JUNGBRUNNEN1, a Reactive Oxygen Species–Responsive NAC Transcription Factor, Regulates Longevity in *Arabidopsis*

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The transition from juvenility through maturation to senescence is a complex process that involves the regulation of longevity. Here, we identify JUNGBRUNNEN1 (JUB1), a hydrogen peroxide (H_2O_2)-induced NAC transcription factor, as a central longevity regulator in *Arabidopsis thaliana*. JUB1 overexpression strongly delays senescence, dampens intracellular H_2O_2 levels, and enhances tolerance to various abiotic stresses, whereas in *jub1-1* knockdown plants, precocious senescence and lowered abiotic stress tolerance are observed. A JUB1 binding site containing a RRYGCCGT core sequence is present in the promoter of *DREB2A*, which plays an important role in abiotic stress responses. JUB1 transactivates *DREB2A* expression in mesophyll cell protoplasts and transgenic plants and binds directly to the *DREB2A* promoter. Transcriptome profiling of *JUB1* overexpressors revealed elevated expression of several reactive oxygen species-responsive genes, including heat shock protein and glutathione S-transferase genes, whose expression is further induced by H_2O_2 treatment. Metabolite profiling identified elevated Pro and trehalose levels in *JUB1* overexpressors, in accordance with their enhanced abiotic stress tolerance. We suggest that JUB1 constitutes a central regulator of a finely tuned control system that modulates cellular H_2O_2 level and primes the plants for upcoming stress through a gene regulatory network that involves DREB2A.

INTRODUCTION

In plants, the transition from juvenility through maturation to senescence is a physiologically complex process that involves a large number of molecular and physiological events regulated by genetically determined and environmentally modified regulatory networks (e.g., Smart et al., 1995; Hinderhofer and Zentgraf, 2001; He and Gan, 2002; Gepstein et al., 2003; Buchanan-Wollaston et al., 2005; van der Graaff et al., 2006; Lim et al., 2007). Senescence is triggered by adverse environmental conditions, such as high salinity, low light intensity, drought, pathogen attack, nutrient deficiency, and other stresses (Dwidedi et al., 1979; Dhindsa et al., 1981; Bohnert et al., 1995; Balazadeh et al., 2010a, 2011). Many genes, including those that control

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[™]Open Access articles can be viewed online without a subscription. www.plantcell.org/cgi/doi/10.1105/tpc.111.090894 transcription, undergo expression changes during senescence (e.g., Gepstein et al., 2003; Guo et al., 2004; Buchanan-Wollaston et al., 2005; van der Graaff et al., 2006; Balazadeh et al., 2008b; Parlitz et al., 2011). Recently, high-resolution temporal profiling of gene expression during leaf senescence in Arabidopsis thaliana revealed clusters of coexpressed genes and a distinct chronology of senescence-associated processes (Breeze et al., 2011). Among the transcription factors (TFs), the NAC (for NAM, ATAF1, 2, and CUC2) and WRKY families are particularly rich in senescence-regulated TFs in many plant species (Andersson et al., 2004; Guo et al., 2004; Lin and Wu, 2004; Buchanan-Wollaston et al., 2005; Gregersen and Holm, 2007; Balazadeh et al., 2008b), suggesting they play important roles in leaf senescence. Although more than 20 of the 106 known NAC genes in Arabidopsis exhibit senescence-dependent expression (and, thus, represent senescence-associated genes [SAGs]), a distinct regulatory function with respect to senescence has only been reported for some members so far, including At NAP (for NAC-LIKE, ACTIVATED BY APETALA3/PISTILLATA; also called Arabidopsis NAC [ANAC] 029; Guo and Gan, 2006), ORESARA1 (ORE1; ANAC092, At NAC2; Kim et al., 2009; Balazadeh et al., 2010a), and ORESARA1 SISTER1 (ORS1; ANAC059; Balazadeh et al., 2011). Inhibiting either NAC individually delays senescence, identifying them as nonredundant positive regulators of

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senescence (Guo and Gan, 2006; Kim et al., 2009; Balazadeh et al., 2010a, 2011). A dual-function NAC gene, VASCULAR-RELATED NAC-DOMAIN INTERACTING2 (VNI2; ANAC083), was recently reported to integrate abscisic acid (ABA) signaling with leaf senescence (Yang et al., 2011) and to negatively regulate xylem vessel formation (Yamaguchi et al., 2010). Genes downstream of ORE1 and ORS1 have been identified; from these studies, it became apparent that both TFs exert their function through regulatory networks that include many known SAGs (Balazadeh et al., 2010a, 2011). Of the 170 genes upregulated after expression of ANAC092 is induced, 102 genes were also enhanced by senescence in the time-course experiment reported by Breeze et al. (2011). The majority of these genes (75%) fell into clusters whose gene members were enriched for promoterlocalized NAC binding sites. This observation supports the conclusion that ANAC092/ORE1 and most likely other NAC TFs play an important role in regulating senescence.

