

A Rice Ca²⁺ Binding Protein Is Required for Tapetum Function and Pollen Formation¹[OPEN]

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In flowering plants, successful male reproduction requires the sophisticated interaction between somatic anther wall layers and reproductive cells. Timely degradation of the innermost tissue of the anther wall layer, the tapetal layer, is critical for pollen development. Ca²⁺ is a well-known stimulus for plant development, but whether it plays a role in affecting male reproduction remains elusive. Here we report a role of Defective in Exine Formation 1 (OsDEX1) in rice (*Oryza sativa*), a Ca²⁺ binding protein, in regulating rice tapetal cell degradation and pollen formation. In *osdex1* anthers, tapetal cell degeneration is delayed and degradation of the callose wall surrounding the microspores is compromised, leading to aborted pollen formation and complete male sterility. *OsDEX1* is expressed in tapetal cells and microspores during early anther development. Recombinant OsDEX1 is able to bind Ca²⁺ and regulate Ca²⁺ homeostasis in vitro, and *osdex1* exhibited disturbed Ca²⁺ homeostasis in tapetal cells. Phylogenetic analysis suggested that OsDEX1 may have a conserved function in binding Ca²⁺ in flowering plants, and genetic complementation of pollen wall defects of an Arabidopsis (*Arabidopsis thaliana*) *dex1* mutant confirmed its evolutionary conservation in pollen development. Collectively, these findings suggest that OsDEX1 plays a fundamental role in the development of tapetal cells and pollen formation, possibly via modulating the Ca²⁺ homeostasis during pollen development.

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J.Y. performed most of the experiments; Z.M. conceived the original screening of mutant identification; S.B. and J.K. analyzed the Ca²⁺ concentration; J.K. analyzed the Ca²⁺ binding activity; M.R.T. performed the callose immunolabeling; Z.L. and M.C. generated F2 population for mapping; D.X., G.Z., J.W., and S.Z. performed map-based cloning; Y.J.K. contributed for preparation of the manuscript; D.Z. and W.L. supervised the project; D.Z. complemented the writing.

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Higher plants alternate their life cycle between sporophytic and gametophytic generations that result from two sequential processes: sporogenesis and gametogenesis (Goldberg et al., 1993). Formation of the male gametophyte is a complex process that starts with anther morphogenesis, followed by microspore formation via meiosis and mitosis (Ma, 2005; Gómez et al., 2015; Zhang and Liang, 2016). The somatic cell layers surrounding the microsporocytes include the epidermis, endothecium, middle layer, and tapetum, which are required for normal pollen development. The differentiation of the innermost tapetal cell layer is crucial for pollen formation (Kelliher and Walbot, 2012; Fu et al., 2014; Zhang and Yang, 2014). After the formation of tapetal cells, subsequent tapetal cell and callose degradation, as well as primexine formation, are vital for pollen development (Ma, 2005; Li et al., 2006; Wu and Cheun, 2000; Paxson-Sowders et al., 2001; Ariizumi et al., 2004; Li et al., 2011a; Chang et al., 2012; Ji et al., 2013; Niu et al., 2013; Sun et al., 2013).

During tapetum development, regulated tapetal cell death is of vital importance for primexine formation, sporopollenin synthesis, and exine formation (Shi et al.,

2015; Zhang and Liang 2016). Until now, several transcription factors and their associated targets have been reported to play role in tapetal cell death (Sorensen et al., 2003; Jung et al., 2005; Li et al., 2006; Aya et al., 2009; Xu et al., 2010; Li et al., 2011a; Niu et al., 2013; Ji et al., 2013; Fu et al., 2014; Ko et al., 2014). In rice (*Oryza sativa*), mutations in the basic helix-loop-helix (bHLH) transcription factor TDR Interacting Protein2 (TIP2), also called bHLH142, show compromised inner anther wall layer differentiation and defects in microspore development (Fu et al., 2014; Ko et al., 2014). A mutant of the bHLH transcription factor, Undeveloped Tapetum1 (UDT1), displays abnormal development and degeneration of the tapetum and middle layer (Jung et al., 2005). *gamyb*, which encodes a MYB transcription factor, shows aborted tapetum degradation (Aya et al., 2009). Another bHLH transcription factor, Eternal Tapetum1 (EAT1), also called DTD, as well as one PHD finger protein, Persistent Tapetal Cell1 (PTC1), regulate microspore development via controlling tapetal cell death (Li et al., 2011a; Ji et al., 2013; Niu et al., 2013). TDR and EAT1 directly regulate the expression of proteases, which are regarded as executors for programmed cell death (PCD) in animals (Li et al., 2006; Niu et al., 2013; Woltering, 2010). For instance, EAT1 regulates the two aspartic proteases *OsAP25* and *OsAP37*, which trigger PCD in both yeast and plants (Niu et al., 2013). As a positive tapetal PCD determinant, TDR regulates the expression of *OsCPI1*, a Cys protease that is involved in tapetal degradation (Lee et al., 2004; Li et al., 2006; Niu et al., 2013). In addition, *OsCPI1* is also regulated by two ATP-dependent RNA helicases, AIP1 and AIP2 (Li et al., 2011b).

Tapetal cell death is also associated with the degradation of callose and the formation of primexine on the surface of the microspore, which is the first step of the pollen cell wall formation (Ariizumi and Toriyama, 2011; Shi et al., 2015). Mutants defective in callose degradation in rice also have defects in exine formation and pollen development (Wan et al., 2011). In Arabidopsis (*Arabidopsis thaliana*), Defective in Exine Formation1 (DEX1), No Exine Formation1 (NEF1), Ruptured Pollen Grain1 (RPG1), Ruptured Pollen Grain2 (RPG2), and No Primexine and Plasma Membrane Undulation (NPU) are required for primexine formation, and their corresponding mutants are defective in exine formation (Paxson-Sowders et al., 2001; Ariizumi et al., 2004; Chang et al., 2012; Sun et al., 2013). After the formation of the primexine, sporopollenin was synthesized in the tapetum and transported to the microspore. Several lipid metabolism-related genes, such as *Defective Pollen Wall* (DPW), *CYP704B2*, and *CYP703A3* have been reported to be essential for rice exine formation (Shi et al., 2011; Li et al., 2010; Yang et al., 2014b; Shi et al., 2015; Zhang et al., 2016). Additionally, sporopollenin precursor transport-related genes, such as *Postmeiotic Deficient Anther1* (*OsABCG15/PDA1*), *OsABCG26*, and *OsC6* (Zhang et al., 2010; Qin et al., 2013; Zhu et al., 2013; Zhang and Li 2014; Zhao et al.,

2015), are also required for pollen exine formation (Supplemental Fig. S1).

Up to now, the progress made toward understanding tapetal cell death is mainly restricted to the level of transcription factors and their related regulatory effects (Sorensen et al., 2003; Li et al., 2006; Aya et al., 2009; Xu et al., 2010; Li et al., 2011a; Niu et al., 2013; Ji et al., 2013; Fu et al., 2014; Ko et al., 2014). Consequently, further approaches are needed to reveal additional details of tapetal cell death regulation. Ca^{2+} is an essential component regulating a wide range of biological processes including cell division, differentiation, motility, and PCD (Poovaiah and Reddy, 1993; Trewavas and Malhó, 1998; Zielinski, 1998; Reddy, 2001; Lopez-Fernandez et al., 2015). Ca^{2+} affects cell death, including apoptosis, autophagy, and autolysis (Groover and Jones, 1999; Giorgi et al., 2008) by triggering the increase of cytosolic Ca^{2+} ($[\text{Ca}^{2+}]_{\text{cyt}}$; McConkey and Orrenius, 1997; Ferrari et al., 2002; Orrenius et al., 2003; Smaili et al., 2003; Giorgi et al., 2008). The endoplasmic reticulum (ER) is considered as an important Ca^{2+} -rich organelle, and ER-localized proteins have been reported to influence Ca^{2+} homeostasis in cells (Lam et al., 1994; Foyouzi-Youssefi et al., 2000; Kowaltowski et al., 2000; Pinton et al., 2000; Vanden Abeele et al., 2002). In Arabidopsis, BCL2-Associated X protein inhibitor1 acts as an inducer of ER stress-mediated PCD, which may play a role in affecting Ca^{2+} homeostasis on ER stress-mediated PCD in plants (Watanabe and Lam, 2008). Overexpression of ER-localized Bcl-2 protein promoted Ca^{2+} leakage from the ER and suppressed Ca^{2+} reuptake by the ER, causing the consequent increase of $[\text{Ca}^{2+}]_{\text{cyt}}$ and subsequent apoptosis (Hofer et al., 1998; Foyouzi-Youssefi et al., 2000; Vanden Abeele et al., 2002; Ferrari et al., 2002).

