Mutations of genes in synthesis of the carotenoid precursors of ABA lead to pre-harvest sprouting and photo-oxidation in rice

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Summary

Pre-harvest sprouting (PHS) or vivipary in cereals is an important agronomic trait that results in significant economic loss. A considerable number of mutations that cause PHS have been identified in several species. However, relatively few viviparous mutants in rice (Oryza sativa L.) have been reported. To explore the mechanism of PHS in rice, we carried out an extensive genetic screening and identified 12 PHS mutants (phs). Based on their phenotypes, these phs mutants were classified into three groups. Here we characterize in detail one of these groups, which contains mutations in genes encoding major enzymes of the carotenoid biosynthesis pathway, including phytoene desaturase (OsPDS), f-carotene desaturase (OsZDS), carotenoid isomerase (OsCRTISO) and lycopene β -cyclase (β -OsLCY), which are essential for the biosynthesis of carotenoid precursors of ABA. As expected, the amount of ABA was reduced in all four phs mutants compared with that in the wild type. Chlorophyll fluorescence analysis revealed the occurrence of photoinhibition in the photosystem and decreased capacity for eliminating excess energy by thermal dissipation. The greatly increased activities of reactive oxygen species (ROS) scavenging enzymes, and reduced photosystem (PS) II core proteins CP43, CP47 and D1 in leaves of the Oscrtiso/phs3-1mutant and OsLCY RNAi transgenic rice indicated that photo-oxidative damage occurred in PS II, consistent with the accumulation of ROS in these plants. These results suggest that the impairment of carotenoid biosynthesis causes photo-oxidation and ABA-deficiency phenotypes, of which the latter is a major factor controlling the PHS trait in rice.

Keywords: rice, pre-harvest sprouting, photo-oxidation, carotenoid biosynthesis, abscisic acid.

Introduction

The phenomenon of germination of cereal grains in the ear or panicle, usually under wet conditions shortly before

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harvest, is termed pre-harvest sprouting (PHS) or vivipary. Pre-harvest sprouting of cereal grains not only causes reduction of grain yield but also affects the quality of the grain. In Southeast Asia, PHS frequently occurs in rice due to

the long spell of rainy weather in early summer and autumn (Wan et al., 2006). In south China alone, heavy PHS sometimes occurs in >6% of the rice acreage, which could be up to 20% for hybrid rice (Guo et al., 2004).

Previous work demonstrated that mutants impaired in abscisic acid (ABA) biosynthesis or responsiveness, such as maize viviparous (vp), Arabidopsis ABA-deficient (aba) and ABA-insensitive (abi) mutants, often produce precociously germinating seeds (McCarty, 1995). At least 10 viviparous mutants have been identified from maize (Zea mays L.), most of which (vp2, vp5, vp7, vp9, w3, y3, and y9) were blocked in biosynthesis of the carotenoid precursors for de novo ABA synthesis (Figure S1; Singh et al., 2003). In the early steps of carotenoid biosynthesis, the head-to-head condensation of C₂₀ geranylgeranyl diphosphate (GGPP) molecules to produce a C_{40} carotenoid phytoene (colorless) is mediated by a soluble enzyme called phytoene synthase (PSY), which is the first committed step in carotenoid synthesis. Subsequently the phytoene undergoes four desaturation reactions with the production of lycopene (Cunningham and Gantt, 1998). Then, a series of steps including cyclization and hydroxylation reactions take place to yield a-carotene, β -carotene, lutein, xanthophyll, and zeaxanthin. The C₄₀ carotenoid precursor is cleaved and followed by a two-step conversion of the intermediate xanthoxin to ABA via ABA aldehyde (Schwartz et al., 2003; Xiong and Zhu, 2003).

The maize $Vp5$ gene was found to encode a phytoene desaturase (PDS), and transgenic rice plants harboring the PDS–RNAi construct showed a clear albino phenotype (Hable et al., 1998; Miki and Shimamoto, 2004). The maize vp9 mutant and the non dormant-1 (nd-1) mutant of sunflower (Helianthus annuus L.) have mutations in the gene coding for zeta-carotene desaturase (ZDS; Conti et al., 2004; Matthews et al., 2003). The carotenoid isomerase (CRTISO) gene was cloned from the tangerine mutant of tomato and ccr2 mutant of Arabidopsis (Isaacson et al., 2002; Park et al., 2002). The Vp7/Ps1 gene encodes a lycopene-b-cyclase and is necessary for the accumulation of both ABA and carotenoid zeaxanthin in mature maize embryos; the mutant is easily discernible as it has pink kernels because of lycopene accumulation (Singh et al., 2003). These mutants containing defects in carotenoid precursor synthesis exhibit pleiotropic phenotypes, such as albino or pale green, non-viable seedlings and vivipary, due to deficiencies of carotenoid and ABA.

Within the thylakoid membranes of chloroplasts, carotenoids are found to be bound to specific protein complexes of photosystem I (PS I) and photosystem II (PS II), where they augment the light-harvesting capacity by absorbing light in the blue–green range of the visible spectrum (450–550 nm) and transferring the energy to chlorophylls (Holt et al., 2005; Moise et al., 2005). Carotenoid deficiency often causes aberrations in plastid ultrastructure, such as etioplasts from the cotyledon of dark-grown ccr2 mutants (carotenoid

isomerase, CRTISO mutation) which lack prolamellar bodies (PLBs; Park et al., 2002). In addition, carotenoids play an essential role in photoprotection in plants. During photosynthesis, the excess absorbed energy can be eliminated as heat by de-exciting ¹Chl through the process of non-photochemical quenching of chlorophyll fluorescence (NPQ), minimizing the generation of harmful reactive oxygen species (ROS; Ma et al., 2003; Niyogi, 1999; Tracewell et al., 2001).