An apparent signaling element for the regulation of senescence is hydrogen peroxide (H₂O₂). Diverse environmental and developmental stimuli, such as heat stress, salinity, cold, and pathogen attack, are known to trigger an accumulation of intracellular H₂O₂ and through this regulate the expression of many genes, including TFs of, for example, the NAC, WRKY, and APETALA2 (AP2)/ethyleneresponsive element binding protein (EREBP) families (Vanderauwera et al., 2005; Gadjev et al., 2006). Notably, the expression of at least 15 senescence-associated NAC TFs increases rapidly after H₂O₂ treatment, including, for example, At NAP, ANAC092/ORE1, ORS1, ARABIDOPSIS TRANSCRIPTION ACTIVATION FACTOR1 (ATAF1), and ANAC032 (Balazadeh et al., 2010b, 2011). However, the H₂O₂ and senescence-dependent gene regulatory networks of these TFs are just emerging and are far from being fully understood.

DEHYDRATION-RESPONSIVE ELEMENT BINDING PRO-TEIN2A (DREB2A) belongs to the EREBP family of TFs. Although DREB2A expression increases during senescence, its biological role in this process has not been analyzed. Expression of DREB2A is induced by dehydration, salinity, heat, and different oxidative stress treatments, including H₂O₂ stress (Gadjev et al., 2006; Sakuma et al., 2006a; Suzuki et al., 2011). DREB2A regulates the water deficit-inducible expression of target genes and requires posttranslational modification for its activation. Two C3HC4 RING domain-containing proteins, including DREB2A-INTERACTING PROTEIN1 (DRIP1) and DRIP2, have been discovered and shown to interact with DREB2A in the nucleus. DRIPs act as E3 ubiguitin ligases that mediate the ubiguitination of DREB2A, thereby targeting the TF to 26S proteasome degradation. DRIP1 and DRIP2 thus function as negative regulators of drought-responsive gene expression (Qin et al., 2008). Recently, Suzuki et al. (2011) identified a new heat stress regulon in Arabidopsis regulated by multiprotein bridging factor 1c (MBF1c). DREB2A is one of the 36 genes whose expression during heat stress is regulated by MBF1c, and genetic studies demonstrated that DREB2A is required for plant survival under heat stress (Suzuki et al., 2011). Overexpression of a constitutively active form of DREB2A (35S:DREB2A CA) resulted in the upregulation of nearly 500 genes, including drought- and saltresponsive genes, and also heat shock-related genes (Sakuma et al., 2006b). The heat shock TF gene HsfA3 is a known direct target of DREB2A (Schramm et al., 2008).

Here, we report the identification of JUNGBRUNNEN1 (JUB1), a NAC TF that negatively regulates senescence. *JUB1* is rapidly and strongly induced by H_2O_2 treatment. Overexpression of *JUB1* markedly extends leaf longevity and promotes tolerance to various abiotic stresses. We found that increased tolerance to abiotic stress correlates with reduced levels of H_2O_2 in *JUB1* overexpressors, whereas the opposite is observed in *jub1-1* knockdown plants, suggesting that JUB1 participates in regulating the cellular H_2O_2 homeostasis network. We further determined the binding site of the JUB1 TF by in vitro binding site selection and discovered *DREB2A* as one of its direct downstream target genes. Thus, JUB1 links H_2O_2 signaling to senescence regulation and the downstream activation of *DREB2A* and its direct target *HsfA3*.

