Triticum Aestivum L. cv. Guizi 1 ANS-6D Positively Regulates Leaf Senescence Through the Abscisic Acid Mediated Chlorophyll Degradation in Tobacco

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Abstract

Anthocyanidin synthase (ANS) is involved in the synthesis of anthocyanins, which are important phytonutrients because of their beneficial effects on human health. Here, we identified ANS-6D of purple-colored *Triticum aestivum* L. cv. Guizi 1 (Gz) that is involved in leaf senescence through the abscisic acid (ABA) mediated chlorophyll degradation pathway in tobacco. After characterizing the leaf-senescence phenotype in GzANS-6D overexpression (*OxGzANS-6D*) lines, we found that the increased anthocyanin accumulation and decreased chlorophyll content in *OxGzANS-6D* lines were closely correlated with the expression levels of anthocyanin synthesis-related structural genes and senescence marker genes, as well as the accumulation of reactive oxygen species. The endogenous ABA content increased and ethylene content decreased in *OxGzANS-6D* transgenic lines compared with wild type. Additionally, the levels of the abscisic acid-responsive transcription factors *ABF1* and *ABF2*, as well as those of chlorophyll degradation-related genes (*PAO, NYC, SGR* and *CHL*), were significantly higher in *OxGzANS-6D* transgenic lines than in wild type. Furthermore, we found that GzABF1 and NtABF1 binds to the promoter of *GzANS-6D*, and NtABF2 binds to the promoter of *NtSGR*. Thus, *GzANS-6D* participated in leaf senescence through ABA-mediated chlorophyll degradation, and *ABF1/2* play important role in *GzANS-6D* functions.

Introduction

Anthocyanins, which are responsible for the colors (e.g., purple, red and blue) of cereal grains, are water-soluble pigments belonging to the flavonoid group (Chen et al. 2013). Purple-colored wheat (*Triticum aestivum* L.) has purple-colored grains that contain significant levels of anthocyanins (Zeven 1991; Liu et al. 2010). The anthocyanins benefit human health by improving vision and neurological health, and having anticarcinogenic and anti-inflammatory roles owing to their antioxidant activities (Cavalcanti et al. 2011; Khoo et al. 2017). The anthocyanin content is an important indicator of phytonutrients (Wang et al. 2019). Therefore, researches have focused on analyzing anthocyanin synthesis in, and the breeding of, purple-colored wheat.

Structural genes control anthocyanin accumulation through the regulation of the anthocyanin biosynthetic pathway. Phenylalanine ammonia-lyase (*PAL*), 4-coumarate-CoA ligase (*4CL*), chalcone synthase (*CHS*), chalcone isomerase (*CHI*), flavanone 3-hydroxylase (*F3H*), dihydroflavonol 4-reductase (*DFR*) and anthocyanidin synthase (*ANS*) are important structural genes in the anthocyanin biosynthetic pathway, among which the *ANS*-encoded anthocyanidin synthase enzyme catalyzes the conversion of leucoanthocyanidin to anthocyanidin (Gonzalez et al. 2008; Jaakola et al. 2013; Li et al. 2018). Transcriptome analyses of genes involved in anthocyanin biosynthesis have demonstrated that *ANS* plays important roles in anthocyanin biosynthesis in cherry (*Prunus avium* L.), mulberry (*Morus alba* L.) and wheat (Wei et al. 2015; Huang et al. 2020; Li et al. 2018). However, only limited studies on *ANS* functions have demonstrated that overexpressing *ANS* enhances anthocyanin accumulation, leading to an increased antioxidant potential in rice (Reddy et al. 2007) and reduced salvianolic acid B (SAB) and rosmarinic acid (RA) biosynthesis in *Salvia miltiorrhiza* and *S. miltiorrhiza* Bge f. alba (Li et al. 2019).
plantlets, respectively. Additionally, *Fedia graciliflora* the ANS knockdown mutants develop larger flowers with wider corolla tube openings than the wild type (WT) (Berger et al. 2017).

