The molecular cloning and clarification of a photorespiratory mutant, oscdm1, using enhancer trapping

Jinxia Wu†, Zhiguo Zhang†, Qian Zhang, Xiao Han, Xiaofeng Gu* and Tiegang Lu*

Biotechnology Research Institute/National Key Facility for Genetic Resources and Gene Improvement, The Chinese Academy of Agricultural Sciences, Beijing, China

Enhancer trap systems have been demonstrated to increase the effectiveness of gene identification in rice. In this study, a chlorophyll-deficient mutant, named oscdm1, was screened and characterized in detail from a T-DNA enhancer-tagged population. The oscdm1 plants were different from other chlorophyll-deficient mutants; they produced chlorotic leaves at the third leaf stage, which gradually died with further growth of the plants. However, the oscdm1 plants were able to survive exposure to elevated CO\textsubscript{2} levels, similar to photorespiratory mutants. An analysis of the T-DNA flanking sequence in the oscdm1 plants showed that the T-DNA was inserted into the promoter region of a serine hydroxymethyltransferase (SHMT) gene. OsSHMT1 is a key enzyme that is ubiquitous in nature and structurally conserved across kingdoms. The enzyme is responsible for the interconversion of serine and glycine and is essential for cellular one-carbon metabolism. Full-length OsSHMT1 complemented the oscdm1 phenotype, and the downregulation of OsSHMT1 in wild-type plants by RNA interference (RNAi) produced plants that mimicked the oscdm1 phenotype. GUS assays and quantitative PCR revealed the preferential expression of OsSHMT1 in young leaves. TEM revealed serious damage to the thylakoid membrane in oscdm1 chloroplasts. The oscdm1 plants showed more extensive damage than wild type using an IMAGING-PAM fluorometer, especially under high light intensities. OsSHMT1-GFP localized exclusively to mitochondria. Further analysis revealed that the H\textsubscript{2}O\textsubscript{2} content in the oscdm1 plants was twice that in wild type at the fourth leaf stage. This suggests that the thylakoid membrane damage observed in the oscdm1 plants was caused by excessive H\textsubscript{2}O\textsubscript{2}. Interestingly, OsSHMT1-overexpressing plants exhibited increased photosynthetic efficiency and improved plant productivity. These results lay the foundation for further study of the OsSHMT1 gene and will help illuminate the functional role of OsSHMT1 in photorespiration in rice.

Keywords: rice, serine hydroxymethyltransferase, photorespiratory, mitochondria, hydrogen peroxide
Introduction

Leaves are the most important organs for photosynthesis in plants; indeed, the photosynthetic efficiency of leaves determines crop productivity. Improving photosynthesis in leaves is a reliable method for breeding cultivars with high photosynthetic efficiency (Govaerts et al., 1996; Zhang et al., 2009). The key to this method is cloning and analyzing the function of genes involved in leaf photosynthesis. Leaf color mutants are ideal genetic materials for studying the molecular mechanisms that regulate leaf photosynthesis and photorespiration in plants. To date, a considerable number of leaf color mutations have been reported in rice (http://www.shigen.nig.ac.jp/rice/oryzabaseV4), and some of them have been cloned successfully. For example, OsCHLH was the first gene from rice to be identified from a T-DNA insertion mutant using gene trapping methods. OsCHLH encodes the OsChlH subunit of magnesium chelatase, and its mutation produces plants that lack chlorophyll in their thylakoids (Jung et al., 2003; Goh et al., 2004). YGL1 encodes a chlorophyll synthase; its mutation reduces chlorophyll accumulation and delays chloroplast development (Wu et al., 2007). Chl1 and Chl9 encode key enzymes for chlorophyll synthesis and chloroplast development, respectively. Ultrastructural analyses have revealed that the grana of Chl1 and Chl9 mutants are poorly stacked, resulting in underdeveloped chloroplasts (Zhang et al., 2006). OsPPRI encodes a pentatricopeptide repeat protein that may play an essential role in chloroplast biogenesis; its mutant exhibits typical phenotypes of chlorophyll-deficient plants (Gothandam et al., 2005). OsHAP3 regulates normal chloroplast development and chlorophyll biosynthesis, and its mutant has pale green leaves (Miyoshi et al., 2003).

The mutants and genes described above are mostly related with chlorophyll synthesis or degradation in rice. In recent years, several genes from other pathways affecting leaf color have been identified in monocotyledonous plants. For example, several tie-dyed mutants, including tie-dyed1, tie-dyed2, camouflauge1, psychedelic, and sucrose export defective1, have been described that show chlorotic/variegated leaves and carbon hyperaccumulation in their leaf blades (Braun et al., 2006; Baker and Braun, 2007, 2008; Ma et al., 2008; Huang et al., 2009; Slewinski and Braun, 2010; Slewinski et al., 2012; Baker et al., 2013). Tie-dyed1 (Tdy1) is a novel transmembrane protein that is present only in grasses. Tdy1 functions in carbon partitioning by promoting phloem loading; accordingly, it is expressed exclusively in the phloem cells of both source and sink tissues. The tie-dyed2 (tdy2) mutant exhibits variegated green and yellow leaves. Tie-dyed2 is a callose synthase that functions in vein development and affects symplastic trafficking within the phloem of maize leaves. Several other maize leaf color genes, including Oil Yellow1 (Oy1), those in certain lesion mimic mutants, and Ioiap (ij), have also been cloned. The semi-dominant Oy1 mutants of maize are deficient in the first committed step of chlorophyll biosynthesis: the conversion of protoporphyrin IX to magnesium protoporphyrin IX. The Oy1 gene encodes subunit I of magnesium chelatase (Sawers et al., 2006). The lethal leaf spot 1 lesion mimic locus of maize (2mLls1) encodes pheophorbide a oxygenase. This locus corresponds to gene At3g44880 on chromosome 3 of Arabidopsis thaliana (Gray et al., 2002; Yang et al., 2004a). Iojap (ij) is a recessive striped mutant of maize in which plastid development is locally altered in a position-dependent manner on the leaves; ij-affected plastids can be transmitted to progeny even when the function of the nuclear gene is restored. The ij mutant is characterized by a number of independent transposon insertion mutations created using Robertson's Mutator. The ij gene encodes a 24.8-kDa protein with no significant sequence similarity to proteins in public databases (Han et al., 1992; Byrne and Taylor, 1996; Silhavy and Maliga, 1998). In addition to these leaf color mutants, the barley mutants tigrina-d (Lee et al., 2003; Khandal et al., 2009) and chlorina have been well characterized (Knoetzel et al., 1998; Krol et al., 2009).