RESULTS

JUB1 Promotes Leaf Longevity

To identify novel transcription regulators that modulate leaf senescence, we systematically screened NAC gene T-DNA insertion mutants and cauliflower mosaic virus (CaMV) 35Sdriven overexpression lines for extended longevity compared with wild-type Arabidopsis plants kept under identical growth conditions. We found that plants overexpressing At2g43000 developed senescence considerably later than the wild type and retained fully green leaves for a much longer period (see below). By contrast, a knockdown mutant of At2q43000 (Salk ID 036474) developed leaf senescence earlier than the Columbia-0 (Col-0) wild type (see below). Similarly, we observed earlier senescence in transgenic plants upon suppression of At2g43000 transcript abundance by means of a genome-inserted artificial microRNA construct (see Supplemental Figure 1 online). These observations identify At2g43000 as a novel genetic regulator of plant senescence. As elevated expression of At2g43000 extended longevity, we designated it JUNGBRUNNEN1 (JUB1; German for "Fountain of Youth").

JUB1 encodes a 275–amino acid protein of a calculated molecular mass of 31.5 kD. JUB1 contains a NAM domain (pfam02365) at its N terminus. Its coding region consists of three exons, interrupted by two introns. Phylogenetic analyses revealed the presence of *JUB1* orthologs in other plant species, including rice (*Oryza sativa*; ONAC066; Ooka et al., 2003) and *Populus trichocarpa* (PNAC080-083; Hu et al., 2010). Expression of a JUB1–green fluorescence protein (GFP) fusion in *Arabidopsis* showed its predominant accumulation in the nucleus (see Supplemental Figure 2 online), consistent with its function as a transcription regulator.

RNA gel blot analysis and quantitative RT-PCR (qRT-PCR) confirmed elevated *JUB1* transcript level in multiple independent *35S:JUB1* transformants compared with the wild type (Figures 1A and 1B). When grown in soil under long-day conditions, leaf senescence and bolting were strongly delayed in *35S:JUB1* plants compared with the wild type and empty vector (EV)–transformed control lines (Figure 1C). Overexpressors started bolting 10 to 14 d later than the EV lines. Chlorophyll levels of 12 individual rosette leaves of overexpression plants were higher than those in age-matched EV plants (Figure 1D). Mortality

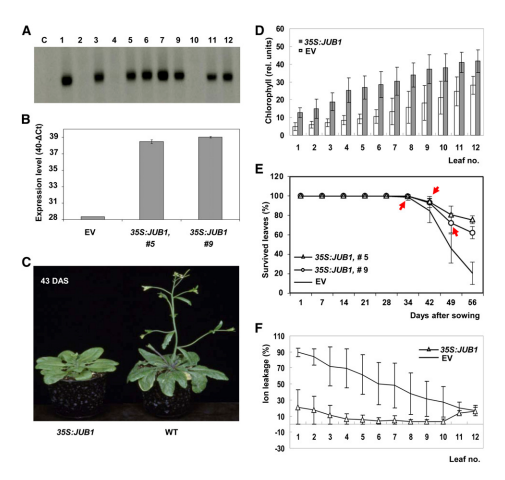


Figure 1. Physiological and Molecular Characterization of JUB1 Overexpression Plants.

(A) RNA gel blot analysis of plants transformed with the 35S: JUB1 construct. Radiolabeled JUB1 cDNA was used as hybridization probe. Numbers indicate individual transformants; C, nontransformed wild-type (Col-0) control. Elevated JUB1 expression compared with wild-type plants is observed in transgenic lines 1, 3, 5, 6, 7, 9, 11, and 12.

(B) Increased JUB1 expression in lines 5 and 9, as confirmed by qRT-PCR, compared with EV control.

(C) Delayed bolting in 35S:JUB1 overexpression line compared with the wild type (WT) at 43 d after sowing (DAS).

(D) Elevated chlorophyll content in leaves 1 to 12 of 35S:JUB1 overexpressors compared with EV control plants at \sim 60 DAS (n = 6).

(E) Percentage of survived leaves at different plant ages (given as DAS). Bolting time points are indicated by red arrows. Note steeper slope for curve of EV plants compared with 35S: JUB1 overexpressors between days 34 and 56 (*n* = 14 to 17).

(F) Ion leakage of leaves 1 to 12 of 35S: JUB1 and EV control plants at \sim 60 DAS.

