Hydrogen Sulfide-Linked Persulfidation Maintains Protein Stability of ABSCISIC ACID-INSENSITIVE 4 and Delays Seed Germination

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Abstract: Hydrogen sulfide (H\textsubscript{2}S) is an endogenous gaseous molecule that plays an important role in the plant life cycle. The multiple transcription factor ABSCISIC ACID INSENSITIVE 4 (ABI4) was precisely regulated to participate in the abscisic acid (ABA) mediated signaling cascade. However, the molecular mechanisms of how H\textsubscript{2}S regulates ABI4 protein level to control seed germination and seedling growth have remained elusive. In this study, we demonstrated that ABI4 controls the expression of L-CYSTEINE DESULFHYDRASE1 (DES1), a critical endogenous H\textsubscript{2}S-producing enzyme, and both ABI4 and DES1-produced H\textsubscript{2}S have inhibitory effects on seed germination. Furthermore, the ABI4 level decreased during seed germination while H\textsubscript{2}S triggered the enhancement of the persulfidation level of ABI4 and alleviated its degradation rate, which in turn inhibited seed germination and seedling establishment. Conversely, the mutation of ABI4 at Cys250 decreased ABI4 protein stability and facilitated seed germination. Moreover, ABI4 degradation is also regulated via the 26S proteasome pathway. Taken together, these findings suggest a molecular link between DES1 and ABI4 through the post-translational modifications of persulfidation during early seedling development.

Keywords: hydrogen sulfide; persulfidation; DES1; ABI4; protein stability

1. Introduction

Growing bodies of reports have demonstrated that H\textsubscript{2}S, a gaseous signaling molecule, is involved in a variety of physiological processes during plant growth and development, such as autophagy, flowering, and stomatal closure [1–4]. The multiple H\textsubscript{2}S generation pathways have been correspondingly identified, including L/D-cysteine desulphydrase, β-cyanoalanine synthase, and O-acetylserine(thiol)lyase [5,6]. Particularly, cytoplasmic L-Cys desulphydrase DES1 was considered as one of the most critical H\textsubscript{2}S-producing enzymes in Arabidopsis. DES1 exhibited a higher affinity to L-cysteine and a strong catalytic capacity to degrade L-cysteine to sulfide, ammonia, and pyruvate [7]. In parallel, endogenous H\textsubscript{2}S levels reduced by 30% in the Arabidopsis des1 mutant [7], which has brought great convenience for us to investigate the physiological effects of H\textsubscript{2}S in plants [8,9]. A greater understanding of the mechanism for H\textsubscript{2}S biological functions in plants has been achieved, accompanied by the key feature of H\textsubscript{2}S at protein persulfidation, one type of post-translational modification of oxidized cysteine residues to form a persulfide group (RSSH) in target proteins [10,11]. To date, persulfidated proteins in plants have been widely reported and the function of persulfidation has exhibited great complexity and diversity in different tissues and organs of plants [12,13]. Persulfidation significantly impacts protein biological functions, such as enzymatic activity, subcellular localization, and protein interaction [14–16]. At least 5214...
proteins, 13% of the entire annotated proteome in *Arabidopsis* roots, were susceptible to persulfidation [17]. However, our current understanding of persulfidation remains relatively limited compared to other protein modifications such as phosphorylation and acetylation.

Seed germination is an essential stage in the whole life cycle of plants, which is crucial for plant propagation and requires the precise coordination of multiple external and internal signals [18]. A sequence of mechanisms has been adopted to fine-tune intercellular homeostasis during seed germination in plants due to its high vulnerability to biotic and abiotic stresses [19,20]. ABSCISE ACID-INSENSITIVE 4 (ABI4), an APETALA2 (AP2)/ETHYLENE-RESPONSIVE ELEMENT BINDING PROTEIN (EREBP) domain-containing transcription factor (TF), acts as a positive modulator in the ABA signaling pathway [21]. During seed germination and post-germination process, ABI4 is involved in regulating a wide range of important biological events, including stress response, chloroplast and mitochondria retrograde, and lipid metabolism [22–25]. ABI4 integrates with various phytohormones signaling during seed germination and flowering [23,26,27]. Interestingly, ABI4 acts as either activator or repressor to control the transcription of downstream responsive genes by recognizing coupling 1 elements (CE1) (CACCG and CCAC motif) within their promoters [21,28]. For example, ABI4 could directly control the MAPK cascade by specifically binding to the CE1 motif of the *MAPKK18* promoter [16]. However, ABI4 inhibits the transcript abundance of the mitochondrial retrograde signal gene ALTERNATIVE OXIDASE1a (*AOX1a*) by targeting the CGTGAT element in the promoter [29].