Plant photorespiration is an essential prerequisite for oxygenic photosynthesis. This metabolic repair pathway b estrides four compartments and requires several metabolite transporters for pathway function and the well-studied enzymatic steps of the core photorespiratory cycle (Eisenhut et al., 2014). Photorespiration dissipates excess photochemical energy to provide protection against oxidative damage under stressful conditions in which CO\textsubscript{2} assimilation is reduced. In addition, photorespiration provides metabolites for protection against stress, including glycine for the synthesis of glutathione (Lakshmanan et al., 2013). Mutants with photorespiration pathway deficiencies exhibit chlorotic and lethal phenotypes when grown in ambient CO\textsubscript{2} (Somerville and Ogren, 1981; Jamai et al., 2009). The enzymatic steps involved in the photorespiratory pathway have been well established. For example, mitochondrial serine hydroxymethyltransferase (SHMT) has been reported to be an essential component of the pathway; it plays an important role in one-carbon metabolism. Specifically, the enzyme produces N\textsubscript{5},N\textsubscript{10}-methylene tetrahydrofolate (THF) glycine (Gly) from Ser and THF, and generates one-carbon units for cellular use (Mouillon et al., 1999). Seven SHMT genes have been found in Arabidopsis and other species. The Arabidopsis SHMT mutant shm1 was one of the first photorespiratory mutants described by Somerville and Ogren (1981); the affected gene was later cloned by map-based cloning (Voll et al., 2006). AtSHMT1 was reported to play a critical role in controlling cell damage caused by abiotic stress, whereas its mutant shm1-1 has a conditional lethal photorespiratory phenotype (Moreno et al., 2005). Thus, SHMT has acquired a new or altered function. A soybean (Glycine max L. Merr.) SHMT conferring resistance to nematodes plays an important role in plant resistance mechanisms against the pathogen (Liu et al., 2012).

T-DNA tagging methods, including gene trapping, promoter trapping, and enhancer trapping, have been shown in rice to be more effective and faster than positional cloning. These techniques are also advantageous when dealing with early embryogenic or gametophytic lethality. Several genes have been cloned by gene trapping or enhancer trapping, including OsCHLH (Jung et al., 2003), Wda1 (Jung et al., 2006), UDT1 (Jung et al., 2005), Rip1 (Han et al., 2006), and MIT (Bashir et al., 2011). We previously reported the establishment of a large T-DNA enhancer trapping population (Yang et al., 2004b; Peng et al., 2006).
et al., 2005; Wan et al., 2009). The vector used to establish the pool carried a minimal cauliflower mosaic virus (CaMV) 35S promoter containing only the TATA box and transcription start site, which was placed adjacent to the T-DNA border. Only when the insertion site is next to an enhancer can the minimal promoter drive reporter gene expression. The GAL4/VP16-UAS enhancer trap was used in our insertion pool; this system improves the efficiency of enhancer trapping greatly. Insertion of the T-DNA into a gene in the proper orientation enables fusion between the endogenous gene and the GUS reporter gene; thus, T-DNA-tagged lines can be verified by a GUS assay. Screening of our insertion lines revealed that 40% were positive for GUS in the leaves or seeds (Peng et al., 2005).

To understand the molecular mechanism controlling leaf color, we first screened for leaf color mutants in the T1 generation. Leaf color phenotypes co-segregated with GUS staining in nine independent lines. One line, oscdm1, showed co-segregation between its mutant phenotype and the T-DNA; thus, it was selected for further analysis. The oscdm1 plants had yellow-green chlorotic leaves; this trait was found to be controlled by a single nuclear gene. The oscdm1 plants displayed a lethal phenotype under natural conditions. Amplification of the T-DNA insertion-flanking sequence by PCR-based genome walking and subsequent sequencing resulted in the identification of an SHMT gene, OsSHMT1, which is predicted to encode the largest subunit of SHMT. The mutant phenotype was complemented by transformation with the wild-type gene, and plants in which OsSHMT1 expression was reduced by RNA interference (RNAi) exhibited the oscdm1 mutant phenotype. Moreover, the oscdm1 plants exhibited extensive thylakoid membrane damage compared to wild type, especially at high light intensities. This damage may have resulted from excessive H2O2 levels. The identification of OsSHMT1 using enhancer trapping and the characterization of oscdm1 will aid future studies of the exact role of OsSHMT1 in photorespiration in rice.

Materials and Methods

Plant Materials

The enhancer trapping population used in the present study for the mutant screen was described previously (Peng et al., 2005). The functional regions of the enhancer trap vector pFX-E24.2-15R are shown in Additional File 1: Figure S1. Within the T-DNA, the reporter gene was comprised of −48CaMV, GAL4/Vp16, 6×UAS, BoGUS, and EGFP; the reporter was located at the right border (RB) of the T-DNA. The minimal promoter −48CaMV cannot drive expression of the reporter gene alone; however, when the reporter gene is inserted into a gene or nearby enhancer it can be activated by the minimal promoter with the help of these enhancer elements.

Leaf Color Mutant Isolation

About 30 T1 seeds per line were sterilized in 1% sodium hypochlorite for 30 min, washed with running tap water, and then imbibed in tap water for 3 days at 20°C. The seeds were then sown on a plastic net floating on nutrient fluid (Yoshida et al., 1976). The pH was adjusted every 2 days, with replacement of the nutrient fluid every week. The plants were grown for 14–30 days under greenhouse conditions, with natural sunlight and a minimum nighttime temperature of 25°C.

Histochemical Staining for GUS Activity in Plant Tissues

GUS assays were performed essentially as described by Jefferson et al. (1987). Rice tissues were removed, transferred to Eppendorf tubes containing GUS staining solution (50 mM sodium phosphate, pH 7.0, 10 mM EDTA, 0.1% Triton X-100, 1 mg/ml X-gluc, 0.1 mM potassium ferricyanide, and 10% methanol), and incubated at 37°C for 10–12 h. The staining solution was then removed and the tissues were stored in 70% ethanol. The stained tissues were examined under a zoom stereomicroscope (Nikon SMZ1000; Tokyo, Japan) and photos were taken with a digital camera (Nikon DXM1200F).

Amplification and Analysis of the T-DNA Flanking Sequences

Genomic DNA from the enhancer trap mutants was isolated using an improved version of the CTAB method (Murray and Thompson, 1980), and the T-DNA left border (LB) flanking regions were rescued using a modified version of the PCR-based genome walking protocols of Siebert et al. (1995) and Sallaud et al. (2003). The method consisted of three steps: digestion of genomic DNA using a blunt-end restriction enzyme and ligation of an asymmetrical adaptor, amplification by PCR using primers specific for the T-DNA and the adaptor, respectively, and successive PCR using two nested, specific primers. The adaptor (ADAR, 50 mM) was prepared by heating a mixture comprised of equal volumes of the complementary oligonucleotides ADAR1 (100 mM) and ADAR2 (100 mM) to 80°C for 10 min, and then allowing it to cool gradually to room temperature for annealing. The specific primers for the adaptor were APR1 (20 mM) and APR2 (20 mM); the specific primers for the T-DNA LB in pFX-E24.2-15R were LB1 (20 mM) and LB2 (20 mM).

