Characterization of the Arabidopsis thaliana E3 Ubiquitin-Ligase AtSINAL7 and Identification of the Ubiquitination Sites

Diego A. Peralta¹, Alejandro Araya³, Cristina F. Nardi², Maria V. Busi¹*, Diego F. Gomez-Casati¹*

¹ Centro de Estudios Fotosintéticos y Bioquímicos (CEFobi-CONICET), Universidad Nacional de Rosario, Rosario, Argentina, ² Instituto de Investigaciones Biotecnológicas, Instituto Tecnológico de Chascomús (IIB-INTECH-CONICET), Universidad Nacional de San Martín, Chascomús, Argentina, ³ Centre National de la Recherche Scientifique and UMR 1332 – Biologie du Fruit et Pathologie, Institute National de la Recherche Agronomique (INRA) Bordeaux Aquitaine, Villenave D’Ornon, France

Abstract

Protein ubiquitination leading to degradation by the proteasome is an important mechanism in regulating key cellular functions. Protein ubiquitination is carried out by a three step process involving ubiquitin (Ub) activation by a E1 enzyme, the transfer of Ub to a protein E2, finally an ubiquitin ligase E3 catalyzes the transfer of the Ub peptide to an acceptor protein. The E3 component is responsible for the specific recognition of the target, making the unveiling of E3 components essential to understand the mechanisms regulating fundamental cell processes through the protein degradation pathways. The Arabidopsis thaliana seven in absentia-like 7 (AtSINAL7) gene encodes for a protein with characteristics from a C3HC4-type E3 ubiquitin ligase. We demonstrate here that AtSINAL7 protein is indeed an E3 protein ligase based on the self-ubiquitination in vitro assay. This activity is dependent of the presence of a Lys residue in position 124. We also found that higher AtSINAL7 transcript levels are present in tissues undergoing active cell division during floral development. An interesting observation is the circadian expression pattern of AtSINAL7 mRNA in floral buds. Furthermore, UV–B irradiation induces the expression of this transcript indicating that AtSINAL7 may be involved in a wide range of different cell processes.

Introduction

Protein turnover through the ubiquitin-mediated proteasome system plays a pivotal role in many regulatory pathways such as growth, cell differentiation, cell cycle control, stress response and apoptosis [1–3]. First described in Drosophila melanogaster, Seven in absentia (SINA) proteins are E3 ubiquitin ligases with a characteristic N-terminal RING (Really Interesting New Gene) finger domain, linked to a conserved C-terminal domain required for oligomerization and binding to target proteins [4]. The D. melanogaster SIN A regulates photoreceptor differentiation by targeting the transcription factor Tramtrack for proteasomal degradation [5–7]. The RING finger motif is defined as a 40-60 cysteine rich domain coordinating two Zinc ions which can fold into a compact domain comprising a small central β-sheet and an α-helix [8] (http://www.ncbi.nlm.nih.gov/Structure/cdd/cddsrv.cgi?uid=pfam00097).

The RING finger domain from many E3 ubiquitin-ligases is required for interaction with an E2 ubiquitin-conjugation protein, leading to the transfer of ubiquitin to the target protein. E3 ubiquitin-ligases are able to recognize a large number of targets through adaptor proteins, which provides precise functional specificity [9,10]. Besides the proteolytic pathways, protein ubiquitination can also regulate protein functions [11,12], making these enzymes key mediators of post-translational protein regulation. Homologous to the Sina superfamily, which is composed of about 35 highly conserved proteins, have also been found throughout eukaryotes [13,14].

The Arabidopsis genome contains more than one thousand genes encoding for E3 ubiquitin ligases, 469 of them predict proteins presenting one or more of the various types of RING domains [15]. Eighteen Siah homologous, SIN A, SINAT1–T5, SINAL1–11 and PEX14 are encoded by the A. thaliana genome (www.arabidopsis.org). The participation of Sina in plant resistance to pathogens and plant growth has been
described [16,17]. Other studies revealed the existence of a link between hormone response on floral development and the ubiquitination pathway [18]. However, very little is known about the function of Siah protein counterparts in plants.

