Genome-wide identification and analysis of the SGR gene family in *Cucumis melo* L.

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ABSTRACT. Chlorophyll (CHL) is present in many plant organs, and its metabolism is strongly regulated throughout plant development. Understanding the fate of CHL in senescent leaves or during fruit ripening is a complex process. The stay-green (SGR) protein has been shown to affect CHL degradation. In this study, we used the conserved sequences of STAY-GREEN domain protein (NP_567673) in *Arabidopsis thaliana* as a probe to search SGR family genes in the genome-wide melon protein database. Four candidate SGR family genes were identified in melon (*Cucumis melo* L. Hetao). The phylogenetic evolution, gene structure, and conserved motifs were subsequently analyzed. In order to verify the function of *CmSGR* genes in CHL degradation, *CmSGR1* and *CmSGR2* were transiently overexpressed and silenced using different plasmids in melon. Overexpression of *CmSGR1*
or CmSGR2 induced leaf yellowing or fruit ripening, while silencing of CmSGR1 or CmSGR2 via RNA interference delayed CHL breakdown during fruit ripening or leaf senescence compared with the wild type. Next, the expression profile was analyzed, and we found that CmSGR genes were expressed ubiquitously. Moreover, CmSGR1 and CmSGR2 were upregulated, and promoted fruit ripening. CmSGR3 and CmSGR4 were more highly expressed in leaves, cotyledon, and stem compared with CmSGR1 or CmSGR2. Thus, we conclude that CmSGR genes are crucial for fruit ripening and leaf senescence. CmSGR protein structure and function were further clarified to provide a theoretical foundation and valuable information for improved performance of melon.

Key words: Melon; Stay-green; Bioinformatic analysis; Chlorophyll degradation; Leaf senescence

INTRODUCTION

As a photoreceptor, chlorophyll (CHL) is a key component of the photosynthesis machinery and is required for the absorption of sunlight. CHL exists in chloroplast membranes where it is bound in CHL-protein complexes (Markwell et al., 1979). The protein components of the two photosystems are structurally organized into morphologically distinct membrane subunits (Arntzen, 1978), including photosystem I (PSI) and photosystem II (PSII) reaction center complexes (Kusaba et al., 2007). The final step of leaf and fruit development is senescence and degradation, which is an active process that salvages nutrients from the fruit and leaf (Matile, 2000; Pružinská et al., 2005; Hörtensteiner and Kräutler, 2011). Leaf senescence is a complex process in which multiple cellular events proceed in parallel or sequentially. Furthermore, this process requires a lot of energy, and involves a sequence of physiological and biochemical events (Lim et al., 2007; Procházková and Wilhelmová; 2007). These events are characteristically followed by the breakdown of nuclei, plastids, vacuoles, and mitochondria, and ultimately lead to cell death (Buchanan-Wollaston, 1997).

Mutants that retain green pigments during senescence are collectively called stay-green (or non-yellowing) mutants, and have been isolated from several plants (Grassl et al., 2012). Stay-green or non-yellowing mutants have been of high interest in the determination of genetic and biochemical mechanisms of CHL breakdown during leaf senescence and fruit ripening. Earlier stay-green mutants have been reported to maintain greenness in leaf or fruit longer than in wild-type plants during senescence (Thomas and Smart, 1993; Spano et al., 2003; Hörtensteiner, 2009). The breakdown of CHL is catalyzed by several CHL catabolic enzymes in an enzymatic process (Rong et al., 2013). Several genes (SGR, RCCR, PAO, PPH, NYC1, and NOL) regulating the process of leaf senescence and fruit ripening have been identified (Lim et al., 2003; Hörtensteiner and Kräutler, 2011; Luo et al., 2013). Senescence-associated gene (SGR) plays a critical role in the regulation of CHL degradation and senescence. In addition, SGR is typically upregulated during senescence (Ren et al., 2007; Hörtensteiner, 2009). Chloroplast-located proteins encoded by SGR genes in different species are well conserved in higher plants, and it is expected that SGR interacts with subunits of the light harvesting complex of PSII (Park et al., 2007; Hörtensteiner, 2009). In recent years, the functions of the products of SGR genes have been identified in Arabidopsis thaliana (Sakuraba
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et al., 2012), Oryza sativa (Jiang et al., 2007), Pisum sativum (Sato et al., 2007), Solanum lycopersicum (Akhtar et al., 1999; Luo et al., 2013), Triticum aestivum (Spano et al., 2003), and other species. Expression of SGR in Arabidopsis and tobacco induces the degradation and senescence of leaf CHL (Park et al., 2007; Ren et al., 2007; Grassl et al., 2012). Silencing of SGR by RNA interference (RNAi) leads to reduced CHL degradation in leaves and fruits (Hu et al., 2011; Zhou et al., 2011; Luo et al., 2013).