The distance between the LB binding site and the LB was about 180 bp. Unwanted amplification resulting from the two adaptors was inhibited by the asymmetry of the adaptor and its NH2-blocked internal 3’ end; the second round of nested PCR also reduced non-specific amplification reactions (Siebert et al., 1995). Each DNA sample (40–100 ng) was digested using DnaI, EcoRV, or PvuII (2 U; Takara, Dalian, China) and ligated with ADAR (0.1 ml) using T4 DNA ligase (70°C; Takara) at 25°C in a total volume of 10 μl (1 μl of 10X ligase buffer). After digestion/ligation for 10-12 h, aliquots (2 μl each) were used for the first round of PCR (MJ Research PTC-100 DNA Engine; MJ Research Inc., Quebec, Canada) in a total volume of 20 μl (0.3 ml of LB1, 0.3 ml of APR1, 2 U of Takara r-Taq, 100 mM dNTPs, and 2 μl of 10X PCR buffer). The program was as follows: 3 min at 94°C followed by 35 cycles of 94°C for 30 s, 67°C for 45 s, and 72°C for 150 s, with a final elongation step at 72°C for 5 min. Next, 2 μl of a 1:50 dilution of the product was used for nested PCR in a total volume of 20 μl (0.5 μl of LB2, 0.5 μl of APR2, 0.8 U of r-Taq, 200 mM dNTPs, and 2 μl of 10X PCR buffer) under the same conditions. All of the products from the second round of PCR were loaded onto a 1.5% agarose
Leaves (300 mg) were ground to a powder in the sequencer's capability of >1 kb. LB2 was used as the primer for sequencing according to the protocol included with the BigDye Terminator v3.1 Cycle Sequencing Kit (Life Technologies).

**Co-segregating Genotype Analysis of oscdm1 by PCR**

An analysis of the oscdm1 plants was performed by PCR using two gene-specific primers flanking the insertion site (P1 and P2) and another T-DNA LB-specific primer (P3). PCR was carried out in a total volume of 20 µl containing 20 ng of plant DNA, 10× PCR buffer, 0.2 mM dNTPs, 0.5 U of rTaq polymerase, and 1 mM each primer. The DNA was denatured at 95°C for 4 min, followed by 35 cycles of 94°C for 1 min, 58°C for 1 min, and 72°C for 2 min. The products were loaded onto a 1.5% agarose gel for electrophoresis.

**OsSHMT1 cDNA Amplification, Complementation Analysis, and Overexpression Line Construction**

Total RNA was extracted using TRizol reagent (Invitrogen, Carlsbad, CA, USA) from wild-type plants. The RNA was quantified using a UV spectrophotometer (Beckman Coulter Inc., Brea, CA, USA). First-strand cDNA was synthesized by reverse transcription using a cDNA synthesis kit (Takara) in a total volume of 20 µl containing 1 µg of total RNA, 10 ng of oligo(dT)14 primer, 2.5 mM dNTPs, 1 µl of AMV, and 0.5 µl of RNAsin. Amplification of the full-length cDNA sequence of OsSHMT1 (1542 bp) was performed in a total of 20 µl containing a 1/20 aliquot of the cDNA reaction, 0.5 µM gene-specific primers (OP1, which contained a SalI digestion site, and OP2, which contained a SmaI digestion site), 10 mM dNTPs, 1 U of rTaq DNA polymerase, and 2 µl of 10× reaction buffer. The reaction protocol was as follows: denaturation at 94°C for 3 min followed by 25 cycles of 94°C for 30 s, 60°C for 45 s, and 72°C for 1 min, with a final step at 72°C for 10 min. An aliquot (1 ¸ tl) of the reaction was loaded onto a 1.5% agarose gel (regular; Bio-Rad, Hercules, CA, USA) and analyzed by electrophoresis. The PCR product was cloned into the pEASY-Blunt simple cloning vector (Beijing TransGen Biotech Co., Ltd.) and sequenced. The vector was inserted into pTCK309 using the SacI, SpeI, KpnI, and BamHI sites in inverted orientations. The pUbi-RNAi309-SHMT1 construct was transferred into Agrobacterium strain AGL1 by electroporation and then introduced into wild-type rice calli through A. tumefaciens-mediated transformation. The regenerated T0 plants were grown in a paddy field at the Chinese Academy of Agricultural Sciences in Beijing, China. Leaves of the pUbi-RNAi309-SHMT1 transgenic plants showing the chlorina phenotype were selected and analyzed.

**Reverse Transcription (RT)-PCR and Quantitative Real-time RT-PCR (qRT-PCR)**

To study the OsSHMT1 expression level, total RNA was extracted using TRizol reagent (Invitrogen) from the leaves of wild-type and oscdm1 plants. Total RNA (2 µg) from each sample was reverse-transcribed with oligo(dT) primer and PrimeScript RT Enzyme (Takara) according to the manufacturer's instructions. For RT-PCR, the primers used to amplify OsSHMT1 were Q1f and Q1r, while the primers used to amplify actin were Actinf and Actinr. The PCR conditions were as follows: preincubation at 94°C for 2.5 min, then 30 cycles of 94°C for 30 s, 52°C for 30 s, and 72°C for 1 min. qRT-PCR was performed using an iQTM5 Mutescolor Real-Time PCR Detection System (Bio-Rad, Hercules, CA, USA) with SYBR® Green Real-Time PCR Master Mix (Life Technologies). The primers used to amplify OsSHMT1 were qF and qR.

**Transmission Electron Microscopy (TEM)**

Leaves were harvested from wild-type and oscdm1 plants at the third leaf stage that had been grown in a greenhouse under a medium light intensity (∼150 µmol of photons/m²). Leaf sections were fixed in 2% glutaraldehyde and further fixed in 1% OsO4. The tissues were then stained with uranyl acetate, dehydrated in ethanol, and embedded in Spurr’s medium prior to thin sectioning. The samples were then stained again and examined with a JEOL 100 CX electron microscope (JEOL Ltd., Tokyo, Japan). The total chlorophyll content was determined spectrophotometrically according to the method of Arnon and Whatley (1949). Leaves (300 mg) were ground to a powder in liquid nitrogen and then transferred to a 15-ml Falcon tube. Next, 5 ml of 80% acetone were added to the tube and the mixture was thoroughly combined then left in the dark overnight. Centrifugation was performed at 4°C for 15 min (3000 rpm). The supernatant was transferred to a new centrifuge tube and the chlorophyll absorbance (hereafter termed A) was measured by spectrophotometry. The chlorophyll concentrations were calculated as follows (using 80% acetone as a blank control): C_{a+b} (mg/g) = [8.026A_{663} + 20.206A_{645}] × V/10000 × W, where V is the volume of the extract (ml) and W is the fresh leaf weight (g).