Several lines of evidence reveal that floral development requires proper mitochondrial function [19–21]. We recently demonstrated that plants with a mitochondrial dysfunction were affected in Arabidopsis thaliana SINA like 7 (AtSINAL7, At5g37890) mRNA levels [22,23]. AtSINAL7, predicted to be expressed in 21 plant structures most of the growth stages, seem to be involved in multicellular organismal development such as protein ubiquitination and ubiquitin-dependent protein catabolic process (http://www.arabidopsis.org/servlets/TairObject?id=131882&type=locus). However, the precise function of this protein remains unknown.

We decided to characterize biochemically the AtSINAL7 encoded protein and to study their expression. AtSINAL7 is a protein characterized by the presence of a canonical C3HC4 RING-type cysteine-rich domain able to coordinates two zinc atoms, homologous to the RING-finger E3 ubiquitin ligase protein, SINA [9]. Here, we show that AtSINAL7 functions, indeed, as an ubiquitin ligase using a self-ubiquitination assay. We present evidence that this activity depends of a Lys residue. Moreover, we study the expression of AtSINAL7 in different tissues and during daily light cycle and under UV–B exposure. These results strongly argue that AtSINAL7 is an E3 ubiquitin ligase and suggests that it plays an important role in cell processes, particularly during flower development.

Results

Expression and purification of recombinant AtSINAL7

To characterize the seven in absentia like 7 protein from Arabidopsis thaliana, the DNA fragment containing the AtSINAL7 coding sequence (286 codons) was fused to an N-terminal His6-tag when cloned onto pRSETb expression vector. The recombinant protein was purified using a HisTrap chelating affinity chromatography after expression in E. coli (BL21) pLys strain. The purified recombinant AtSINAL7 protein of 32 KDa was successfully induced (Figure 1, lane 2) and purified to homogeneity as shown by protein staining (lane 3). The presence of recombinant AtSINAL7 was assessed by Western blot analysis using with the anti-His antibodies (lane 4).

Transcript levels of AtSINAL7 in different tissues of Arabidopsis

The expression of AtSINAL7 was determined in several tissues from wild-type Arabidopsis by qRT-PCR (Figure 2A). RNA from Root, Rosette leaves, Inflorescence stage 6, Inflorescence stage 12, and Siliqua were isolated from 28 days old A. thaliana Col 0 plants grown under long-day condition greenhouse (see Methods). cDNA was synthesized from total RNA and was quantified spectrophotometrically. Identical amounts of different cDNA samples were used for qPCR amplifications using Arabidopsis sина like 7 with specific primers (Table 1). The data presented concern relative values obtained from the average of three biological and technical replicates using the cbp gene (cap binding protein At5g44200) as a house-keeping control [24] (Figure 2A). AtSINAL7 transcript levels are highly expressed in siliques (2.3770) and stage 12 inflorescences (0.7053), and roots (0.1524), being lower at Stage 6 inflorescences (0.08) and rosette leaves (0.06). Thus, AtSINAL7 seems to be expressed in a tissue-specific manner with a strong induction of 12 and 40 times in stage 12 flowers and siliqua respectively, compared to rosette leaves.