Data in (B) and (D) to (F) are the means of at least three biological replicates \pm SD.

curves confirmed later senescence in overexpressors than in EV lines (Figure 1E). In accordance with this, ion leakage as an indicator of senescence was much less pronounced in overexpression plants than in EV controls (Figure 1F). We also expressed *JUB1* from the *RD29A* promoter, which shows basal activity in nonstressed plants, but enhanced activity upon abiotic stress (Kasuga et al., 1999). We observed extended longevity, accompanied by delayed bolting, in *RD29A:JUB1* plants, which was dependent on *JUB1* expression level (see Supplemental Figure 3 online).

Next, we characterized a SALK T-DNA insertion line (Col-0 background). The T-DNA insertion in the second intron, 1034-bp downstream of the start codon (Figure 2A), was confirmed by PCR on genomic DNA of *jub1-1* plants (see Supplemental Figure

4 online). RT-PCR analysis revealed a reduction in *JUB1* transcript abundance in fully expanded *jub1-1* mutant leaves compared with leaves of equivalently aged wild-type plants (Figure 2B). Phenotypic analysis of the *jub1-1* mutant revealed earlier bolting than in the wild type (up to 3 d; Figure 2C). A >95% reduction of *JUB1* transcript level was detected by qRT-PCR in *jub1-1* plants (Figure 2D), which was accompanied by an earlier loss of chlorophyll (Figure 2E) and precocious senescence (Figures 2F).

As our data indicated a longevity-extending effect of elevated *JUB1* expression, we also tested its role in dark-induced senescence. Leaves were detached from 39-d-old soil-grown plants and kept for up to 6 d on moist filter paper in the dark; we observed that 35S:*JUB1* overexpressors retained chlorophyll

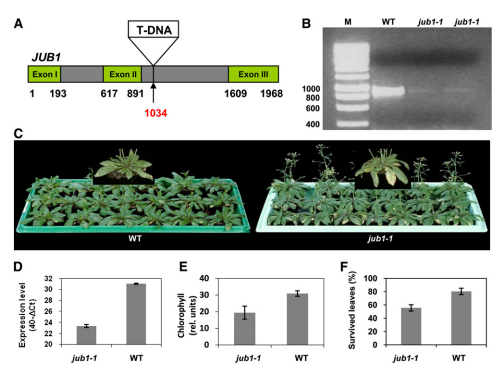


Figure 2. Physiological and Molecular Characterization of the jub1-1 Mutant.

(A) T-DNA inserted at nucleotide position 1034 downstream of the start codon (indicated by arrow) in the second intron.

(B) Downregulation of *JUB1* transcripts in *jub1-1* mutant shown by RT-PCR with primers annealing to the start and stop regions of the coding sequence. M, molecular size marker (sizes in base pairs); WT, wild type.

(C) Comparison of jub1-1 plants with wild-type plants at 47 DAS. Note early bolting and early senescence in the mutant plants.

(D) Downregulation of *JUB1* transcript abundance in *jub1-1* line, as confirmed by qRT-PCR.

(E) The jub1-1 mutant contains less chlorophyll than the wild type in the five biggest leaves at 47 DAS.

(F) The *jub1-1* mutant exhibits a lower percentage of survived leaves than the wild type at 47 DAS. Data in graphs are the means of at least three biological replicates \pm sD.

better than the *jub1-1* knockdown and EV control lines, reminiscent of a delay in senescence (see Supplemental Figure 5 online).

Expression Profiling of SAGs in JUB1 Transgenic Plants

To further substantiate the role of JUB1 for the regulation of leaf senescence, we analyzed the expression of 168 SAGs (including 49 TFs) in wild-type and JUB1-modified plants by gRT-PCR. The SAGs included in our expression profiling platform were previously shown to be highly upregulated during natural senescence in wildtype plants (see Supplemental Data Set 1 online; Buchanan-Wollaston et al., 2005; van der Graaff et al., 2006; Balazadeh et al., 2008b; Parlitz et al., 2011). The platform included SAG12, a wellknown senescence marker gene (Noh and Amasino, 1999). At a twofold cutoff, the expression of 89 SAGs (including 21 TFs and SAG12), representing \sim 53% of all SAGs tested, was downregulated in 35S:JUB1 plants compared with the wild type, while only six SAGs (including JUB1) were upregulated (see Supplemental Data Set 1 online; Figure 3A). By contrast, expression of 97 SAGs (including 19 TFs), representing \sim 58% of all SAGs tested here, were upregulated in jub1-1 compared with the wild type, and only two SAGs (including JUB1) were downregulated in jub1-1 plants (Figures 3B and 3C; see Supplemental Data Set 1 online). Collectively, our data support the model that JUB1 constitutes a negative regulator of leaf senescence and, hence, a driver of longevity.