Ca^{2+} signal is sensed by a series of Ca^{2+} binding proteins, calmodulin (CaM), calcineurin B-like proteins, and Ca^{2+} -dependent protein kinases, and activates a number of transcriptional regulators by a kinase cascade or by direct interaction of CaM in a Ca^{2+} -dependent manner (Hiraga et al., 1993; Corneliusen et al., 1994; Enslin et al., 1995; Tokumitsu et al., 1995; Shi et al., 1999). Many Ca^{2+} sensors contain domain E and F (EF)-hand domains for Ca^{2+} binding (Snedden and Fromm, 1998). The typical EF-hand is a helix-loop-helix structure, in which the residues with $+X^*+Y^*+Z^*-Y^*-X^{**}-Z$ are associated with Ca^{2+} binding (Day et al., 2002; Derbyshire et al., 2007; Rigden et al., 2011). In addition, other motifs such as helix-loop-strand, helix-loop-turn, strand-loop-helix, strand-loop-strand, and several structural contexts without regular secondary structure elements either before or after the Dx Dx DG-containing loop can provide affinity for binding Ca^{2+} (Rigden and Galperin, 2004).

In this work, we characterized a Ca^{2+} binding protein, OsDEX1, which is required for tapetal function and pollen development in rice. OsDEX1 is conserved in flowering plants, and it is able to complement the mutant of its Arabidopsis counterpart. These findings provides the first evidence for an essential role of the Ca^{2+} homeostasis in plant pollen development.

RESULTS

Phenotypic Analysis of *osdex1*

To identify genes that are required for rice male reproduction, we isolated the *defective in exine formation1* in rice (*osdex1*) mutant (see below) from a rice mutant library generated in *ssp. japonica* cv 9522 background by ^{60}Co γ -ray irradiation (Chen et al., 2006), which showed a complete male sterility phenotype. Compared with wild-type plants, *osdex1* mutant displayed morphologically normal vegetative and nonreproductive floral organs, but smaller and pale yellow anthers lacking normal mature pollen grain, leading to complete male sterility (Fig. 1). All of the F1 progeny between wild type and *osdex1* were fertile, and the segregation rate of F2 generation was approximately 1:3 (sterility/fertility = 24:81), suggesting *osdex1* in caused by a single recessive mutation.

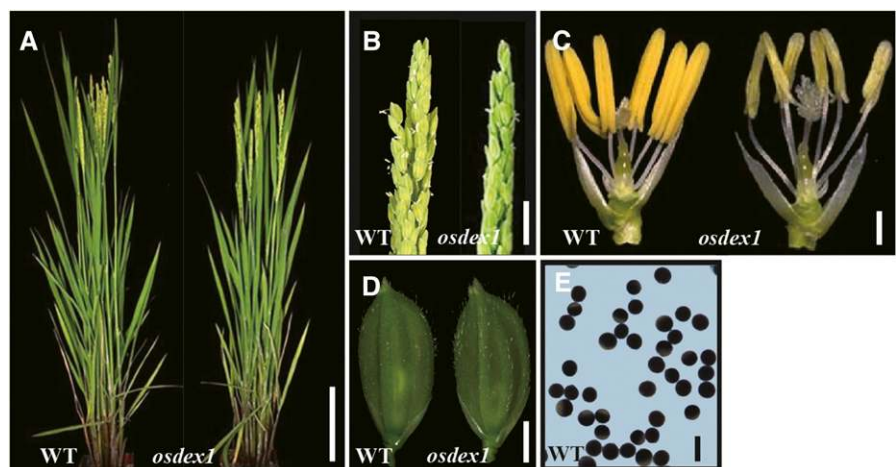
OsDEX1 Affects Tapetal Cell Death and Primexine Formation

Transverse sectioning was used to investigate the cellular morphological alterations of *osdex1* during pollen development, which was delineated based on a previous report (Zhang et al., 2011). No detectable morphological defect was observed in *osdex1* anthers until stage 8b when the microspore mother cells of both wild type and *osdex1* had undergone the second meiosis to produce ellipsoidal-shaped dyads, and tapetal cells were darkly stained with a vacuolated shape (Fig. 2, A and E). At stage 9, the wild-type microspores had separated from the tetrads and were distributed in anther lobes (Fig. 2B), while *osdex1* microspores were not separated (Fig. 2F). At stage 10, the wild-type microspores were covered in a thick layer of exine, displayed a round shape due to vacuolization, and the tapetum were degenerated into a thin layer (Fig. 2C). In contrast, *osdex1* microspores were smaller, lacked a clear layer of exine compared with the wild type, and the tapetal cells were not degenerated but appeared expanded in some

regions (Fig. 2G). At stage 11, the wild-type tapetal cells were completely degraded and sickle-shape microspores exhibited storage starch accumulation (Fig. 2D). However, *osdex1* displayed a persistent tapetal layer, and the microspores showed a similar morphology to previous stages (Fig. 2H). These observations indicate that *osdex1* exhibits delayed tapetal degradation and aborted exine formation.

Transmission electron microscopy (TEM) was used to further investigate the defects in *osdex1* pollen formation and tapetum degeneration. Consistent with the observations by light microscopy, no visible morphological differences were observed between the wild type and *osdex1* (Supplemental Fig. S2) until stage 8a. At stage 8b, wild-type tapetal cells became largely vacuolated with a thin layer of cell wall (Fig. 3, A and E), and the plasma membrane (PM) of wild-type microspore showed regular undulations and primexine deposition (Fig. 3E). In *osdex1* tapetal cells, where vacuoles were reabsorbed, their cytoplasm appeared condensed with a large number of mitochondria and they displayed a thicker cell wall (Fig. 3, I, M, and Q). PM undulation was not observed in mutant microspores and primexine deposition did not occur (Fig. 3Q). At stage 9, wild-type tapetal cells displayed a waved shape accompanied by an enrichment of organelles, such as mitochondria and ER, and small bubbles that likely represented the early stages of orbicule development (Fig. 3, B and F). Correspondingly, a thin layer of exine (including probacula and nexine) was present on the wild-type microspore surface (Fig. 3F). Consistent with this observation, sporopollenin precursor synthesis and transport-related genes such as *DPW*, *CYP703A3*, *CYP704B2*, *OsC6*, and *PDA* were highly expressed in wild-type anthers (Supplemental Fig. S3; Shi et al., 2011; Yang et al., 2014b; Li et al., 2010; Zhu et al., 2013; Zhang et al., 2010). By contrast, in *osdex1*, at the same stage, tapetal cells were flat with a thicker cell wall and contained a larger number of vacuoles with variable size compared to wild type (Fig. 3, J and R). Notably, the vacuoles in the mutant tapetal cells were of irregular

Figure 1. Phenotypic comparison between the wild type and the *osdex1* mutant. A, Wild-type plant (left) and the *osdex1* mutant plant (right) after heading. B, Part of the wild-type panicle showing the dehiscent anther (left) and part of the *osdex1* panicle (right) showing a smaller anther at the pollination stage. C, Wild-type (left) and *osdex1* (right) flower organs after removal of the palea and lemma. D, Wild-type (left) and *osdex1* (right) flowers before anthesis. E, Wild-type stained pollen at stage 13. Bars = 10 cm (A), 2 cm (B), 5 mm (C and D), and 100 μm (E).



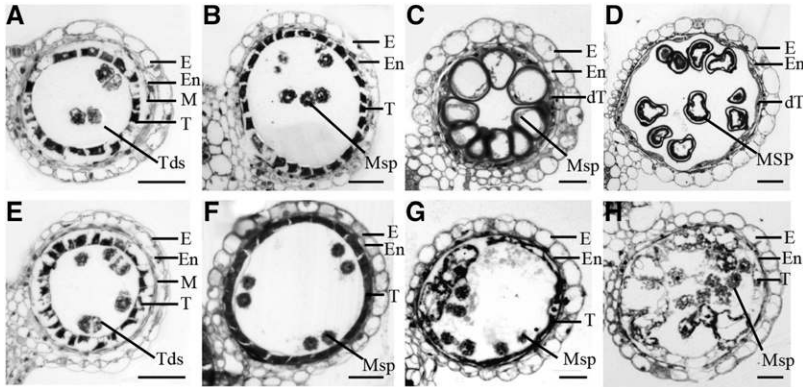


Figure 2. Bright field microscopy of transverse sections showing anther and microspore development in wild type and *osdex1*. Locules from the anther section of the wild type (A–D) and *osdex1* (E–H) from stage 8 to stage 11. dT, degenerated tapetal layer; E, Epidermis; En, endothecium; M, middle layer; Msp, microspores; T, tapetal layer; Tds, tetrads. Bars = 15 μ m. A and E, Stage 8b. B and F, Stage 9. C and G, Stage 10. D and H, Early stage 11.

shape and were attached to each other exhibiting membrane ablation (Fig. 3N). Furthermore, the structures of nucleus and ER seemed disrupted in the mutant tapetal cells (Fig. 3, O and P). No intact exine structure was formed on the *osdex1* microspore surface, which only showed fragmented primexine deposition (Fig. 3R). Furthermore, no obvious expression of *DPW*,

CYP703A3, *CYP704B2*, *OsC6*, and *PDA* was detected in the mutant, suggesting the defective synthesis and transport of sporopollenin precursors (Supplemental Fig. S3). At stage 10, wild-type tapetal cells were thin and contained a large number of mature orbicules on the inner surface (Fig. 3, C and G). As a result, wild-type microspores possessed a thick layer of exine on the