AtSINAL7 gene expression show a light-dependent behavior in stage 12 flowers

Flowering behavior is strongly dependent from different environmental signals, particularly light. Day length, a major regulator of flowering, allows sexual reproduction to proceed at an appropriate time [25]. As the higher expression was obtained in floral tissues, a time course profile of the study of AtSINAL7 transcript levels was performed on Arabidopsis stage 12 inflorescences throughout 24 hours under long day growth conditions (see Methods). The expression of cbp gene (At5g44200) was used as an internal standard ( [24]). The mRNA levels decrease after 4 h darkness reaching a minimum value (0.365). At the beginning of daylight, the mRNA levels increase to reach a maximum value (0.7053) at 20 hours of the experimental period (12 h after light onset). Thus, a two-fold...
Figure 2. Analysis of AtSINAL7 mRNA levels (A) AtSINAL7 transcript expression profile in different organs from A. thaliana. AtSINAL7 transcript levels relative to CBP control gene expression were determined by RT-qPCR in several A. thaliana organs. Data shown represent at least three independent experiments (significant statistical difference was determined using t test, P <0.05). (B) Time-course expression of AtSINAL7 transcripts. Determination of the transcription levels of AtSINAL7 at 0, 4, 8, 12, 16, 20, and 24 hours under long day growth conditions. Transcripts levels are plotted as relative values using the cap binding protein (At5g44200) mRNA as an internal control. Data shown represent at least three independent experiments (significant statistical difference was determined using t test, P <0.05). (C) Effect of the UV–B treatment on the AtSINAL7 transcript levels in wild type A. thaliana inflorescences. The AtSINAL7 transcript expression in A. thaliana inflorescences (stage 12) relative to cpk3 (calcium-dependent protein kinase 3, At4g23650) was determined after 4 h of UV–B exposure. Control plants were protected from UV–B irradiation using polyester filters (see Methods). Data shown represent at least three independent experiments (significant statistical differences determined using t test, P <0.05).

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Table 1. Primers Used in qPCR experiments.

| Gene       | Primer Name | Sequence                  |
|------------|-------------|----------------------------|
| At5g44200  | CBPFw       | CCG GCC TAT TCG TGT GGA TTT TGA |
| At5g44200  | CBPRv       | CAT AAT TCG TTG GCG CAG CTT GAG |
| At4g23650  | CPK3Fw      | AAT CCA CGG ATG ATT TAG CA   |
| At4g23650  | CPK3Rv      | ATC TGG AGT GCT GGT GTG AT   |
| At5g37890  | RTAtSINAL7Fw| GCT ACG AAG CTT TCA CAA TCC C |
| At5g37890  | RTAtSINAL7Rv| GTA CAG ATC CTT GTA GCT A   |

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Characterization of AtSINAL7 Ubiquitin Ligase

variation of AtSINAL7 transcript levels is observed over a daily basis (Figure 2B).

AtSINAL7 transcript levels increase after UV–B treatment

Ubiquitination plays an important role in DNA damage signal amplification. Upon UV-induced DNA damage, several ubiquitin-tagged proteins are degraded by proteasome [26]. To investigate the behavior of AtSINAL7 gene in plants under UV–B exposure, we determined the transcript levels by qRT-PCR at stage 12 inflorescence. The experiment was carried out as follows: wild type Arabidopsis plants were grown in a long-day greenhouse during 4 weeks and then irradiated with UV–B light at 16 hours (8h dark plus 8h light) after the onset of the experiment. Control plants, present in the same environment, were protected with a plastic membrane able to exclude the UV–B light. The inflorescence stage 12 cDNA from control and UV-treated plants were submitted to qRT-PCR analysis (Figure 2C) using cpk3 (calcium-dependent protein kinase 3, At4g23650) as a house-keeping control gene whose expression remains unchanged after UV–B treatment [27]. The results allowed us to conclude that SINAL7 gene expression augments almost 2-fold in inflorescences after UV–B treated plants. Similar results were obtained in rosette leaves, (data not shown), proving that UV–B irradiation triggers AtSINAL7 transcript expression.

AtSINAL7 undergoes self-ubiquitination

The in vitro ubiquitination assay was carried out using heterologous partners: a yeast ubiquitin-activating (E1) enzyme and the human recombinant UbcH2 ubiquitin-conjugating (E2) enzyme, in the presence of cMyc-tagged ubiquitin (cMyc-ubiquitin) as described in the Methods section. Purified recombinant AtSINAL7 was used as ubiquitin-ligating (E3) enzyme. As expected, no signal was detected in lanes lacking cMyc-ubiquitin (Figure 3A, lanes 1, 2, 3, 5, and 6), indicating that the anti-cMyc antibodies do not cross react with other protein components present in the reaction mixture. Lanes 4 and 7, containing only the tagged ubiquitin and the reaction mixture lacking the E3 component respectively, show four bands characteristic of the monomer (9.3 kDa) and oligomers corresponding to cMyc-ubiquitin. Interestingly, the complete reaction mixture shows a protein band corresponding to ubiquitinated AtSINAL7 migrating with an apparent molecular weight approximately 72 kDa (Figure 3A, lane 8). To assess
that AtSINAL7 was indeed self-ubiquitinated the samples 5 and 
8 containing the E3 component alone and the complete 
reaction mixture respectively were analyzed by Western blot 
using anti-His6 (Santa Cruz Biotechnology) and anti-
recombinant AtSINAL7 antibodies (Figure 3B). The same band 
is revealed by both anti-His and anti-AtSINAL7 (Figure 3B , 
lanes 2 and 3). It is interesting to note that AtSINAL7 identified 
by anti-AtSINAL7 antibody (lane 3) migrates at 34 kDa (lane 4) 
while the self-ubiquitinated form show a mobility shift to near 70 
kDa, suggesting that more than one ubiquitin molecules is has 
been incorporated to the AtSINAL7 protein.