Estradiol-Inducible JUB1 Overexpression

We next tested the effect of estradiol (EST)-inducible JUB1 expression on bolting and leaf senescence using the system described by Zuo et al. (2000). qRT-PCR revealed induction of JUB1 expression in EST-treated JUB1-IOE seedlings already after 1 h of EST treatment and expression further increased at longer incubation times (Figure 4A). We sowed JUB1-IOE lines on half-strength Murashige and Skoog (MS) medium containing 1% Suc and 10 μ M EST (or 0.1% ethanol in control experiments). After stratification at 4°C for 3 d, plates were transferred to a growth chamber at long-day conditions. Bolting was delayed in EST-treated JUB1-IOE plants compared with mock-treated controls (Figure 4B). When grown in liquid medium, JUB1-IOE seedlings remained green (nonsenescent) longer in the presence of EST (15 µM) than in mock-treated seedlings (Figure 4C). Thus, high JUB1 expression suppressed bolting and delayed senescence under in vitro conditions, similar to plants grown in soil.

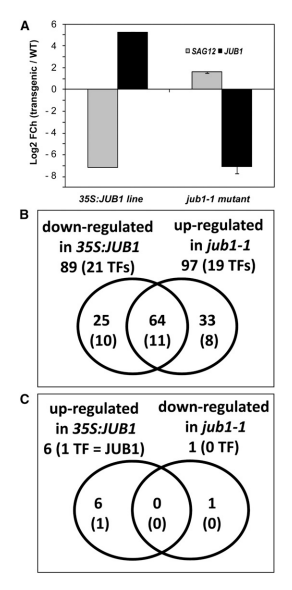


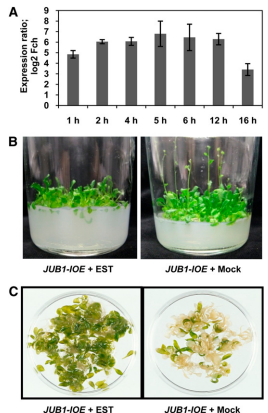
Figure 3. Transcript Profiling of SAGs.

(A) Expression of JUB1 and the late-senescence marker gene SAG12 in 35S:JUB1 and jub1-1 plants compared with the wild type (WT) (numbers on the y axis indicate log2 fold-change (FCh) expression ratio compared with the wild type).

(B) and (C) Venn diagrams of SAGs differentially expressed in 35S:JUB1 and jub1-1 plants compared with the wild type at 47 DAS. Numbers in parentheses indicate senescence-associated TFs. See also Supplemental Data Set 1 online.

Age-Dependent JUB1 Expression

We next analyzed the JUB1 expression pattern in Arabidopsis (Col-0) and tobacco (Nicotiana tabacum cv Samsum NN) plants transformed with a JUB1 promoter-β-glucuronidase (GUS) reporter construct (ProJUB1:GUS). JUB1 expression was observed in various tissues throughout plant development. In Arabidopsis seedlings, GUS activity was preferentially detected in roots, cotyledons, and the tips of young leaves. GUS staining was generally more pronounced in leaf tips and margins than the central part of the leaf blade (Figures 5A and 5B). Expression was also observed in floral tissues, preferentially in old sepals, petals, stamens, mature anthers, and pollen grains, while immature floral tissue did not show GUS activity (Figures 5C and 5D). GUS activity was also observed in the abscission zone of open flowers (data not shown). In the half expanded leaves of soil-grown plants, GUS staining was observed in the tip region only; in fully expanded leaves, strong GUS staining was observed in the senescent regions (Figures 5E and 5F). Moreover, GUS activity was detected in primary and lateral roots (Figures 5A, 5G, and



JUB1-IOE + EST

Figure 4. Phenotypic Analysis of JUB1-IOE Lines.

(A) JUB1 expression is induced in leaves of JUB1-IOE seedlings after treatment with 10 μ M EST compared with mock treatment (0.1% ethanol). Treatment times are indicated. Data are the means of three biological replicates \pm sp. Fch, fold change.

