During leaf senescence, plants degrade chlorophyll to colorless linear tetrapyrroles that are stored in the vacuole of senescing cells. The early steps of chlorophyll breakdown occur in plastids. To date, five chlorophyll catabolic enzymes (CCEs), NONYELLOW COLORING1 (NYC1), NYC1-LIKE, pheophytinase, pheophorbide a oxygenase (PAO), and red chlorophyll catabolite reductase, have been identified; these enzymes catalyze the stepwise degradation of chlorophyll to a fluorescent intermediate, pFCC, which is then exported from the plastid. In addition, STAY-GREEN (SGR), Mendel’s green cotyledon gene encoding a chloroplast protein, is required for the initiation of chlorophyll breakdown in plastids. Senescence-induced SGR binds to light-harvesting complex II (LHCII), but its exact role remains elusive. Here, we show that all five CCEs also specifically interact with LHCII. In addition, SGR and CCEs interact directly or indirectly with each other at LHCII, and SGR is essential for recruiting CCEs in senescing chloroplasts. PAO, which had been attributed to the inner envelope, is found to localize in the thylakoid membrane. These data indicate a predominant role for the SGR-CCE-LHCII protein interaction in the breakdown of LHCII-located chlorophyll, likely to allow metabolic channeling of phototoxic chlorophyll breakdown intermediates upstream of nontoxic pFCC.

INTRODUCTION

Leaf senescence is a genetically determined and highly ordered process that constitutes the final stage of leaf development. It remobilizes nutrients, in particular nitrogen and phosphorus, to sink organs, such as storage tissues or seeds. On the cellular level, the most significant early changes occur in the chloroplasts, where grana membranes are unstacked and overall thylakoid membrane density is reduced. This catabolic process is accompanied by the massive degradation of chloroplast proteins, which constitute ~70% of total cellular protein (Hörtensteiner and Feller, 2002). By contrast, the chloroplast envelope remains intact late into senescence, indicating that cellular compartmentalization is maintained during senescence (Matile et al., 1999; Thomas et al., 2003). This is in agreement with the observation that leaf senescence can be experimentally reversed (Zavaleta-Mancera et al., 1999a, 1999b), up to a “point of no return,” beyond which senescence is followed by death (Guilboineau et al., 2010). Hence, cellular senescence is seen as a transdifferentiation rather than a cell death process, and viability needs to be maintained to allow senescence initiation and progression (Thomas et al., 2003).

Loss of green color is the visible symptom of leaf senescence and is caused by the degradation of chlorophyll. In recent years, a pathway has been elucidated that is active during senescence and converts chlorophyll to colorless linear tetrapyrroles, so-called nonfluorescent chlorophyll catabolites (NCCs), as end products of chlorophyll breakdown (Hörtensteiner, 2006; Kräutler, 2008; Hörtensteiner and Kräutler, 2011). The early, plastid-localized reactions of the pathway end with the formation of a primary fluorescent chlorophyll catabolite (pFCC). After export from the plastid, several peripheral side chains of pFCC are modified in the cytosol to produce a species-specific variety of FCCs. Finally, after import into the vacuole, these modified FCCs are isomerized to their respective NCCs in a nonenzymatic reaction driven by the acidic vacuolar pH (Oberhuber et al., 2003). Interestingly, all except one of the NCCs identified to date are derived from chlorophyll a, and conversion from chlorophyll b to chlorophyll a was shown to be a prerequisite for chlorophyll breakdown (Hörtensteiner et al., 1995; Hörtensteiner and Kräutler, 2011). Consequently, mutants deficient in chlorophyll b reductase, catalyzing the first of two consecutive reactions of chlorophyll b to chlorophyll a reduction, develop a stay-green phenotype and
retain large quantities of chlorophyll, in particular chlorophyll b. In *Arabidopsis thaliana* and rice (*Oryza sativa*), chlorophyll b reductase is encoded by two orthologous genes each, *NONYELLOW COLORING1* (NYC1) and *NYC1-LIKE* (NOL) (Kusaba et al., 2007; Horie et al., 2009; Sato et al., 2009). Recently, 7-hydroxymethyl chlorophyll a reductase (HMCOR), catalyzing the second step of conversion of chlorophyll b to chlorophyll a, has been identified at the molecular level (Meguro et al., 2011).

The plastid-located part of the chlorophyll degradation pathway starts with the removal of the central Mg atom by a metal chelating substance, whose molecular nature is as yet unknown, and is followed by phytol hydrolysis yielding phaeophorbide (Pheide) a. Dephylation was for a long time considered to be catalyzed by chlorophyllase (i.e., to precede Mg dechelation and to yield chlorophyllide as an intermediate) (Takamiya et al., 2000). However, recent investigation of leaf senescence in *Arabidopsis* and rice showed that, instead, phaeophytinase (PPH) is active, which specifically dephylates phaeophytin (Mg-free chlorophyll), but does not accept chlorophyll as substrate (Morita et al., 2009; Schelbert et al., 2009; Ren et al., 2010). Next, the chlorin macrocycle of Pheide a is oxygenolytically opened by a Rieske-type monooxygenase, termed Pheide a oxygenase (PAO) (Pružinská et al., 2003, 2005). The product of this reaction, red chlorophyll catabolite (RCC), is then reduced to pFCC in a regio- and stereoselective manner catalyzed by RCC reductase (RCCR) (Pružinská et al., 2007). Biochemical and two-hybrid experiments indicated interaction between PAO and RCCR as well as metabolic channeling of RCC (Rodoni et al., 1997; Pružinská et al., 2007). PAO activity provides the structural basis for all further breakdown products (i.e., RCCs, FCCs, and NCCs). Therefore, this pathway is termed the PAO pathway (Hörtensteiner and Kräutler, 2011).