Our current understandings of the mechanism of *H*₂*S*-regulated signal transduction have been achieved in genetic and molecular studies. Our recent study has illustrated a DES1-ABI4 loop molecular regulatory mechanism to control plant responses to ABA. ABA-triggered accumulation of *H*₂*S* leads to persulfidation of ABI4 at Cys250 [16]. Meanwhile, ABI4 binds specifically to the CE1 motif of the *DES1* promoter to control *DES1* expression and further *H*₂*S* production. However, ABI4 transcript and protein levels are fairly low and even undetectable during vegetative growth in mature overexpressing transgenic plants [30], implying that ABI4 is precisely regulated by a complex network while exercising its function. The general mechanism behind the protein stability of ABI4 controlled by protein persulfidation and the underlying signaling pathways remain to be characterized.

In this study, we show that ABI4 controls the expression of *DES1*, and both ABI4 and DES1-produced *H*₂*S* have inhibitory effects on seed germination. *H*₂*S*-linked persulfidation on ABI4 Cys250 enhances ABI4 stability, which in turn inhibits seed germination and post-germination development in *Arabidopsis*. We establish a molecular mechanism for DES1 and ABI4 synergistically in the regulation of seed germination and seedling establishment through ABI4 protein stability.

2. Results

2.1. Inhibition of Germination and Seedling Growth in *Arabidopsis* by Exogenous *H*₂*S*

Our previous studies have revealed that persulfidation plays an indispensable role in regulating plant ABA responses, including the induction of stomatal closure [14] and the inhibition of primary root growth [16]. To determine whether *H*₂*S* is involved in the *Arabidopsis* germination process, we measured the relative expression of *DES1* during the WT seed germination period. Quantitative PCR (qPCR) results showed that the relative expression of *DES1* decreased in a time-dependent manner from dormant seeds to 6-day-old seedlings (Figure 1A). Meanwhile, the enzyme activity of L-cysteine desulphhydrase (L-CDes) was also detected. Mutation of *DES1* resulted in a marked decrease of L-CDes activity in germinating seeds (Figure 1B). Moreover, we observed that the enzyme activity of total L-CDes decreased significantly during the progress of germination in WT but not in *des1* mutant (Figure 1B). These results further proved that L-CDes activity decrease was closely related to DES1, which may function as a negative regulator during seed germination.
of total L-CDes decreased significantly during the progress of germination in WT but not in des1 mutant (Figure 1B). These results further proved that L-CDes activity decrease was expected, the inhibitory effects of NaHS on Arabidopsis seed germination were observed over a wide range of NaHS concentrations from 0.05 to 0.2 mM (Figure 1C). These results indicated that NaHS treatment could delay the beginning of germination in Arabidopsis and prolong the full germination period in a concentration-dependent manner. However,

To further analyze the effect of H2S on Arabidopsis seed germination, NaHS, a well-known H2S donor, was applied into an MS medium with different concentrations. As expected, the inhibitory effects of NaHS on Arabidopsis seed germination were observed over a wide range of NaHS concentrations from 0.05 to 0.2 mM (Figure 1C). These results indicated that NaHS treatment could delay the beginning of germination in Arabidopsis and prolong the full germination period in a concentration-dependent manner. However,
NaHS-treated seeds can fully germinate at 96 h after treatment (Figure 1C). Subsequently, we examined NaHS affect the post-germination process. We observed that Arabidopsis cultured in different NaHS concentrations displayed growth inhibition as shown by a lower greening degree and smaller leaves of the seedlings. NaHS strengthened this inhibitory effect in a concentration-dependent manner (Figure 1D). Primary root length was measured to evaluate the effect of NaHS on post-germination growth. Accordingly, NaHS treatment inhibited primary root growth of 5-day-old Arabidopsis seedlings, suggesting H$_2$S inhibits seedling growth (Supplementary Figure S1). Taken together, these results revealed that exogenous H$_2$S delayed the initiation of seed germination and inhibited post-germination growth.