The residue K124 is involved in self-ubiquitination of 
AtSINAL7

To determine the residues involved in the AtSINAL7 self-
ubiquitination, in silico analysis was conducted using the program UbPred: Predictor of protein ubiquitination sites [28]. 
Two out of 19 Lys, residues on AtSINAL7 were predicted as 
susceptible to accept the ubiquitin molecule by the bio-
computer analysis (http://www.ubpred.org/), at position K23 
and K124. To confirm this hypothesis, we construct mutant 
proteins where the residues K23 and K124 were replaced by 
alanine (see methods). Three constructs K23A, K124A and the 
double mutant K23A/K124A were used to produce the recombinant proteins SL7K23A, SL7K124A and SL7K23AK124A for in vitro ubiquitination experiments (Figure 4). Only the mutant SL7K23A was able to sustain self-
ubiquitination (Figure 4, lane 2) which is comparable to result 
obtained with the wild-type control (Figure 4, lane 7). In 
contrast, the mutant SL7K124A (lane 4) and the double mutant 
SL7K23AK124A (Figure 4, lane 6) were unable to underwent 
self-ubiquitination.

Mutations K23A and K124A do not affect protein 
folding

Since the mutation of Lys residues may induce structural 
changes on the protein, we decided to study the secondary 
structure of the recombinant AtSINAL7 and the respective K-to-
A mutants. The correct folding of recombinant 
AtSINAL7(K23A), AtSINAL7(K124A) and 
AtSINAL7(K23AK124A) proteins was analyzed using circular 
dichroism (CD) (Figure 5). The percentage of the secondary 
structure was estimated from CD spectra using the K2D 
algorithm [29]. The content of alpha helix, beta sheet and 
random coil were 26.76, 30.03 and 43.21% respectively for the 
recombinant AtSINAL7. Interestingly, all three mutants 
presented identical CD spectra to the wild type AtSINAL7 
protein, indicating that no changes on the secondary structure 
were induced when the residues K23, K124 or both 
simultaneously were changed for Ala on the protein.

Considering that the average error observed from CD spectra 
was lower than the values obtained in secondary structure 
prediction using K2D (0.08; 0.09), we conclude that the lack of 
self-ubiquitination of AtSINAL7(K124A) and 
AtSINAL7(K23AK124A) was not resulting from a loss of
secondary structure, but to the replacement of the Lys-124 residue.

Discussion

AtSINAL7 is a RING-finger protein homologous to the Drosophila’s seven in absentia (sina) protein. The Drosophila sina superfamily possesses important functions in development control and protein degradation by the proteasome. These proteins are an essential part of the ubiquitin protein modification machinery found in all eukaryotic organisms [9,18]. This particular protein modification participates in many different cell process by affecting gene expression through the protein turnover pathways or by modification of the protein function [2]. The important role played by these proteins is resulting from both the E3 ubiquitin ligase activity, and the capacity of E3 ligases to recognize specifically the target protein. Thus, the knowledge of E3 RING-finger ubiquitin ligases is crucial to understand the physiological mechanisms involved in plant development and adaptation.