Screening for stay-green mutants in many species uncovered a novel chloroplast-located protein, termed STAY-GREEN (SGR) (Hörtensteiner, 2009), whose function is considered to be related to chlorophyll breakdown, but is not a chlorophyll catabolic enzyme (CCE) itself. SGR was shown to specifically interact with light-harvesting complex subunits of photosystem II (LHCII) but not with core complexes or LHCI subunits (Park et al., 2007). It is assumed that SGR interaction with LHCII may trigger destabilization of these chlorophyll-apoprotein complexes as a prerequisite for the subsequent degradation of both chlorophyll and apoproteins (Park et al., 2007; Hörtensteiner, 2009). In line with this is the observation that besides retention of chlorophyll, sgr mutants in various plants also retain large quantities of LHCII subunits (Jiang et al., 2007; Park et al., 2007; Aubry et al., 2008). The same is true for other sgr mutants caused by a deficiency in either NYC1 or PPH (Kusaba et al., 2007; Horie et al., 2009; Morita et al., 2009; Schelbert et al., 2009), and it has been assumed that the concerted activity of these three proteins is required for the initiation of LHCII protein degradation during leaf senescence (Schelbert et al., 2009). By contrast, deficiency in PAO or RCCR results in an accelerated cell death phenotype, which is caused by the accumulation of the substrates of respective reactions, Pheide a or RCC (Mach et al., 2001; Pružinská et al., 2003, 2005, 2007). These colored intermediates of chlorophyll breakdown are potentially phototoxic, and tight control of the PAO pathway has been considered important to prevent premature cell death during senescence (Hörtensteiner, 2004, 2006).

Using different complementary methods, including yeast two-hybrid analysis, in vitro and in vivo pull-down assays, and bimolecular fluorescence complementation (BiFC), we provide evidence that SGR and five CCEs, involved in the conversion of chlorophyll to pFCC, localize to LHCII and molecularly interact with each other. Hence, during active chlorophyll breakdown, dynamic SGR-CCE-LHCII protein interaction occurs at the thylakoid membrane. The likely role of these interactions is to metabolically channel chlorophyll breakdown pigments to minimize the risk of photodamnism of these light-excitable intermediates and, thus, to prevent accelerated cell death during leaf senescence.

**RESULTS**

**Arabidopsis Plants Expressing Epitope-Tagged SGR or CCEs Exhibit Enhanced Chlorophyll Breakdown during Senescence**

During leaf senescence, SGR and five CCEs (RCCR, PAO, PPH, NYC1, and NOL) have been identified as essential components of chlorophyll degradation (Hörtensteiner and Kräutler, 2011).

![Figure 1](image-url)

*Figure 1. Accelerated Leaf Yelllowing of Arabidopsis Plants Constitutively Expressing GFP-Tagged SGR or CCEs during Dark-Induced Senescence.*

Three-week-old plants grown under long-day conditions were used in this study. Photographs were taken from whole plants ([A]) or detached leaves ([B]) before (0 DDI) or after incubation in darkness for 4 d (4 DDI; [A] and [B]). WT, wild type. Bar = 5 cm.
For this study, we produced Arabidopsis transgenic lines that constitutively expressed SGR or one of the five CCEs as fusion proteins with green fluorescent protein (GFP), tandem affinity purification (TAP), or glutathione S-transferase (GST) tags (see Supplemental Table 1 online). Transgenic lines with the highest levels of transgene expression were selected, and correct sizes of fusion proteins were verified by immunoblot analysis using antibodies against GFP (α-GFP), myc (α-myc; for TAP detection), and GST (α-GST). All the GFP-tagged SGR and CCEs were mostly detected in membrane-enriched fractions and barely in soluble fractions of total protein extracts (see Supplemental Figure 1 online).

To examine the effects of constitutive expression of GFP-tagged CCEs on chlorophyll degradation during leaf senescence, we used 3-week-old plants to dark-induce senescence in whole plants (Figure 1A) or detached leaves (Figure 1B). Before dark incubation (0 d of dark incubation [DDI]), chlorophyll levels of chlorophyll a/β ratios (Table 1) of these transgenic plants were almost indistinguishable from the wild-type plants. However, accelerated leaf yellowing (Figure 1) and reduced chlorophyll a levels and chlorophyll a/β ratios (Table 1) were observed at 4 DDI in both whole plants and detached leaves compared with the wild-type plants. In addition, because of an assumingly enhanced chlorophyll b reductase activity, 35S:PPH-GFP and 35S:NOL-GFP plants exhibited higher chlorophyll a/β ratios (Table 1). These plants exhibited a similar phenotype under natural senescence-induced leaves (see Supplemental Figure 3 online). After senescence induction, SGR, RCCR, PAO, and NY1C1, but not NOL, were significantly higher expressed in the PPH-GFP overexpressing plants than in the wild-type plants. These results indicate that constitutive expression of GFP-tagged CCEs is not sufficient to activate chlorophyll degradation during vegetative growth, but significantly accelerates chlorophyll degradation during leaf senescence, likely through transcriptional coactivation of other genes of the pathway.