2.2. DES1-Produced H$_2$S Inhibits Arabidopsis Seed Germination and Post-Germination Growth

To test whether DES1-produced H$_2$S impacted seed germination, the germination performance of the des1 mutant was compared with WT. Under normal conditions, des1 mutant germinates faster compared to WT, especially 36 to 48 h after germination. However, both WT and des1 mutant completely germinated after 72 h (Figure 1E). Subsequently, the des1 mutant was treated with NaHS. As shown, the germination rate of both WT and des1 displayed a similar, overall downward trend with the treatment of NaHS (Figure 1F). Long treatment with NaHS can lead to sulfide oxidation and thus form polysulfides. In this study, the treatment of chemical H$_2$S donor GYY4137 resulted in a similar inhibitory effect on both WT and des1 as NaHS, suggesting that the effects on seed germination observed are related to H$_2$S directly (Supplementary Figure S2). Meanwhile, both WT and des1 mutant seed germination was significantly accelerated by hypotaurine (HT), an H$_2$S scavenger (Figure 1G).

The post-germination phenotype was also investigated. des1 mutant displayed a faster growth trend than WT during the post-germination process (Figure 1H). The greening degree of des1 was significantly higher than that of WT, and the leaves were larger after five days of germination under control conditions. Meanwhile, the application of NaHS inhibited the development of the leaf and delayed the greening of both des1 and WT. By contrast, treatment of HT accelerated seedling growth and abolished the phenotypic differences between WT and des1 (Figure 1H). Therefore, these experimental data revealed that DES1-produced H$_2$S has an inhibitory effect on both germination and post-germination growth of Arabidopsis.

2.3. Functional of DES1 and ABI4 on Seed Germination and Post-Germination Growth

Our previous studies demonstrated that ABI4 can activate DES1 transcription by binding to its promoter, and Cys250 from ABI4 is critical for its binding [16]. In this study, the qPCR analysis showed that the expression of DES1 was significantly blocked in the 5-day-old abi4 mutant seedling (Figure 2A). Correspondingly, DES1 expression was higher in transgenic plants overexpressing ABI4 but not the ABI4Cys250Ala variant (Figure 2A), implying that Cys250 is crucial for ABI4-activated DES1 expression in Arabidopsis seedlings. Subsequently, the total L-CDes activity was also measured in Arabidopsis seedlings of WT, abi4, 35S:ABI4 abi4, and 35S:ABI4Cys250Ala abi4. abi4 and 35S:ABI4Cys250Ala abi4 exhibited decreased L-CDes activities compared with WT, whereas L-CDes activity was increased in 35S:ABI4 abi4 (Figure 2B). These results demonstrated that DES1 expression was controlled by functional ABI4 in Arabidopsis seedlings.
Time-course changes of germination percentage were examined in WT and abi4 Arabidopsis seedlings upon NaHS treatment. While the germination speed of abi4 is faster than wild-type, NaHS-triggered inhibitory response in germination is weakened by the mutation of ABI4 (Figure 2C,D). The germination rates of des1 and abi4 mutant lines were further compared. Mutation of either DES1 or ABI4 reduced the time to have 50% germinated seeds (T50; Figure 2E). Meanwhile, the T50 index is significantly shortened in des1 abi4 double mutant than its parental lines, all of which were significantly increased by NaHS treatment. Furthermore, DES1 and ABI4 also showed additive effects with regard to seedling primary root growth (Supplementary Figure S3). Taken together, these results demonstrated that H2S and ABI4 have additive inhibitory effects during seed germination and the post-germination stages.
2.4. ABI4 Protein and Its Persulfidation Level Decreases during Germination and Post-Germination Stages

ABI4 is a positive regulator of ABA regulation, and ABA has an inhibitory effect on the germination of Arabidopsis [21]. However, the expression of ABI4 was very low in vegetative growth [30]. In this study, 35S:ABI4-GFP transgenic seedlings were used. Meanwhile, the effect of NaHS on ABI4 protein level during seed germination (2 days) and post-germination (6 days) stages were compared. The results showed that the level of ABI4 protein in the post-germination stage decreased by half compared with that of the germination stage (Figure 3A,B). However, compared to the control sample, treatment with NaHS caused a 40% increase in ABI4 protein level after two days of sampling. Meanwhile, the decreased ABI4 protein level at 6-days after germination increased by 100% after the addition of NaHS treatment. In conclusion, the ABI4 protein level decreased markedly during seed germination and post-germination growth, which can be future attenuated by the addition of H2S.

