%, 70%, 80%, 90%, and 95% for 10 min each and twice with 100% ethanol), and dried by critical-point drying with CO<sub>2</sub>. Alteration on the morphology of the seed were analyzed by using the Smart-grain software (Tanabata et al., 2012). At least 100 seeds per genotype (for three independent biological replicates) were spread uniformly on the glass with a black background, and images were captured with HP Scanner at 300 dpi. The software determined seed shape parameters such as seed length, width, perimeter, and area, and also calculated length-to-width ratio and circularity.

## Statistical Analyses

All statistical analyses are shown in Supplemental Table S6. Statistical significance of each experiment was determined by using GraphPad Prism 7 (<https://www.graphpad.com/scientific-software/prism>). For flowering time phenotype, we chose one-way ANOVA; multiple comparisons were then corrected with Dunnett's or Dunn's tests. For other kind of data, we compared two columns by using unpaired Student's two-tailed *t* test, with confidence interval at 95%.

Following instructions of GraphPad Prism, we first transformed ratios in log of ratios ( $Y = \log(Y)$ ), for Figure 5 and Supplemental Figure S11. Then we created a column data table and entered two columns of data (control – *OsTEM1*, +*OsTEM1*) with matched values on the same row. We chose *t* tests from the list of column analyses and selected conditions as follows: Experimental design: Paired; Assume Gaussian distribution: Yes; Choose test: Ratio paired *t* test. On the second tab of the *t* test dialog, we chose to compute +*OsTEM1* versus –*OsTEM1*.

## Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers. AtRAV locus identifiers: AT1G13260 (AtRAV1), AT3G25730 (AtRAV1-like), At1g25560 (AtRAV2-like/TEM1), AT1G68840 (RAV2/TEM2), At1g50680 (AtRAV3), At1g51120 (AtRAV3-like). *OsRAV* locus identifiers: LOC\_Os10g39190 (OsRAV2), LOC\_Os08g06120 (OsRAV3), LOC\_Os03g02900 (OsRAV4), LOC\_Os04g49230 (OsRAV5), LOC\_Os02g45850 (OsRAV6), LOC\_Os11g05740 (OsRAV7), LOC\_Os01g04750 (OsRAV8), LOC\_Os01g04800 (OsRAV9/OsTEM1), LOC\_Os01g49830 (OsRAV11), LOC\_Os05g47650 (OsRAV12). *OsMADS* locus

identifiers: LOC\_Os03g54160 (OsMADS14), LOC\_Os07g01820 (OsMADS15), LOC\_Os07g41370 (OsMADS18), LOC\_Os03g54170 (OsMADS34), LOC\_Os03g03100 (OsMADS50/OsSOCI), LOC\_Os01g69850 (OsMADS51).

## SUPPLEMENTAL DATA

The following supplemental materials are available.

**Supplemental Figure S1.** Phylogenomic analysis of *TEM* homologs in sequenced land plants.

**Supplemental Figure S2.** Analysis of genes encoding AP2-B3 (RAV) proteins.

**Supplemental Figure S3.** Similarity clustering based on distribution patterns of known conserved motifs in RAV proteins.

**Supplemental Figure S4.** Similarity clustering based on distribution patterns of conserved features related to posttranslational modification.

**Supplemental Figure S5.** Analysis of RAV genes in *O. sativa* ssp. *japonica*.

**Supplemental Figure S6.** GO analysis of genes strongly coexpressed with *OsRAV9*.

**Supplemental Figure S7.** GO test of genes negatively correlated with *OsRAV9* and expression analysis of genes acting in apical meristems at floral transition in rice.

**Supplemental Figure S8.** Inferred expression profiles of *OsRAV* genes during plant development.

**Supplemental Figure S9.** Analysis of T1 and T2 transgenic lines ectopically expressing *OsRAV9* and *OsRAV11* in *Arabidopsis*.

**Supplemental Figure S10.** Molecular and phenotypic analysis of T1 transgenic rice lines silencing *OsRAV9*.

**Supplemental Figure S11.** Transactivation activity of *OsTEM1* on floral activators in rice protoplasts.

**Supplemental Figure S12.** Expression analyses of genes involved in the floral transition in *OsRAV9-i* rice lines.

**Supplemental Figure S13.** Analyses of *OsRAV11* and *OsRAV12*.

**Supplemental Figure S14.** Molecular characterization of *OsRAV11* and *OsRAV12*.

**Supplemental Figure S15.** Analysis of RAV proteins from *Arabidopsis* and *O. sativa*.

**Supplemental Table S1.** List of RAV genes in *O. sativa* reported by Swaminathan et al. (2008).

**Supplemental Table S2.** List of genes encoding B3-domain TFs down-regulated in *OsMADS1* knock-down panicles (modified from Khanday et al., 2013).

**Supplemental Table S3.** List of primers used for genotyping.

**Supplemental Table S4.** List of primers used for cloning.

**Supplemental Table S5.** List of primers used for expression analyses.

**Supplemental Table S6.** Statistical analyses.

**Supplemental Dataset S1.** Coexpression analyses

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