Phytohormones play important roles in regulating the anthocyanin content. Both abscisic acid (ABA) and ethylene regulate anthocyanin metabolism (Peng et al. 2011). Early in 1992, Hattor et al. demonstrated that ABA activates the *C1* regulatory gene for anthocyanin biosynthesis during seed maturation in maize (Hattor et al. 1992). In addition, ABA plays positive roles in inducing anthocyanin accumulation by regulating the expression of structural and/or regulatory genes in rice (*Oryza sativa* L.) (Hung and Kao 2005; Hung et al. 2008) and *Arabidopsis thaliana* (Zhou et al. 2009; Watanabe et al. 2018). The MYB96 mutant (*myb96-1d*) shows obvious anthocyanin accumulations in leaves after simultaneous stimulation with both ABA and salicylic acid (SA) in *Arabidopsis* (Seo and Park 2010). González-Villagra et al. (2017) further indicated that ABA participates in anthocyanin synthesis through the regulation of microRNA156, augmenting the expression levels of anthocyanin synthesis-related genes (such as *DFR* and *ANS*), resulting in anthocyanin accumulation. Recently, overexpressing the *More Axillary Branches 2* gene from *Sapium sebiferum* in *Arabidopsis* resulted in higher expression levels of ABA biosynthetic genes, increased anthocyanin accumulations and a higher retained chlorophyll content (Wang et al. 2019).

Interestingly, the ABA content is elevated along with anthocyanin accumulation in the *OsPL6* (Purple Leaf) mutant; however, the chlorophyll content is significantly reduced at the tillering stage (Khan et al. 2020). Additionally, the ABA and anthocyanin contents increase, while the chlorophyll content decreases in torenia (*Torenia fournieri*) and allotetraploid rapeseed (*Brassica napus* L.) under sucrose and severe salinity treatments, respectively (Nagira et al. 2006; Feng et al. 2020). Ethylene signaling plays a negative role in Pi starvation- (Lei et al. 2011), sucrose- (Jeong et al. 2010; Kwon et al. 2011; Meng et al. 2018) and light-induced (Jeong et al. 2010) anthocyanin accumulation in *Arabidopsis*. Sewelam et al. demonstrated that ETHYLENE RESPONSE FACTOR6 insertion mutant plants showed increased H$_2$O$_2$ and anthocyanin levels in *Arabidopsis* (Sewelam et al. 2013). Under nitrogen starvation conditions, the ROOT HAIR DEFECTIVE3 mutant shows an anthocyanin overaccumulation phenotype similar to ethylene signaling mutants *etr1*, *ein2* and *ein3/eil1* in *Arabidopsis*. Additionally, the ethylene precursor ACC strongly suppresses anthocyanin accumulation in a partially ROOT HAIR DEFECTIVE3-dependent manner (Wang et al. 2015). Recently, ethylene was reported to repress anthocyanin biosynthesis and accumulation in pericarps in the dark, and the aminoethoxyvinylglycine hydrochloride (an ethylene biosynthesis inhibitor) treatment results in significantly higher transcript levels of biosynthetic genes (e.g., *DFR*, *CHS* and *ANS*) in rice (Kumar et al. 2019). Carvalho et al. (2010) reported that ABA and anthocyanin biosynthetic genes in the serine/arginine-rich 45 mutant were over-induced by glucose; however, there is no effect on ethylene signaling (Carvalho et al. 2010). Most reports have shown that ABA and ethylene participate in regulation anthocyanin biosynthesis in *Arabidopsis* and rice, however, the effects of ABA on anthocyanin biosynthesis in wheat still require research.

Based on the transcriptome sequencing results (Li et al. 2018), we found that the expression levels of the structural *GzANSs* (especially *GzANS-6D*) underwent positive dynamic changes along with anthocyanin accumulation. Here, we characterized the functions of *GzANS-6D* in tobacco. We first analyzed the
dynamic changes in GzANS-6D in three important developmental periods of ‘Guizi 1’ (Gz) wheat. Then, we analyzed the phenotypes of transgenic tobacco lines overexpressing GzANS-6D (OxGzANS-6D) grown under normal conditions. The OxGzANS-6D lines showed early senescence phenotypes with higher anthocyanin contents and lower chlorophyll contents than WT. The OxGzANS-6D lines had higher anthocyanin biosynthetic and senescence-related gene expression levels than the WT. Consistently, these transgenic lines displayed accumulations of reactive oxygen species (ROS) content and losses in antioxidant enzyme activity levels. Furthermore, the endogenous contents of ABA and ethylene were detected, and the OxGzANS-6D lines possessed higher ABA content and lower ethylene content compare with WT. Finally, compared with the WT plants, the OxGzANS-6D lines exhibited higher expression levels of ABFs and chlorophyll degradation-related genes. These results demonstrated that GzANS-6D participated in leaf senescence through the regulation of endogenous ABA signaling.