![Figure 3](image-url) The stability and persulfidation level of ABI4 during germination and post-germination growth. (A) NaHS affects the ABI4 protein level In vivo. 35S:ABI4-GFP transgenic seedlings grown on 1/2 MS medium with or without NaHS (0.1 mM) were harvested at the indicated times. Total proteins were extracted and then checked by immunoblot analysis. ABI4 proteins were detected using an anti-GFP antibody. Relative amounts of proteins were determined by densitometry normalized to actin. (B) Quantification of the relative band intensity shown in (A). (C) Effect of the NaHS on the persulfidation of ABI4 In vivo. Sample treatments were as described for (A). Total proteins were extracted and then subjected to the biotin-switch assay to analyze persulfidation levels (ABI4-GFP-SSH). Persulfidated ABI4-GFP protein was detected with anti-GFP antibody after tag-switch labeling and streptavidin purification. The bottom panels show the total ABI4-GFP used for the tag-switch assay as a loading control. (D) Quantification of the ABI4-GFP-SSH level is shown in (C). For (B,D), the bars indicate the relative abundance of the corresponding protein (B) or persulfidated protein (D) compared with that of the control un-treated sample (set to 1.0). Signals from two independent experiments were quantified. Different letters indicate significant differences at p < 0.05 according to one-way ANOVA (post-hoc Tukey’s HSD test).

To verify whether the above NaHS-driven responses were related to ABI4 persulfidation, we further checked the persulfidation level of ABI4 by a biotin-switch method in the following experiment [15]. As shown in Figure 3C, the ABI4 persulfidation level of seedlings decreased by approximately 50% after six days of growth, suggesting the ABI4 presulfidation level decreased during seed germination and seedling establishment. However, the declined level of ABI4 persulfidation was reversed to a higher level in the presence of NaHS, compared with that of the 2-day sample (Figure 3D). Besides, the level of ABI4 persulfidation increased dramatically in both 2-day and 6-day samples after exposure.
to NaHS. These results suggested that both protein and corresponding persulfidation level of ABI4 was regulated by H₂S.

2.5. Persulfidation at Cys250 of ABI4 Inhibits Its Degradation

We hypothesized that persulfidation of ABI4 may regulate its protein level during germination. To test whether persulfidation affects ABI4 stability, we set up a cell-free assay using ABI4-His recombinant protein purified from *Escherichia coli*, treating with either NaHS or dithiothreitol (DTT) followed by dialysis. The level of ABI4-His recombinant protein was detected after 3–18 h incubation with total protein extract from WT, followed by using immunoblotting with anti-His antibody. With an increased incubation period, the ABI4-His protein level was gradually decreased, which was further attenuated or exacerbated by NaHS or DTT treatment, respectively (Figure 4A,B). For example, the ABI4-His protein level decreased by 60% after 9 h of incubation, while this value is 20% or 85% after treatment of NaHS or DTT, respectively. Moreover, the decreased tendency of ABI4-His protein level became faster when its Cys250 was mutated to Ala (Figure 4C,D), which showed a maximum decrease of 40% at 6 h of incubation. Meanwhile, treatment of NaHS or DTT enhance or decrease the persulfidation level of recombinant ABI4 protein, whereas Cys250Ala mutation almost fully abolished the persulfidation of recombinant ABI4 protein (Supplementary Figure S4). These results further indicated persulfidation on Cys250 is critical for ABI4 stability.

![Figure 4](image.png)
control (Con). During incubation, samples were collected at indicated time points for immunoblot analysis using anti-His antibody. Samples collected before the addition of protein extracts were set as internal control, as normalized to load equal amounts determined by Coomassie brilliant blue staining (CCB). (B) Quantification of the relative band intensity shown in (A). (C) Cell-free degradation analysis of ABI4 mutant recombinant proteins. Purified ABI4-His and ABI4Cys250Ala-His recombinant protein were dialyzed and then incubated with extracts of 10-day-old Arabidopsis seedlings. Sample treatments and measurements were as described for (A). (D) Quantification of the relative band intensity shown in (C). (E) The effect of MG132 on the stability of ABI4 in vitro. Purified ABI4-His recombinant proteins were dialyzed and then incubated with extracts containing 50 µμM MG132 of 10-day-old Arabidopsis seedlings. Sample treatments and measurements were as described for (A). (F) Quantification of the relative band intensity shown in (E). For the quantification of relative band intensity, the data indicate the relative abundance of the corresponding protein compared with that of the control sample (set to 1.0). Signals from two independent experiments were quantified. Asterisks represent significant differences between treatments according to Student's t-test (* p < 0.05, ** p < 0.01, *** p < 0.001).