**Chlorophyll Fluorescence Measurement**

The middle portion of the first leaves from wild-type and oscdm1 plants at the third leaf stage were collected and used to measure the leaf photosynthetic efficiency. Chlorophyll fluorescence measurements were performed with an IMAGING-PAM 2000 chlorophyll fluorometer (Walz, Effeltrich, Germany). A small
piece of the mutant leaf blade was removed and placed in the well of a PCR plate filled with water; after 15 min of dark preadaptation, the measurements were performed. Changes in the chlorophyll fluorescence of the chlorophyll-deficient mutants were documented based on F₀, the steady level of chlorophyll fluorescence; Fₘ/F₀ = (Fₘ₋F₀)/F₀, the maximum quantum yield of photosystem (PS)II photochemistry; Φₛₛₚᵢₐₜ = (Fₘ′−F)/Fₘ′, the effective quantum yield of PSII; qP = (Fₘ′−F)/(Fₘ′−F₀), photochemical quenching; and qN = 1−(Fₘ′−F₀)/[Fₘ′−F₀]), non-photochemical quenching. The measurements were recorded after 5 min under actinic light. F₀ was calculated as F₀ = F₀/(Fₘ′/Fₘ′+F₀/Fₘ′) because of the limitations of the fluorometer.

**Blue Native (BN) Test**

BN polyacrylamide gel electrophoresis for the isolation of membrane protein complexes in an enzymatically active form was performed mostly as described (Schagger et al., 1994; Yan et al., 2007). Thylakoids were resuspended in medium A (25 mM Bis-Tris HCl, pH 7.0, 20% [w/v] glycerol, and 0.25 mg/ml Pefabloc) to a final concentration of 0.5 mg/ml chlorophyll, and the chlorophyll fluorescence of the chlorophyll-deficient mutants was recorded after 5 min under actinic light. F₀ was measured at the full heading stage using a portable photosynthetic chamber, and 1800 umol/s light quantum flux density.

**Phylogenetic Analysis**

The BLAST search program (http://blast.ncbi.nlm.nih.gov/) was used to identify homologs of OsSHMT1 using its amino acid sequence as the query. The resulting sequences were aligned using ClustalX1.83 in multiple alignment modes, and a neighbor-joining phylogenetic tree was generated using MEGA 4.0 (Tamura et al., 2007). The bootstrap values for nodes in the phylogenetic tree were from 1000 replications. The handling gap option was pairwise deletion; the numbers at the branch points indicate the bootstrap values. The peptide identities among the different proteins were calculated using GeneDoc software. The OsSHMT1 nucleotide/amino acid sequence has been deposited in the NCBI database with accession number: Os03g0738400. The OsSHMT1 full cDNA clone AK063056 has been deposited in KOME.

**Subcellular Localization**

Full-length OsSHMT1 cDNA was amplified using the primers F1 and R1, after which the product was cut with Sma1 and Xba1 and then fused with PAN580. The 35S::OsSHMT1-GFP expression construct was transfected into rice protoplasts as described previously (Zhang et al., 2011). Briefly, for each sample, 10 µg of plasmid DNA were mixed with 200 µl of protoplasts (~1 × 10⁶ cells). Freshly prepared polyethylene glycol solution [220 µl; 40% (w/v) polyethylene glycol 4000, 0.2M mannitol, and 0.1 M CaCl₂] was added, and the mixture was incubated at room temperature for 20 min in the dark. After incubation, 1 ml of W5 solution was added slowly to the samples. The resulting solution was mixed well by gently inverting the tube, and the protoplasts were pelleted by centrifugation at 1500 rpm for 3 min. The protoplasts were resuspended gently in 1 ml of W5 solution. Finally, the protoplasts were transferred to multi-well plates and cultured under light or in the dark at room temperature for 16 h. MitoTracker® Red CMXRos was purchased from Invitrogen. The samples were observed with a confocal laser scanning microscope (Leica TCS SP5; Leica Microsystems, Wetzlar, Germany).

**Measurement of the H₂O₂ Levels**

The H₂O₂ levels in the plants were measured according to the method of Dagnar et al. (2001). Briefly, 500 mg of leaves were homogenized in a cold mortar with a pestle and 0.2 g of silicon dioxide in pre-cooled acetone (5 ml). The homogenate was centrifuged at 12,000 × g for 5 min, after which 1 ml of the supernatant was mixed with 0.1 ml of 5% Ti(SO₄)₂ and 0.2 ml of 19% ammonia. After the formation of a precipitate, the reaction mixture was centrifuged at 12,000 × g for 5 min. The resulting pellet was dissolved in 3 ml of 2 M H₂SO₄ and the absorbance of the solution was recorded at 415 nm. The H₂O₂ content was calculated according to a standard curve of H₂O₂ ranging from 0 to 10 mM.

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**Statistical Analysis**

All the experiments were carried out at least in triplicate. The values shown in the figures are mean values ± SD. For multiple comparisons, means were compared by one-way analysis of variance and Duncan’s multiple range test with a 5% level of significance.

**Primer Sequence**

All the primers for gene cloning, complementation analysis, over-expression construction, sub-cellular localization, reverse transcription (RT)-PCR and quantitative real-time (qRT-PCR) and etc in our paper are listed in Table S1.

**Results**

Isolation of Chlorophyll-deficient Mutants from the T-DNA Insertion Population

To identify new genes modulating the photosynthetic mechanism in rice, we conducted a genetic screen of an enhancer trapping population containing more than 10,000 individual lines. In total,
420 lines with leaf blade color defects were identified, including 251 albino, 132 pale green, 35 chlorotic (yellow), 25 stripe, and 13 zebra. The mutant lines varied in phenotype from albino to pale green according to the degree of chlorophyll deficiency (Figure 1A). Of the 420 leaf color mutants tested, 34 lines were positive for GUS staining. However, only in nine lines were all of the mutants positive for GUS staining. These lines were selected for further study. To test for the co-segregation of GUS staining and the hygromycin resistance gene HptII in the T1 generation, we amplified DNA from each mutant line using HptII-specific primers. The results were consistent, as expected, for all mutants of the nine lines. The GUS expression patterns in the mutant leaves are shown in Figure 1B.

Analysis of T-DNA Insertion Sites in the Rice Genome and Prediction of Tagged Genes

PCR-based genome walking has been shown to be a highly efficient way of isolating T-DNA flanking sequences (Peng et al., 2005). Using this technique, 10 bands were amplified from the nine mutant lines. Of all of the bands sequenced, only six contained rice genomic sequences. The other four contained vector sequences, indicating border read-through or the termination of sequencing. Of the 10 flanking sequences rescued, six sequences originating from five mutants showed a unique hit to rice artificial chromosome (BAC/PAC) clones based on homology searches against sequences obtained from the NCBI and KOME databases. Of the six sequences, four flanking sequences were identified in genic regions and the other two in intergenic regions, based on a public annotation database for the rice genome. The locations of the T-DNA insertions are shown in Table 1.

Identification of a Mutation in the SHMT Promoter

Sequence analyses using the NCBI database identified one line, P239, also named oscdm1, as having a T-DNA insertion upstream of a gene with high homology to SHMT genes from several plant species. OsSHMT1 encodes the largest subunit of SHMT. OsSHMT1 is 3962 bp in length and consists of 16 introns and 17 exons, with a complete coding sequence of 1542 bp encoding 513 amino acids and a molecular mass of 56.3 kDa; it encodes a putative serine methyltransferase domain, which spans from residues 51–449 at the C-terminus (Figure S2).