Materials And Methods

Plant materials and plant growth conditions

Triticum aestivum L. cv. Guizi 1 (Certificate No. Qian2015003) was planted on the experimental farm, and the field management was performed in accordance with Li et al. (Li et al. 2018). The developing caryopses were selected from spikes at the identical flowering stages at 10-, 15- and 25-days post anthesis (dpa), and samples were stored at −80°C for RNA extraction. The coding sequence of GzANS-6D was obtained from 25 dpa caryopses of ‘GZ1’, placed into the pBI121 vector, and then transformed into the tobacco (Nicotiana benthamiana). From the regenerated plants, transgenic GzANS-6D (OxGzANS-6D) overexpression lines were selected (Luo et al. 2006). The OxGzANS-6D transgenic lines were confirmed using PCR. Seeds were surface sterilized with 75% [v/v] ethanol for 2 min and then 1% [v/v] sodium hypochlorite for 10 min, followed by washing four times for 5 min each time with distilled water. Seeds were vernalized at 4°C for 3 days in the dark. Seedlings were grown vertically on MS medium. Two-week-old plants were transferred to soil. The plants were grown at 25°C/22°C (day/night) under long-day conditions (16-h light/8-h dark). The 7th and/or 8th leaves (cotyledons were excluded) were harvested from 10-week-old tobacco plants for RNA extraction, and the determination of anthocyanin, chlorophyll and phytohormone contents, ROS accumulation levels and enzyme activities.

RNA isolation and quantitative real-time PCR

Total RNAs from wheat and tobacco were extracted using an EASYspin Plus Complex Plant RNA Kit (Aidlab, Beijing, China), and then, first-strand cDNA was synthesized from 1 μg of RNA using a TUREscript 1st Strand cDNA Synthesis Kit gDNA Eraser (Aidlab, Beijing, China). The β-actin gene was used as the internal control, and a qRT-PCR analysis was performed using LightCycler480 SYBR Green (Bio-Rad, CFX 96 Touch, USA). The primers for qRT-PCR are listed in Table S1.

Extraction and measurement of total anthocyanins
The extraction and total content determination of anthocyanins were performed in accordance with Li et al. (Li et al. 2018). Briefly, 0.5 g frozen samples were ground into powder in liquid nitrogen, then 4 mL of acidified ethanol was added and mixed with each sample for anthocyanin extraction. The total anthocyanins level was determined by measuring the absorbance at 527 with an Evolution 220 (Thermo Fisher, USA) and calculated using the following formula: anthocyanin content (mg/Kg) = \( (C \times V) / m \), where \( C \) represents anthocyanin concentration, \( V \) represents volume and \( m \) represents sample weight.

**Extraction and determination of total chlorophyll**

The extraction and determination of chlorophyll were performed in accordance with previously described protocols (Zhang and Guo 2018; Kong et al. 2014). Briefly, 0.25 g fresh leaf samples were used for chlorophyll extraction, and 12.5 mL extraction solution was added at room temperature in darkness until the leaf tissue was completely bleached. The leaves were incubated in the dark for 24 h at room temperature. The absorbance levels of the solution were measured at 665 and 649 nm using an Evolution 220 (Thermo Fisher) and calculated using the following formulae:

- Chlorophyll a content (\( C_a \)) (\( \mu \)g/mL) = 13.95\( \times A_{665} \)− 6.88\( \times A_{649} \);
- Chlorophyll b content (\( C_b \)) (\( \mu \)g/mL) = 24.96\( \times A_{649} \)− 7.32\( \times A_{665} \); and
- Total chlorophyll (mg/g) = \( (C_a + C_b) \times V / W \), where \( V \) represents the volume of the extraction solution; \( W \) represents the weight of freshly detached leaves.