To further investigate whether the decrease of ABI4 level is also attributed to the ubiquitin-26S proteasome pathway, MG132, a specific 26S proteasome inhibitor of ubiquitin-mediated protein degradation, was used. In this study, ABI1 protein, an important component of ABA signaling, where its degradation was regulated by ubiquitination [31,32], was used as a positive control. In vitro and in vivo experiments demonstrated that MG132 sufficiently blocks ABI1 degradation (Supplementary Figure S5). As expected, the decrease level of ABI4-His recombinant protein was relieved by the application MG132 to some extent, with maximum mitigation effect at 6 h by 25% (Figure 4E,F). These results indicated that the ABI4 Cys250 persulfidation regulates its degradation In vitro, and this process is partially related to 26S proteasome activity.

2.6. The Stability of ABI4 Is Regulated by Its Persulfidation

To investigate the effect of persulfidation on protein stability of ABI4 in vivo, a protoplast-based time course degradation assay was performed. The protoplasts from WT plants were transfected with equal amounts of plasmids expressing either ABI4-GFP or ABI4Cys250Ala-GFP, respectively. The protoplasts expressing ABI4-GFP were treated with or without NaHS before lysed. Cycloheximide (CHX) was used to inhibit protein synthesis during protein extraction. As shown in Figure 5A,B, NaHS treatment attenuated the decrease of ABI4-GFP level during the whole detection period compared with the control sample, indicating that NaHS enhanced the stability of ABI4-GFP In vivo. However, the possibility that NaHS might inhibit proteasome activity and thus indirectly enhance the stability of ABI4 cannot be ruled out.

Cys250 mutation almost completely abolished ABI4 persulfidation (Supplementary Figure S4). Next, the degradation rate between ABI4-GFP and ABI4Cys250Ala-GFP were compared. Notably, our results exhibited that the ABI4Cys250Ala-GFP mutant version caused a decrease rate which increased to approximately 80% at 9 h, whereas ABI4-GFP only decreased by 50% (Figure 5C,D). These results demonstrated that Cys250 residue is necessary for maintaining the ABI4-GFP level. Subsequently, the following experiment demonstrated that MG132 attenuated the decreased tendency of the ABI4-GFP level in a time-dependent manner (Figure 5E,F), indicating the involvement of the 26S proteasome-mediated degradation pathway. The persulfidation level of ABI4-GFP was increased by NaHS, regardless of the presence of MG132 (Supplementary Figure S6). Furthermore, the degradation rate of ABI4-GFP was further inhibited by the application of NaHS in the presence of MG132, especially during 0–3 h of treatment. After 9 h of incubation, the degradation rate was inhibited by 40% with the treatment of both MG132 and NaHS (Figure 5E,F). Taken together, these results indicated that persulfidation regulates ABI4 protein stability, in which the pathway of 26S proteasome may involve.
were detected using an anti-GFP antibody, and relative amounts of proteins were determined by densitometry normalized to actin. (B) Quantification of the relative band intensity shown in (A). (C) The stability of wild-type ABI4 or its Cys250Ala mutated version In vivo. Arabidopsis protoplasts isolated from the wild-type lines were transfected with ABI4-GFP or ABI4Cys250Ala-GFP expressing plasmid constructs, respectively. Sample harvest and measurements were as described for (A). (D) Quantification of the relative band intensity shown in (C). (E) The effect of MG132 and NaHS on the stability of ABI4 In vivo. Protoplasts were treated with 50 µM MG132 or 50 µM MG132 and 0.1 mM NaHS, respectively (or distilled water, as a control) for 1 h, and other sample treatments and measurements were as described for (A). (F) Quantification of the relative band intensity shown in (E). For the quantification of relative band intensity, the data indicate the relative abundance of the corresponding protein compared with that of the control sample (set to 1.0). Signals from two independent experiments were quantified. Asterisks represent significant differences between treatments according to Student’s t-test (* p < 0.05, ** p < 0.01, *** p < 0.001).