Besides, as accessory pigments in photosynthesis and photoprotectors preventing photo-oxidative damage, carotenoids can also be the precursors to the hormone ABA. Analysis of mutant and transgenic plants has provided strong evidence that ABA biosynthesis and responses to this phytohormone are clearly involved in the onset and maintenance of dormancy and inhibition of PHS (Bewley, 1997). Abscisic acid biosynthetic mutants fail to induce seed dormancy and exhibit a vegetative wilty phenotype. These phenotypes have been used to define and prove the function of ABA in seed dormancy and water relations. The seeds of ABA-deficient mutants of tomato (Solanum lycopersicum L., e.g. sitiens), Arabidopsis (e.g. aba) and maize treated with fluridone (a carotenoid biosynthesis inhibitor) germinate readily when placed in water and sometimes show vivipary (Fong et al., 1983; Groot and Karssen, 1992; Leon-Kloosterziel et al., 1996a).

Another phytohormone, gibberellin (GA), is also involved in the release from dormancy of various species. It has been shown that GA-deficient mutants of Arabidopsis and tomato are dependent on exogenous GA for germination (Koornneef et al., 2002). Further work in maize showed that inhibition of GA biosynthesis mimics the effect of exogenous ABA in suppressing vivipary (White and Rivin, 2000; White et al., 2000). These results suggested that the ABA/GA ratio and not the absolute hormone content controls germination (Finch-Savage and Leubner-Metzger, 2006).

In contrast to the intensive molecular and genetic studies of seed dormancy in maize and Arabidopsis, the molecular mechanism of seed dormancy in rice is poorly understood, mainly because of the lack of available mutants with reduced dormancy. So far most research has focused on the quantitative trait locus (QTL) analysis of the natural differences in seed dormancy characteristics (Gonzalez-Guzman et al., 2004; Gu et al., 2004, 2006; Takeuchi et al., 2003; Wan et al., 2006). Only a few reports have been published on phs mutants and only one gene (OsABA1) has been cloned which was involved in ABA biosynthesis (Agrawal et al., 2001).We performed a large-scale screening of a rice T-DNA/ Tos17 insertion mutant population to identify the genes involved in PHS. The warm and damp weather, a good elicitor of PHS, during the harvest season in Zhejiang Province on the southeast coast of China, enables us to isolate phs mutants efficiently. In an intensive screening of approximately 16 000 rice T_1 mutant lines, we obtained 12 viviparous mutants. In this paper, four genes involved in carotenoid precursors of ABA biosynthesis were cloned. Our results suggested that the impairment in synthesis of the carotenoid precursors of ABA leads to photo-oxidation and PHS in rice, which will definitely be helpful for elucidating the molecular mechanisms of PHS in other crops such as wheat and barley that are susceptible to PHS.

Results

Identification and genetic analysis of the rice phs mutants

To identify rice phs mutants, we have screened a T-DNA/ Tos17-mutagenized population (Nipponbare background) under field conditions in Hangzhou, downstream of the Yangtze River with a relatively high degree of humidity. Approximately 16 000 transgenic T_1 lines were screened prior to harvest by visual inspection in the paddy field, and 27 putative mutants were identified with a viviparous phenotype. A representative mutant is shown in Figure S2(a). T_2 seeds of these 27 putative *phs* mutants were then grown in Beijing with a lower degree of humidity for a second round of screening. From the secondary screening, 12 mutants showing a viviparous phenotype were recovered (Figure S2b), which could be simply categorized into three groups based on phenotypes besides vivipary. Mutants from category I exhibit an albino or photobleaching phenotype (Figure S2c); while mutants from categories II and III do not show an albino or photobleaching phenotype but have an enhanced wilty phenotype under conditions of water stress (category II; Figure S2d) or with embryo/seedlinglethal phenotypes (category III; Figure S2e). We present here a detailed characterization of six mutants which belong to category I (Table 1).

To further characterize these mutants the viviparous seedlings were rescued, and five viviparous mutant lines showed albino seedlings (Figure 1a), these homozygous plants eventually died at about 4 weeks after germination. Interestingly, the homozygous seeds of two mutant lines among them had pink embryos (Figure 1b). Moreover, one of the viviparous mutant lines, T09, developed alternating green and yellow crossbands on the leaf blades at the tillering stage (Figure 1c), like rice zebra mutants previously described (Kusumi et al., 2000), and finally showed pale green on entire plants when mature (Figure 1d).

Except for the T09 mutant, all the other viviparous mutants (T_1 plants) are lethal due to lack of pigments, therefore only two genotypes – heterozygous and wild type in the T_1 mutant seeds – could germinate after sowing. Statistical analysis of segregation ratios of viviparous and non-viviparous plants demonstrated that the segregation of viviparous and non-viviparous plants is in line with the expected 2:1 ratio (Table S2). Furthermore, in $T₂$ progeny (seeds in the ear) obtained from the viviparous plants, the mutant phenotype segregated in 3:1 (Wt: vivipary, data not shown), indicating that each mutant phenotype was caused by a single recessive mutation. Subsequently genetic analysis suggested that these six mutants could be assigned to four loci and were consecutively designated as phs1 through phs4, respectively, of which two mutant alleles were identified in phs2 (phs2-1 and phs2-2) and phs4 (phs4-1 and phs4-2; see Table 1). In addition, we later identified another two phs3 alleles by the zebra phenotype (see below).