**Histochemical staining of ROS and an antioxidant enzyme activity analysis**

\( \text{H}_2\text{O}_2 \) and \( \text{O}_2^- \) were detected using a 3,3′-diaminobenzidine (DAB) and nitroblue tetrazolium chloride (NBT) staining kit (Solarbio, Beijing, China). Then, the leaves were imaged and used to calculate the relative gray value with ImageJ software (black is 0 and white is 255). The activity levels of antioxidant enzymes (POD, CAT and SOD) were detected using the appropriate corresponding kit (Suzhou Grace Biotechnology Co.Ltd).

**Determination of phytohormones**

Endogenous ABA levels were measured using a plant ELISA kit (Shanghai Enzyme-linked Biotechnology Co. Ltd., Shanghai, China) following the producer's instructions. Leaves of \( \text{OxGzANS-6D} \) transgenic plants and WT were ground in liquid nitrogen and used to measure the ABA content. The calibration standards and samples were simultaneously determined by measuring the absorbance at 450 nm using a microplate reader (Infinite F50, TECAN). The standard curve was constructed using the concentrations of calibration standards against optical density (OD). Then, the sample ABA contents were established by tracking the standard curve and establishing similar trends using previously reported sample values (Khan et al., 2020). The endogenous ethylene content was measured in accordance with Li et al. (2014). Briefly, 0.2 g of \( \text{OxGzANS-6D} \) transgenic plants and WT leaves were placed separately in 1-mL gas-tight glass vessels. A drop of distilled water was added, the caps tightened and the samples incubated at room temperature for 24 h. For ethylene determination, 5 \( \mu \)L sample of gas was removed and analyzed using a gas chromatograph (GC7890, Agilent, USA) with a chromatographic column (hp-5; 30 \( \times \) 0.25 mm). The oven, injector and detector temperatures were 50, 150 and 230°C, respectively.
Yeast one-hybrid (Y1H) assay

The Y1H was performed using the Matchmaker Gold Yeast One-Hybrid Library Screening System (Clontech). The cDNA sequence was cloned into the prey vector pGADT7 and transformed into the yeast strain Y1H Gold. The \textit{GzANS-6D} and \textit{NtSGR} promoter was inserted into the pABAI-bait vector.

Results

Phylogenetic tree, multiple alignment and expression levels of \textit{GzANS} genes

Based on the transcriptome sequencing results of three important periods (10 dpa, 25 dpa and 35 dpa) of ‘Guizi 1’ for anthocyanin biosynthesis, we found that the structural \textit{GzANS} genes’ expression levels positively changed along with anthocyanin accumulation (Fig. 1A). The expression levels of \textit{GzANS}s were significantly higher at 25 dpa and 35 dpa than at 10 dpa. They exhibited dynamic increases from 10 to 25 dpa and then decreased from 25 to 35 dpa. Among the \textit{GzANS}s, \textit{GzANS-6D} exhibited a remarkable change. Then, qRTPCR validation was carried out using \textit{GzANS-6D} in ‘Guizi 1’ and ‘Chinese Spring’ wheat (Fig. 1B). In the former, \textit{GzANS-6D} was significantly higher at 25 dpa and 35 dpa than at 10 dpa. It exhibited a dynamic increase from 10 to 25 dpa and then decreased from 25 to 35 dpa, which coincided with the transcriptome sequencing results. However, in the latter, there were no significant differences in \textit{GzANS-6D} expression at the different developmental periods. A phylogenetic tree analysis was carried out for ANS proteins having protein homology levels greater than 70% compared with \textit{GzANS-6D} (Fig. S1). The ANSs clustered into several subgroups, with \textit{GzANS}s and \textit{TaANS}s being in the same group. Multiple alignments of ANSs in wheat revealed that the protein sequences of \textit{GzANS-6D} and \textit{GzANS-6A1} were identical to those of \textit{TaANS-D1} and \textit{TaANS-A1}, respectively, and the protein sequences of \textit{GzANS-6B} were different by one amino acid from \textit{TaANS-B1} (Fig. S2). The ANSs are involved in the regulation of anthocyanin biosynthesis.