3. Discussion

In this study, we focused on the effects of H2S and ABI4 on seed germination and post-germination growth under normal physiological conditions. While ABA levels decreased gradually during seed germination, we found that ABI4 and DES1-produced H2S...
have an inhibitory effect on seed germination synergistically. We established a molecular framework for H₂S-linked persulfidation maintaining protein stability of ABI4 in the regulation of *Arabidopsis* seed germination and post-germination process. During this process, ABI4 protein was degraded concurrently and its persulfidation level decreased, which was closely related to H₂S/DES1. ABI4 controls the expression of DES1, and both ABI4 and DES1-produced H₂S have inhibitory effects during seed germination and post-germination growth. We further discovered persulfidation regulates ABI4 stability during seed germination and post-germination growth in *Arabidopsis*.

Multiple lines of evidence obtained from genetic and physiological studies demonstrate that DES1-produced H₂S has an inhibitory effect on both germination and post-germination growth of *Arabidopsis*. This idea is supported by the observation that seed germination and post-germination development was inhibited by NaHS in a concentration-dependent manner (Figure 1D). Consistently, the des1 mutant showed faster germination compared with WT. Our data further suggested that ABI4 regulates DES1 expression, as DES1 expression and L-CDes activities in abi4 mutants was severely reduced from that in the WT, respectively. These results may be supported by the fact that ABI4 activates DES1 transcription by binding to its promoter [16]. Interestingly, the germination of WT, des1, abi4, and des1 abi4 seeds were all significantly inhibited by the treatment of NaHS (Figure 2E). Based on these findings, this study provides new evidence for the notion that ABI4 integrates with DES1 as a negative regulator to participate in seed germination coordinately.

ABI4 plays an essential role as one of the positive regulation transcription factors mediating ABA-dependent stress response [25]. The ABI4 protein level is precisely regulated by a complex network under a vegetative state [30]. In this study, protein degradation analysis revealed that MG132, a specific 26S proteasome, relieved the decrease level of ABI4 protein, indicating that ABI4 degradation was also regulated via the 26S proteasome pathway, which was also found in ABI1 and ABI5 [32,33]. However, a recent study has found that phytomias SAP05 (secreted AY-WB proteins) mediates the concurrent degradation of SPL and GATA developmental regulators via hijacking the plant ubiquitin receptor RPN10 independent of substrate ubiquitination [34]. Thus, the degradation of ABI4 may also not be dependent on ubiquitination entirely. Interestingly, the accumulation of ABI4 in seeds was observed at the early stage of germination, and its steady-state mRNA levels dropped sharply a few days after germination [35]. Importantly, the ABI4 level in the NaHS treated seedlings was evidently higher than in the control. These results further implied that H₂S might also be involved in the mechanism of regulation of ABI4 stability to control seed germination.

H₂S is emerging as a potential messenger molecule involved in the modulation of various aspects of physiological processes in plants [36,37]. Signaling by H₂S is proposed to occur via persulfidation, a post-translational modification of protein Cys residues (R-SHs) to form persulfides (R-SSSH). Our results revealed that the ABI4 presulfidation level decreased during seed germination (Figure 3C), and the ABI4 protein stability could be attenuated or accelerated by NaHS or DTT (Figure 4A). Furthermore, mutation of Cys250 to Ala accelerated ABI4 decrease tendency (Figures 4C and 5C). Collectively, these results illustrated that the persulfidation of ABI4 was also linked to ABI4 stability.

In conclusion, we discovered that H₂S could greatly trigger the enhancement of the persulfidation level of ABI4 and alleviate its degradation rate, which in turn inhibits *Arabidopsis* germination. The discovery of ABI4 protein stability regulated by its persulfidation expands our understanding of H₂S role in plant signal transduction networks and establishes a molecular framework for the crosstalk between different post-translational modifications during seed germination.

4. Materials and Methods

4.1. Plant Materials and Growth Conditions

*Arabidopsis* mutants of des1 (SALK_103855; Col-0) and abi4 (SALK_080095; Col-0) mutants were obtained from the Arabidopsis Biological Resource Center (http://www.
Cross abi4 with des1 to obtain the double mutant des1 abi4. 35S:ABI4-GFP transgenic materials were obtained from Dr. Wei Chi (Institute of Botany, The Chinese Academy of Sciences, Beijing, China). Seeds were disinfected with sodium hypochlorite for 20 min and then washed three times with sterile water. They were cultured in Petri dishes on semi-solid Murashige and Skoog (1/2 MS) medium (pH 5.8). The plants were grown in a 16 h/8 h (23 °C/18 °C) growth chamber using a bulb-type fluorescent lamp with a light intensity of 100 mmol photons m$^{-2}$ s$^{-1}$ irradiation.