Molecular cloning of PHS1 through PHS4 genes

Genetic analysis suggested that T-DNA was not co-segregated with the mutant phenotype in all these mutants (data not shown). We then made a rough mapping to identify candidate genes in these mutants, and the PHS genes were located to different chromosomes (Table 1 and Figure S5).

Because only mutants specifically blocked in the carotenoid biosynthetic pathways can manifest as a combination of vivipary with albino phenotype (Wurtzel et al., 2001), we presumed that these mutants may carry lesions in key genes for biosynthesis of carotenoids or ABA. So the predicted amino acid sequences of ABA biosynthesis genes from Arabidopsis were used as probes to screen in silico all available rice databases. In these four regions where mutations phs1 through phs4 mapped, four candidate genes appearing to be orthologs of Arabidopsis genes were found in the carotenoid biosynthesis pathway (Table 1), including a PDS-like gene (phs1), a ZDS-like gene (phs2-1 and phs2-2), a carotenoid isomerase-like gene ($phs3-1$) and a β -LCY-like gene (phs4-1 and phs4-2). To examine whether these candidate genes carried mutations in the respective

Table 1 Rice pre-harvest sprouting mutants and corresponding genes

Category	Original line	Gene	Arabidopsis homolog	Mutation	Chromosome location	Protein (a.a.)	Accession no.
	T01	<i>OsPDS</i>	AT4G14210	Insertion	Chr.3, 20.3cM	566	LOC 0s030g08570
	HF807 HG4123	<i>OsZDS</i>	AT3G04870	Insertion Deletion	Chr.7, 41.7cM	578	AK065213
	T09	OsCRTISO [*]	AT1G06820	Substitution	Chr.11, 84.6cM	586	EF417892
	HC2621 HD1449	B-OsLYC	AT3G10230	Deletion Deletion	Chr.2, 23.3cM	493	OC 0s02g09750

* The nucleotide sequence of OsCRTISO reported in this paper has been submitted to GenBank under accession number EF417892.

Figure 1. The phenotypes of pre-harvest sprouting (phs) mutants in category I.

(a) The albino phenotype of phs seedlings.

(b) The pink-embryo seeds of phs4-1 and phs4-2 lines.

(c) Phenotypes of the wild type (left) and phs3-1 mutant (right) at the early tillering stage. The inset shows the magnified leaves of the wild type and phs3-1 mutant, respectively.

(d) The phs3-1 plant shows pale green over the whole leaves at the mature stage in the field.

mutants, we analyzed the corresponding candidate genes by polymerase chain reaction (PCR) and sequencing; the results revealed that a Tos17 was inserted in intron 4 of the putative OsPDS in phs1 (Figure 2a). As expected, no OsPDS expression was detected by reverse transcription (RT)-PCR analysis in phs1 (Figure 2b). The phs2-1 carried a Tos17 insertion in exon 3 of OsZDS, whereas phs2-2 had a 1.7-kb deletion started from intron 3 spanned to exon 7 (Figure 2c). No OsZDS expression was detectable in either mutant allele (Figure 2d).

In phs3-1, a G-to-C transition was identified at position 1995 (the putative translation start codon is referred to as $+1$) of the OsCRTISO gene, which is predicted to be the donor site of intron 6 (Figure 2e). This mutation presumably causes the utilization of a novel splicing donor-like site (GT) 24 bp upstream from the mutated site. Consistent with this speculation, RT-PCR analysis revealed the transcript in phs3-1 was slightly shorter than that in the wild type (Figure 2f), which was confirmed by direct DNA sequencing. In addition, another two phs3 alleles were collected by the zebra phenotype. Sequence analysis revealed that mutations of various types were identified in the candidate gene CRTISO. A Tos17 retrotransposon inserted in exon 7 of phs3-2 (NG0489 from the Rice Tos17 Insertion Mutant database http://tos.nias.affrc.go.jp/~miyao/pub/tos17/index.html.en) completely disrupts the function of CRTISO (Figure 2f). phs3-3 has a 30-bp deletion in exon 6. The deletions in phs3-1 and phs3-3 do not cause a frame-shift in the open reading frame (ORF).

The rice β -OsLCY gene has no intron, and sequence analysis of phs4-1 mutant showed a 1-bp deletion at the 1314 bp starting from ATG, and the *phs4-2* allele has a 25-bp deletion in 473–497 bp (the putative translation start codon is referred to as +1). The deletions caused a frame-shift and subsequently created a new stop codon that might result in immature translation (Figure 2g) and disrupt the normal function of the β -LCY gene. Expression analysis also showed that the β -OsLCY transcript was barely detectable in phs4 alleles compared with the wild type (Figure 2h).

To confirm the identity of these candidate genes, we also performed genetic complementation experiments. Since homozygous mutant seedlings from three genes (OsPDS, $OsZDS$, and β -OsLCY) are non-viable the homozygous seeds were screened out simply from the heterozygous lines at an early stage of callus induction using mature embryos. The embryos of the homozygous mutant initiated an albino bud after 2 weeks' culture on Murashige and Skoog medium (Murashige and Skoog, 1962) with addition of 2 mg I^{-1} 2,4dichlorophenoxyacetic acid (2,4-D) in the light, and their calli derived from homozygous seeds regenerated the albino shoot during later regeneration (Figure 3a). Genomic DNA fragments covering the respective 5'-upstream regions, entire genes and 3¢-downstream regions of respective candidate genes were introduced into corresponding mutants by Agrobacterium-mediated transformation. In all cases, the mutant phenotypes were rescued when genomic fragments containing the candidate genes were introduced (Figure 3b,c, and Table S3). In contrast, the transgenic plants containing the empty vectors failed to rescue the *phs* mutants (data not shown).