(B) Induction of JUB1 expression by EST in JUB1-IOE plants delays bolting when grown in vitro. Plants were grown for 6 weeks in glass jars on medium containing 10 µM EST (0.1% ethanol for control experiment). In this experiment, five independent transgenic lines were tested; the photographs shown represent a typical result.

(C) Delayed senescence in JUB1-IOE plants grown in vitro. Two-weekold JUB1-IOE seedlings were transferred to flasks with liquid medium containing 15 μ M EST, or 0.15% ethanol as control, and kept on a rotary shaker (slow motion) under continuous light for 1 week. Note the delayed senescence upon JUB1 induction. Three independent transgenic lines were used to confirm the observation made here.

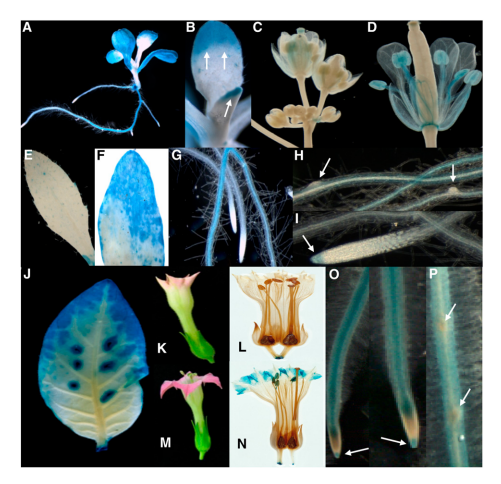


Figure 5. GUS Activity in ProJUB1:GUS Lines.

(A) to (I) Arabidopsis.

(A) Ten-day-old seedling. GUS staining is mainly localized to cotyledons, the tip regions and margins of leaves representing the oldest but not yet senescent leaf regions, and primary and secondary roots.

(B) Ten-day-old seedling. Strong GUS staining located in tips of newly emerging leaves (arrows).

(C) and (D) Flowers at different development stages. GUS staining is virtually absent in unopened young flowers. In open flowers (D), GUS staining appears in mature anthers, filaments, and the stamen abscission zone.

(E) GUS staining is weak to absent in \sim 50% expanded leaves from soil-grown plants.

- (F) Strong GUS staining is located in senescent regions of a partially senescent leaf from soil-grown plants.
- (G) GUS staining in roots; note the absence of GUS activity in the meristematic zone.
- (H) GUS staining is absent from emerging lateral roots (arrows).
- (I) GUS staining in root cap (arrow). Staining was for \sim 1 h.

(J) to (P) Tobacco.

- (J) Leaf, with more intense staining in the tip and margins. GUS activity is also visible around wound sites.
- (K) and (L) Young flower without GUS staining.
- (M) and (N) Open flower showing GUS staining in petal tips and anthers.
- (O) Roots. Note the absence of GUS activity in the meristematic zone, whereas the root cap shows GUS staining (arrows).
- (P) Absence of GUS activity from emerging lateral roots (arrows). Staining was for ${\sim}6$ h.

5H), including the root cap of extended roots (Figure 5I). However, *JUB1* promoter activity was absent from cells of emerging lateral roots (Figure 5H) and from root meristematic zones that faded into the elongation zones (Figures 5G and 5I).

In transgenic *Pro_{JUB1}:GUS* tobacco leaves, GUS staining was only observed in older parts (i.e., the tips and margins), consistent with an age-dependent upregulation of *JUB1* expression (Figure 5J). No GUS staining was observed in young flowers

when corolla limbs began to open (Figures 5K and 5L); however, in open flowers, intense GUS staining was evident in the tip regions of corolla limbs, but not corolla tubes. Additionally, *JUB1* promoter activity was visible in anthers (Figures 5M and 5N). Moreover, GUS staining was observed in roots (Figures 5O and 5P), including the root tip of extended, but not young, emerging roots. As in *Arabidopsis*, no *JUB1* promoter activity was detected in the root meristematic zone (Figure 5O).