Characterization of the oscdm1 Mutant

There was no significant difference between 10-day-old (first leaf stage) oscdm1 and wild-type plants (Figure 2A). However, the oscdm1 plants appeared chlorotic at the third leaf stage (Figure 2B), and they gradually died off from the fourth leaf stage onward. We measured the chlorophyll contents of both the oscdm1 plants and wild-type plants at the third leaf stage. The ratio of chlorophyll a to chlorophyll b in the wild-type plants was about 3.5. The chlorophyll a, chlorophyll b, and total chlorophyll contents in the oscdm1 plants at the third leaf stage were drastically lower than in the controls (Figure 2C). These results suggest that the chlorotic phenotype of the oscdm1 seedlings was probably caused by a reduction in the total chlorophyll content, rather than a reduction in a particular pigment.

Light microscopic observations of cross-sections of a chlorina mutant leaf blade did not show any significant change in the size or number of mesophyll cells (data not shown). To examine whether the lack of chlorophyll in the oscdm1 plants was accompanied by ultrastructural changes in the chloroplasts, we compared the ultrastructure of chloroplasts in the oscdm1 and control plants at the third leaf stage using TEM. The analysis revealed no great change in chloroplast structure at the first leaf stage between the wild-type and oscdm1 plants; however, at the third leaf stage, substantial changes in the number of chloroplasts, lamellar structures, and organization of the grana in the oscdm1 chloroplasts (Figures 3B,D) were noted compared with wild type (Figures 3A,C). In addition, the mutant chloroplasts exhibited vacuolation (Figure 3E). The BN test was used to assess thylakoid membrane integrity. A BN gel analysis of the thylakoid membrane showed some differences in five bands between the mutant and wild-type plants; in particular, bands III and IV were slightly more diffuse in the mutant plants. We concluded that the light harvesting complexes (LHCs) in the thylakoid membrane were altered and that chlorophyll synthesis was affected in the mutants beginning at the third leaf stage (Figure S3).

Co-segregation of the T-DNA and GUS Activity with the Chlorina Phenotype

The oscdm1 plants appeared chlorotic at the third leaf stage. These plants eventually died both under greenhouse conditions and in a paddy field because of insufficient photosynthesis. In the T2 generation, 80 plants were wild type and 30 were oscdm1. In the
TABLE 1 | Locations of the T-DNA insertion and tagged putative genes.

| Line | GenBank accession number | Insertion site | BLASTn E-value | Chr. | Insertion region (putative genic or intergenic) | Localization of putative protein |
|------|--------------------------|----------------|----------------|------|-----------------------------------------------|--------------------------------|
| A33  | AL731638                 | 10450          | 2E-33          | 4    | Intergenic region                             |                                |
| A167 | AP002837                 | 46557          | 9E-07          | 6    | ORF of a putative Fe-SOD gene                 | Chloroplast                    |
| P239 | AC117986                 | 110492         | 9E-07          | 3    | ORF of a putative glycine hydroxymethyltransferase gene | Mitochondrion                  |
| A304 | AP004809                 | 45787          | 3E-07          | 6    | Intergenic region                             |                                |
| A401-1| AC125782                | 158205         | 3E-07          | 11   | ORF of a putative GTP-binding gene             | Chloroplast                    |
| A401-2| AP004182                | 53463          | 1E-17          | 7    | ORF of a putative TPR-like domain-containing gene | No distinct localization       |

ORF, open reading frame; SOD, superoxide dismutase; TPR, tetratricopeptide repeat.

T2 and T3 seedlings were genotyped to test whether the chlorotic phenotype co-segregated with the T-DNA insertion (Figures 4A,B). Two randomly selected lines, 4 and 18, exhibited a wild-type phenotype, and T3 seedlings of those lines were wild type in appearance. However, the progeny of plants 3 and 19 exhibited wild-type and mutant phenotypes at a ratio of 3:1. In contrast, plants 4 and 7 displayed the chlorina phenotype. These results demonstrate that the chlorina phenotype co-segregated with the T-DNA insertion. Furthermore, RT-PCR using RNA from the homozygous oscdm1 plants did not detect full-length transcripts for OsSHMT1. Thus, the oscdm1 plants were deemed to be loss-of-function mutants (Figure 4C).

Functional Complementation of osshmt1

To further demonstrate that the oscdm1 phenotype was due to insertion of the T-DNA into the candidate gene, functional complementation of the oscdm1 mutant with wild-type OsSHMT1 was performed. Given the lethality of the homozygous mutation, homozygous mutant individuals were identified by genotyping the progeny of a heterozygous plant.
that was allowed to self-fertilize. These homozygous mutant
individuals were used to initiate a callus in tissue culture, which
was used for Agrobacterium-mediated transformation with the
complementation construct. The sequence of OsSHMT1 was
used to search for the full-length cDNA sequence in the KOME
database (http://cdna01.dna.affrc.go.jp/cDNA/). Fortunately,
the full-length cDNA sequence of OsSHMT1 was available.
We cloned the full-length cDNA sequence by RT-PCR. The
full-length cDNA sequence of OsSHMT1, which is 1542 bp
in length, was found to be identical to that of the annotated
gene Os03g0738400 in the TIGR database. This cDNA was
cloned into a binary overexpression vector containing the actin
promoter and subsequently transformed into oscdm1 calli via
A. tumefaciens-mediated transformation with G418 selection
resistance (Figure 5A).

The resulting G418-resistant lines were confirmed using
specific PCR primers (p1, p2, and p3) and cDNA-amplifying
primers (OP1 and OP2). In addition, quantitative PCR revealed
that the exogenous OsSHMT1 gene was overexpressed in 50
transgenic lines (CP1, CP2, and CP3) (Figure 5C). Interestingly,
these 50 lines displayed a wild-type phenotype even though
their background was homozygous oscdm1 (Figure 5B). The
chlorophyll content in these lines was rescued to the wild-type
level (Figure 5D).

**RNAi Against OsSHMT1 Reproduced the oscdm1
Mutant Phenotype**

To further confirm that OsSHMT1 was the gene associated
with the observed mutant phenotype, an OsSHMT1 RNAi vector was
constructed and transformed into japonica variety Nipponbare
(Figure 6A). About 400 independent transgenic lines were
obtained and confirmed by PCR, and more than 100 of them
displayed visibly chlorotic leaves during the early growth period
(Figure 6B); however, after further growth and development the
plants eventually died. Real-time PCR revealed that OsSHMT1
transcription was decreased in the RNAi transgenic plants
compared to that in the Nipponbare controls (Figure 6D). The
chlorophyll contents were decreased to 11 and 37% of that
in Nipponbare using five representative T0 lines (RNAi-1–5,
respectively) (Figure 6E). Those RNAi plants showing a severe
chlorotic phenotype eventually died in the field (Figure 6C).