Carotenoid profiles of phs mutants

To analyze carotenoid metabolism in the phs mutants, highperformance liquid chromatograph (HPLC) analysis was performed on the extracts prepared from light-grown wildtype and mutant seedlings. As shown in Figure 4a, carotenoids were almost undetectable in phs1 (Ospds mutants) and phs2-1 or phs2-2 (Oszds mutants) seedlings at 460 nm. However, lycopene accumulated in phs4-1or phs4-2 (B-Oslcy mutants). Carotenoids, especially lutein, were dramatically reduced in light-grown phs3-1 (Oscrtiso mutants) and an alltrans lycopene precursor, prolycopene, was accumulated in dark-grown phs3-1 seedlings (data not shown). The Ospds mutant contained a peak at 287 nm with a retention time of 25.54 min, which was absent in the wild type (Figure 4b). The derived spectrum of the peak showed characteristics of

Figure 2. The sites of mutation and the expression of the pre-harvest sprouting (PHS) genes. (a) The position of the mutation in the OsPDS gene of phs1 mutant.

(b) Expression of the OsPDS gene in the wild type and mutant.

(c) The positions of the mutation in the OsZDS gene of phs2-1 and phs2-2 mutants.

(d) Expression of the OsZDS gene in the wild type and mutants.

(e) The positions of the mutation in the OsCRTI-SO gene of phs3-1, phs3-2/NG0489 and phs3-3 mutants.

(f) Expression of the OsCRTISO gene in wild type and mutants.

(g) The positions of the mutation in the β -OsLCY gene of phs4-1 and phs4-2 mutants.

(h) Expression of the β -OsLCY gene in the wild type and mutant.

Arrowheads indicate the insertion sites of a rice retrotransposon, Tos17. Boxes and lines indicate exons and introns, respectively.

phytoene with three main peaks (data not shown). In phs2 seedlings only small peaks of ξ -carotene and its isomers were detected at 430 nm, which were absent in the wild type (Figure 4c).

These results were consistent with the mutated genes in the biosynthetic pathway of carotenoids, and carotenoid biosynthesis is indeed impaired in these mutants. The albino phenotypes of the phs mutants (Ospds, Oszds, and β -Oslcy) were due to chlorophyll and carotenoid deficiency. The pink embryo in phs4 suggested lycopene accumulation, and the phenotype of phs3 (Oscrtiso mutant) seemed to result from photobleaching.

Cellular characters of the phs mutants

Carotenoids are essential components of photosynthetic systems, which occur in thylakoid membranes in chloroplasts in association with chlorophyll–protein complexes

(Green and Durnford, 1996; Moise et al., 2005). To assess the effect of the PHS mutations on chloroplast development, the leaves of 2-week-old seedlings of all phs mutants mentioned above were examined by transmission electron microscopy. In wild types, chloroplasts were well-developed and organized in the mesophyll cells (Figure 5a), in contrast, no plastid- or chloroplast-like structures were found in the mesophyll cells of albino mutants (phs1, phs2, and phs4; Figure 5b–d). In the mesophyll cells of Oscrtiso mutants, irregularly shaped and abnormally developed chloroplasts were observed, with reduced and irregularly organized thylakoid membranes (Figure 5e,f).

OsLCY-RNAi plants and phs mutants suffered from photo-oxidative damages

Due to the photoprotective function of carotenoids, plants deprived of these pigments suffered from photobleaching

Figure 3. Complementation test.

(a) Plant regeneration from calli derived from embryos of the heterozygous phs4 or wild-type seeds (left) and the homozygous phs4 seeds (right). (b) The calli derived from embryos of homozygous phs4 seeds were used for transformation.

(c) The phenotypes of panicles in mature plants of wt (left), heterozygous mutant (middle) and complementation transgenic plant (right).

Figure 4. HPLC chromatograms of the wild type and homozygous phs mutants grown in light.

Chl a, chlorophyll a; Chl b, chlorophyll b.

damage, especially under high light conditions (Niyogi, 1999; Sagar et al., 1988). Since the homozygous mutant seedlings from three genes (OsPDS, OsZDS, and β -OsLCY)

Figure 5. Mesophyll cells and chloroplasts from the *phs* mutants and wild type.

Wild type (a), Ospds (b), Oszds (c), β-Oslcy (d), Oscrtiso (e); the chloroplast from wild type (left) and Oscrtiso mutant (right; f).

CP, chloroplast; CW, cell wall; Thy, thylakoid. Scale bar = $1 \mu m$.

are nonviable, we generated transgenic rice plants harboring the β -OsLCY-RNAi construct. It was demonstrated that β -OsLCY-RNAi plants and Oscrtiso mutants showed a distinctly photobleached phenotype on leaves (Figure 6a,b), which was in accordance with the accumulation of ROS as revealed by nitro blue tetrazolium (NBT) staining (Figure 6c,d). Furthermore, compared with that in the wild type, higher activities of ROS-scavenging enzymes were found in both β -OsLCY-RNAi plant and phs3-1 mutant (Figure S3), suggesting that these two plants were under oxidative stress.