\textit{GzANS-6D} overexpression promotes anthocyanin accumulation and leaf senescence in tobacco

To further understand physiological functions of \textit{GzANS-6D} in plant growth and development, we generated \textit{GzANS-6D}-overexpression (\textit{OxGzANS-6D}) lines in tobacco. The rosette diameters and the seventh-leaf sizes (including the length and width) of 5-week-old seedlings of \textit{OxGzANS-6D} lines were significantly smaller than those of the WT (Fig. S3). \textit{OxGzANS-6D} lines exhibited an early senescence phenotype in 10-week-old transgenic plants (Fig. 2A). Subsequently, the anthocyanin and chlorophyll contents of the seventh leaves were detected. The former was significantly higher in \textit{OxGzANS-6D} lines than in WT (Fig. 2B); however, the latter was significantly lower in \textit{OxGzANS-6D} lines than in WT (Fig. 2C).

Our laboratory previously determined that anthocyanin accumulation is often accompanied by the up-regulated expression of anthocyanin synthesis-related structural genes (Li et al. 2018). In our study, RNA was extracted from the seventh leaves, and the transcript levels of six structural genes, \textit{PAL}, \textit{4CL}, \textit{CHS}, \textit{CHI}, \textit{F3H} and \textit{DFR}, were detected using qPCR. The transcript levels of these genes were significantly higher in \textit{OxGzANS-6D} transgenic leaves than those in WT leaves (Fig. 3A–F). The senescence marker...
genes glutamate dehydrogenase (GDH1), din, cysteine protease (CP1) and senescence-associated gene (SAG12) are upregulated during leaf senescence (Li et al., 2016; Liu et al., 2016). Here, the transcript levels of these four genes were significantly higher in OxGzANS-6D transgenic leaves than in WT leaves (Fig. 4A–D), with the transcript levels of CP1 and SAG12 being increased by more than 40-fold (Fig. 4C) and 90-fold (Fig. 4D) in OxGzANS-6D transgenic leaves compared with WT leaves, respectively. Thus, the overexpression of GzANS-6D, which results in anthocyanin accumulation, correlates with plant senescence. Furthermore, the yield traits of the WT and OxGzANS-6D transgenic lines revealed that the latter had significantly lower caryopsis numbers and thousand grain weights than the WT (Table 1), which coincided with their early senescence phenotype.

![Table 1](https://example.com/table1.png)

**Table 1**

Analysis of yield traits in WT and transgenic lines

<table>
<thead>
<tr>
<th>Yield traits</th>
<th>WT</th>
<th>OE6</th>
<th>OE7</th>
<th>OE9</th>
</tr>
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<tr>
<td>Caryopsis number</td>
<td>43.00 ± 1.41</td>
<td>33.75 ± 1.71**</td>
<td>34.00 ± 1.83**</td>
<td>32.25 ± 2.87**</td>
</tr>
<tr>
<td>Thousand grains weight (mg)</td>
<td>65.33 ± 1.49</td>
<td>58.67 ± 1.70**</td>
<td>60.67 ± 1.11**</td>
<td>59.33 ± 1.11**</td>
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Note: ‘**’ means extremely significant level (P < 0.01).

**Overexpression of GzANS-6D increases ROS accumulation and reduces antioxidant enzyme activity levels**

To verify the function of GzANS-6D in leaf senescence, we performed assays to measure the accumulation of ROS (H_2O_2 and O_2·−) in the leaves of the WT and OxGzANS-6D plants. The eighth fresh leaves of 10-week-old plants were used for DAB and NBT staining to detect the accumulations of H_2O_2 and O_2·−, respectively. Leaves from OxGzANS-6D transgenic lines exhibited greater accumulations of H_2O_2 (Fig. 5A and 5C) and O_2·− (Fig. 5B and 5D) than leaves from the WT. ROS levels in plants are controlled by biosynthetic processes and antioxidant systems (Mittler et al. 2004). Therefore, we detected the activity levels of the antioxidant enzymes POD (Fig. 6A), CAT (Fig. 6B) and SOD (Fig. 6C). All three antioxidant enzymes’ activity levels were significantly decreased in OxGzANS-6D transgenic lines compared with WT. These results indicated that GzANS-6D participates in leaf senescence by regulating ROS accumulation-associated biological processes.