In this study, we performed a predictive computer analysis for SGR genes in melons, and four putative SGR genes were identified from the MELONOMICS (https://melonomics.net) database, which includes CM_protein_v3.5, melon_genome pseudomolecules v3.5. Phylogenetic analyses were performed and gene structures were compared to reveal the evolutionary relationships among SGRs. The transient expression of SGR by agro-infiltration in melon fruit may enhance chlorophyll degradation, while silencing of SGR by RNAi with agro-infiltration in melon fruit resulted in reduced chlorophyll degradation. In addition, to demonstrate the spatial patterns of SGR family gene expression, gene expression in seven tissues was analyzed.

MATERIAL AND METHODS

Identification of the SGR family in melon

To identify members of the SGR gene family in melon, multiple database searches were performed. An A. thaliana protein with a typical STAY-GREEN domain structure was selected as the query sequence in the National Center for Biotechnology Information (NCBI) database (accession No. NP_567673), using BLASTp (Altschul et al., 1990) at E values ≤ 10^{-3} to avoid false positives, and SGR family genes were searched in the melon genome. CM_protein_v3.5 (https://melonomics.net/files/Genome), which was used as a search database for local BLASTp. BioEdit 7.0.9 (Hall, 1999) was used to analyze homologs of SGR family genes in local BLASTp. We also performed database searches using amino acid sequences of the STAY-GREEN domain of other members of the A. thaliana SGR family as the query to confirm the completeness of the collection. To confirm the putative SGR family genes, amino acid sequences were searched for the STAY-GREEN domain using the Simple Modular Architecture Research Tool (SMART) (Schultz et al., 1998; Letunic et al., 2004). The Pfam (http://pfam.sanger.ac.uk/search) and SMART (http://smart.embl-heidelberg.de/) databases were used to confirm each predicted melon SGR protein sequence.

Gene structure analysis of the melon SGR genes

Information on melon SGR genes, including accession number, open-reading frame (ORF) length, and exon-intron structure, was retrieved from CM_protein_v3.5 (https://melonomics.net/files/Genome).

Phylogenetic analysis and characterization of melon SGR proteins

Multiple alignments of SGR proteins were carried out using the Clustal W (http://bioinformatics.ubc.ca/resources/tools/clustalw) program (Chenna et al., 2003). DNAMAN software (Version 7.0) was also used as a secondary tool to align sequences and calibrate
the results. Phylogenetic trees were constructed using MEGA 6.0 software (http://www.megasoftware.net/mega.html) (Tamura et al., 2013) with the neighbor-joining (NJ) method (Saitou and Nei, 1987) and the 1000-bootstrap replicates. The phylogenetic trees were drawn with MEGA 6.0, and the Multiple Em for Motif Elicitation (MEME) program v4.9.0 (http://meme-suite.org/tools/meme) (Bailey et al., 2009) was used to predict potential motifs in the putative SGR family gene sequences with the following parameters: optimum width of 6-50 amino acids, and the maximum number of motifs was set to five. The significance of the resulting motifs was checked in the NCBI and SMART databases (Sharma et al., 2010).

Plant materials and growth conditions

Melon (Cucumis melo L. cv Hetao) plants were used for the experiments and were grown in the field. Self-pollination was performed by hand and the pollination time was recorded and controlled between 9:00 and 11:30 a.m. Each plant retained only one fruit. The injected tissues were sampled and harvested with phenotypes. The mesocarp was frozen in liquid nitrogen after collection and then stored at -80°C for subsequent extraction of total RNA.