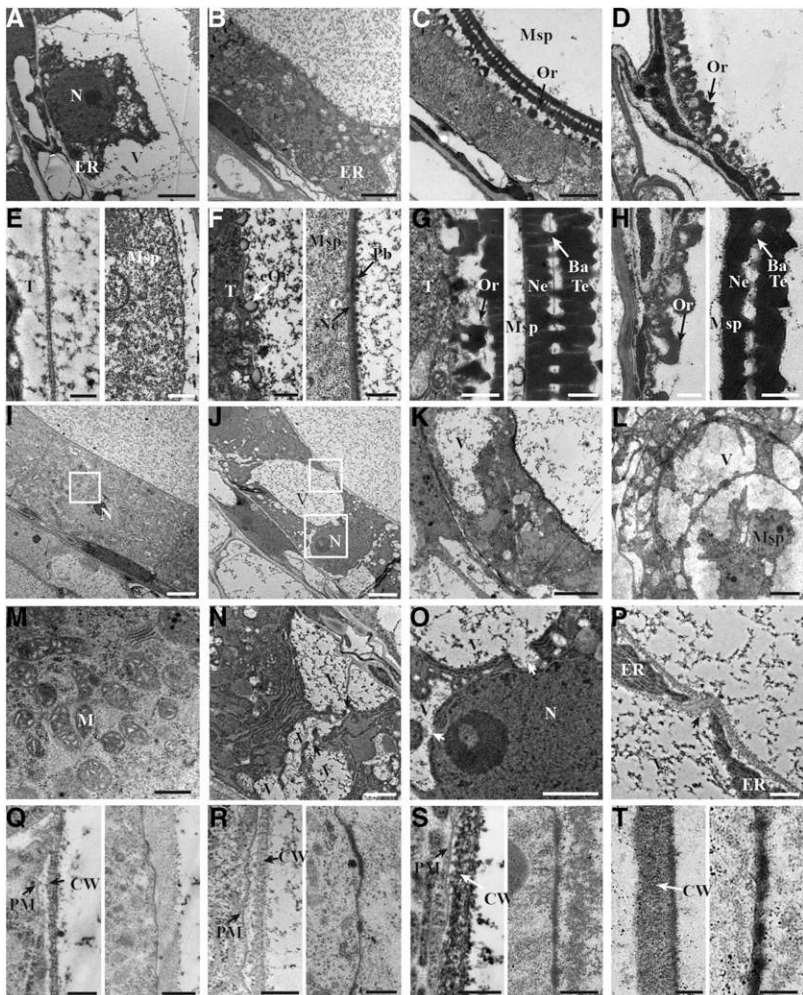


Figure 3. TEM images of the anthers from the wild type and *osdex1*. A to D, TEM observation showing tapetal cells of wild-type anthers at stage 8b (A), stage 9 (B), stage 10 (C), and stage 11 (D). E to H, Wild-type tapetal cell wall (left) and microspore cell wall (right) at stage 8b (E), stage 9 (F), stage 10 (G), and stage 11 (H). I to L, TEM observation showing tapetal cells of *osdex1* anthers at stage 8b (I), stage 9 (J), stage 10 (K), and stage 11 (L). M, Higher magnification of the highlighted region in (I) showing details in tapetal cells. N, Vacuoles fusion in *osdex1* tapetal cells at stage 9. Black arrows show the attachment of vacuoles. O and P, Higher magnification of the highlighted region in (J) showing details in tapetal cells. White arrows show the breakage of vacuoles. Black arrow shows the breakage of ER. Q to T, *osdex1* tapetal cell wall (left) and microspore cell wall (right) at stage 8b (Q), stage 9 (R), stage 10 (S), and stage 11 (T). Ba, Bacula; CW, cell wall; eOr, early stage of orbicules; Pb, probacula; M, mitochondria; N, nucleus; Ne, nexine; Or, orbicules; T, tapetal cells; Te, tectum; V, vacuoles. Bars = 2 μ m (A–C and I–K), 1 μ m (D and N), 10 μ m (L), and 0.5 μ m (E–H, M, O–P, and Q–T).

surface (Fig. 3G). By contrast, *osdex1* tapetal cells were expended in several regions because of large vacuoles (Fig. 3K). The persistent tapetal cells were covered by a thick cell wall without any orbicules (Fig. 3S). Moreover, mutant microspores were covered by a thin layer of darkly stained material, lacking the normal structure of exine (Fig. 3S). At stage 11, the tapetal layer in wild-type anthers was almost completely degenerated and orbicules were present around microspores forming a thick exine (Fig. 3, D and H), while *osdex1* exhibited expanded tapetal cells with large vacuoles, and extremely thick cell walls (Fig. 3, L and T) without sculptured exine on the microspores (Fig. 3T). These results suggest that OsDEX1 affects the normal tapetal cell death, synthesis of sporopollenin precursors, and the formation of pollen wall ranging from the primexine to multiple-layer exine during male development.

The persistence of tapetal cells in *osdex1* mutants was investigated using the terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling (TUNEL) assay in which the fluorescein-12-dUTP-labeled DNA is catalytically incorporated into fragmented DNA, and can be visualized by confocal laser scanning microscope (Li et al., 2006). The wild-type tapetum displayed DNA fragmentation from stage 8b until stage 9, and there was no DNA fragmentation at stage 7 and stage 8a. In contrast, the DNA fragmentation in *osdex1* tapetal cells could be observed starting from stage 7 until stage 8a, and declined at stage 8b and stage 9 (Fig. 4). Therefore, DNA fragmentation appears to be activated early in *osdex1* tapetal cells and disrupted at the stages when wild-type tapetal cells show DNA fragmentation. To further assess tapetal cell death, the expression pattern of tapetal cell death-related genes was determined. *OsAP25*, *OsAP37* and *OsCP1* were not induced in *osdex1* anthers compared to wild-type anthers undergoing cell death (Supplemental Fig. S4), suggesting that the normal protein degradation pathway was disrupted in the *osdex1* mutant during tapetum development.

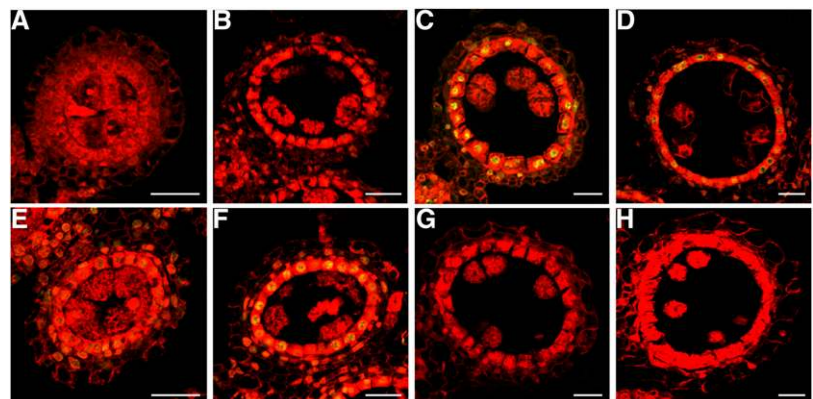
Our observations using TUNEL and TEM suggest abnormal tapetal cell degeneration in *osdex1*. Consistent with the earlier induction of DNA fragmentation in *osdex1* tapetal cells, in *osdex1* we observed

up-regulation of five endonuclease-encoding genes (*LOC_07g45100*, *LOC_01g03740*, *LOC_04g58850*, *LOC_02g50040*, and *LOC_08g29700*) that are down-regulated in the tapetal cell death-deficient mutants *gamyb* and *tip2* at stage 8 (Aya et al., 2009; Fu et al., 2014). Similarly at stage 7, the genes were either up-regulated or showed the opposite expression pattern compared to *gamyb* and *tip2*. These findings are in agreement with the opposite cell death phenotype. At stage 9 when the DNA fragmentation signal was declined, the expression of these genes was down-regulated (Supplemental Fig. S4).

OsDEX1 Affects Callose Degradation

The TEM analysis indicated that *osdex1* microspores were covered in an electro-dense matrix at later stages (Fig. 5, A to D), likely indicative of abnormal callose degradation. To further characterize the role of OsDEX1 in callose dynamics during microspore development, aniline blue staining and immunolabelling with a callose antibody were performed (Figs. 5, E to L; S5, E to L). At stage 7, both wild-type and *osdex1* tetrads were labeled with aniline blue, indicating the relatively normal synthesis of callose in *osdex1* (Supplemental Fig. S5, E and I). A similar pattern was observed using callose immunolabeling, although in this case staining was weaker around *osdex1* tetrads than in wild type at stage 8b (Fig. 5, F and J). This is supported by the TEM observations showing a looser electron-dense matrix around *osdex1* tetrads (Supplemental Fig. S5, A and B). Also at stage 8b, both aniline blue and callose antibody stained cell-wall material in the center and the border region surrounding the wild-type tetrads (Figs. 5, F and J; S5, F and J). Remarkably, in *osdex1*, aniline blue staining was restricted mainly to the inner tetrad walls, suggesting some changes in cell-wall composition of the outer walls, possibly because of precocious yet incomplete callose degradation (Supplemental Fig. S5F). Specific callose labeling was detected by aniline blue and callose antibody directly adjoining *osdex1* microspores, and weakly in the diffuse halos surrounding them, until stage 10 (Figs. 5, K and L; S5, K and L) when

Figure 4. DNA fragmentation is initiated earlier and subsequently blocked in *osdex1* mutant. A to H, DNA fragment signal at stage 7, stage 8a, stage 8b, and stage 9 in wild type (A–D) and *osdex1* mutant (E–H). The red fluorescence shows the propidium iodide staining of anther cells using confocal laser scanning microscope; the yellow fluorescence shows the TUNEL-positive nuclei staining in confocal laser scanning microscope overlays of fluorescein staining and propidium iodide staining. Bars = 15 μ m.