Eighteen genes homologous to Siah from Sina superfamily, SINA, SINAT1–T5, SINA like 1–11, and PEX14 are present in the Arabidopsis genome (www.arabidopsis.org). Several functions have been reported for plant Sina genes such as resistance to pathogens, plant growth [16,17], hormone response and floral development [18]. Interesting, floral development depends on proper mitochondrial function. Consistent with this observation, we found that mitochondrial dysfunction affected AtSINAL7 expression in young flowers [22,23]. Here, we show that AtSINAL7 transcripts are more abundant in tissues engaged in active cell division in different organs of A. thaliana, with higher levels in flowers and siliques (Figure 2A), these results are in agreement with the transcript level pattern shown in Arabidopsis eFP Browser at bar.utoronto.ca [30]. An important result of this study is the fact that the transcript levels oscillate daily with the lower level at 4 hours in dark, and attains the higher levels after 12 hours of light onset in a long-day growing cycle. This behavior is in agreement with the presence of putative circadian control boxes in the

![Figure 5. Far-UV CD Spectra of recombinant AtSINAL7 and AtSINAL7 mutants.](doi:10.1371/journal.pone.0073104.g005)
promoter region of the AtSINAL7 gene, predicted by using the Genomatix Software Suite (http://www.genomatix.de/solutions/genomatix-software-suite.html). In addition, an increase of AtSINAL7 level was induced by UV–B irradiation in Arabidopsis. During DNA damage response generated by UV light, several proteins undergo polyubiquitylation to be processed by the proteasome [26], indicating that important transcriptional and metabolic processes are affected by UV–B irradiation [31]. It is interesting to note that the expression of the multifunctional E3 ubiquitin ligase, COP1 gene, is controlled by a combinatorial regulation of FHY3 and HY5 in response to UV–B [32]. Thus, considering that AtSINAL7 has the signature of a RING-type E3 ubiquitin ligase, the increase of transcript upon UV–B exposure is in agreement with a potential role in complex signalling pathways.

We demonstrate that SINAL7 is able to participate as an ubiquitin ligase (E3 component) in an E1-E2-E3 assay (Figure 3A). Moreover, we found that the Lys24, located in the DNA domain, is required for this function. However, we tested this function in a self-modification assay; if this residue acts as an acceptor of ubiquitin, plays a role in catalysis or both, remains to be elucidated.

Self-ubiquitination of E3 ubiquitin ligases has been proposed to regulate their activity, the recruitment of substrates and to participate in non-catalytic functions of these proteins [1,33]. Considering that AtSINAL7 contains a TRAF-like domain, self-ubiquitination may be a mechanism to recruit substrates with ubiquitin-ubiquitin properties as shown for the TRAF6, a RING domain E3 ligase that has a crucial role in the initial activation of signaling cascades [34–36]. Thus, it is possible to speculate that AtSINAL7 undergo self-ubiquitination as a target for degradation by the proteasome complex, but also to accomplish different activities in plant cells. In vivo studies will be necessary to validate this hypothesis.

Conclusions

We show here that AtSINAL7, a predicted RING-domain ubiquitin E3 ligase, is able of undergo self-ubiquitination in vitro. This activity is dependent of the presence of a Lys residue in position 124. In addition, we show that AtSINAL7 transcript levels are high in Arabidopsis thaliana tissues undergoing active cell division, suggesting a role during floral development. An interesting observation is the circcadian expression pattern to be elucidated.

Materials and Methods

Plant Material and Bacterial Strains

Arabidopsis thaliana (var. Columbia Col-0), grown in a greenhouse under long day conditions (16 day/8 night), were used in this study. Escherichia coli BL21(DE3) pLysS strain (E. coli B F- dcm ompT hsdS(rB–mB–) gal K1(DE3) [pLysS CamR]) was used as bacterial hosts in cloning and expression experiments.