Vector construction and Agrobacterium-based transient transformation

Total RNA was isolated from melon fruit mesocarp. The full-length cDNA of CmSGR1 and CmSGR2 was amplified, using the forward primer: 5'-TTAATTTTATAGATGGCGAGAT-3' and 5'-TCTGTGAGAAATGAGGGTT-3', and the reverse primer: 5'-GGGAGCCAAAT AATCAAT-3' and 5'-TAAGGGGGAGCCAAATAAATC-3', respectively. Amplified fragments of these genes were cloned into the pEASY® Blunt Cloning Vector (TransGen, China), according to the manufacturer instructions, and sequenced. These were then cloned into the pPZP221 (35S-NOS) and pART27, using the primers detailed in Table S1. All vectors were transferred into Agrobacterium tumefaciens AGL1 by the freeze-thaw method, and subsequently the Agrobacterium were used for agroinfiltration into melon fruit.

Agrobacterium cultures (5 mL) were grown overnight from single colonies in Agrobacterium rhizogenes culture (YEB) medium plus selective antibiotics (rifampicin 50 mg/L, spectinomycin 20 mg/L), at 28°C, 175 rpm, which were then transferred into 50-mL induction medium (0.5% peptone, 0.5% beef extract, 0.1% yeast extract, 0.5% sucrose, 2 mM MgSO₄, 20 μM acetosyringone, 10 mM MES, pH 5.6) plus antibiotics (rifampicin 50 mg/L, spectinomycin 20 mg/L), and grown overnight to the log phase (OD ≈ 0.8), at 28°C, 180 rpm. Agrobacterium cultures were recovered by centrifugation and resuspended in MMA medium (10 mM MgCl₂, 10 mM MES, 200 μM acetosyringone, pH 5.6) plus 100 mg/L 2, 4-dichlorophenoxyacetic acid and 0.005% Tween-20. The Agrobacterium suspension was incubated at room temperature with gentle agitation (20 rpm) for a minimum of 2 h and then used for infiltration medium and injected in the fruits. Agroinjection was performed as follows: melon fruits at different stages of development were infiltrated using a 1-mL syringe with a 0.7 x 30-mm needle, which was introduced 5 to 7 mm in depth. The infiltration medium was slowly injected into the fruit. The total volume of solution injected varied at different developmental stages, with a maximum volume of 1000 μL injected into mature green melons.
RNA isolation and real-time quantitative RT-PCR expression analysis

Total RNA was extracted using RNAiso plus for polysaccharide-rich plant tissue (TaKaRa, Japan) according to the manufacturer instructions. Total RNA extracts were analyzed by UV spectrophotometry and agarose gel electrophoresis. Contaminated DNA was removed with RNase-free DNase I (TaKaRa) treatment First-strand cDNA was reverse transcribed from 0.5 μg total RNA from melon fruits at various stages of development and ripening and other tissues, and was used as a template in a 10-μL reaction mixture using PrimeScript™ RT Master Mix (Perfect Real Time) (TaKaRa) according to the manufacturer instructions. The primer sequences are listed in Table S2. Amplification of melon glyceraldehyde 3-phosphate dehydrogenase (GAPDH) gene was used as an internal control to calculate the relative fold differences based on the comparative Ct method. cDNA was amplified with SYBR® Premix Ex Taq™ (Perfect Real Time) (TaKaRa) using the Opticon 3 real-time system (BioRad, USA) in a 25-μL volume. Melting curves were generated immediately after the last cycle to exclude any influence of primer dimers. Cycle numbers, representing the point at which the fluorescence passed the cycle threshold (Ct), were analyzed, and the relative expression was calculated by the 2^−ΔΔCt method. At least three technical and three biological replicates were performed for each reaction. Data points in the quantitative real-time PCR (qRT-PCR) time course are reported as means ± SE of three biological replicates.