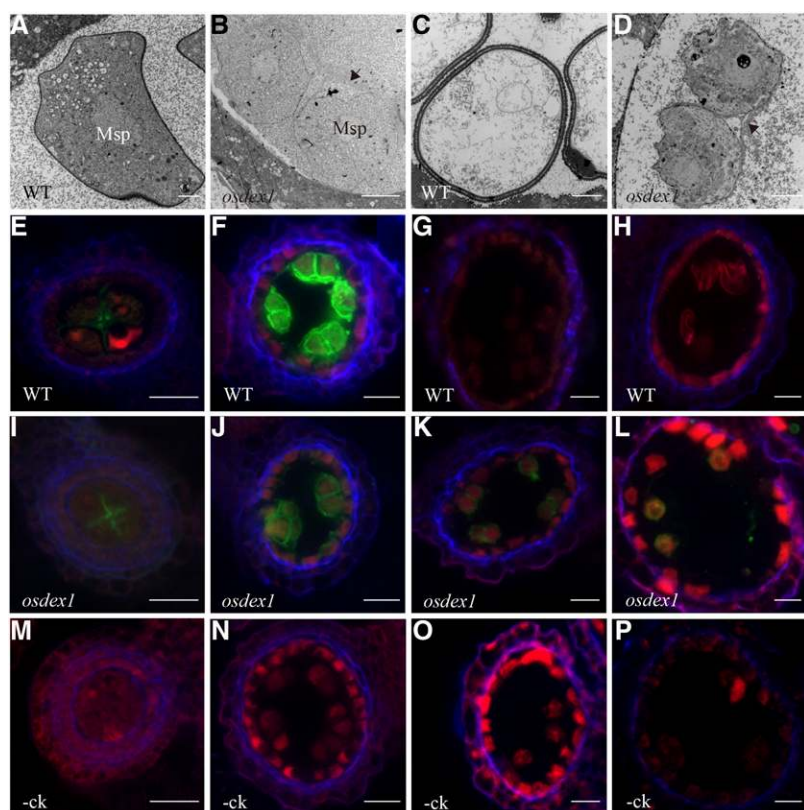


Figure 5. Callose degradation is retarded in *osdex1*. A to D, TEM of extracellular materials from the wild type and *osdex1* at stage 9 (A and B), and stage 10 (C and D). Black arrows show the site of callose deposition. E to L, Immunolabeling of wild-type (E–H) and *osdex1* (I–L) anther sections from stage 7 to stage 10 observed by epifluorescence microscopy. M to P, Negative controls of immunolabeling. In (E) to (P), the green channel shows immunostaining with callose antibody; blue counterstaining shows 1,4- and 1,3;1,4-glucan polymers stained with 0.01% calcofluor white; and red staining shows background autofluorescence. Bars = 2 μm (A), 5 μm (B–D), and 15 μm (E–P).

staining was absent around the wild-type microspores. These results suggest that OsDEX1 affects callose degradation during microspore development.

Cloning and Expression Analysis of *OsDEX1*

To identify the gene responsible for the *osdex1* phenotype, we employed a map-based cloning approach. *OsDEX1* was mapped to chromosome 3 between the markers of Os315 and Os315-2, which incorporated a 26 kb DNA fragment and seven putative genes (Supplemental Fig. S6A). After sequencing anther-expressed candidate genes within this region, a 1 bp deletion was identified in the coding region of a gene corresponding to *LOC_Os03g61050* (<http://www.gramene.org/>, also annotated as Os03g0825700 by <http://rapdb.dna.affrc.go.jp/>), which deletion resulted in a frame shift and subsequently an earlier termination of the translation of this gene (Supplemental Fig. S6B). Moreover, three additional alleles of *OsDEX1* were identified, and we re-named *osdex1* as *osdex1-1*, and the additional three alleles were designated as *osdex1-2*, *osdex1-3*, and *osdex1-4* (Supplemental Fig. S6B). *osdex1-2* has a one-nucleotide G to A substitution at an intron boundary (position 4296), causing the alternative splicing of *OsDEX1*. *osdex1-3* had a 374 bp deletion in the 10th exon leading to a 286 bp deletion of the mRNA. *osdex1-4* had a 4 bp deletion in the sixth exon leading to an earlier termination of *OsDEX1*. All of

the mutants exhibited similar male sterile phenotype and expanded tapetal cells (Supplemental Fig S7). Allelic tests confirmed that the four mutants were indeed alleles of a single locus. The *OsDEX1* transcript is 3241 bp in length and contains a 2556 bp coding sequence with 12 introns, a 180 bp 5' untranslated region, and a 505 bp 3' untranslated region (Supplemental Fig. S6B). The predicted *OsDEX1* protein encompasses 851 amino acids and harbors one putative N-terminal signal peptide (amino acids 1 to 23), one integrin α -N-terminal domain (amino acids 69 to 582) containing two phenylalanyl-glycyl-glycyl-alanyl-prolyl (FG-GAP) domains (amino acids 460 to 486 and 554 to 579; PF01839; <http://pfam.xfam.org>), and one transmembrane domain (amino acids 816 to 826; Supplemental Fig. S6C). The integrin α -N-terminal domain is predicted to mediate cell adhesion, and the FG-GAP domain has been shown to be important for ligand binding (Loftus et al., 1994).

To gain insights into the function of *OsDEX1*, quantitative RT-PCR (qRT-PCR) analysis was used to further investigate the spatio-temporal expression pattern of *OsDEX1*. In the wild type, *OsDEX1* expression was detectable in roots, shoots, and leaves, with a preferable expression in the anther, starting from stage 6, reaching an expression maximum at stage 8 and stage 9. *OsDEX1* expression showed a dramatic reduction in *osdex1-1*, particularly in anthers (Supplemental Fig. S6D). Although the *OsDEX1* expression was high in wild-type leaves and much reduced in the mutant, no visible

phenotypic change was observed in mutant leaves under normal growth conditions, suggesting no obvious role of OsDEX1 or redundant gene function during leaf development. Furthermore, in situ hybridization also showed that *OsDEX1* was expressed in tapetal cells and microspores from stage 8a to stage 9, and low level at stage 10 (Supplemental Fig. S6, E to L).

OsDEX1 Has a Conserved Function in Pollen Development

To elucidate the evolutionary conservation and distribution of OsDEX1, the full-length OsDEX1 protein was used as the query to search for its closest relatives in public databases, including the National Center for Biotechnology Information, The Arabidopsis Information Resource, Phytozome (<http://www.phytozome.net/>), and PLAZA (<http://bioinformatics.psb.ugent.be/plaza/>), which yielded 25 sequences from 24 different species, and all of which were then included in the phylogenetic analysis (Fig. 6). Among the 24 species, besides rice, there are monocots such as *Oryza brachyantha*, dicots such as Arabidopsis, basal angiosperm such as *Amborella trichopoda*, Pteridophyta such as *Selaginella moellendorffii*, Bryophyta such as *Physcomitrella patens*, Chlorophyta such as *Chlamydomonas reinhardtii*, Protozoa such as *Phytomonas sp. isolate Hart*, Mollusca such as *Strongylocentrotus purpuratus*, and Prokaryotes such as *Firmicutes bacterium*. These data

suggested that OsDEX1 belongs to an ancient protein subfamily, present in both prokaryotes and eukaryotes, which was designated as DEX1 (PTHR21419:SF23) by the panther database (<http://www.pantherdb.org/>). Notably, proteins in this family only exist in lower animals, while they are present in both higher plants and lower plants. OsDEX1 clustered together with other monocot DEX1 sequences in the same clade, while proteins from dicots, including the Arabidopsis homolog DEX1, clustered in a separate clade. In addition, the homologous eukaryotic sequences all contained a transmembrane domain, suggesting their similar membrane localization in the cell. The FG-GAP domain was only detectable in the sequences of flowering plants, which has been shown to be important for ligand binding (Loftus et al., 1994). These results suggest that OsDEX1 might have an evolutionarily conserved ancient function in male gametophyte development that, however, may have obtained the ability ligand binding during the formation of angiosperms.