Conversely, other transgenic lines lacking the pUbi-RNAi309-
SHMT1 vector (data not shown) failed to exhibit the chlorotic
phenotype. These complementation and RNAi results strongly
indicate that Os03g0738400 corresponds to OsSHMT1.

**Sequence Analysis Of OsSHMT1**

BLAST searches of all available genomic sequences revealed
that the rice genome contains five genes encoding an SHMT
Whether OsSHMT1 is localized to mitochondria, the complete OsSHMT1 coding region was fused to the gene encoding GFP under the control of the CaMV 35S promoter. Vectors to express the 35S::OsSHMT1-GFP fusion protein and 35S::GFP (as a control) were introduced into rice protoplasts. MitoTracker® Red was used as a positive control for mitochondria. The 35S::OsSHMT1-GFP fusion protein co-localized with the MitoTracker® Red signal (see the merged image), suggesting that OsSHMT1 is a mitochondrial protein (Figures 7E–H).

**OsSHMT1 Expression Pattern**

An analysis by qRT-PCR revealed the expression of OsSHMT1 mainly in young leaves, and not in root, leaf sheath, or flower, implying that OsSHMT1 functions in seedlings or young leaves. To investigate the expression pattern of the gene in greater detail, total RNA was extracted from leaves from the first leaf stage to the mature stage. A qRT-PCR analysis of these RNA samples showed low-level expression of OsSHMT1 at the first leaf stage; however, the expression level increased with leaf development and reached its highest level at the third leaf stage. Expression of the gene remained at a relatively high and stable level until the fifth leaf stage, suggesting that OsSHMT1 is expressed at an early stage of growth and may play an important role in leaf development (Figure 7I).

Because the GUS gene in the enhancer trapping vector and the OsSHMT1 gene shared the same orientation, we expected that the expression pattern in oscdm1 would be consistent with the OsSHMT1 expression pattern. Thus, we analyzed heterozygous plant tissues from an oscdm1 segregating population for GUS staining. In vegetative shoots, strong GUS activity was noted in young leaf and leaf ligule, whereas relatively weak GUS activity was detected in root and flower. This result indicates that OsSHMT1 is expressed mainly in young leaves, consistent with our qRT-PCR results (Figures 7A–D,I).

**Comparison of Chlorophyll Fluorescence Parameters between Wild-type and oscdm1 Plants**

Chloroplasts are the organelles that perform photosynthesis in plants; thus, they play an essential role in plant growth. To test whether the photosynthetic apparatus in the oscdm1 plants was defective, we compared some key parameters of PSI and PSII between oscdm1 and wild-type plants using an IMAGING-PAM fluorometer. The results of the comparison are shown in Table 2. Leaves from wild-type and oscdm1 plants were dark-adapted for 10 min prior to fluorescence recording, and then a saturating light pulse was applied. \( F_v/F_m \) is widely considered to be a sensitive indicator of the maximum quantum efficiency of PSII. The maximum quantum yield (\( F_v/F_m \)) of PSII in the oscdm1 plants was ~0.6, which is considerably lower than the value of 0.83 found in wild-type plants under a low light intensity, and which is the normal value for the mesophyll cells of most plant species. During illumination with actinic light, the maximal fluorescence yields (\( F_m' \)) were assessed in a time-dependent manner. The results indicate that the PSII quantum yield ratio \( \Delta F/F_m' = (F_m' - F)/F_m' \) was remarkably low in the

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**FIGURE 5** | Complemented expression of OsSHMT1 rescued the mutant phenotypes of oscdm1. (A) Schematic structure of the actin-OsSHMT1 complementation vector. NPTII was used as a marker in the transformation test. (B) Complementation of the oscdm1 mutant by the wild-type gene. Japonica rice oscdm1 callus was used for transformation (see the "Materials and Methods"). Images of the wild-type plants, oscdm1 mutant plants, and independent complemented plants (cp)1, cp2, and cp3 were taken after 1 month. Bar = 2 cm. (C) qRT-PCR analyses of OsSHMT1 expression in wild type, oscdm1, and in cp1, cp2, and cp3 showed that the expression level in cp1, cp2, and cp3 was near the wild-type level. The experiments are biological replicates with the SD. (D) The leaf pigment contents in wild type, oscdm1, and in cp1, cp2, and cp3 were measured. The pigment contents of the complemented lines were rescued to the wild-type level. Data were compared by one-way analysis of variance and Duncan’s multiple range test. Different letters (a–b) indicate significant differences (P<0.05) between lines.
**oscdm1** plants as compared with wild-type plants (Figures 8A,B). These results indicate that the **oscdm1** plants exhibited low photosynthetic activity because of damage to the reaction center of PSII.

**The oscdm1 Mutant was MoreSensitive at a High Light Intensity**

To detect the influence of different light intensities on the photosynthetic efficiency of the plants, **oscdm1** and wild-type plants were grown at a low (57 µmol/m²s) or high light intensity (103 µmol/m²s) and a chlorophyll fluorescence response curve was produced. Our results demonstrate no significant change in qP and qN between the wild-type and mutant plants up to 120 s of treatment at a low light intensity. However, at a high light intensity the change occurred earlier. This result suggests that the photochemical and non-photochemical quenching capacities of the mutant were affected, and that the reaction speed became slower, especially at a high light intensity. Similarly, a large change in qN was noted in the mutant plants at a high light intensity. The above results further demonstrate that the photosynthetic

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**FIGURE 6 | RNAi against OsSHMT1 in wild type mimicked the mutant phenotypes of oscdm1.**

(A) Schematic structure of the pUbi-RNAi309-SHMT1 vector, including the inverted repeat at the 3’ end of the SHMT DNA sequence. (B) Phenotypes of 1-month-old wild-type plants and five independent RNAi lines (Ri-1–5). Bar = 2 cm. (C) Photographs of independent RNAi lines in the paddy field after 40 days. (D) qRT-PCR analyses of OsSHMT1 expression in wild type (WT-1 and -2) and five independent RNAi lines (Ri-1–5). The results indicate that the expression of OsSHMT1 in the five independent RNAi lines had decreased to some extent. The experiments are biological replicates with the SD. (E) The pigment contents in the leaves of wild-type plants and independent complemented lines (Ri-1–5) were measured.
capacity of the mutant was abolished at a high light intensity (Figures 8C–F).

**Oscdm1 is a Photorespiratory Mutant with a Chlorotic Phenotype Caused by Excessive H₂O₂**

The *Arabidopsis* mutant *shm1-1* was one of the first photorespiratory mutants to be identified. The affected gene in *shm1-1* encodes an SHMT isozyme. The *shm1-1* mutant displays a lethal photorespiratory phenotype when grown at ambient CO₂ levels, but it is virtually unaffected by elevated CO₂ levels (Somerville and Ogren, 1981; Moreno et al., 2005; Jamai et al., 2009). Given the strong homology between *OsSHMT1* and *AtSHM1*, we examined whether the oscdm1 mutant could survive exposure to a high (1%) CO₂ concentration. Thus, oscdm1 mutant (leftmost plant in Figure 9A) plants were grown under ambient CO₂ levels while wild-type and oscdm1-hu (two rightmost plants in Figure 9A) plants were grown under elevated CO₂ levels in an artificial climate chamber. After 1 month at 1% CO₂, the oscdm1 mutant plants were able to grow as well as wild type under ambient conditions (Figure 9A). Since the plants were kept in an artificial climate chamber for an extended period, some senescing leaves were observed on the wild-type and oscdm1 plants. The pigment content of the oscdm1-hu plants was restored to the wild-type level (Figure 9B). This result is similar to that obtained for *Arabidopsis shm1-1*.