**Endogenous ABA and ethylene content changes in OxGzANS-6D transgenic plants**

The ABA and ethylene contents were determined in the leaves of WT and OxGzANS-6D plants. The seventh fresh leaves of 10-week-old plants were used to analyze endogenous ABA and ethylene contents. The ABA content significantly increased in OxGzANS-6D transgenic lines compared with WT (Fig. 7A); however, the ethylene content significantly decreased in OxGzANS-6D transgenic lines compared with WT (Fig. 7B). Thus, ABA and ethylene may play different roles during anthocyanin accumulation in OxGzANS-6D plants.
ABFs and chlorophyll degradation-related genes are upregulated in OxGzANS-6D transgenic plants

To further investigate the role of ABA in the leaf senescence of OxGzANS-6D transgenic lines, we performed an RT-qPCR analysis on ABA-responsive element (ABRE)-binding transcription factors genes, such as ABF1 and ABF2, and several chlorophyll degradation-related genes, such as pheophorbide a oxygenase (PAO), non-yellow coloring (NYC), non-yellowing (SGR, also known as NYE) and chlorophyllase (CHL) (Fig. 8). The ABFs are direct activators of PAO, NYC and SGR, which are involved in the regulation of chlorophyll degradation (Gao et al. 2016; Hu et al. 2019). The expression levels of ABF1 (Fig. 8A) and ABF2 (Fig. 8B) were significantly higher in OxGzANS-6D transgenic plants than in WT. In addition, the expression levels of four chlorophyll degradation-related genes (PAO, NYC, SGR and CHL) were significantly higher in OxGzANS-6D transgenic lines compared with WT (Fig. 8C–F).

GzABF1 and NtABF1 binds to the promoter of GzANS-6D, and NtABF2 binds to the promoter of NtSGR

As indicated in Fig. 9A and 9B, a ABRE sequence of GzANS-6D promoter was identified as the possible binding region by ABFs (Fig. 9A). Y1H assay indicated GzABF1 and NtABF1 binds to the GzANS-6D promoter, whereas NtABF2 cannot binds to the promoter of GzANS-6D (Fig. 9B). Meanwhile, Y1H assay indicated that NtABF2 binds to the NtSGR protomer (Fig. 9C). The results demonstrated the expression of GzANS-6D regulated by GzABF1 and NtABF1, and expression of NtSGR regulated by NtABF2 (Fig. 9).

Discussion

Transcriptome analyses of genes involved in anthocyanin biosynthesis have demonstrated the importance of ANS (Zhang et al. 2016; Li et al. 2018; Huang et al. 2020). Additionally, limited studies demonstrated that ANS plays important roles in antioxidant potential, phenolic acids biosynthesis and floral morphology (Reddy et al. 2007; Li et al. 2019; Berger et al. 2017). Therefore, more studies are needed to explore functions of ANS. Here, our transcriptome sequencing (Fig. 1A) and qRTPCR validation (Fig. 1B) results demonstrated that GzANS-6D positively changed with anthocyanin accumulation in wheat, which is in agreement with our previous study (Li et al. 2018).