Based on the phylogenetic analysis, OsDEX1 was identified as a putative ortholog of Arabidopsis DEX1. *dex1* was also reported as a male sterile line due to its delayed primexine formation, and aborted pollen wall development, but lacked characterization of its tapetal and biochemical function (Paxson-Sowers et al., 2001). The amino acid sequence of OsDEX1 was overall 66.5%

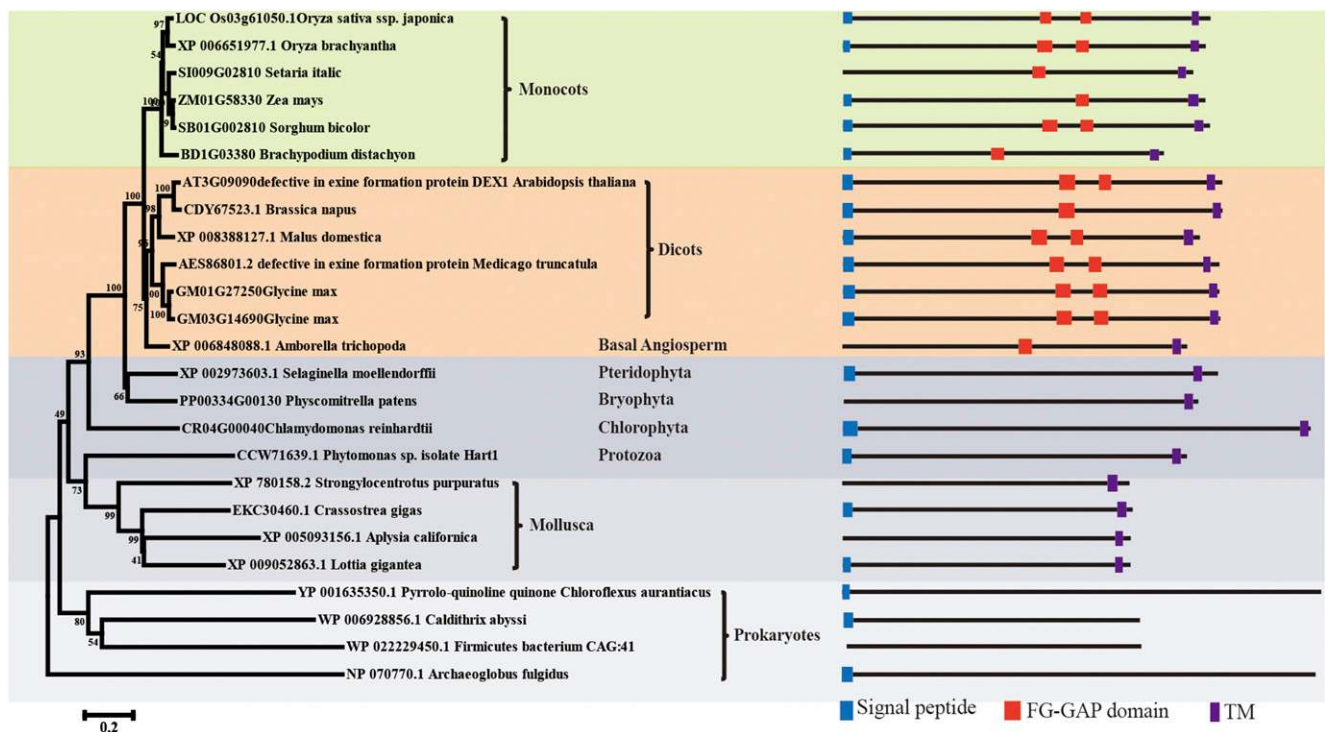


Figure 6. Phylogenetic analysis of OsDEX1 and its related proteins. Neighboring-joint analysis was performed using MEGA 6.1 (see “Materials and Methods”) based on the alignment given in Supplemental Data Set S1 online of OsDEX1 with the most similar OsDEX1 sequences from the species shown. The species were classified by evolutionary relationship.

identical to that of DEX1 (Supplemental Fig. S8), suggesting that they may have similar function. To test this hypothesis, *OsDEX1* cDNA driven by *DEX1* promoter (1.7 Kb) was introduced into *dex1* heterozygous plants. More than 100 T1 transformants were identified, and among the *dex* homozygous lines containing the transgene, 42% displayed full fertility and 25% showed partial fertility. Notably, Alexander staining showed that in the fully fertile lines, all the pollen grains in the anther were stained red, indicating that these pollen grains are viable (Fig. 7C). In the partially fertile lines, some anthers showed all pollen grains stained in red, while others showed a reduced amount of red-stained pollen grains (Fig. 7D). Altogether, these results demonstrate that OsDEX1 fulfills a conserved function in pollen development.

OsDEX1 Is a Calcium Binding Protein

The phylogenetic analysis suggested the importance of the FG-GAP domain in flowering plants, and FG-GAP domain has also been also reported to mediate calcium binding (Tuckwell et al., 1992; Loftus et al., 1994; Springer, 1997). To investigate whether the conserved FG-GAP domain of OsDEX1 can indeed bind to Ca^{2+} , a three-dimensional (3D) structure prediction of OsDEX1 was conducted using SWISSMODEL (<http://www.swissmodel.expasy.org/>), which revealed that the 3D structure of OsDEX1 was rich in β -sheet linked by loops (Supplemental Fig. S9), implying its potential ability to bind Ca^{2+} . Structural analysis showed the oxygen atoms from the side chains of the first, third, and fifth residues in the EF-hand motif that may form a pocket suitable for calcium atom binding (Fig. 8, A and B).

An in vitro Ca^{2+} binding assay (Gregersen et al., 1990; Ling and Zielinski, 1993; Libich and Harauz, 2002) showed that the truncated protein of OsDEX1 (amino acids 336 to 634), containing 3 EF-hands motifs, was shifted in an acrylamide-SDS gel after incubation with 1 mM CaCl_2 . However, when Ca^{2+} was chelated by

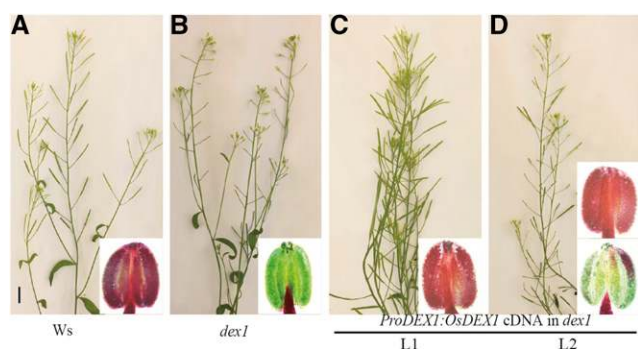


Figure 7. A to D, Transgenic lines containing Pro:*DEX1*:*OsDEX1* in the Arabidopsis *dex1* mutant display rescued male fertility. Insets show the pollen tested by Alexander staining using bright field microscopy. Ws, wild-type plant of Wassilewskija ecotype. Bar = 10 mm.

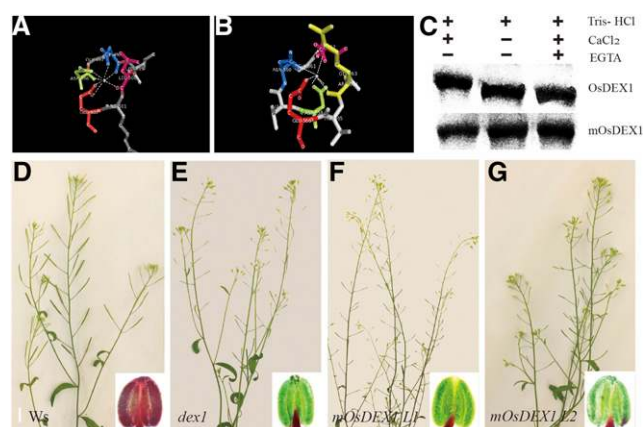


Figure 8. Recombinant OsDEX1 has Ca^{2+} binding activity. A, 3D structure of EF-hand motif of YesW. B, 3D structure of EF-hand motif of OsDEX1. C, In vitro Ca^{2+} binding assay shows that OsDEX1 has Ca^{2+} binding activity. D to G, Failure in complementation by OsDEX1 with the mutated Ca^{2+} binding sites in *dex1*. Insets show pollen tested by Alexander staining using bright field microscopy. Bar = 10 mm.

adding EGTA, no shift in fragment migration was detected. In addition, when an isoform of OsDEX1 the first, third, fourth, fifth, and sixth residues of the conserved calcium binding domain were muted was incubated with CaCl_2 , no shift was observed (Fig. 8C). Importantly, when the mutated *OsDEX1* cDNA driven by *DEX1* promoter (1.7 Kb) was introduced into the Arabidopsis *dex1* mutant, the transgenic plants homozygous for *dex1* failed to produce viable pollen grains (Fig. 8, D to G). These results identify OsDEX1 as a Ca^{2+} binding protein and provide evidence for a pivotal role of Ca^{2+} binding in the function of OsDEX1.

OsDEX1 Is Required for Adequate Ca^{2+} Homeostasis in Tapetum

The abnormal tapetal cell death phenotype and the alteration of the Ca^{2+} binding ability of a truncated OsDEX1 protein suggested that OsDEX1 might regulate tapetal cell death by affecting Ca^{2+} distribution in the tapetal cells, as suggested by previous reports (Wyllie, 1980; Cohen and Duke, 1984). To test this, yellow cameleons 3.6 (YC3.6) expressed under the control of the constitutive *UBI10* promoter, a powerful tool monitoring the spatio-temporal dynamics of Ca^{2+} fluxes in different tissues of Arabidopsis and rice (Krebs et al., 2012; Nehera et al., 2015), was used to monitor Ca^{2+} homeostasis in rice tapetum. There are three detectable somatic cell layers in the rice anther at stage 9, i.e. the epidermis, endothecium, and tapetum. In Figure 9, the anther images show the morphology of tapetal cells, which is consistent with a previous report (Zhao et al., 2015). PM-YC3.6, which is a plasma membrane targeted version of YC3.6, is able to distinguish the faint Ca^{2+} signal at the cytosolic side of the plasma membrane from the strong autofluorescence of anthers.