Given that the *Arabidopsis* mutant *atshmt1-1* accumulates reactive oxygen species (ROS) under natural conditions (Moreno et al., 2005), we investigated the H₂O₂ contents among wild-type, oscdm1 mutant, and OsSHMT1-OV plants. As shown in Figure 10A, for all of the lines the H₂O₂ level in the oscdm1 mutant plants was significantly higher than that in wild type at the fourth leaf stage. By contrast, the level of accumulation of H₂O₂ in the OsSHMT1-OV transgenic lines was slightly lower than that in wild type (Figure 10A). The above analyses indicate that OsSHMT1 encodes an SHMT that is required for photorespiration and which plays a role in attenuating ROS production in chloroplasts to mitigate photoinhibition.

**The OsSHMT1-OV Transgenic Lines Exhibited Increased Photosynthetic Efficiency and Improved Productivity**

To evaluate whether OsSHMT1 could improve photosynthetic efficiency, several photosynthetic parameters were measured using a portable LCpro® photosynthesis measurement instrument (ADC BioScientific). The OsSHMT1-OV transgenic lines had a higher photosynthetic rate, stomatal conductance, intercellular CO₂ concentration, and transpiration rate than wild type at the heading stage (Figure 10B and Table 3). Interestingly, the productivity (grain number per plant) of the OsSHMT1-OV lines was 5% greater than that of wild type (Figures 10C,D). Thus, OsSHMT1 overexpression can improve the photosynthetic efficiency of plants and could be applied to breeding programs in the future.

**Discussion**

**Enhancer Trapping is a Valuable Tool for Gene Identification**

GAL4/VP16-UAS, the improved binary T-DNA vector used to construct our study population (Wu et al., 2003; Peng et al., 2005), contains a promoter-less GUS reporter gene next to the RB. This enhancer trapping vector is designed to detect fusion between the GUS gene and an endogenous gene tagged by the T-DNA. Insertion of the construct not only destroys normal gene function, it also activates expression of the reporter gene. In...
FIGURE 8 | Chlorophyll fluorescence analysis in oscdm1 and wild-type plants. (A) Chlorophyll fluorescence induction curve was produced using third leaf stage plants under actinic light (A), low light intensity \( [57 \mu \text{mol/m}^2 \text{s}] \); (B), high light intensity \( [103 \mu \text{mol/m}^2 \text{s}] \). The induction curve did not differ between wild type and the mutant up to 120 s of treatment under a low light intensity, but after 120 s the wild-type plants exhibited a greater fluorescence quenching capacity than the oscdm1 plants. Moreover, under a high light intensity this change occurred at an earlier time. (C–F) Changes in qP and qN in wild-type and oscdm1 plants. (C,D), qP and qN changed under a low actinic light intensity \( (57 \mu \text{mol/m}^2 \text{s}) \). (E,F), qP and qN changed under a high actinic light intensity \( (103 \mu \text{mol/m}^2 \text{s}) \). The data show that qP and qN were not overly different between the wild-type and oscdm1 mutant plants up to 120 s of treatment at a low light intensity. However, under a high light intensity the change occurred earlier. This result suggests that the photochemical and non-photochemical quenching capacities of the mutant were affected and that the reaction speed became slower, especially under the higher light intensity. The data are the means \( \pm \text{SD} (n = 5) \). These experiments were repeated more than three times with similar results.

rice, at least 40% of the insertion events led to GUS activation in various tissues (e.g., roots, leaves, flowers, and seeds) (Peng et al., 2005). Since T-DNA insertion lines do not all carry a single copy of the insert, positive GUS staining results do not fully confirm co-segregation between the mutant phenotype and gene. Still, this method has a number of advantages. First, it can be used to identify a candidate gene from several mutant lines in a short amount of time. Jung et al. (2003) used gene trapping technology to clone OsCHLH; since then, a number of genes, including Wd1 (Jung et al., 2006), have been successfully cloned using this method. Second, enhancer trap vectors can be used to determine candidate gene expression patterns. GUS expression data can provide helpful information for functional gene analyses, including spatial-temporal expression specificity and clues as to a gene’s function (Johnson et al., 2005; Peng et al., 2005; Liang et al., 2006). The cell division-related gene CYCD3;2 was successfully cloned by reporter gene trapping. The GUS staining result in that case was similar but slightly different from the temporal patterns in T-DNA lines owing to the opposing orientation of the T-DNA insertion and CYCD3;2. This result may have been influenced by neighboring genes (Swaminathan et al., 2000). In the present study, the GUS gene was in the same orientation as OsSHMT1. Thus, the technique provided accurate OsSHMT1 organ expression patterns and yielded helpful data for our bioinformatic analysis and prediction of gene function.
The OsSHMT1 Gene has a Conserved Regulatory Function in Monocots and Dicots

In this report, a leaf color mutant with a chlorina phenotype was identified from a rice T-DNA-tagged pool and the gene was trapped by GUS staining. The mutated gene was identified as OsSHMT1, which encodes the largest subunit of SHMT. Rice OsSHMT1 is highly homologous to SHMT from Arabidopsis. The T-DNA was integrated into the OsSHMT1 promoter, 99 bp away from the start codon (ATG). This generated a fusion transcript between the OsSHMT1 promoter and GUS. Moreover, because of the insertion, the wild-type OsSHMT1 transcript was not detectable in plants homozygous for the T-DNA. T-DNA insertion led to a loss-of-function mutation in OsSHMT1. The rice genome includes five genes that are homologous to OsSHMT1. RNai against four other SHMT family genes caused no change in photorespiration compared with wild type (data not shown). This suggests that OsSHMT1 is only one of multiple photorespiration-related genes in the rice SHMT family. SHMT in Arabidopsis is encoded by seven SHM genes, two of which encode mitochondrial isoforms (Jamai et al., 2009). However, only SHM1 is necessary and sufficient to specify photorespiratory SHMT activity (Voll et al., 2006). The oscdm1 plants produced chlorotic leaves, accumulated excessive H$_2$O$_2$, and subsequently died, similar to the mutant Atshm1. Moreover, the oscdm1 plants were able to survive at a high concentration of CO$_2$ (1% CO$_2$). Therefore, SHMT family genes appear to have a conserved regulatory function in photorespiration in both monocots and dicots.