Cloning and Expression of AtSINAL7

The total RNA extracted from Arabidopsis leaves was used as template for cDNA synthesis using random hexamers. The cDNA fragment (860bp) containing the AtSINAL7 coding region was PCR amplified using GoTaq Polymerase (Promega, WI, USA) and the following primers: SINAL7up: 5’-AAAGGATCCATGGGTGCCGAGTTTTG-’3 and SINAL7down: 5’-CTCGAGTTACTTCTTTGTTCAACCTTGACGAC-’. The PCR product was cloned into pGEMT-Easy (Catalog# A1360 Promega, WI, USA). The AtSINAL7 coding sequence was obtained by double digestion with Xhol and BamHI restriction enzymes from the recombinant plasmid. The purified fragment was inserted into prSETb expression plasmid (Invitrogen, Carlsbad, CA, USA), previously digested with the same restriction enzymes, using TDNA ligase (Promega, WI, USA). The resulting recombinant plasmid, pRSETbSL7, containing 286 codons from the Arabidopsis thaliana seven in absentia like 7 (AtSINAL7) containing a N-terminal His-tag sequence, under the control of phage T7 promoter, was used to transform E. coli BL21 (DE3) pLysS cells.

Purification of AtSINAL7, SL7K23A, SL7K124A and SL7K23AK124A

E. coli BL21 (DE3) pLysS cells harboring plasmid pRSTEbSL7 were grown at 37°C in TB medium, containing 100 mg ml⁻¹ ampicillin to an OD₆₀₀ = 0.6. AtSINAL7 production was induced by the addition of 1 mM IPTG and subsequent incubation at 30°C for 8 h. Cells were harvested and re-suspended in 20 mM Tris-HCl, pH 7.4, containing 1 mM phenylmethylsulfonyl fluoride (PMSF), disrupted by sonication and centrifuged at 7000 x g for 15 min at 4°C. The supernatant was loaded onto a HiTrap chelating column (GE Healthcare). After washing with 20 ml of 20 mM Tris-HCl, pH 7.4, 20 mM imidazole, the recombinant protein was eluted using a 20–500 mM imidazole gradient in buffer 20 mM Tris-HCl, pH 7.4. The elution of the recombinant protein was monitored by enzyme activity and SDS–PAGE analysis of chromatography fractions. The purified enzyme was pooled and concentrated to >1 mg ml⁻¹ and used immediately after purification process.

Protein analyses

SDS–PAGE was performed using 12% (w/v) gels as described by Laemmli [37]. Gels were revealed by Coomassie blue staining or after electro-blotting onto nitrocellulose membranes (Bio-Rad). Membranes were incubated with peroxidase-linked anti-mouse IgG or anti-rabbit IgG, followed by staining with BCIP and NBT [38]. Total protein concentration was determined as described by Bradford [39].

E3 ubiquitin ligase activity assay

In vitro ubiquitination assay was adapted from the protocol described by Wertz et al. [40]. The reaction mixture (25 μl) contained 20 mM HEPES, 100 mM NaCl, 5 mM MgCl₂, 5 mM ATP, 10 mM DTT, 5 μg cMyG-ubiquitin (Boston Biochem), 150
nM yeast E1 (Bostom Biochem), 200 mM human recombinant Ubch2 (Bostom Biochem) and 5 µg of purified His-AtSINAL7. The reaction mixture was incubated at 30°C. After 2 h, the reaction was stopped by adding 5X SDS-PAGE Sample buffer (125 mM Tris-HCl pH 6.8, 20% Glycerol, 4% SDS and 10% β-mercaptoethanol) and boiled at 100°C for 5 min. Protein samples were analyzed by SDS-PAGE electrophoresis followed by protein gel blotting. Blots were immersed using anti-cMyc antibodies (Bostom Biochem), followed by incubation with anti-mouse Alkaline Phosphatase conjugated antibodies (Sigma).