RESULTS

Genome-wide identification of the SGR family in melon

To detect candidate SGR homologs in the melon genome, the SGR domain of a melon SGR amino acid sequence was used as a protein BLAST query sequence using the MELONOMICS database. Four candidate SGR sequences were identified as potentially containing the SGR domain (Table 1). In order to verify its reliability, the *A. thaliana* SGR family was used as the query (accession No. NP_567673), and the results were consistent with the initial search. Four equivalent sequences were obtained using the MELONOMICS database by hidden Markov model analysis with PF12638, which contains a typical SGR domain (Table 1). SMART analysis indicated that the SGR domain of each of the four CmSGR sequence was typical in order to verify their reliability. Using those methods, five SGRs from *A. thaliana* (B3H593, Q66WT5, 891373, Q94AQ9, and O82741) and three SGRs from *S. lycopersicum* (NP_001234723, XP_004252642, and XP_004237702), two SGRs from the *Cucumis sativus* genome (Cucsa.302630 and Cucsa.077260) were detected (Table S3).

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<th>Stop</th>
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<th>Exon number</th>
<th>Intron number</th>
<th>Length (aa)</th>
<th>MW (kDa)</th>
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<td>2</td>
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*CmSGR*: *Cucumis melo* stay green protein; ORF: open reading frame; MW: molecular weight; pI: isoelectric point.
Structure analysis of the melon SGR genes

CmSGR ORF lengths ranged from 504 (CmSGR4) to 774 bp (CmSGR2), molecular weights ranged from 19.02 (CmSGR4) to 29.48 kDa (CmSGR2), and pI values ranged from 6.19 (CmSGR4) to 8.72 (CmSGR2) (Table 1). CmSGR genes were distributed on all chromosome of the melon genome. CmSGR1 and CmSGR2 genes were located on chromosome 9, and the another two were detected on chromosome 3. Based on the exon and intron structures, the number of introns differed. The number of introns in the CmSGR2 or CmSGR3 genes was three, while CmSGR1 and CmSGR4 genes contained two (Table 1 and Figure 1). Although, the protein analysis programs SMART and Pfam (Finn et al., 2014) could be used to analyze the major domains encoded by the CmSGR1, those programs were unable to identify smaller individual motifs. Hence, we used the MEME database to recognize further motifs and conserved domains shared within the SGR gene family. The motifs identified by MEME were between 6 and 50 amino acids in length. Five common motifs were observed, but the biological functions of these motifs were not known. Structure analysis revealed that the STAY-GREEN domain at the N-terminal was highly conservative. A chloroplast transit peptide was conserved motif near the N-terminal. The domains I, II, and III formed the STAY-GREEN domain. The C-terminal containing domain III possessed 2 to 6 conserved glutamines (Q) (Figure 2). Motifs 1, 2, 3, and 5 formed the configurations of the SGR domain I. Motif 4 was distinctively detected in CmSGR1 and CmSGR2, and formed domain II. Motif 4 determined C-X3-C-X-C2-F-P-X5-P domain in C-terminal conservative amino acid sequence, which was the highly homologous core region of SGRs, but the Motif 4 was deleted in CmSGR3 and CmSGR4. (Table 2 and Figure 3).

Figure 1. Structure of the CmSGR genes. Exons and introns are depicted by filled yellow boxes and single lines, respectively.

Phylogenetic and structural analyses

To explore the classifications and phylogenetic relationships of SGR proteins, we carried out phylogenetic analyses on different SGR proteins from different sequenced genomes of plants. Comprising A. thaliana, S. lycopersicum, and C. sativus, information on the SGR family was listed in Table S3. Full-length protein sequences from plants were used to construct an unrooted NJ phylogenetic tree, which was generated from the aligned full-length protein sequences of all four CmSGRs, five AtSGRs, three SlSGRs, and two CsSGRs. Arabidopsis, cucumber, tomato, and melon SGR proteins were grouped into two subfamilies via analysis of phylogenetic tree with well-supported bootstrap values (1000 replicates). The resulting
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Figure 2. Sequence alignment of SGR proteins from melon and other higher plant species. Arabidopsis (B3H593, Q66WT5, 891373, Q94AQ9, Q82741), tomato (NP_001234723, XP_004252642, XP_004237702), and cucumber (Cucsa.302630, Cucsa.077260) proteins belonging to SGR protein alignments base on the highly conserved central core of the STAY-GREREN domain were performed using DNAMAN Version 7.0. Sequences consisting of the cysteine-rich motif (C-X3-C-X-C2-F-P-X5-P) defined by Aubry et al. (2008) are shown in the red frame, Black shading with white letters, pink shading with white letters, and blue shading with black letters indicate 100, 75, and 50% sequence identity, respectively.