### Table 1. Chlorophyll Levels of 3-Week-Old Transgenic Arabidopsis Plants Expressing Tagged Versions of SGR and CCEs

<table>
<thead>
<tr>
<th>Arabidopsis Transformants</th>
<th>Before Dark Incubation (0 DDI)</th>
<th>After 4 DDI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total Chlorophyll&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Chlorophyll a/β Ratio</td>
</tr>
<tr>
<td>Whole plants</td>
<td>Sample No.</td>
<td>Mean</td>
</tr>
<tr>
<td>Wild type</td>
<td>7</td>
<td>1885</td>
</tr>
<tr>
<td>35S:SGR-GFP</td>
<td>6</td>
<td>1821</td>
</tr>
<tr>
<td>35S:RCCR-GFP</td>
<td>5</td>
<td>1877</td>
</tr>
<tr>
<td>35S:NYC1-GFP</td>
<td>6</td>
<td>1865</td>
</tr>
<tr>
<td>35S:NOL-GFP</td>
<td>5</td>
<td>1760</td>
</tr>
<tr>
<td>35S:PPH-GFP</td>
<td>5</td>
<td>1843</td>
</tr>
<tr>
<td>35S:PAO-GFP</td>
<td>5</td>
<td>1885</td>
</tr>
<tr>
<td>Detached leaves</td>
<td>Sample No.</td>
<td>Mean</td>
</tr>
<tr>
<td>Wild type</td>
<td>5</td>
<td>1865</td>
</tr>
<tr>
<td>35S:SGR-GFP</td>
<td>7</td>
<td>1818</td>
</tr>
<tr>
<td>35S:RCCR-GFP</td>
<td>7</td>
<td>1892</td>
</tr>
<tr>
<td>35S:NYC1-GFP</td>
<td>7</td>
<td>1954</td>
</tr>
<tr>
<td>35S:NOL-GFP</td>
<td>6</td>
<td>1734</td>
</tr>
<tr>
<td>35S:PPH-GFP</td>
<td>5</td>
<td>1865</td>
</tr>
<tr>
<td>35S:PAO-GFP</td>
<td>7</td>
<td>1834</td>
</tr>
</tbody>
</table>

<sup>a</sup>Unit of total chlorophyll is nmol mg<sup>-1</sup> fresh weight.

### SGR and CCEs Specifically Interact with LHCII at the Thylakoid Membrane

Previously, we showed that SGR interacts with LHCII in vitro and in vivo (Park et al., 2007). Using α-GFP– and α-GST–conjugated beads for in vivo pull-down assays with membrane-enriched fractions of nonsenescent (0 DDI) or senescence-induced (3 DDI) 35S:SGR-GFP plants, we found that SGR-GFP, which is constitutively present in this line, interacts with LHCII regardless of the senescence conditions (see Supplemental Figure 4 online). We extended this analysis by testing whether CCEs also interact with LHCII and/or other photosystem proteins. For this, we performed in vivo pull-down assays with nonsenescent (0 DDI) GFP- or GST-tagged transgenic plants using α-GFP– and α-GST–conjugated beads, respectively, followed by immunoblot analysis with antibodies against three photosystem proteins (α-Lhcb1, α-Lhca1, and α-CP43). Like SGR (Figure 2A), all five CCE proteins were communoprecipitated with Lhcb1, but not with Lhca1 or CP43 (Figures 2B to 2F), indicating that not only SGR but also chloroplast-located CCEs bind to LHCII at the thylakoid membrane.

### Simultaneous Pull-Down of SGR and CCEs at the Thylakoid Membrane in Senescing Chloroplasts

The interaction of SGR and the five CCEs with LHCII indicated the formation of a large chlorophyll catabolic complex for chlorophyll breakdown during leaf senescence. To investigate this possibility, 35S:PPH-GFP plants were employed for in vivo pull-down assays. To this end, intact plants were senescence-induced by 3 DDI, and

*Chlorophyll Detoxification* 3 of 12
and the membrane-enriched fractions were treated with α-GFP–conjugated beads. As revealed using α-SGR, α-RCCR, α-PAO, α-NYC1, and α-NOL, we found that all tested endogenous proteins (i.e., SGR, RCCR, PAO, NYC1, and NOL) were coimmunoprecipitated with PPH-GFP (Figure 3).

Pairwise Interactions among SGR and CCEs in Yeast Two-Hybrid and in Vitro Pull-Down Assays

Although the results shown in Figure 3 supported the possibility of multiprotein complex formation, they left the question open, whether the observed coimmunoprecipitation between SGR and CCEs solely occurred through their interaction with LHCII or through sole bilateral interaction between pulled PPH and the other proteins, or whether more complex patterns of direct interaction might exist among SGR and CCEs. For example, in vitro interactions of NYC1-NOL and PAO-RCCR have been reported (Pruzinská et al., 2007; Morita et al., 2009). To address this, we first examined pairwise interactions among them by measuring β-galactosidase activity in yeast two-hybrid assays (see Methods for further details). We found significant interactions between SGR and each of the five CCEs and between RCCR and the other four CCEs (Figure 4). Among the latter CCEs (PAO, PPH, NYC1, and NOL), only interactions of PAO-PPH and NYC1-NOL were significant. To eliminate possible interference of the N-terminal chloroplast-targeting sequences of the proteins investigated in the yeast two-hybrid assays, we further examined pairwise interactions by in vitro pull-down assays using the membrane-enriched fractions of GFP-tagged transgenic lines (Figure 5; see Supplemental Figure 5 online). The results were consistent with the yeast two-hybrid interactions (Figure 4). Together, these data strongly suggested that SGR (and possibly RCCR) may act as key players to recruit other CCEs into a possible multiprotein complex for rapid and safe chlorophyll breakdown during leaf senescence.