The maximal efficiency of PS II photochemistry (F_v/F_m) is often considered to be an indicator of PS II function (Demmig-Adams and Adams, 1992; Krause and Weis, 1991). Therefore, we further investigated PS II photochemistry in $phs3-1/O$ scrtiso mutants and β -OsLCY-RNAi plants, as well as the wild type. The F_v/F_m ratios in wild-type plants are above 0.8. However, phs3-1/Oscrtiso mutant and β -OsLCY-RNAi plants show a decrease in the F_v/F_m ratio, which is about 0.65 (Figure 7a,b).

The $phs3-1/O$ scrtiso mutants and β -OsLCY-RNAi plants also show different responses of NPQ to increasing photosynthetic photon flux density (PPFD). When PPFD was $<$ 500 µmol m⁻² sec⁻¹, the *phs3-1/Oscrtiso* mutants and β -OsLCY-RNAi plants showed almost the same value of NPQ in comparison with the wild type. With further increase in PPFD, NPQ increased more rapidly in wild-type plants than in the Oscrtiso/phs3-1mutants and β -OsLCY-RNAi plants, leading to much higher values of NPQ than for the latter two plants (Figure 7c,d).

⁽a) HPLC chromatograms recorded at 460 nm (most carotenoids). (b) HPLC chromatograms recorded at 287 nm (phytoene).

⁽c) Chromatograms recorded at 430 nm (č-carotene).

Pre-harvest sprouting in rice 183

Figure 6. Photobleaching phenotype (a, b) and the detection of ROS (c, d) of O scrtiso and β -LCY-RNAi plants and the corresponding wild types.

To get further evidence for the photo-oxidative damage occurring in PS II, we examined the levels of essential PS II core proteins CP43, CP47 and D1 (Green and Durnford, 1996), which are the targets of ROS (Niyogi, 1999), in the leaves from the $phs3-1$ mutant and β -OsLCY-RNAi plants, respectively. Western analysis showed that the level of these proteins in the $phs3-1$ mutant and β -OsLCY-RNAi plants decreased to 64–77% of that in the wild type, indicating the occurrence of photo-oxidative damage in PS II core proteins (Figure 8).

ABA and GA synthesis are altered in phs mutants

To investigate whether the reduced level of carotenoids in phs mutants causes a decreased level of ABA we measured the ABA content in the respective *phs* mutants by immunoassay. As shown in Figure 9a, the amount of ABA was reduced in all four mutants, but much more significantly in Ospds, Oszds, and Oslcy mutants compared with that in the wild type. Since ABA is also directly involved in the regulation of stomatal aperture, we examined the water loss characteristics of phs3-1/Oscrtiso plants. Wild-type and mutant plants were grown in the soil under a normal irrigation regime. The water loss analysis was carried out at the tillering stage, and the results revealed a water loss of about 53% of fresh weight within 120 min in *phs* mutants, whereas wild-type leaves showed only a 38% loss, indicating that the rate of water loss in *phs* mutants was faster than in wild-type plants (Figure 9b).

To test whether phs mutant can sense or respond to ABA, we cultured seeds [30 days after pollination (DAP)] from phs3-1 mutant and the wild type in water with or without ABA (of different concentrations) treatment (Figure S4). In all treatments, the *phs3-1* mutant and the wild type grew with similar kinetics, indicating that the *phs3-1* mutant has a normal response to ABA.

Abscisic acid and another important phytohormone, GA, which can directly antagonize ABA signaling during induction of dormancy, are two important regulators of seed dormancy. Therefore, we measured the ABA and GA content in homozygous *phs4-1*/ β -Oslcy seeds 30 DAP (*phs4* homozygous seeds were easily discernible by their pink embryo). As expected, the ABA level in $phs4-1/B-Oslcy$ was lower than

Figure 7. The maximal efficiency of PS II photochemistry $(F_v/F_m; a, b)$ and non-photochemical quenching of chlorophyll fluorescence (NPQ; c, d) of $phs3$ -1mutant and β -OsLCY-RNAi plants. Standard deviations were obtained from six measurements.

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Figure 8. Western analysis of the PS II core proteins CP47, CP43 and D1.

Figure 9. ABA content and water loss assay of phs mutants and wild types. (a) ABA content in wild-type and mutant seedlings. (b) Water loss assays for the leaves of the wild type and Oscrtiso mutant (T09/ phs3-1) were performed within 120 min. Standard deviations were obtained from five measurements.

that in the wild type; however, GA in $phs4-1/B-Oslcy$ is higher than that in the wild type (Figure 10a). We further examined the expression of two ABA-regulated genes, rab16B and TRAB1, as well as a GA-induced gene, the α -amylase gene RAmy1A (AK073487), which were involved in embryo development and seed maturation (Hobo et al., 1999; Mitsui et al., 1996; O'Neill et al., 1990; Ono et al., 1996). The results demonstrated that the transcripts of rab16B and TRAB1 were significantly reduced in $phs4-1/\beta$ -Oslcy seeds while the transcript of RAmy1A was increased (Figure 10b), suggesting that the ABA/GA ratio in $phs4-1/\beta-Oslcy$ seeds is probably a switch for PHS.