Table 2. Analysis of conserved motifs in the CmSGR family in melon.

<table>
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<td>Motif 5</td>
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</table>

phylogenetic tree demonstrated that CmSGRs share and evolutionary close relationship with CsSGRs. CmSGR1 and CmSGR2 were grouped into the first subfamily, and these two melon genes were clustered with CsSGR based on high sequence similarity with Cucsa.302630. The two external nodes at the end of the same clades of the phylogenetic tree were likely to represent the closest homologous gene pairs. The phylogenetic tree showed that CmSGR1 and CmSGR2 in the same clade were duplicated several times before the formation of melon species, and that this occurred after repeated events, which was similar to AtSGRs (Q66WT5...
Figure 3. Distribution of conserved motifs in four CmSGR proteins. The distribution of five conserved domains, represented by the colored boxes, was identified using CmSGR protein sequences in MEME suite 4.10.1. The order of the motif corresponds to their position in the individual protein sequences.

and O82741, 891373 and Q94AQ9) and SISGRs (NP_001234723 and XP_004252642) in Arabidopsis and tomato gene families. CmSGR3 and CmSGR4 were grouped in the second subfamily, and were most homologous to CsSGR (Cucsa.077260). Melons and cucumbers belong to the cucurbit family, and SGRs were clustered together and were less evolved following speciation. However, in Arabidopsis, those genes were generated through long-term evolution and might have species-specific functions. The combined phylogenetic analysis revealed that the ancestral gene of the SGR family originated from the divergence of other species and repeated events (Figure 4).

Figure 4. Phylogenetic relationships among melon, tomato, cucumber, and Arabidopsis SGR proteins. Neighbor-joining tree was created using the MEGA6.0 program (bootstrap value set at 1000) with full-length sequences of four melon SGR proteins (MELO3C005616P1, MELO3C005616P2, MELO3C008435P1, MELO3C008435P2) with yellow shading, three tomato (NP_001234723, XP_004252642, XP_004237702), two cucumber (Cucsa.302630, Cucsa.077260), and five Arabidopsis (B3HS93, Q66WT5, 891373, Q94AQ9, O82741) proteins.

Infiltration of melon fruit tissues with Agrobacterium

To confirm the function of the SGR gene in melon fruits, Oe-CmSGR (overexpressing) (p35S-SGR1, p35S-SGR2) and CmSGR-RNAi (pART27-SGR1, pART27-SGR2) vectors were constructed and introduced into melon fruit by Agrobacterium-based transient transformation. Agrobacterium cultures containing p35S-SGR1, p35S-SGR2, pART27-SGR1, pART27-SGR2, pPZP221 (35S-NOS), and pART27 were syringe-infiltrated into the mesocarp (mature green), respectively. About 5-7 days after infiltration, approximately 50% of the infiltrated fruit (total 50 pieces) was discarded because of decay and other factors, but the phenotype of the Oe-CmSGR1 (16%), and Oe-CmSGR2-infiltrated fruit (14%) turned a yellow color. However, control fruit infiltrated with pPZP221-35S-NOS and Agrobacterium cultures alone remained green, with a similar phenotype to the wild type (WT) fruit (Figure 5A, 5B, 5C). There was
increased expression of SGR at the mature green stage, and the highest level was reached at
the Breaker stage, where expressed induced by ripening was observed. Compared with non-
injection fruit, at about B+7 (7 days after breaker), the pART27-SGR1 (16%) and, pART27-
SGR2-infiltrated fruit (12%) exhibited a delay in turning yellow at the injection site (stay green),
and a green phenotype was retained for at least 7-10 days (Figure 5D, E, F). Approximately 40%
of the infiltrated fruit (total 50 pieces) was discarded. Those results suggested that CmSGR-
RNAi suppressed CHL degradation and exhibited a stay-green phenomenon, whose function
was similar to that of the LeSGR1 gene in tomato plants (Hu et al., 2011).