In Vivo Interactions among SGR and CCEs in Senescing Chloroplasts

Next, we used BiFC as an alternative method to analyze pairwise interactions among SGR and CCEs in vivo (Figure 6; see Supplemental Figure 6 online). Different combinations of SGR-CCE proteins that were fused to either the N- or C-terminal half of yellow fluorescent protein (YFPn or YFPc, respectively) were cotransformed into mesophyll protoplasts isolated from 0 DDI green or 4 DDI senescent leaf tissues. As a positive control for protein–protein interaction that is unrelated to chlorophyll breakdown, we used the two halves of YFP fused to either phosphoribulokinase (PRK) or chloroplast protein 12 (CP12), two proteins that have been shown to form a complex in chloroplasts (Scheibe et al., 2002) (Figure 6B). Most positive interactions among SGR and CCEs described above (Figures 4 and 5; see Supplemental Figure 5 online), including SGR-PAO and NYC1-NOL, also gave positive BiFC fluorescence signals (Figure 6A; see Supplemental
However, we were unable to verify the in vitro interaction of SGR-PPH or SGR-NOL, notably combinations that resulted in rather weak interaction in yeast two-hybrid assays (Figure 4). Surprisingly, YFP fluorescence signals were only obtained in protoplasts isolated from senescent leaves (4 DDI) but not in nonsenescent protoplasts (0 DDI) (Figure 6A).

Based on yeast two-hybrid and in vitro pull-down analysis, we suggested that SGR may act as a key player for protein interaction. In order to address this possibility, we analyzed in vivo interaction between PAO and RCCR in the Arabidopsis sgr

nye1-

1 mutant, by BiFC. In contrast with the wild type, YFP fluorescence was absent in senescent protoplasts of nye1-

1. Functionality of the nye1-

1 protoplasts was confirmed by positive YFP fluorescence when using the PRK/CP12 control. These results indicated that presence of SGR in senescing chloroplasts is a prerequisite for CCE protein interaction.

Taking all the results together, we propose that SGR and CCEs interact directly and indirectly with each other to possibly form a large SGR-CCE-LHCII multiprotein complex at the thylakoid membrane during leaf senescence.

**Figure 3.** Coimmunoprecipitation of All CCEs and SGR in Senescing Chloroplasts.

35S:GFP and 35S:PPH-GFP transgenic plants grown for 3 weeks under long-day conditions were transferred to darkness and sampled at 3 DDI. Membrane-enriched fractions were used for in vivo pull-down assays. For this, GFP was immunoprecipitated (GFP-IP) with α-GFP–conjugated beads. Native SGR, RCCR, PAO, NYC1, and NOL in the input samples (left panel) and the pulled fractions (right panel) were detected using respective antibodies. The expression of GFP (negative control) and PPH-GFP were confirmed by α-GFP.

**PAO Localizes to the Thylakoid Membrane**

Based on their primary structures, all CCEs except PAO and NYC1 are soluble proteins. NYC1 and NOL have been shown to localize at the thylakoid membrane in rice (Sato et al., 2009). By contrast, PAO activity was attributed to the chloroplast envelope in barley (Hordeum vulgare; Matile and Schellenberg, 1996), and proteome analyses also favored envelope localization for PAO in Arabidopsis (Joyard et al., 2007). However, this proposed envelope localization of PAO conflicted with its proposed interaction with LHCII, SGR, and other CCEs at the thylakoid membrane as described here. Therefore, we readdressed the subchloroplast localization of PAO in mesophyll protoplasts by transiently expressing a PAO-GFP fusion protein (Figure 7). As a positive control for inner envelope localization, we used a GFP-tagged translocon at the inner chloroplast envelope 110 (TIC110-GFP), a component of the chloroplast protein import machinery. PAO-GFP fluorescence signals entirely overlapped with chlorophyll autofluorescence, whereas TIC110-GFP specifically labeled the chloroplast envelope (Figure 7A). In addition, using the wild-type plants, we separated chloroplast membranes by Suc density gradient centrifugation and investigated the distribution of PAO along with chloroplast membrane marker proteins by immunoblot
analysis of individual density gradient fractions (Figure 7B). PAO clearly comigrated with chlorophyll a/b binding proteins (CAB) but not with an envelope (TOC75) or a plastoglobule (PGL35) marker. Together, these data indicate a thylakoid rather than an envelope localization of PAO, which is in agreement with the finding presented here that PAO is a component of a possible chlorophyll degrading protein complex at the thylakoid membrane.