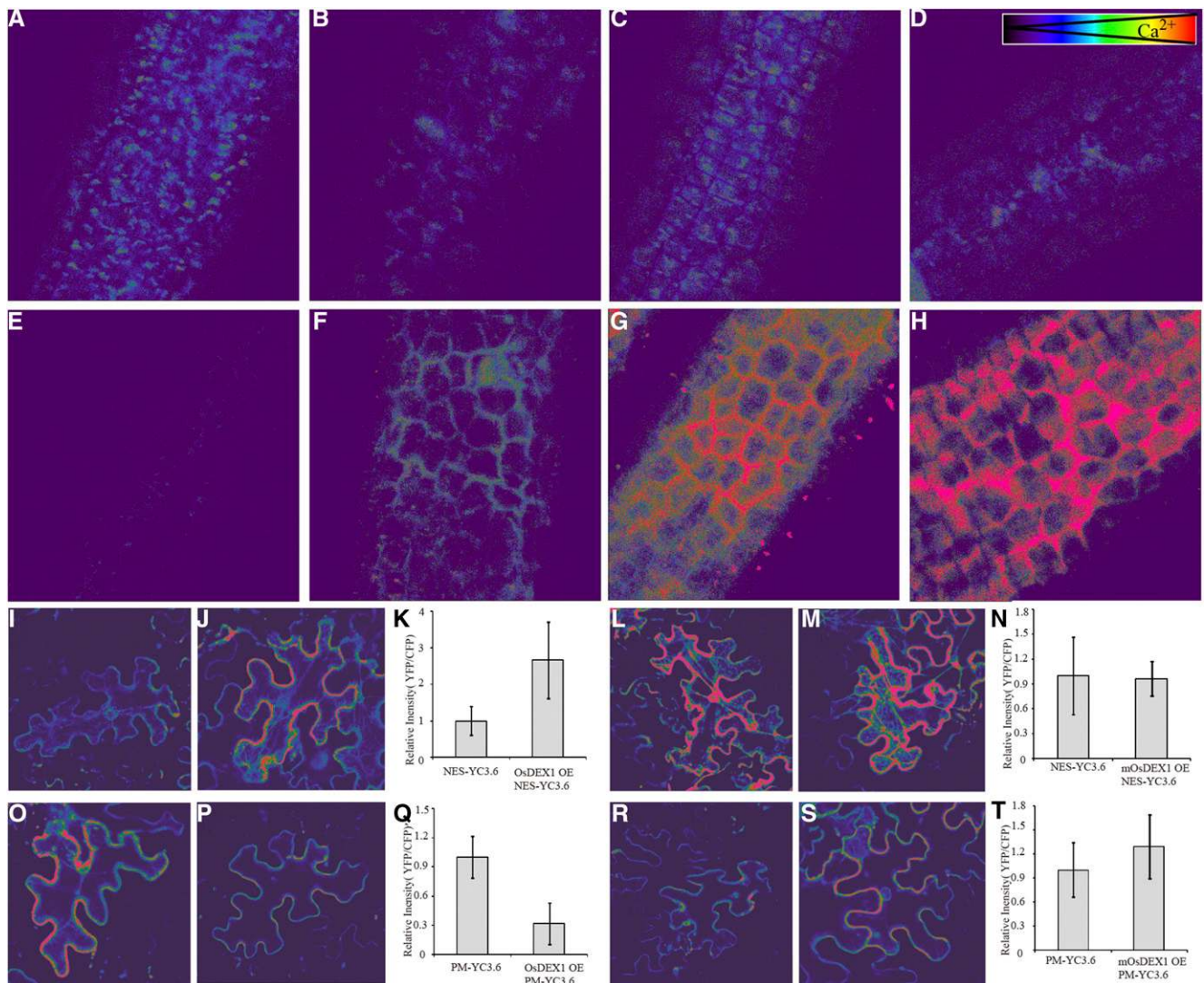


Figure 9. The Ca^{2+} concentration calculated by emission ratios between YFP and CFP intensities using confocal laser scanning microscope in rice anthers and tobacco epidermal cells. A to H, Ca^{2+} concentration on plasma membrane in tapetal cells of wild type (A–D) and *osdex1* (E and F) from stage 8 to stage 11. I to T, Comparison of cytosolic (I to N) and plasma membrane (O–T) Ca^{2+} concentration in tobacco epidermal cells overexpressed with OsDEX1 (J and P) or mOsDEX1 (M and S). I, L, O, and R, Control of YC3.6 overexpression. K, N, Q, and T, Statistical analysis of ratios between YFP and CFP intensities in the cells (K and N) and on the plasma membrane (Q and T).

During anther development from stage 8 to stage 10, only a faint Ca^{2+} signal was observed in the wild-type anthers. In sharp contrast, in the *osdex1* mutant, the plasma membrane-localized Ca^{2+} reporter indicated a dramatic increase of intracellular Ca^{2+} concentration in the tapetal cells specifically initiating from stage 9, that was even further enhanced at stage 10 (Fig. 9, A to H). These results identify rapid and strong elevations of intracellular Ca^{2+} concentration that occur stage specific only in *osdex1* and the coincide with the establishment of tapetal phenotypes in this mutant. These findings support the conclusion that the increase in $[\text{Ca}^{2+}]$ specifically observed in *osdex1* represents an important facet of the cell death ongoing in these cells and there by

attribute a crucial function for maintaining appropriate cellular $[\text{Ca}^{2+}]$ homeostasis to OsDEX1.

In another assay for cytosolic $[\text{Ca}^{2+}]$ homeostasis, cytoplasmic NES-YC3.6 or plasma membrane targeted PM-YC3.6 were expressed alone or coexpressed together with either OsDEX1 protein or non- Ca^{2+} binding mOsDEX1 protein transiently in tobacco epidermal cells. Consistent with the results in rice tapetal cells, $[\text{Ca}^{2+}]_{\text{PM}}$ was lower when PM-YC3.6 was coexpressed with OsDEX1 compared to cell that did only express PM-YC3.6 while $[\text{Ca}^{2+}]_{\text{cyt}}$ was elevated when NES-YC3.6 was coexpressed with OsDEX1 compared to cells that only expressed NES-YC3.6. Importantly, the Ca^{2+} concentration was not affected when PM-YC3.6 or NES-YC3.6

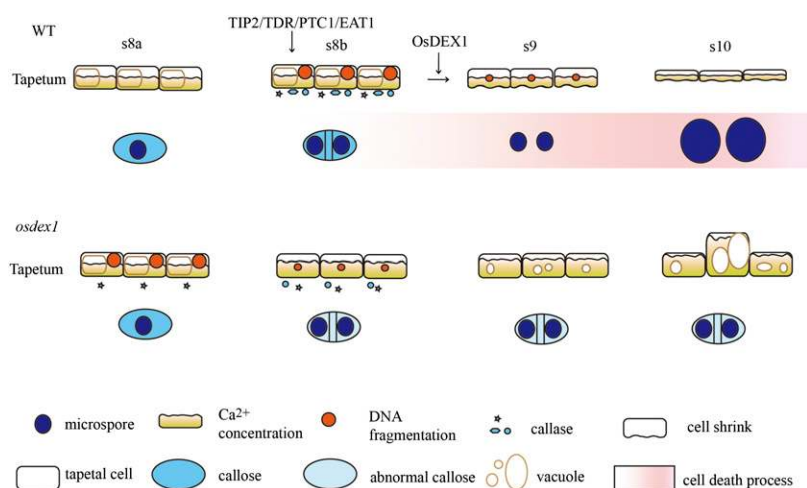


Figure 10. Model for OsDEX1 in anther development. Transcription factors such as TIP2, TDR, PTC1, and EAT1 function as master valves to switch cell death signaling on or off; OsDEX1 buffers the Ca^{2+} concentration in the cells to function as a component of cell death signaling.

were coexpressed with mOsDEX1, which lacks Ca^{2+} binding activity (Fig. 9). These results reveal that the expression of OsDEX1 modulates the Ca^{2+} homeostasis in the cells, which is considered as a key mechanism for cell death regulation (Giorgi et al., 2008).

To determine the subcellular localization of OsDEX1 at which this protein impacts on cellular $[\text{Ca}^{2+}]$ dynamics, the full-length OsDEX1 coding region was fused with the GFP at its C terminus, and transiently expressed in tobacco epidermal cells. The OsDEX1-GFP signal was detected on the ER (Supplemental Fig. S10), which is in agreement with previous reports suggesting that the ER is a main storage site for Ca^{2+} in the cell (Pozzan et al., 1994). Considering the results described above, we therefore hypothesize that OsDEX1 may regulate tapetal cell death via modulating Ca^{2+} homeostasis between the ER and other cellular compartments.