**RNA Preparation and Quantitative Real-Time PCR**

Total RNA was isolated from 30 mg of tissue using SV Total RNA Isolation System (Promega, WI, USA) as described in the manufacturer’s protocol. cDNA synthesized using 5X M-MLV buffer (250 mM Tris-HCl, 375 mM KCl, 15 mM MgCl2, 50mM DTT), dNTPs MIX (dATP, dCTP, dGTP and dTTP 10 mM each), 2 µg/µL random hexamers pd(N) 6 (Amersham #27-2166-01) as primers, Recombinant RNAsin® Ribonuclease Inhibitor (25 units), and 200 units of MMLV reverse transcriptase (USB Corp., Cleveland, OH, USA), incubating 1 h at 37 °C. After quantification by UV absorption at 260 nm, 2 µg of cDNA was used as a template for qPCR amplification in a MiniOPTICON2 apparatus (Bio-Rad), using the intercalation dye SYBRGreen I (Invitrogen) as a fluorescent reporter and GoTaq Polymerase (Promega). Primers, able to amplify unique 150-200 bp products, were designed using the online primer design tool Primer-BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/) (Table 1). Amplification conditions were : 2 min denaturation at 94°C; 40–45 cycles at 94°C for 15 s, 57°C for 20 s, and 72°C for 20 s; followed by 10 min extension at 72°C. Three technical replicates were performed for each sample. RNA from each sample was obtained from pools of at least three plants. Melting profile for each PCR was determined by measuring the decrease of fluorescence with increasing temperature (from 65°C to 98°C). The size of the amplification products was verified on 2% (w/v) agarose gel. Gene expression was normalized to the housekeeping gene [27] in UV–B treatment experiments.

**Site-directed Lys-to-Ala SINAL7 mutants**

The codons for Lys 23 and Lys 124 on the AtSINAL7 coding region were changed to Ala triplets using the QuickChange II XL site-directed mutagenesis kit (Stratagene, La Jolla, CA, USA). The pRSETbSL7 vector was used as template for PCR amplification. The primers used were: SL7K23A, GATCTAAACGACTTTCTCAGCAAgcGAGACAACCTTTCTCAGTGAT; SL7K124A, TTGCAAAAAGAATGTATCTTATGGGgcAGAGTTAACTCATGTG; and their respective complementary oligonucleotides. Base substitutions are indicated by lowercase letter. The resultant, single, SL7K23A, SL7K124A, and double SL7K23AK124A mutant vectors were verified by DNA sequencing and used to transform E. coli BL21 (DE3) pLysS cells.

**UV–B treatment of Arabidopsis plants**

UV–B irradiation of plants was performed as described by Lario [41]. Arabidopsis plants were exposed for 4 h to UV–B radiation in a growth chamber using UV–B bulbs (Bio-Rad, Hercules, CA, USA). UV–B lamps were covered with cellulose acetate filters to exclude the wavelengths below 280 nm (100 nm extra clear cellulose acetate plastic; Tap Plastics, Mountain View, CA, USA) and placed 30 cm above the plants. The UV radiation measured with a UV–B/UVA radiometer (UV203 AB radiometer; Macam Photometrics, Scotland, UK) was 2 W m$^{-2}$ for UV–B and 0.65 W m$^{-2}$ for UV-A. Control plants, were exposed for the same period of time to the light sources described above covered with a polyester filter (100 µm clear polyester plastic; Tap Plastics) to absorb both UV–B (0.04 W m$^{-2}$) and wavelengths <280 (UV-A radiation intensity was 0.4 W m$^{-2}$). Immediately after irradiation, samples from at least three independent biological replicates were collected, frozen in liquid nitrogen, and stored at -80°C until its use for RNA isolation.

**Circular Dichroism (CD) studies**

Far-UV CD spectra were obtained using a Jasco J-810 spectropolarimeter (Jasco International Co.) over the wavelength range from 200 to 250 nm, at 25°C. Measurements were performed in a 0.2 cm quartz cuvette at rate of 100 nm.min$^{-1}$, bandwidth of 1 nm, response time of 2 s, data pitch of 1 nm, and accumulation of 10. CD data are shown as the mean residue ellipticity (deg cm$^{2}$ dmol$^{-1}$) obtained after subtracting the baseline, smoothing, and data normalization. CD spectra for AtSINAL7, AtSINAL7 K23A, AtSINAL7 K124A and AtSINAL7 K23AK124A (0.1-1 mg ml$^{-1}$) were recorded in 20 mM Sodium phosphate buffer, pH 7.4. Secondary structure analysis from CD spectra data was performed using the K2d algorithm [29].

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**Author Contributions**

Conceived and designed the experiments: DAP AA MVB DGC. Performed the experiments: DAP CFN. Analyzed the data: DAP AA MVB DGC. Wrote the manuscript: DAP AA MVB DGC.
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