DISCUSSION

Chlorophyll breakdown is an integral process of senescence, the final part of leaf development. In this respect, loss of green color visually marks the initiation of dramatic metabolic changes that occur during this final phase of development (Lim et al., 2007). Among other processes, senescence is accompanied by a loss of photosynthetic capacity and the massive degradation of cellular proteins. These processes remobilize nutrients from senescing leaves and occur in living cells (i.e., before the ultimate death of the cell). However, chlorophyll breakdown is seen as a detoxification rather than a remobilization process (Hörtensteiner, 2009; Hörtensteiner and Kräutler, 2011). This view is supported by the fact that NCCs, identified as final products of chlorophyll breakdown (Kräutler et al., 1991; Kräutler, 2008), do not absorb visible light and thus are photodynamically safe; however, they still contain the four moles of nitrogen that are also present in chlorophyll. Furthermore, downstream steps of chlorophyll breakdown (i.e., FCC hydroxylation, conjugation, and excretion to the vacuole) resemble the three-step process of plants that is active for the detoxification of toxic endogenous and xenobiotic compounds, such as herbicides (Kreuz et al., 1996). Finally, mutants that are defective in several steps of chlorophyll breakdown develop an accelerated cell death phenotype, which has been attributed to the accumulation of respective phototoxic chlorophyll breakdown intermediates (Pružinska et al., 2003, 2005, 2007; Tanaka et al., 2003). Among the chlorophyll breakdown intermediates generated in the PAO pathway, all the ones upstream of pFCC have the potential to generate singlet oxygen in light, causing toxicity. All this information implies that the steps of chlorophyll breakdown required to produce pFCC need to be tightly controlled and accumulation of chlorophyll intermediates upstream of pFCC must be prevented or minimized during senescence.

Here, we show that such a control could, at least in part, be accomplished by metabolic channeling of chlorophyll to pFCC through dynamic interaction between SGR and CCEs, thereby likely forming a multiprotein complex, which specifically interacts with LHCII (Figure 8).

Possible Formation of a CCE Complex at the Thylakoid Membrane and Specific Interaction with LHCII

The results of these experiments, which included different in vitro and in vivo methods, are summarized in Supplemental Figure 7...
online. Not all tested pairwise protein combinations resulted in positive interaction, and different methods yielded partially different results. It remains unclear, however, whether this was due to limitations of the used respective methods or whether this might indicate rather dynamic and possibly only transient interaction among SGR and CCEs in vivo. Nevertheless, in vivo pull-down experiments (Figure 3) confirmed coimmunoprecipitation of all tested CCEs and SGR, indicating that they indeed might form a multiprotein complex, possibly with varying protein composition.

We considered the thylakoid membrane as the likely site of protein interaction because SGR had been demonstrated to interact with LHCII (Park et al., 2007), and NYC1, in complex with NOL, had been suggested to localize to thylakoids (Sato et al., 2009). Here, we provide evidence that also PAO, which had been proposed to localize to the chloroplast envelope (Matile and Schellenberg, 1996; Joyard et al., 2009), resides in thylakoid membranes (Figure 7). Using in vivo pull-down experiments, we demonstrate that the SGR/CCE complex components specifically interact with LHCII (Figure 2), in agreement with the presence of GFP-tagged SGR/CCE proteins in membrane-enriched fractions rather than soluble fractions (see Supplemental Figure 1 online). This specificity for LHCII is surprising because during senescence, chlorophyll in both LHCI and LHCII (and probably also core complexes) is degraded. In accordance with this, sgr mutants retain both LHCI and LHCII subunits during senescence (Park et al., 2007; Sato et al., 2007), suggesting the involvement of SGR in chlorophyll degradation in the antenna of both photosystems. Nevertheless, SGR specifically binds to LHCII (Figure 2; see Supplemental Figure 4 online) (Park et al., 2007). By contrast, other sgr mutants that are deficient in PPH or NYC1 specifically retain LHCII subunits, with comparably minor alterations of LHCI compared with the wild-type plants (Kusaba et al., 2007; Horie et al., 2009; Morita et al., 2009; Sato et al., 2009; Schelbert et al., 2009). This together with a particularly high retention of chlorophyll b in these mutants, which is indicative of a LHCII-related defect, challenges their role in degradation of chlorophyll from LHCI. However, we cannot rule out the possibility that SGR/CCEs might interact with other LHCI subunits, which were not tested in this work. In summary, published data are conflicting with respect to the specificity of SGR/CCEs for particular LHCs, but the data presented here indicate an interaction of SGR/CCEs specifically with LHCII, leaving open the question whether degradation of LHCI-located chlorophyll involves CCEs without direct contact to LHCI or whether other, so far unknown, enzymes or localizing proteins are required.