4.2. Molecular Cloning
4.2.1. For Expression in Escherichia coli

The fragment of ABI4 amplified by PCR was introduced into the PET28a vector (for His fusion) using a homologous recombination technique (Vazyme, Nanjing, China) with the enzyme digestion sites Ndel and XhoI. Site-directed mutagenesis was performed using a Mut Express II Fast Mutagenesis Kit (Vazyme). All procedures followed the manufacturer’s manual. The specific primers used for ABI4 are listed in Supplementary Data 1.

4.2.2. For Transient Expression in Arabidopsis Protoplasts

Homologous recombination technology (Vazyme) was used to transfer PCR amplified fragments into the PAN580 vector at XbaI and BamHI. Arabidopsis protoplasts were extracted according to the method previously described by Yoo et al. [38] with some modifications. The constructed transient expression vector PAN580 was transformed into Arabidopsis protoplasts combined with PEG calcium-mediated method and cultured in the dark at 22 °C for 16 h. The protoplasts expressing target GFP were identified by fluorescence microscopy and used for transient expression analysis.

4.2.3. For Expression in Planta

ABI4 fragments were cloned into the pCAMBIA 1302 vector. The constructed plasmid was transferred into a competent cell of Agrobacterium tumefaciens and then transformed into Arabidopsis using the inflorescence infection method. 1/2 MS medium with 50 mg/mL hygromycin B was used to select transgenic plants. PCR, fluorescence observation, and western blot analysis were combined to identify transgenic plants.

4.3. Real-Time RT-PCR Analysis

Mature Arabidopsis seedlings were collected for RNA extraction. According to the manufacturer’s instructions, seedlings were ground using a mortar and pestle in liquid nitrogen until a fine powder appeared, and then separated total RNA using RNA-easy Isolation Reagent (Vazyme, Nanjing, China). 1000 ng RNA from seedlings was used to synthesize the first-strand cDNA in a 20 µL reaction volume (Vazyme, Nanjing, China) using 1 µM of primers. According to the manufacturer’s instructions, the AceQ qPCR SYBR green master mix (Vazyme) was used for real-time RT-PCR in the Mastercycler ep realplex real-time PCR system (Eppendorf, Germany). The specific primers used are listed in Supplementary Data 1.

4.4. Determination of Activity of L-Cysteine Desulfhydrase

The method for determining L-cysteine desulfhydrase activity was described by Riemenschneider et al. [39] with some modifications. 0.2 g of seedlings was collected and ground with liquid nitrogen and the soluble protein was extracted using 1 mL 20 mM pH 8.0 Tris-HCl. The protein concentration was determined using a BCA kit (Takara, Dalian, China) and calibrated to be consistent. The release of H$_2$S was determined to evaluate L-cysteine desulfhydrase activity. 100 mM Tris HCl (pH 9.0), 2.5 mM DTT, 0.8 mM L-cysteine and 10 µg protein solutions were mixed to 1 mL. The reaction was initiated by the addition of L-cysteine and terminated by 100 µL 30 mM FeCl$_3$ and 100 µL 20 mM N, N-dimethyl-p-phenylenediamine dihydrochloride after incubation at 37 °C for 15 min in the dark. The content of H$_2$S was measured by colorimetric at 670 nm. Taking the known
Na$_2$S concentration as the standard curve, the activity of L-cysteine desulphydrase was expressed as nmol g$^{-1}$ FW min$^{-1}$.

4.5. Seed Germination and Green Open Cotyledon Assays

The seeds were germinated and grown on $1/2$ MS medium with or without NaHS, if there were no other instructions in the text. The percentage of germinated seeds and green cotyledons were recorded, and the seedlings were photographed at the designated time points.

4.6. Expression and Purification of Recombinant Protein

The recombinant protein was expressed and purified in BL21 competent cells (Vazyme). 0.1 mM IPTG was added, and the bacteria were grown to OD$_{600}$ = 0.4 to 0.6 at 16°C for 12 h. After enriching the bacterial solution, suspend the pellet in PBS buffer and use an ultrasonic breaker. The protein was broken and centrifuged at 12,000× g for 30 min, and then the extract was collected for purification. NI-NTA pre-packaged gravity column (Sangon Biotech, Shanghai, China) was used to purify the His-labeled protein, and the protein purification procedure was performed in accordance with the column specifications.