The oscdm1 Mutant Exhibits Damage Caused by Photooxidation

Leaf color mutants may result from a variety of problems, including defects in chlorophyll synthesis, abnormal degradation pathways, and light damage. The oscdm1 mutant is different from other reported leaf color mutants of rice. Almost all leaf color mutants show the mutant phenotype at the start of the second to the third leaf stage; at that stage, the oscdm1 plants began to show slight chlorosis, which increased gradually. Structurally, the chloroplasts in the oscdm1 plants developed well initially, but as time progressed the chloroplasts were destroyed and the plants eventually died. This chain of events is similar to that observed for photorespiration mutants of Arabidopsis. Photorespiration dissipates excess light energy as a protective process in plants. Based on our results, the phenotype of the oscdm1 plants was due to photooxidation and changes in chloroplast structure. When the leaves of the mutant were subjected to stress, the chloroplasts exhibited ultrastructural changes, including irregularly stacked grana, and symptoms of vacuolization. The ultrastructural changes observed in the chloroplasts of the oscdm1 plants are similar to the ultrastructural changes observed in the leaves under conditions of stress. When the plants were under stress, the level of ROS increased and caused cellular injury due to membrane oxidation. A number of physiological processes were affected, including photosynthesis, programmed cell death, hormone action, growth and development, and mitochondrial membrane structure (Figure S5). In higher plants, multiple SHMT genes predicted to localize to different compartments have been described (Zhang et al., 2010). Apart from its photorespiratory role in mitochondria, the physiological roles of SHMT are not well characterized. Given the finding that the OsSHMT1-OV lines had a lower H$_2$O$_2$ concentration compared with wild type, OsSHMT1 may participate in biotic and abiotic stress responses. Genetic engineering using OsSHMT1 represents a potential way to improve the tolerance of crop plants to stress.

Differences between the Current Study and a Recently Published Report

As we were submitting the present study for review, a report entitled “Characterization and molecular cloning of a serine
TABLE 3 | Comparison of photosynthesis parameters between WT and OsSHMT1-OV rice leaves (n = 15), the asterisks shows significantly different between the OsSHMT1-OV and wild type at P ≤ 0.05.

| Parameter                  | WT          | OsSHMT1-OV lines |
|----------------------------|-------------|------------------|
| Stomatal conductance (mol/m²/s) | 0.159 ± 0.03 | 0.194 ± 0.05*    |
| Intercellular CO₂ concentration (µmol/mol) | 237.1 ± 5.2 | 333.6 ± 7.3* |
| Transpiration rate (mmol/m²/s) | 1.77 ± 0.21 | 2.62 ± 0.25*    |

The OsSHMT1-OV Lines Exhibited Increased Photosynthetic Efficiency and Improved Productivity

A particularly intriguing finding of our study is that OsSHMT1 transcript accumulation increased the photosynthetic efficiency and productivity of the plants. This phenomenon can be explained as follows. The negative impact of photorespiration on plant growth and yield has been demonstrated by doubling the CO₂ concentration in a glasshouse environment; it can dramatically increase the performance of several crops. However, such a strategy cannot be implemented in the field, where photorespiratory losses are exacerbated by high temperatures and suboptimal water supplies. Under these conditions, the stomata close and the intercellular O₂ concentration increases through the release of O₂ from the light reactions of photosynthesis. Despite these disadvantages, photorespiration is important because it recovers 75% of the carbon from phosphoglycolate and efficiently removes potent inhibitors of photosynthesis. Moreover, photorespiration dissipates excess photochemical energy under high light intensities, thus protecting chloroplasts from overreduction. Kebeish et al. (2007) introduced the Escherichia coli glycolate catabolic pathway into A. thaliana chloroplasts to reduce the loss of fixed carbon and nitrogen that occurs in C3 plants when phosphoglycolate, an inevitable by-product of photosynthesis, is recycled by photorespiration. In our study, the transgenic OsSHMT1-OV lines grew faster, produced more shoots and a greater panicle biomass, and contained more soluble sugars, reflecting reduced photorespiration and enhanced photosynthesis that correlated with an increased chloroplastic soluble sugars, reflecting reduced photorespiration and enhanced photosynthesis that correlated with an increased chloroplastic CO₂ concentration in the vicinity of ribulose-1,5-bisphosphate carboxylase/oxygenase. The greater panicle biomass of the plants indicates that the overexpression of OsSHMT1 enabled the plants to overcome at least some of the photorespiration-induced limitations on growth. This result indicates that the increase in biomass production of the transgenic plants coincided with improved photosynthesis.
In conclusion, we showed that the oscdm1 chloroplast thylakoid membrane damage may be from the excessive reactive oxygen (H₂O₂). The reactive oxygen (H₂O₂) was mainly produced in mitochondria. Mitochondria and chloroplasts are thought to have originated from prokaryotes during endosymbiotic evolution in eukaryotic cells. Intimate communication among organelles is necessary to coordinate their activities during growth, development, and other physiological processes. It is reasonable to assume that an intracellular mobile signal is transported from mitochondria to chloroplasts to trigger chloroplast change. This signal should be clarified in future investigations by taking advantage of oscdm1 plants.

Acknowledgments

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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fgene.2015.00226

Figure S1 | Schematic map of the enhancer trap vector. The upper and lower portions of the map show the T-DNA and vector backbone, respectively. LB,

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Figure S2 | Sequence alignment of SHMT homologs from soybean, rice, and Arabidopsis. Identical residues are boxed in black; similar residues are highlighted in gray. The pyridoxal phosphate binding site is marked using a red line. The OsSHMT1 SHMT domain spans residues 51–449.

Figure S3 | BN gel analysis of thylakoid membrane protein complexes. Lane 1, wild type; lane 2, oscdm1. Band I, PSI monomer and PSII dimer; II, ATP synthase; III, monomer; IV, CP43-less PSI; V, LHC II.

Figure S4 | A phylogenetic tree showing the predicted relationships among soybean, rice, and Arabidopsis. The full-length amino acid sequences of each protein were aligned using CLUSTALW and revised manually. The tree was constructed using the neighbor-joining method with full-length protein sequences from soybean, rice, and Arabidopsis. The results indicate that the SHMT protein sequences were divided into three subfamilies: SUBI [Oryza sativa: OsSHMT1 and OsSHMT3; soybean: GmSHMT1 and GmSHMT4; and Arabidopsis: ASHMT3, ASHMT4, and ASHMT5]; SUBII [Oryza sativa: OsSHMT2 and OsSHMT4; soybean: GmSHMT3 and GmSHMT5; and Arabidopsis: ASHMT1 and ASHMT2]; and SUBIII [Oryza sativa: OsSHMT5; soybean: GmSHMT2; and Arabidopsis: ASHMT6 and ASHMT7]. The bar represents the genetic distance in the phylogenetic tree.

Figure S5 | Transmission electron microscopic images of mitochondria from 1-month-old wild-type and oscdm1 mutant plants. (A) A wild-type plant. (B) An oscdm1 plant. The wild-type mitochondrial membranes were well-ordered, while the mutant mitochondrial membranes were abnormal and became irregular. Bar = 100 nm.

Table S1 | Primer sequence for test.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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