Figure 5. Recombinant vectors were used to infect melon fruit by Agrobacterium-based transient transformation. pPZP221-35S-NOS (A) and pART27 (D) plasmids were used as the control. Detached mature green melon fruits were infiltrated with Agrobacterium and transformed with p35S-SGR1 (B) and p35S-SGR2 (C) (overexpressing), pART27- SGR1 (E), and pART27-SGR2 (F) (RNAi). Ov-CmSGR was upregulated and promoted fruit ripening, which exhibited a yellow color phenotype, while CmSGR-RNAi inhibited chlorophyll degradation and exhibited a stay-green phenotype.

SGR gene expression in different tissues and at developmental stages

To determine the involvement of SGR genes in growth and development during the
melon life cycle, the expression profile of SGR family genes under normal growth conditions
was investigated by qRT-PCR in various tissues, including the root, stem, leaf, and fruit at
10, 20, 30, 40 (breaker stage), and 45 days after pollination (DAP). The results indicated
that SGRs were expressed in all tissues (Figure 6). CmSGR1 and CmSGR2 were expressed
at relatively high levels in cotyledon and at 45 DAP, with fruit maturation. Expression of
CmSGR1 and CmSGR2 was increased, and the highest expression was observed at 45 DAP.
We speculated that CmSGR1 and CmSGR2 might be related to fruit development and leaf
senescence. However, CmSGR3 and CmSGR4 were more highly expressed in leaves and
cotyledon than in fruit and root (5-fold increase). As the fruits develop, the expression levels
of CmSGR3 and CmSGR4 continuously decreased from 10 to 30 DAP, and slightly increased
from 30 to 45 DAP. The results indicated that expression of CmSGR3 and CmSGR4 were
increasing during fruit ripening and CHL degradation.

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DISCUSSION

CHL degradation is an important process of senescence and is a characteristic of fleshy fruit, in combination with anthocyanin accumulation and carotenoid retention (Park et al., 2007). CHL degreening is associated with the upregulation of CHL degradation and downregulation of CHL biosynthesis. In several fruits including melon, fruit color is used as an index of maturity. SGR genes have been observed in sgr mutants and are upregulated during senescence. In many crops, stay-green genotypes have also been shown to enhance resistance to disease and drought and possess leaves. In our study, we used SGR gene and protein sequences from A. thaliana as probes to identify CmSGR genes from C. melo L. Hetao, which were used to investigate the molecular mechanism of color development during ripening. Finally, four non-redundant SGR genes were identified and characterized in the melon genome. Although the C. melo genome is approximately 3-fold larger than the Arabidopsis genome (375 and 125 Mb, respectively) (Arabidopsis Genome Initiative, 2000; Garcia-Mas et al., 2012), the gene number in C. melo is less than that in Arabidopsis (4:7). Those results suggested that gene loss had occurred during genome duplication, the gene number in C. melo is 2-fold the SGR genes of C. sativus (4:2), and it probably support the idea that these SGR genes was highly conserved in monocot and dicot plant species.