Discussion

Analysis of phs mutants in rice

Under normal circumstances, germination in the ear of developing cereal grain is prevented by physiological mechanisms that include dormancy. Only under exceptional circumstances (e.g. rain late in grain maturation) may PHS occur. Since dormancy is a complex trait which is controlled

(b) Expression of rab16B (LEA protein gene), TRAB1 (a bZIP factor gene) and RAmy1A (α -amylase gene) genes in wild-type and β -Oslcy seeds at 30 DAP.

by a large number of genes, not only with strong intergenic interactions but also strong interactions between genes and the environment. Thus, QTL analysis has often been the tool of choice for the dissection of dormancy and germination, and a large number of the QTLs controlling seed dormancy and germination have been identified in Arabidopsis, rice, wheat, and barley (Clerkx et al., 2004; Li et al., 2004; Prada et al., 2004). Comparative genomics approaches were also used to identify candidate gene(s) controlling dormancy and PHS based on the availability of the rice genome sequence, a large number of barley and wheat ESTs, and also the high co-linearity among rice, wheat, and barley (Li et al., 2004; Wilkinson et al., 2002).

Since PHS is a complex trait controlled by both environmental and genetic factors, we carried out intensive screening of the rice mutant population in south China, where long spells of rainy weather often occur in early summer and autumn and successfully isolated 27 rice phs mutants in the paddy fields.

In this study, we focused on *phs* mutants which belong to category I; using both map-based cloning and comparative genetics approaches, we have identified that the mutated genes in category I mutants encode major enzymes (OsPDS, OsZDS, OsCRTISO, and β -OsLCY) in the carotenoid biosynthesis pathway. The candidate genes were all confirmed via functional complementation. Further work with these rice *phs* mutants will provide novel insights into the role of these PHS genes in dormancy and germination.

Carotenoid metabolism, plastid development and photo-oxidation in phs mutants

In plant chloroplasts, carotenoids serve important roles in the photosynthetic apparatus as structural components and photosynthetic and photoprotective pigments (Green and Durnford, 1996; Niyogi, 1999; Niyogi et al., 2001). Just as expected, according to the enzymatic steps blocked, phs1 and $phs2$ accumulated phytoene and ξ -carotene in the lightgrown seedlings, respectively. The results are in accordance with maize vp5 and vp9 mutants with mutations in PDS and ZDS (Hable et al., 1998; Matthews et al., 2003). In case of $phs4$, the mutation in lycopene β -cyclase leads to high lycopene accumulation in both seedlings and embryos from homozygous phs4 seeds. The accumulation of lycopene may be explained by the presumption that lycopene $\epsilon\text{-c}$ yclase and lycopene b-cyclase constitute an enzyme complex in the thylakoid membranes of plant chloroplasts; in the absence of b-cyclase, the complex is destabilized, resulting in the loss of both ϵ - and β -cyclase activity and so lycopene accumulates (Cunningham and Gantt, 1998). The results are consistent with 2-(4-chlorophenylthio)-triethylamine hydrochloride (CPTA)-treated barley seedlings (CPTA is a specific inhibitor of the cyclization of lycopene, especially lycopene b-cyclase; La Rocca et al., 2007). In addition, PSY mRNA is virtually absent in rice endosperm, which does not provide the substrate for these downstream enzymes and is consequently unable to form downstream products (Schaub et al., 2005). Therefore, the phs4 mutant accumulates lycopene (showing a pink color) in the embryo but not in the endosperm.

Unlike phs1, phs2, and phs4 mutants, phs3 mutant show a non-lethal and 'variegated' phenotype. The mutation in the PHS3 gene results in a dramatic reduction of lutein in lightgrown leaves and accumulation of prolycopene in etiolated seedlings. This result was in agreement with other crtiso mutants from tomato and Arabidopsis (Isaacson et al., 2002; Park et al., 2002). Interestingly, the CRTISO activity could partially be substituted by light, i.e. photoisomerization, which may account for the survival of the *phs3* mutant (Isaacson et al., 2002; Park et al., 2002).

Since carotenoids are located in the photosynthetic membrane in the form of chlorophyll–carotenoid–protein complexes and some carotenogenic enzymes are membrane-associated (Cunningham and Gantt, 1998; Green and Durnford, 1996), changes in carotenoid composition or the carotenogenic enzyme itself always lead to abnormal plastid development (Park et al., 2002; Welsch et al., 2000). It has been demonstrated that the location of PSY protein influenced the formation of PLBs (Welsch et al., 2000), CRTISO is required in PS II assembly in cyanobacteria, and loss of CRTISO activity results in the absence of PLB in the ccr2 Arabidopsis mutant (Masamoto et al., 2004; Park et al., 2002). All the phs mutants (phs1 through phs4) showed dramatic changes in carotenoid composition and had abnormal chloroplasts or even none at all, suggesting the importance of these enzymes for plastid development.

Carotenoids could also act as antioxidants to quench excessive free radicals and ROS generated from photooxidation (Hirayama et al., 1994; Niyogi, 1999; Niyogi et al., 1998; Woodall et al., 1997). The increasing activities of ROS-scavenging enzymes in β -OsLCY RNAi plants and

Oscrtiso mutants suggested that these two plants suffered from oxidative damage. Non-photochemical quenching of chlorophyll fluorescence is one of the major indicators which are closely associated with the onset of harmless dissipation of the excess energy present in the pigment of light-harvesting complexes as heat (Gilmore, 1997; Niyogi, 1999). The significant decrease of NPQ in β -OsLCY RNAi plants and Oscrtiso mutants indicated that excessive absorbed light could not be efficiently dissipated as heat and the excessive energy might result in ROS generation. Nitro blue tetrazolium staining further confirmed that considerable superoxide accumulated in both β -OsLCY RNAi plants and Oscrtiso mutants. The value of F_v/F_m is normally in the range of 0.8–0.85 in healthy leaves independent of plant species, and a lower value indicated that a proportion of PS II reaction centers were damaged (Demmig-Adams and Adams, 1992). The decreased F_v/F_m indicated that photoinhibition occurred in the β -OsLCY RNAi plants and Oscrtiso mutants under normal conditions. Moreover, the decrease of PS II core proteins such as CP43, CP47, and D1 in b-OsLCY RNAi plants and Oscrtiso mutants may provide further evidence for the occurrence of photo-oxidative damage in PS II. Taken together, these results further confirm that the investigated PHS genes are involved in not only synthesis of the carotenoid precursors of ABA, but also chloroplast development and photoprotection. However, in the dark or in weak light the mechanism by which carotenoid deficiency influences plastid development needs to be further elucidated.