### Regulatory Role of SGR

Among the different experiments that we performed to demonstrate SGR/CCE interactions, BiFC analysis yielded particularly interesting results because, despite the use of a constitutive 35S promoter for expression, none of the positive interactions observed in senescent protoplasts were found in the wild-type protoplasts before senescence induction (Figure 6). By contrast, interactions of SGR/CCEs with LHCII were senescence independent when constitutively expressing particular

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**Figure 6.** In Vivo Interactions among SGR and CCEs Analyzed by BiFC.  
(A) For BiFC assays, construct pairs expressing fusions between SGR, PAO, NYC1, or NOL, and the N- or C-terminal half of YFP (YFPn or YFPc, respectively), were coexpressed in Arabidopsis mesophyll protoplasts isolated from 0 DDI green or 4 DDI senescing leaves. Confocal microscopy analysis was performed after 24 h. Note that YFP fluorescence was not detected at 0 DDI. Further positive BiFC interactions among SGR and CCEs are shown in Supplemental Figure 6 online. Auto, chlorophyll autofluorescence. Bars = 10 μm.  
(B) As a positive control for chloroplast-located BiFC interaction, PRK and CP12 fusions were used. Note that positive interaction was detected in both 0 DDI and 4 DDI protoplasts. Bars = 10 μm.  
(C) BiFC interaction between PAO and RCCR was positive in the wild-type protoplasts at 4 DDI but negative in the Arabidopsis sgr mutant nye1-1. As a control, positive interaction of PRK and CP12 was demonstrated in nye1-1. Bars = 10 μm.
SGR/CCE proteins (see Supplemental Figure 4 online). Thus, SGR/CCEs could most likely also bind to LHCII in BiFC experiments at 0 DDI. Despite this, SGR/CCEs interaction was not observed at 0 DDI, implying that additional, senescence-specific components were required to allow pairwise interaction of the chlorophyll catabolic proteins under investigation in each experiment. The nature of such potential factors remains elusive, but SGR was a particularly interesting candidate for interaction regulation because SGR does not exhibit a known catalytic activity in chlorophyll breakdown but could have a structural and/or regulatory function instead (Park et al., 2007; Hörtensteiner, 2009). In addition, modulation of SGR abundance seems to correlate with overall chlorophyll breakdown. Thus, it was shown that during dark-induced senescence of PAO-deficient pao1 and acd1 mutants, SGR expression is severely inhibited (Park et al., 2007), probably with the aim to decrease levels of chlorophyll breakdown if the PAO pathway is blocked. We used nye1-1 to test the role of SGR in CCE protein interaction. Indeed, the well-established interaction between PAO and RCCR (Rodoni et al., 1997; Pružinská et al., 2007) that also gave positive BiFC results in senescent wild-type cells was disabled in senescent nye1-1 protoplasts (Figure 6). This strongly supported the possibility that SGR enables interactions among CCEs after they are bound to LHCII.

Control of Reactive Chlorophyll Metabolites and Mechanism of Chlorophyll-Apoprotein Complex Degradation

The data presented here provide evidence for senescence-related, possibly dynamic, interaction of SGR and CCEs at the thylakoid membrane, specifically interacting with LHCII, that is required for chlorophyll to pFCC conversion. Very likely, such SGR-CCE-LHCII protein interaction allows metabolic channeling of chlorophyll breakdown intermediates, thereby minimizing the risk of chlorophyll intermediate accumulation and potential phototoxicity as seen in the mutants deficient in PAO or RCCR (Tanaka et al., 2003; Pružinská et al., 2005, 2007). In line with this, during normal senescence, none of these phototoxic chlorophyll intermediates accumulates (Pružinská et al., 2005, 2007; Schelbert et al., 2009). Furthermore, induction of chlorophyll breakdown in senescent chloroplasts causes in organello accumulation of pFCC but not of an upstream intermediate of the pathway (Matile et al., 1992; Ginsburg et al., 1994). Surprisingly, however, the mutants deficient in SGR, NYC1, or PPH (i.e., upstream of phytol cleavage) do not show cell death phenotypes (Kusaba et al., 2007; Ren et al., 2007; Schelbert et al., 2009), although in these mutants, the potential of photodynamic effects through excitation of nondegraded chlorophyll increases. It has been argued that in these sgr mutants, chlorophyll is retained.
Do Physical Interaction and Complex Formation Occur during Chlorophyll Biosynthesis?

Metabolome complex formation and metabolic control through channeling likely occurring in chlorophyll breakdown as shown here are known from biosynthetic pathways of several secondary plant compounds (Jørgensen et al., 2005). For chlorophyll biosynthesis, control of metabolite flux is also important. This is seen in the mutants and antisense lines that are impaired in certain steps of chlorophyll synthesis; several of these mutants develop accelerated cell death phenotypes that can be attributed to the accumulation of phototoxic intermediates in chlorophyll synthetic pathway (Tanaka and Tanaka, 2007). However, regulation of chlorophyll intermediate formation during chlorophyll synthesis might be different during chlorophyll degradation. This idea is supported by the fact that besides other regulatory mechanisms, chlorophyll synthesis is under tight metabolic feedback control of δ-aminolevulinic acid synthesis, the rate-limiting step of the pathway (Mochizuki et al., 2010), while direct metabolite feedback control has not been demonstrated for a CCE. Furthermore, although pairwise interaction has been demonstrated for some of the chlorophyll synthetic enzymes (Tanaka and Tanaka, 2007), the existence of a hypothetical megacomplex of chlorophyll synthetic enzymes (Shlyk, 1971) remains to be demonstrated (Tanaka and Tanaka, 2007).