DISCUSSION

Rice is one of the most important foods for the global population. Rice yield improvement is largely dependent on hybrid breeding that employs male sterile lines. Although much progress has been made toward understanding the molecular mechanisms underlying plant male development, the role of Ca^{2+} during anther development has remained elusive. In this study, we have demonstrated that an FG-GAP domain-containing protein, OsDEX1, is required for pollen wall development and tapetum function in a way that has not been elucidated in previous descriptions of the Arabidopsis *dex1* mutant (Paxson-Sowders et al., 2001). The evolutionary importance of the OsDEX1-associated pathway is supported by the fertility restoration of *dex1* by OsDEX1.

OsDEX1 Affects Tapetal Function and Anther Development

The primexine functions as a template for sporopollenin deposition, which is critical for pollen development.

Several genes have been identified that influence primexine formation in Arabidopsis (Paxson-Sowders et al., 2001; Ariizumi et al., 2004; Chang et al., 2012; Sun et al., 2013), while no gene has been identified in rice. In this study, we identified the ortholog of Arabidopsis DEX1 in rice, which shares a similar function with Arabidopsis DEX1 in regulating primexine formation and male fertility (Paxson-Sowders et al., 2001). Similar to *dex1*, *osdex1* showed defects in plasma membrane undulation as well as primexine formation, resulting in the failure of exine formation (Fig. 3). Therefore, DEX1 appears to play an evolutionarily conserved role in male reproduction in model dicot Arabidopsis and monocot rice plants.

Primexine is a structure located at the periphery of the haploid microspores; however, it is believed that primexine formation is controlled by sporophytic cells. Evidence for this comes from reported primexine defective mutants, such as *dex1*, *nef*, *hkm*, and *rpg1*, whose heterozygous mutants showed normal primexine formation (Paxson-Sowders et al., 2001; Ariizumi et al., 2004; Chang et al., 2012; Sun et al., 2013). Tapetal cell death is of vital importance for exine formation (Sorensen et al., 2003; Li et al., 2006; Zhang et al., 2008; Aya et al., 2009; Xu et al., 2010; Niu et al., 2013; Ji et al., 2013). Although its role in primexine formation has not been described previously, we agree that tapetal cell death is also required for primexine formation. In a previous report of Arabidopsis DEX1 (Paxson-Sowders et al., 2001), mutant defects in microspore release from the tetrad, tapetal function as well as a biochemical characterization of DEX1 have not been reported. In this study, we show that the newly produced microspores within the *osdex1* tetrad are covered by an electron-dense matrix, and that microspore release into the lobe is inhibited until at least stage 9. Immunolabeling and aniline blue assays indicate that degradation of microspore callose is abnormal in *osdex1* (Figs. 5 and S5), which has not been reported in Arabidopsis mutants such as *dex1*, *npg1*, *hkm*, *rpg1*, *rpg2*, *npu*, and *nef* (Paxson-Sowders et al., 2001; Ariizumi

et al., 2004; Ariizumi et al., 2005; Guan et al., 2008; Chang et al., 2012, Sun et al., 2013). In addition, diffuse halos that surround *osdex1* microspores at stage 10 only contain very low levels of callose, suggesting that some other polymers are aberrantly deposited onto the *osdex1* microspores. Taken together, the weaker staining of callose by callose antibody and aniline blue at stage 8b, the less compact matrix detected by TEM observation around *osdex1* tetrads, and the persistence of callose until stages 9 and 10 suggest that callose degradation may initiate early in *osdex1*, but then fails to complete due to defects in tapetal development.

As a nutritive tissue, tapetal function such as proper cell death and the biosynthesis of sporollenin precursors is essential for pollen wall formation. Either premature or delayed tapetal degradation frequently causes reduced male sterility (Balk and Leaver, 2001; Ku et al., 2003; Lee et al., 2004; Jung et al., 2005; Luo et al., 2006; Kawanabe et al., 2006; Li et al., 2006; Oshino et al., 2007; Li et al., 2011; Aya et al., 2009; Tan et al., 2012; Niu et al., 2013; Ji et al., 2013; Fu et al., 2014; Ko et al., 2014; Yang et al., 2014; Zhang et al., 2014). We observed a persistent and partially expanded tapetal layer in *osdex1* until stage 11. However, compared to wild-type tapetal cells that are rich in organelles at stage 9, *osdex1* tapetal cells form abnormal vacuoles. The vacuoles expand and encapsulate organelles, leading to degradation of the inclusions in the tapetal cells. These observations suggest that expanding vacuoles may contribute to an alternative form of tapetal cell death in *osdex1*, distinct from normal tapetal cell death (Li et al., 2006; Niu et al., 2013). Consistent with this, the tapetal cell death executors *OsAP25*, *OsAP37*, and *OsCP1* fail to be induced in *osdex1* (Supplemental Fig. S4), suggesting that the normal protein degradation pathway involved in the tapetal cell death is not activated in *osdex1*. Furthermore, compared with the cell death-deficient mutant *tdr* whose tapetum expansion is uniform (Li et al., 2006), the expansion of different tapetal cells in *osdex1* varies (Fig. 3), suggesting a different molecular mechanism of OsDEX1 on regulating tapetal cell death compared with TDR.

Tapetal cell death is characterized by changes such as DNA fragmentation, protease activation, and cell shrinkage (Li et al., 2006; Aya et al., 2009; Xu et al., 2010; Phan et al., 2011; Li et al., 2011a; Niu et al., 2013; Fu et al., 2014; Fig. 10). Notably, DNA fragmentation is observed from stage 8b, while tapetal cell shrinkage and protease induction are observed from stage 9 in the wild-type anthers. The link between DNA fragmentation and cell death remains to be elucidated. DNA fragmentation is observed in the *osdex1* mutant (Fig. 4), but no cell shrinkage or protease induction could be observed, suggesting that OsDEX1 may act as a component required for the tapetal cell death signal transduction (Fig. 10). The early appearance of DNA fragmentation in *osdex1* could be explained by the earlier induction of endonuclease genes (Supplemental Fig. S4). How OsDEX1 regulates the expression of these genes requires further investigation.

OsDEX1 Is a Ca²⁺ Binding Protein

Ca²⁺ concentration in the cytoplasm, organelles, and cell wall is maintained within a certain range (Knight, 2000; White, 2000). Ca²⁺ binding proteins are required for the decoding of transient [Ca²⁺]_{cyt} (Enslin et al., 1995; Snedden and Fromm, 1998; Day et al., 2002; Kudla et al., 2010; Steinhorst and Kudla, 2013), and usually contain EF-hand motifs that are rich in negative amino acid for the coordination with Ca²⁺. EF-hand usually form a helix-loop-helix structure (Day et al., 2002; Derbyshire et al., 2007; Rigden et al., 2011). However, there are some β -propeller structure proteins shown to possess Ca²⁺ binding activity (Cioci et al., 2006). It has been reported that an eight-bladed β -propeller structure protein, RG lyase YesW from saprophytic *Bacillus subtilis*, had the ability for Ca²⁺ binding, despite lacking the canonical EF-hand motif (Ochiai et al., 2007). 3D-structure predictions show that OsDEX1 harbors a β -sheet-rich region linked by loops of negatively amino acids that form a pocket for Ca atom, similar to YesW (Supplemental Figs. S9 and 8). The in vitro Ca²⁺ binding assay indicates that truncated OsDEX1 has the Ca²⁺ binding activity, and the genetic complementation using the mutated OsDEX1 suggests that the Ca²⁺ binding activity is required for pollen development in vivo (Fig. 8). Furthermore, OsDEX1 is an ER-localized protein (Supplemental Fig. S10), and *osdex1* loss-of-function mutant accumulated Ca²⁺ on the plasma membrane, while overexpressing OsDEX1 prevented the Ca²⁺ accumulation on the plasma membrane (Fig. 9). These results suggest a role for OsDEX1 in affecting the level of Ca²⁺ on the plasma membrane (Fig. 10).

OsDEX1 Has a Conserved Function during Plant Male Reproduction

The pollen wall is essential for pollen grains resistant to various biotic and abiotic stresses in flowering plants. In this study, we identified and analyzed 25 DEX1-like proteins from flowering plants, lower plants, lower animals, fungi, and bacteria. Their phylogenetic relationship suggests that these genes probably share a common ancestor and have a conserved function. Notably, the proteins in flowering plants share one or two FG-GAP domains that are required for the Ca²⁺ binding (Fig. 6). The protein structure similarity in flowering plants highlights the conserved and essential role of OsDEX1 and its homologs in pollen wall formation in flowering plants. To our knowledge, OsDEX1 is the first member of the DEX1 family that has been identified as having Ca²⁺ binding activity.

In summary, we have demonstrated that the FG-GAP domain protein OsDEX1 is able to bind Ca²⁺ to modulate cellular Ca²⁺ homeostasis, and regulates tapetal degradation and pollen wall formation. A conserved role for DEX1-like proteins was established by the genetic complementation of *Arabidopsis dex1* by OsDEX1. This finding provides important insights into the role of Ca²⁺ in male reproduction in plants.