It is widely accepted that the pattern of intron/exon positions helps in the understanding of evolutionary relationships (Hu and Liu, 2011). In the SGR genes from melon, CmSGR2 and CmSGR4 contain three introns, CmSGR3 contains four 4 introns, and CmSGR1 contains two introns (Table 1). Phylogenetic analysis showed that melon SGR proteins clustered more...
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closely with those from cucumber than with those from Arabidopsis and tomato, including highest bootstrap values of 100% in cucumber (Figure 4). There was an apparent parallel relationship between the structure of genes and the phylogeny of branches during in the late stages of evolution. The exon/intron pattern of the SGR genes may be stable in melon and cucumber. While lower bootstrap values were found between CmSGR1 and CmSGR2 (54%), higher bootstrap values were found between CmSGR3 and CmSGR4 (96%) (Figure 4) in the joined tree. This may be related to the structures of SGRs and may reflect the functional conservation of plant SGRs. MEME analysis revealed there was extensive conservation in the motif pattern within the SGR; the N-terminal is the putative chloroplast transit peptide region (motif 5). A Cys-rich motif (C-X3-C-X-C2-F-P-X5-P) is present in the C-terminal of the SGR proteins but not in the SGR-like proteins, with a variable region of 12 to 38 amino acids in length (Aubry et al., 2008; Pilkington et al., 2012). However, this motif is absent from the CmSGR3 and CmSGR4 proteins. The four conserved Cys residues, which may account for the activity of the catalytic site, may have functions in intermolecular or intramolecular crosslinking or may be involved in redox regulation (Aubry et al., 2008).

The up- or downregulation of SGR gene expression during leaf senescence or fruit ripening coincides with CHL degradation (Jiang et al., 2007; Kusaba et al., 2007; Park et al., 2007; Tang et al., 2011; Luo et al., 2013). To investigate the function of the CmSGR genes in CHL degradation, we performed a test to determine whether SGR expression could induce leaf yellowing or fruit ripening. We used the Oe-CmSGR plasmid to overexpression of the CmSGR and CmSGR-RNAi vector to repress the activity of endogenous CmSGR in melon. Oe-CmSGR (p35S-SGR1, p35S-SGR2) and CmSGR-RNAi (pART27- SGR1, pART27- SGR2) vectors were constructed and introduced into melon fruit by Agrobacterium-based transient transformation. Agrobacterium cultures containing p35S-SGR or pART27-SGR constructs were syringe-infiltrated into the mesocarp. After 5-7 days infiltration, the fruits were infiltrated with the p35S-NOS vector whether or not the fruits were retained the stay-green phenotype in control group. However, the fruits infiltrated with p35S-SGR1 or p35S-SGR2 developed a yellow color. SGR gene expression analysis showed that SGR increased at the mature green stage, and pART27-SGR1 and pART27-SGR2-infiltrated fruit exhibited suppressed SGR protein activity. We observed that CmSGR-RNAi possessed residual green pigmentation in the epidermis at the breaker +7-day stage when compared to the WT. The SGR gene directly or indirectly affected a specific component of the CHL catabolic pathway and participated in a broad biological processes (Zhou et al., 2011). Expression of SGR increased at the mature green stage and reached its highest level at the Br stage, displayed ripening-induced expression. However, silencing the CmSGR gene using RNAi, delayed CHL breakdown during fruit ripening or leaf senescence; this result is consistent with that obtained using SGR1-RNAi in tomato fruits (Luo et al., 2013). We also performed quantitative PCR analysis to examine expression profiles. These results are consistent with previous observations in the literature. We found SGR to be expressed ubiquitously with melon fruit ripening. CmSGR1 and CmSGR2 were upregulated and promoted fruit ripening, and were not strongly expressed in the root and stem. Compared with other organs, CmSGR3 and CmSGR4 were expressed at lower levels during the course of melon fruit development, while CmSGR3 and CmSGR4 were more highly expressed in leaves, cotyledon, and stem than in fruit and root. CmSGR genes showed tissue-specific expression patterns in melon. Since the roots do not contain chloroplasts, there was minimal expression of the CmSGR genes in the roots. With fruit ripening, CmSGR1 and CmSGR2 play an important role in CHL degradation. CmSGR3 and CmSGR4 exhibit tissue-specific expression in leaves, cotyledon, and stem, but are not strongly expressed in fruit.
Conflicts of interest

The authors declare no conflict of interest.

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REFERENCES


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Identification and analysis of the SGR gene


## Supplementary material

### Table S1
List of primers used in this study.

### Table S2
List of QPCR-primers used in this study.

### Table S3
List of protein sequences used in this study.

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