To further understand the functions of GzANS-6D, we explored the phenotypes of OxGzANS-6D transgenic lines under normal growth conditions. We found that the transgenic lines had smaller rosette diameters (Figure S3) and leaf senescence phenotypes (Fig. 2A) compared with WT. The transgenic lines had higher anthocyanin contents (Fig. 2B) and lower chlorophyll contents (Fig. 2C) compared with WT, which corresponded to GzANS-6D’s function and leaf senescence phenotype, respectively. This result coincides with the functions of LcABFs (Hu et al. 2019) and OsPL6 (Khan et al. 2020), which demonstrated that anthocyanin and chlorophyll have antagonistic effects. Anthocyanin accumulation is often accompanied by the up-regulated expression of anthocyanin synthesis-related structural genes (Li et al. 2018). Out data also show that six structural genes (PAL, 4CL, CHS, CHI, F3H and DFR) involved in anthocyanin synthesis were upregulated in OxGzANS-6D transgenic lines compared with WT (Fig. 3). Leaf senescence is often accompanied by the up-regulated expression of senescence marker genes. Consequently, four senescence marker genes (GDH1, din, CP1 and SAG12) were selected based on the previous reports (Li et
al. 2016; Liu et al. 2016) and their expression levels were determined. qRT-PCR confirmed that these four genes were upregulated in OxGzANS-6D transgenic lines compared with in WT (Fig. 4), which was in agreement with the premature senescence characteristic. Additionally, the early leaf senescence of OxGzANS-6D transgenic lines resulted in lower yields compared with WT, which was consistent with the phenotype of small auxin-up RNA 39 overexpression in rice (Kant et al. 2009). Leaf senescence is often accompanied by ROS accumulation (Cui et al. 2013; Liu et al. 2016; Zhang et al. 2020). Our DAB (for H2O2 content) and NBT (for O2− content) staining results indicated that GzANS-6D positively regulated ROS accumulation (Fig. 5). Antioxidant proteins (such as POD, CAT and SOD) act as ROS-scavenging antioxidants by reducing the ROS level (Nakabayashi et al., 2014; Lotkowska et al., 2015). Here, all three antioxidant enzymes’ activity levels decreased in OxGzANS-6D transgenic lines compared with WT (Fig. 6), which indicated that GzANS-6D participates in leaf senescence by regulating ROS signaling.

ABA promotes, but ethylene inhibits, the accumulation of anthocyanin (Peng et al. 2011) and ROS (Yu et al. 2019). Here, we demonstrated that the endogenous ABA content significantly increased and the ethylene content significantly decreased in OxGzANS-6D transgenic lines compared with WT (Fig. 7), which was in agreement with previous studies (Peng et al. 2011). Therefore, we hypothesized that endogenous ABA and ethylene play different roles in the leaf senescence of OxGzANS-6D transgenic lines. In Arabidopsis, ABF2/3/4 were identified as direct activators of NYE1/2, NYC1 and PAO, which participate in chlorophyll degradation during leaf senescence processes in an ABA-dependent manner (Gao et al. 2016). Recently, in Litchi chinensis, ABF1/2 were identified as recognizing ABA-responsive elements in the promoter regions of PAO and SGR, and they are, therefore, involved in chlorophyll degradation (Hu et al. 2019). qRT-PCR confirmed that the expression levels of ABF1/2 and four chlorophyll degradation-related genes (PAO, NYC, SGR and CHL) were significantly higher in OxGzANS-6D transgenic lines than in WT (Fig. 8). Furthermore, we found GzABF1 and NtABF1 can directly binds to the promoter of GzANS-6D, whereas NtABF1 cannot. Meanwhile, NtABF2 binds to the promoter of NtSGR (Fig. 9). Our results strongly suggest that GzANS-6D participates in leaf senescence through in an ABA-mediated chlorophyll-degradation manner.

Our results demonstrated that GzANS-6D positively regulates anthocyanin synthesis and leaf senescence by increasing the endogenous ABA content and decreasing the ethylene content. ABA-mediated chlorophyll degradation through ABF2’s activation of chlorophyll degradation-related gene (including PAO, NYC, SGR and CHL, mainly SGR) expression in tobacco. In addition, ABA play important regulatory roles in ROS signaling in leaf senescence. Our work reveals a novel regulatory module containing ANS, and ABA that is involved in leaf senescence.

Conclusions

Overexpression of GzANS-6D promote leaf senescence through regulating the ABA-mediated chlorophyll degradation pathway, and ABF1/2 play important role in GzANS-6D functions.

Declarations
Conflict of Interest

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.