ABA and pre-harvest sprouting in phs mutants

Seed dormancy and germination are regulated by a wide range of plant hormones, including ABA, ethylene, GA, and brassinosteroids, of which ABA is the primary mediator of seed dormancy (Koornneef et al., 2002). Abscisic acid-deficient or ABA-insensitive Arabidopsis mutants show reduced seed maturation and dormancy (Finch-Savage and Leubner-Metzger, 2006; Koornneef et al., 2002; Leon-Kloosterziel et al., 1996b). Unlike in Arabidopsis, in cereal plants such as maize embryos from ABA-deficient mutants germinate precociously on the ear. However, the absolute level of ABA in the seeds does not always perfectly correlate with the dormancy and germination events, suggesting that other modulating factors are also involved (White et al., 2000). It has been presumed that dormancy released by after-ripening and stratification caused a switch to ABA catabolism resulting in a decrease in ABA content in the embryo and a corresponding increase in inactive ABA metabolites (Gubler et al., 2005).

The amount of GA also plays an important role in controlling seed dormancy and germination. Previous studies revealed that paclobutrazol treatment of the ear suppresses vivipary in ABA-deficient seeds (White and Rivin,

2000). Here, we have demonstrated that the impaired biosynthesis of the carotenoid led to significant reduction of ABA content in the *phs* mutants. However, these mutants have a normal response to ABA. Interestingly, the amount of GA in phs4-1 seeds was significantly increased; the increased GA might result from a reduced flux of GGPP to carotenogenesis since GGPP is the common precursor for both GA and carotenoid biosynthesis (Rodriguez-Concepcion et al., 2001). It has been demonstrated that the level of GA in PSY-overexpressing plants was reduced due to increasing utilization of GGPP for carotenoid biosynthesis, leading to a dwarf phenotype (Fray et al., 1995). In addition, the ratio of ABA/GA is distinctly reduced in phs4-1 seeds, and the genes involved in seed development were differentially expressed in the seeds of the wild type and mutants. These results suggested that the ABA/GA ratio might play an important role in controlling PHS. However, the contribution of carotenoid synthesis to the regulation of balance between ABA and GA is still elusive.

It was found that some common factors regulated ABA and GA biosynthesis during seed development in Arabidopsis in an opposite manner (Gazzarrini et al., 2004). The viviparous mutants of rice are ideal for elucidating the complex mechanism of germination and dormancy, and a detailed comparison on differential expression patterns of genes between different kinds of phs mutants and wild type by microarray or other technologies will finally help us to identify the major factors controlling PHS and gain more insight into the physiological functions of carotenoids and the molecular mechanism of PHS.

Experimental procedures

Plant materials and growth conditions

Approximately 160 000 rice T-DNA/Tos17 mutant lines (Nipponbare background) were sown in the field in Hangzhou, Zhejiang Province in 2003. The panicles of mature plants were observed in late September to detect the germinated seeds. In 2004, the screened phs mutants (T_2) were sown in Beijing for further screening. Segregation ratios of viviparous and non-viviparous plants were investigated in the T_2 plants. The other two $phs3$ alleles were obtained as follows: phs3-2 was obtained from TOS17 retrotransposon insertion lines (http://tos.nias.affrc.go.jp/~miyao/pub/tos17/phenotype/ 17-NG.html, line number NG 0489) and phs3-3 was derived from a spontaneous mutation.

Map-based cloning of PHS genes

To map the mutated genes, linkage analyses were performed using an F_2 population derived from the cross between mutants (Nipponbare, Japonica) and TN1 or Minghui 63 (Indica) varieties. The simple-sequence repeat (SSR) and sequence-tagged-site (STS) markers used to analyze the polymorphisms between Nipponbare and TN1 (or Minghui 63) were obtained from the DNA bank at the National Institute of Agrobiologica Sciences (http://www.nias. affrc.go.jp).

Sequence analysis

The protein sequences of ABA biosynthetic enzymes from Arabidopsis (http://www.arabidopsis.org/) were compared with the rice database of TIGR Rice Databases (http://www.tigr.org), the Rice Genome Research Program (http://rgp.dna.affrc.go.jp/), the Beijing Genomics Institute (http://btn.genomics.org.cn/rice), and the Knowledge-based Oryza Molecular biological Encyclopedia (KOME; http://cdna01.dna.affrc.go.jp/cDNA/) for searching the rice orthologs.