MATERIALS AND METHODS

Plant Materials, Growth Conditions, and Molecular Cloning of OsDEX1

Rice (*Oryza sativa* ssp. *japonica*, 9522) plants were grown in the paddy field of Shanghai JiaoTong University. Male sterile plants in the F2 progenies generated by a cross between wild-type GuangLuAi species (*indica*) and the *osdex1* mutant (*japonica*) were selected for mapping. Twenty-four pairs of InDel molecular markers were designed based on the polymorphism between *japonica* and *indica* for mapping. Further fine-mapping of OsDEX1 was performed using the previously published method from Li et al. (2006).

Characterization of Mutant Plant Phenotypes

Anthers from different developmental stages, as defined in Zhang et al. (2011), were collected based on the comparison and analysis of wild-type and *osdex1* plants on glume length and morphology. Observation of anther development by semithin section analysis, TEM, and callose staining were performed according to previous studies (Fu et al., 2014; Aditya et al., 2015). For callose immunolabeling, sections were first deparaffinated using xylene before labeling with anticalllose antibody (Meikle et al., 1991). Images were captured on a Zeiss A1 AxioImager using a black and white camera and ZEN2012 software. Identical exposure times were used to enable comparisons between stages and samples. Green immunostaining (Alexafluor-488) was viewed using Zeiss Filter Set 38; blue counterstaining (0.01% calcofluor white) was viewed using Zeiss filter set 49, and red staining (auto-fluorescence) was viewed using Zeiss filter set 43.

TUNEL Assay

Samples from wild-type and *osdex1* anthers at different developmental stages were collected. The TUNEL assay was performed as reported in Fu et al. (2014). The samples were analyzed under a fluorescence confocal scanner microscope (Leica SP5II system). Images were recorded using a HCX PL APO CS 20*0.7 DRY objective. The imaging parameters were as follows: image dimension (1024*1024); pinhole (2.19 airy unit); scanning speed (100 Hz); line average (3). The fluorescein signal was excited using the 488 nm laser line of the Argon laser (30%). The image parameters for this channel were as follows: 488 nm 48%; smart gain (817.0); offset (-2.5%). The propidium iodide signal was excited using the 543 nm laser line of the HeNe 543. The image parameters for this channel were as follows: 543 nm 33%; smart gain (654.0); offset (-2.5%). The overlays of fluorescein signal and propidium iodide signal were shown as the TUNEL-positive signal. All pictures were taken in the same setting.

Cloning of OsDEX1 and Plant Transformation

A 2.5 kb cDNA sequence of OsDEX1 was amplified from the cDNA reverse transcribed by the RNA extracted from anthers of 9522. A 1.7 kb upstream sequence of Arabidopsis *DEX1* was amplified from the genomic DNA of Col-0. The cDNA of mutated EF-hand motif 1 was amplified by the primers mEF1 F1 and mEF1 R1, and mEF1 F2 and mEF1 R2. The PCR products of the amplification were taken as the template for a second PCR, with the primers mEF1 F1 and mEF1 R2. The cDNA of mutated EF-hand motif 2 was amplified by the primers mEF2 F1 and mEF2 R1, and mEF2 F2 and mEF2 R2. The PCR products of the amplification were taken as the template for a second PCR, with the primers mEF2 F1 and mEF2 R2. The two fragments were ligated together by *NcoI* and *XbaI*, which existed in the primer and the pBlueScript SK, and then ligated with the 5' end of the cDNA to a full-length cDNA. Taking the full-length wild-type and mutated full-length cDNA as the templates, the amplified fragments were cloned into the binary vector pCAM-BIA1301. Plasmids were then transferred into *Agrobacterium tumefaciens* GV3101 and Arabidopsis heterozygous *dex1* plants. The transformants were screened for the presence of transgene on hygromycin medium. Over 100 positive transgenic plants in each transformation were obtained and genotyped for the homozygous *dex1* mutant background (primers used are listed in Supplemental Table S1). The PM-YC3.6 (Krebs et al., 2012) were transferred into *A. tumefaciens* EHA105. Calli induced from young panicles of the wild type 9522 and *osdex1* mutant were used for transformation (Li et al., 2006). For transgenic plants, at least three independent lines were obtained for each construct.

qRT-PCR and In Situ Hybridization

Total RNA from corresponding tissues was isolated using TRI reagent from rice tissues. Ninety milligrams of RNA was used to synthesis cDNA in each sample using the Primescript RT reagent kit with genomic DNA eraser (Takara). qRT-PCR was performed with the lightCycler system (Roche) using SYBR Premix Ex Taq GC (Takara) according to the manufacturer's instructions. Amplification was conducted following this protocol: 95°C for 2 min; 40 cycles of 95°C for 5 s; and 55°C for 30 s; 72°C for 30 s. ACTIN (Supplemental Table S1) was used as an internal control, and a relative quantization method (D cycle threshold) was used to quantify the relative expression level of target genes. Three biological replicates with three technique replicates each were included in producing the statistical analysis and the error range analysis. In situ hybridization was performed as described in Fu et al. (2014). Two OsDEX1 cDNA fragments generated by PCR were used for preparing antisense and sense probes.

Phylogenetic Analysis

Multiple alignments were performed using Muscal 3.6 (<http://www.ebi.ac.uk/Tools/msa/muscle/>). A phylogenetic tree was constructed with the aligned sequences from the full-length region of OsDEX1. MEGA (version 6.0; <http://www.megasoftware.net/index.html>) and the Neighbor-Joining methods were used with p-distance model and pairwise deletion and bootstrap (1000 replicates; random seed). The Max Parsimony method of MEGA was also employed to support the Neighbor-Joining tree, using the default parameter.

Heterologous Expression and In Vitro Ca²⁺ Binding Assay

Wild-type and EF-hands-mutated OsDEX1 cDNA was amplified by primers cDNA F and cDNA R fused with GST, expressed in BL₂₁ DE₃ *Escherichia coli* cells. Proteins were purified using amylose and G5SH resin, respectively. The protein extraction was incubated with 1 mM CaCl₂ or 1 mM CaCl₂, together with 1 mM EGTA at 4°C overnight. The mobility was detected in 10% SDS-PAGE.

Calcium Imaging

FRET-based Ca²⁺ imaging was performed as described in Krebs et al. (2012). Anthers at different stages from YC3.6 transgenic lines in wild type or *osdex1* were used for analyses. The samples were analyzed using a fluorescence confocal scanner microscope (Leica SP5II system). The objective, the imaging dimension, and scanning speed were same as above. The pinhole was 2.32 airy unit. The CFP signal was excited using the 458 nm laser line of the Argon laser (30%). The image parameters for the YFP channel were as follows: smart gain (1028.1); offset (-16.1%). The image parameters for this channel were as follows: smart gain (763.0); offset (-12.7%). The ratios between CFP and YFP signal intensity, subtracting the background fluorescence in a region outside the anthers, were calculated. For the ratio calculation, the parameters for image scaling were as follows: Min (0); Max (2).

For the Ca²⁺ imaging in tobacco cells, the epidermis cells transformed with corresponding plasmids for 2 d were observed. The objective, imaging dimension, and scanning speeds were same as above. The pinhole was 1.00 airy unit. The CFP signal was excited using the 458 nm laser line of the Argon laser (50%). The image parameters for this channel were as follows: 458 nm 33%; smart gain (761.0); offset (0.1%). The image parameters for YFP channel were as follows: smart gain (531.0); offset (-1.3%). For the statistical analysis of [Ca²⁺]_{cyt}, the ratio of fluorescence intensity in channel CFP and YFP in the whole region of the cell was measured and calculated as described (Behera et al., 2015). For the statistical analysis of [Ca²⁺]_{PM}, the ratios of fluorescence intensity in channels CFP and YFP around the plasma membrane were measured (*n* > 15 each).

Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers XP_015631235; XP_006651977; XP_008666953; XP_003563417; XP_004981208; XP_002466172; ERN096669; NP_566343; CDY67523; XP_008388127; XP_006573337; XP_006576567; XP_003604604; XP_001782562; XP_002973603; XP_001703106; CCW71639; XP_780158; XP_005093156; EKC30460; XP_009052863; WP_012257615; WP_022229450; NP_070770.

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. The model of anther development from stage 5 to stage 9.

Supplemental Figure S2. Transmission electron micrographs of the anthers from the wild type (A) and the *osdex1* mutant (B) at stage 8a. Bars = 20 μm .

Supplemental Figure S3. Expression of genes associated with sporopollenin synthesis and transport genes in the wild type and *osdex1* during pollen development. Bars = SE.

Supplemental Figure S4. Protease genes and endonuclease genes expression in *osdex1* and tapetum cell death deficient mutants.

Supplemental Figure S5. Callose degradation is abnormal in *osdex1*.

Supplemental Figure S6. Molecular identification and expression pattern of OsDEX1.

Supplemental Figure S7. Phenotype of *osdex1-2* and *osdex1-4*. A, Wild-type flower (left) and *osdex1-2* mutant flower (right) before anthesis.

Supplemental Figure S8. Alignment of OsDEX1 and DEX1.

Supplemental Figure S9. Structure prediction of OsDEX1 and DEX1.

Supplemental Figure S10. Sublocalization analysis using confocal laser scanning microscope of OsDEX1 in tobacco epidermal cell.

Supplemental Table S1. Primers used in this study.

Supplemental Data Set 1. Protein sequences used for phylogenetic tree.

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