Acknowledgments

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References


Figures
Figure 1

Heat map and qRT-PCR validation of GzANS (anthocyanidin synthase) expression levels during different wheat developmental periods. (A) Heat map of GzANS genes. (B) Expression pattern of GzANS-6D at different grain developmental stages at 10, 20 and 35 dpa. Different combinations of numbers and letters (such as 6D, D1, A1, etc.) represent different chromosomal locations. The same chromosome with different numbers indicates corresponding copies of one gene on the same chromosome. Each bar shows the mean ± SD of triplicate assays. ‘**’ indicates a significant difference (P < 0.01).
Figure 2

<p>Phenotypic observations and anthocyanin and chlorophyll content determinations in WT and transgenic plants. (A) Phenotypic observations of WT and 10-week-old <em>OxGzANS-6D</em> transgenic lines. (B) Anthocyanin content analysis. (C) Chlorophyll content analysis. Each bar shows the mean ± SD of triplicate assays. ‘*’ indicates a significant difference (<em>P</em> < 0.05), and ‘**’ indicates a highly significant difference (<em>P</em> < 0.01).</p>

Figure 3
qRT-PCR validation of anthocyanin synthesis-related gene expression levels in WT and transgenic plants. (A–F) Expression levels of phenylalanine ammonia-lyase (PAL), 4-coumarate-CoA ligase (4CL), chalcone synthase (CHS), chalcone isomerase (CHI), flavanone 3-hydroxylase (F3H) and dihydroflavonol 4-reductase (DFR), respectively. Each bar shows the mean ± SD of triplicate assays. ‘*’ indicates a significant difference (P < 0.05), and ‘**’ indicates a highly significant difference (P < 0.01).

Figure 4

qRT-PCR validation of senescence-related gene expression levels in WT and transgenic plants. (A–D) Expression levels of glutamate dehydrogenase (GDH1), (a tobacco senescence-associated gene), cysteine protease (CP1) and senescence-associated gene (SAG12), respectively. Each bar shows the mean ± SD of triplicate assays. ‘**’ indicates a significant difference (P < 0.01).

Figure 5
Analysis of ROS accumulation levels in WT and transgenic plants. (A) Observation of DAB staining for H$_2$O$_2$. (B) Observation of NBT staining for O$_2^−$. (C) Calculation of gray values for DAB staining. (D) Calculation of gray values for NBT staining. Each bar shows the mean ± SD of triplicate assays. ‘*’ indicates a significant difference (P < 0.05), and ‘**’ indicates a highly significant difference (P < 0.01).

Figure 6

Activity levels of antioxidant enzymes in WT and transgenic plants. (A–C) Activity levels of POD, CAT and SOD, respectively. Each bar shows the mean ± SD of triplicate assays. ‘*’ indicates a significant difference (P < 0.05), and ‘**’ indicates a highly significant difference (P < 0.01).
Figure 7

<p>ABA and ethylene contents in WT and transgenic plants. A) Analysis of ABA content. B) Analysis of ethylene content. Each bar shows the mean ± SD of triplicate assays. The ‘*’ represents significant differences (<em>P </em>&lt; 0.05) and the ‘**’ represents significant differences (<em>P </em>&lt; 0.01). </p>

Figure 8

<p>The qRT-PCR validation of chlorophyll degradation-related genes and ABA-related genes in WT and transgenic plants. (A and B) Expression levels of ABRE-binding transcription factors <em>ABF1</em> and <em>ABF2</em>, respectively. (C–F) Expression levels of pheophorbide a oxygenase (<em>PAO</em>), non-yellow coloring (<em>NYC</em>), non-yellowing (<em>SGR</em>, also known as <em>NYE</em>) and chlorophyllase (<em>CHL</em>), respectively. Each bar shows the mean ± SD of triplicate assays. ‘**’ indicates a significant difference (<em>P </em>&lt; 0.01). </p>
Figure 9

Y1H assay. (A) Schematic diagrams of ABRE elements identified in the promoter regions from \textit{GzANS-6D} and \textit{NtSGR}. Triangles, ABRE elements; numbers, distance (bp) from ATG; +, distance from ATG in a positive DNA strand; -, distance from ATG in a negative DNA strand; grey portion, the region cloned into Bait vector in Y1H assay. (B) Y1H assay analyze that NtABF1 and GzABF1 interacted with the promoter of \textit{GzANS-6D}. (C) Y1H assay analyze that NtABF2 interacted with the promoter of \textit{NtSGR}.

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