Construction of complementation and RNAi vectors and plant transformation

A 6.5-kb DNA fragment containing a full-length genomic β -OsLCY gene was obtained by digesting the bacterial artificial chromosome (BAC) clone OSJNBb0031B09 with Pstl and BamHI; the fragment was inserted into a binary vector pCAMBIA2300. Similarly, the complementation vectors for other phs mutants were also constructed, using corresponding BAC clones (Table S3). In addition, empty vectors were transformed into the corresponding phs mutant as the respective controls. For β -OsLCY RNAi construct, an 856-bp fragment from 630 to 1486 bp of the β -OsLCY ORF was inserted as a BamHI/Sall fragment in sense orientation downstream of the potato (Solanum tuberosum L.) GA20 oxidase intron into pUC-RNAi (Luo et al., 2006). The same fragment was inserted in antisense orientation into the Bg/II/Xhol sites of pUC-RNAi already carrying the sense fragment. Subsequently, the fragment comprising sense and antisense fragments of β -OsLCY interspersed by potato GA20 oxidase intron was excised from pUC-RNAi using the flanking Pstl and inserted into a pXQAct plasmid between rice *actin1* promoter and Ocs terminator, yielding the binary construct. The rice transformation was performed as described (Liu et al., 2007).

Carotenoid analysis

The extraction and analysis of carotenoids was performed as previously described (Fraser et al., 2000). Briefly, a reverse-phase C30, 5 µm column (250 \times 4.6 mm) coupled to a 20 \times 4.6 mm C30 guard (YMC Inc.; http://www.ymc.co.jp) with a methanol/tert-methyl butyl ether-based mobile phase were used with a HPLC 10Avp system (Shimadzu, http://www.shimadzu.com/). Throughout chromatography, the elution was monitored continuously from 200 to 600 nm by an online Shimadzu SPD-10Avp PDA detector. Carotenoids were identified by their characteristic absorption spectra, typical retention time, and comparison with authentic standards.

Transmission electron microscopic analysis

Rice leaves were cut into 1 mm squares, fixed in 2.5% (v/v) glutaraldehyde in PBS (pH 7.2) for 24 h and post-fixed in 2% OsO₄ in PBS (pH 7.2) for 2 h. Following ethanol series dehydration, samples were embedded in Epon 812 (Shell Chemicals; http:// www.shellchemicals.com) and polymerized for 24 h at 60°C. Ultrathin sections (50–70 nm) were double stained with uranyl acetate and lead citrate and examined with a transmission electron microscope (TEM; FEI Tecnai 20; http://www.fei.com) at 120 kV.

Histochemical detection of O_2^-

Superoxide accumulation in rice leaves was visualized by 0.1% NBT staining as described (Fitzgerald et al., 2004). Rice leaves were

Chlorophyll fluorescence measurements

Chlorophyll fluorescence measurements were carried out with attached leaves in the greenhouse using a PAM 2100 portable fluorometer (Walz, http://www.walz.com/) as described (Lu et al., 2001).

Determination of ABA, GA, and water-loss assay

Determination of ABA and GA with 2-week-old seedlings was performed as previously described (Agrawal et al., 2001; Luo et al., 2006). For the water-loss assay, plants were grown in a paddy field and the detached leaves (300 mg from 10 seedlings) of 1-month-old seedlings were kept on aluminum foil at room temperature (25°C). The weight of leaves was taken at 10-min intervals until 60 min or longer. Each experiment was performed with five replicates.

Germination and ABA response assay

Seeds (30 days after pollination) from phs mutants, the wild type and complementation transgenic plants were sampled for germination ratio and ABA response assay. Seeds were cultured in a growth chamber at 30°C with a 16-h/8-h cycle for 3 days for the germination assay and 11 days for the ABA response assay. Shoots from seeds without ABA treatment were measured regularly, while shoots from seeds treated with different concentrations of ABA were measured at the 11th day after growth.

Expression analysis

Total RNAs were isolated from various organs of rice as described (Luo et al., 2006). The RT-PCR was performed with DNase-treated total RNAs using the RT-for-PCR Kit (Promega, http://www. promega.com/). The primers used for each gene are listed in Table S1. The products of PCR amplification were separated by electrophoresis on 1.5% (w/v) agarose gels and stained with ethidium bromide.

Western blot analysis

Thylakoid were extracted from the first leaves of **B-OsLCY-RNAi** plants, $phs3-1$ mutant and the wild type when β -OsLCY-RNAi plants and phs3-1 mutant showed a photobleaching phenotype at the tillering stage. Thylakoid membrane preparation and immunodetection of the PS II core proteins CP43, CP47 and D1 were carried out as described (Peng et al., 2006).

Enzyme activity assay

All enzyme activities were measured from 0.5 g first leaves of β -OsLCY-RNAi plants and the *phs3*-1 mutant as well as the wild type when β -OsLCY-RNAi plants and the phs3-1 mutant showed a photobleaching phenotype at the tillering stage. The enzyme activities of SOD, CAT, APX, MDAR, DHAR, and GR were measured as described previously (Chen and Gallie, 2004; Gonzalez et al., 1998; Jiang and Zhang, 2002; Knöraer et al., 1996).

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Supplementary Material

The following supplementary material is available for this article online:

Figure S1. Carotenoid and abscisic acid biosynthetic pathway.

Figure S2. The phenotypes of the rice pre-harvest sprouting mutants.

Figure S3. Activities of ROS-scavenging enzymes of leaves from OsLCY-RNAi plant, phs3-1, and wild type at tillering stage.

Figure S4. Abscisic acid responsiveness test of *phs3-1* mutant.

Figure S5. Map-based cloning of PHS genes.

Table S1. The primers used for expression analysis via RT-PCR. Table S2. Segregation ratios of viviparous and non-viviparous

plants in the T_2 generation.

Table S3. Constructions for genetic complementation of phs mutants and plant transformation events.

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188 Jun Fang et al.

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