METHODS

Plant Materials and Growth Conditions

*Arabidopsis thaliana* wild-type (Columbia-0 ecotype) and transgenic plants were grown on soil in a growth chamber at 21 to 22°C under cool-white fluorescent light (90 to 100 μmol photons m⁻² s⁻¹) under long-day (16 h light/8 h dark) conditions. For protoplast transformation and Suc density gradient centrifugation experiments, wild-type plants were grown for 5 and 8 weeks, respectively, at short-day (8 h light/16 h dark) with fluence rates of 100 to 200 μmol photons m⁻² s⁻¹. Rosette leaves were used for pigment content, immunoblot, and pull-down analyses. For dark-induced senescence of whole plants, 3-week-old plants were transferred to complete darkness. After dark incubation, rosette leaves were sampled at weak green light. For dark treatment of detached leaves, the oldest but still green rosette leaves were incubated on wet filter paper soaked with 3 mM MES, pH 5.7, buffer in complete darkness at 23°C.

Plasmid Construction and *Arabidopsis* Transformation

*Arabidopsis* full-length cDNAs (no stop codon) of SGR, PPH, PAO, RCCR, NYC1, and NOL were PCR amplified. After insertion into the Gateway
entry vector pCR8/GW/TOPO (Invitrogen), inserts were recombined into the binary Gateway vector pEarleyGate 103, thereby introducing a C-terminal GFP-His tag. In addition, using either pEarleyGate 205 (TAP tag) (Earley et al., 2006) or pCAMBIA-GST (GST tag), we introduced alternative C-terminal tags. The primers used for cloning are listed in Supplemental Table 2 online. In all cases, transgene expression was driven by the constitutive 35S promoter. Arabidopsis transgenic plants (see Supplemental Table 1 online) were obtained by Agrobacterium tumefaciens (strain GV3101)–mediated transformation through a floral dipping method (Zhang et al., 2006). As negative controls, transgenic plants transformed with empty pEarleyGate 103 or pEarleyGate 205 or with a chlorophyllase 1 (CLH) (At1g19670)-GST fusion construct were used. Transgenic plants were selected based on the highest expression of tagged proteins in 2 DDI leaf tissues as determined by immunoblot analysis using α-myc (for TAP detection; Santa Cruz Biotechnology), α-GST (Santa Cruz Biotechnology), or α-GFP (Abcam).

Gene Expression Analysis

The mRNA levels of SGR and CCE genes were measured by quantitative real-time PCR (qRT-PCR) analysis. Total RNA was extracted from the rosette leaves using the Total RNA Extraction Kit including RNase-free DNase (iNtRON Biotechnology). First-strand cDNA was synthesized with g of total RNA using M-MLV reverse transcriptase and an oligo(dT)15 primer (Promega) in 20 μl mixture. Then, the reaction was diluted fivefold with water and the cDNA used as template for qRT-PCR. The qRT-PCR mixture (20 μl) contained 2 μl of cDNA template, 10 μl of 2 × LightCycler 480 SYBR Green I Master 1 Master (Roche), and 0.25 μM of forward and reverse primers for each gene (see Supplemental Table 2 online). Reactions were performed using the Light Cycler 2.0 instrument (Roche Diagnostics). The transcript levels of each gene were normalized against those of GAPDH (glyceraldehyde phosphate dehydrogenase; At1g16300) as previously reported (Sakuraba et al., 2010).

Yeast Two-Hybrid Assay

Arabidopsis full-length cDNAs of SGR, NYC1, NOL, PPH, PAO, and RCCR in entry vectors were inserted into the destination vectors pDEST32 (bait) and pDEST22 (prey) (Invitrogen). The yeast strain RCCR in entry vectors were inserted into the destination vectors Arabidopsis Yeast Two-Hybrid Assay reported (Sakuraba et al., 2010). The mRNA levels of SGR and CCE genes were measured by quantitative real-time PCR (qRT-PCR) analysis. Total RNA was extracted from the rosette leaves using the Total RNA Extraction Kit including RNase-free DNase (iNtRON Biotechnology). First-strand cDNA was synthesized with g of total RNA using M-MLV reverse transcriptase and an oligo(dT)15 primer (Promega) in 20 μl mixture. Then, the reaction was diluted fivefold with water and the cDNA used as template for qRT-PCR. The qRT-PCR mixture (20 μl) contained 2 μl of cDNA template, 10 μl of 2 × LightCycler 480 SYBR Green I Master 1 Master (Roche), and 0.25 μM of forward and reverse primers for each gene (see Supplemental Table 2 online). Reactions were performed using the Light Cycler 2.0 instrument (Roche Diagnostics). The transcript levels of each gene were normalized against those of GAPDH (glyceraldehyde phosphate dehydrogenase; At1g16300) as previously reported (Sakuraba et al., 2010).

BiFC Analysis

Full-length cDNAs of SGR and CCE genes were PCR amplified using Pfu polymerase (Promega) with the gene-specific primers listed in Supplemental Table 2 online. The PCR products were digested with BspHI-NotI and cloned into Ncol-NotI into pSY728 and pSY738, respectively (Brachadori et al., 2004), thereby producing C-terminal fusions with the N- and C-terminal halves of YFPs (YFPn and YFPC), respectively. After verifying the inserts by sequencing, constructs were used for BiFC studies.

Arabidopsis mesophyll protoplasts were isolated from either green (0 DDI) or senescent leaves at 4 DDI according to published procedures (Edler et al., 2006). Cell numbers were quantified with a Neubauer chamber and adjusted to a density of 2 × 10^6 protoplasts mL^-1. Protoplasts were cotransformed with each two constructs by 20% polyethylene glycol transformation according to published procedures (Meyer et al., 2006). Twenty micrograms of plasmid of each construct was used. Transformed cells were incubated for 24 h in the dark at room temperature before laser scanning confocal microscopy analysis (DM IRE2; Leica Microsystems). YFP fluorescence was imaged at an excitation wavelength of 512 nm, and the emission signal was recovered between 525 and 565 nm. CP12-YFPC and PRK-YFPn constructs were used as a positive control for plastid colocalization.