4.7. SDS-PAGE and Immunoblotting

Protein extracts were separated by 12.5% SDS-PAGE. The electrophoresis was ended when the bromophenol blue was moved to 5 mm below the gel, and the gel was transferred to a polyvinylidene fluoride membrane (Roche, Shanghai, China) for 60 min at 100 V on ice using a wet transfer method. The membrane was rinsed with deionized water, then immersed in a blocking buffer (5% skim milk), placed on a decolorizing shaker, and incubated slowly for one hour or overnight at 4°C. Place the membrane in TBST with a sufficient amount of primary antibody and incubate at room temperature for 2 h with gentle shaking. After the primary antibody incubation, the membrane was washed three times with TBST for 7 min each time. The appropriate HRP-labeled secondary antibody was labeled according to the source of the primary antibody, it was diluted in the corresponding proportion, and shaken gently at room temperature for 1 h. The horse-radish peroxidase HRP-ECL luminescence method was used to perform the immunoblot analysis. The software ImageJ (https://imagej.nih.gov/ij/, accessed on 31 December 2021) was used to quantify protein abundance, and signals from two independent experiments were quantified. Full-sized membrane scans are presented in Supplemental Figure S7.

4.8. Immunochemical Detection of S-Persulfidated Proteins

The protein persulfidation level was detected with a tag-switch method described by Aroca et al. [15] with modifications. 35S:ABI4-GFP transgenic seedlings were grown in $1/2$ MS medium. Seedling extracts were dissolved with protein extraction buffer (25 mM pH8.0 Tris-HCl, 100 mM NaCl, and 0.2% Triton X-100) containing protease inhibitor (Yeasen, Shanghai, China). The extract was centrifuged at 4°C for 10 min and centrifuged at 12,000× g for 10 min. Take 80 µL supernatant and add 320 µL blocking buffer (50 mM methylsulfonylbenzothiazole was dissolved in tetrahydrofuran), incubated at 37°C for 1 h to block free sulfhydryl groups. The protein was precipitated twice with acetone, the pellet was resuspended with buffer (50 mM pH 8.0 Tris-HCl, 2.5% SDS, and 20 mM HPDP-biotin), and then incubated at 37°C for 3 to 4 h. 33 µL of protein loading buffer was added without β-mercaptoethanol. It was then incubated at 95°C for 5 min, the protein was separated with 12.5%-SDS-PAGE, transferred to the PVDF membrane, and the level of persulfidation was detected with an anti-GFP antibody (Beyotime, Shanghai, China).

4.9. Cell-Free Protein Degradation Assay

The cell-free protein degradation assay was performed as described with some modifications [32]. 10-day-old WT seedlings were harvested, homogenized in liquid nitrogen, and suspended in degradation buffer (25 mM pH 7.4 Tris-HCl, 10 mM MgCl$_2$, 50 mM NaCl,
1 mM DTT, 0.2% Triton X-100, 5 mM ATP, and 1 mM PMSF). The lysis was centrifuged at 12,000 × g at 4 °C for 10 min. 100 ng purified recombinant ABI4-His protein was incubated with 500 µg total proteins in a total volume of 100 µL for each reaction. Samples were kept at 25 °C. The reactions were stopped at indicated times by adding a 5 × SDS loading buffer. The samples were incubated at 95 °C for 5 min and subjected to western blot analysis with anti-His antibody (Beyotime, Shanghai, China).

4.10. Protoplast-Based Protein Degradation Assay

Protoplasts were lysed in 200 µL degradation buffer (25 mM pH 7.4 Tris-HCl, 50 mM NaCl, 1 mM DTT, 1 mM PMSE, 5 mM ATP and 0.2% Triton X-100) with 150 µM CHX. The lysis was divided into 5 tubes (80 µL each) and incubated at 30 °C. The reaction was terminated by adding 20 µL 5 × SDS loading buffer at the indicated time points. Samples were kept on ice until all the reactions were completed, then incubated at 95 °C for 5 min and subjected to Western blot analysis with anti-GFP antibody (Beyotime, Shanghai, China).

4.11. Statistical Analysis

The statistical analysis and graph construction were performed using SPSS v16.0 (https://www.ibm.com/products/spss-statistics, accessed on 30 December 2021). Differences were considered significant at p < 0.001, 0.01, or 0.05.

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