PAO-GFP Fusion Protein Analysis

To examine the localization of PAO in chloroplasts, a full-length cDNA of PAO (Pruzˇinska´ et al., 2003) was PCR amplified using Pfu polymerase (Promega) with the primers listed in Supplemental Table 2 online and cloned into BarnHi-Spel–restricted pUC18-GFP5T-sp (Meyer et al., 2006), thereby producing a C-terminal fusion with GFP (PAO-GFP). Transient transformation of Arabidopsis mesophyll protoplasts and confocal microscopy analysis were performed as described above. GFP fluorescence was imaged at an excitation of 488 nm and emission between 495 and 530 nm. As a control for chloroplast envelope localization, a TIC110-GFP construct was employed (Schelbert et al., 2009).

Pigment Analysis

Chlorophyll was extracted from rosette leaf tissues using ice-cold acetone. Extracts were centrifuged at 15,000 rpm for 10 min at 10°C. The supernatant was diluted with ice-cold water to the final acetone concentration of 80%. Chlorophyll was quantified spectrophotometrically as previously published (Porra et al., 1989).

SDS-PAGE and Immunoblot Analysis

Membrane and soluble proteins were extracted from rosette leaves using the Native Membrane Protein Extraction Kit (Calbiochem). Protein extracts were suspended with an equal volume of 2 × sample buffer (50 mM Tris, pH 6.8, 2 mM EDTA, 10% [w/v] glycerol, 2% SDS, and 6% 2-mercaptoethanol), denatured at 75°C for 3 min, and subjected to SDS-PAGE. For visualization of protein bands, gels were stained with Coomassie Brilliant Blue (Sigma-Aldrich). The resolved proteins were electroblotted onto Immunobilon-P transfer membranes (Millipore). Antibodies against SGR (Park et al., 2007), RCCR (Pruzˇinska´ et al., 2007), PAO (Pruzˇinska´ et al., 2005), NYC1 and NOL (Sato et al., 2009), GFP (Abcam), GST (Santa Cruz Biotechnology), TOC75 and PGL35 (Vidi et al., 2006), and photosystem protein antibodies (CAB [Vidi et al., 2006], Lhcb1, Lhcb1, and CP43; Agrisera, Sweden) were used for immunoblot analysis. Peroxidase activity of secondary antibodies was visualized using the chemiluminescence detection kit WEST SAVE (AbFRONTIER) or ImmunStar WesternC (Bio-Rad) according to the manufacturers’ protocols.

In Vitro and in Vivo Pull-Down Assays

Three-week-old transgenic plants were homogenized with the Native Membrane Protein Extraction Kit (Calbiochem), and membrane-enriched fractions were pulled down using IgG or glutathione agarose beads (Santa Cruz Biotechnology) or α-GFP–conjugated beads (MBL). Precipitated beads were washed at least three times with washing buffer (50 mM Tris-HCl, pH 7.2, 200 mM NaCl, 0.1% Nonidet P-40, 2 mM EDTA, and 10% glycerol). Washed beads were boiled with 20 μl of 2 × SDS sample buffer for 5 min and subjected to SDS-PAGE and immunoblot analysis.

Suc Density Gradient Centrifugation

Intact chloroplasts were isolated and the chloroplast membrane fraction was prepared for density gradient centrifugation as described (Vidi et al., 2006). A linear gradient of Suc between 5 and 45% was employed to separate membrane fractions by centrifugation at 100,000g for 17 h (Vidi et al., 2006). One-milliliter fractions were collected starting from the top of the gradient and fractions were analyzed by SDS-PAGE and immunoblot analysis.
Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: SGR/NYE1, At4g22920; RCCR, At4g37000; NY1; At4g13250; NOL, At5g04900; PPH, At5g13800; PAO, At3g44880; CLH, At1g19670; and GAPDH, At1g16300.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. Subcellular Localization of SGR-GFP and CCE-GFP Fusion Proteins in Transgenic Plants.

Supplemental Figure 2. Natural Senescence of Arabidopsis SGR and CCE Overexpressors.

Supplemental Figure 3. Expression Analysis of Genes Encoding SGR and four CCEs in Wild-Type and 3SS:PPH-GFP Leaves by qRT-PCR.

Supplemental Figure 4. Arabidopsis SGR Interacts with LHCII in Senescing as well as Nonsenescent Chloroplasts.

Supplemental Figure 5. Interactions among CCEs by in Vitro Pull-Down Assays.

Supplemental Figure 6. In Vivo Interactions among SGR and CCEs Analyzed by BIFC.

Supplemental Figure 7. Possible Metabolic Channeling of Chlorophyll Catabolic Intermediates by Interaction of SGR, CCEs, and LHCII at the Thylakoid Membrane.

Supplemental Table 1. Arabidopsis Transformants Expressing a Tagged Version of SGR and CCEs Used in This Study.

Supplemental Table 2. Primers Used in This Study.

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AUTHOR CONTRIBUTIONS


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