H₂O₂ Triggers JUB1 Expression

H₂O₂ plays a central role in plant signaling and stress responses. In microarray hybridization experiments, we previously observed strong induction (~25-fold) of JUB1 transcript level in Arabidopsis seedlings after 5 h of exposure to 10 mM H₂O₂ (Balazadeh et al., 2010b). Here, we confirmed H₂O₂-responsive JUB1 expression by qRT-PCR in 2-week-old Arabidopsis plants treated for 30 min or 2, 4, or 6 h with 10 mM external H₂O₂. JUB1 transcript abundance increased in both whole seedlings and leaves already 30 min after treatment and increased further thereafter (Figures 6A and 6B). Similarly, JUB1 expression increased approximately fivefold within 5 h of pharmacological inhibition of the H₂O₂scavenging enzyme catalase in seedlings (see Supplemental Figure 6A online). In ProjUB1:GUS lines, enhanced GUS activity was observed 30 min or 1 h after treatment with 10 mM or 50 mM H₂O₂ (Figures 6C and 6D), indicating that the response of JUB1 to H₂O₂ is regulated at the promoter level and mediated by one or more currently unknown upstream transcription regulators.

It was previously reported that endogenous H_2O_2 concentration rises during bolting in *Arabidopsis* leaves, and downregulation of catalase (CAT2) activity was suggested to be the initial step of this rise (Zimmermann et al., 2006). To investigate a possible correlation between the level of endogenous H_2O_2 and *JUB1* expression, we measured both H_2O_2 content and *JUB1* expression in leaves of ~35-d-old *Arabidopsis* wild-type plants at bolting (1-cm main flower stalk). Leaves number 2 (second oldest rosette leaf) to 14 were sampled individually; younger leaves were collected in groups A (>25 mm leaf length), B (15 to 25 mm), and C (below 15 mm), respectively (leaf number 1 was too old and deteriorated for measurements). *JUB1* expression was determined by qRT-PCR, and H_2O_2 level was quantified using an Amplex Red assay. We observed that endogenous H_2O_2 and *JUB1* transcript abundance followed similar patterns with higher levels in older leaves (numbers 2 to 12) than in younger leaves (numbers 13 and 14, groups A to C); particularly low H_2O_2 and *JUB1* transcript levels were present in the youngest rosette leaves (see Supplemental Figure 7 online). Thus, our data indicate that *JUB1* expression follows the level of endogenous H_2O_2 dependent on leaf age.

Various abiotic stresses, including salinity, cold, and heat stress, cause the accumulation of endogenous H_2O_2 . In a microarray hybridization experiment, we observed induction of *JUB1* expression when hydroponically grown plants were subjected to salinity stress (150 mM NaCl) over a period of 4 d (see Supplemental Figure 6B online). Salt stress (150 and 200 mM NaCl) slightly induced GUS activity in *Pro_{JUB1}:GUS* plants after 24 h (4-methyl umbelliferyl β -D-glucuronide [MUG] assay; see Supplemental Figure 8A online). Several other treatments that cause an intracellular rise of H_2O_2 level induce *JUB1* expression, including treatment with cellulase R-10 (see Supplemental Figures 8B and 8C online), paraquat (methyl viologen) (see Supplemental Figures 8B and 8C online), ozone (Gadjev et al., 2006), 3-aminotriazole, which blocks the activity of the H_2O_2 scavenging enzyme catalase (Gechev and Hille, 2005), and AAL (*Alternaria alternata* fungal

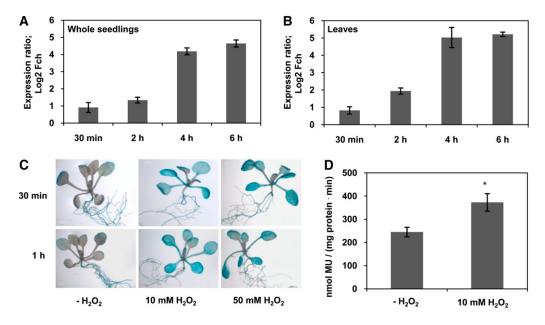


Figure 6. Effect of H₂O₂ on JUB1 Expression.

(A) and (B) JUB1 transcript level in whole seedlings (A) and leaves (B) of wild-type Arabidopsis plants as determined by qRT-PCR after treatment with 10 mM H_2O_2 for 30 min or 2, 4, and 6 h compared with nontreated samples.

(C) *Pro_{JUB1}:GUS* lines treated with H₂O₂. Two-week-old seedlings were transferred to medium containing 10 or 50 mM H₂O₂ and incubated for 30 min or 1 h, respectively. Elevated GUS activity was observed at both concentrations already 30 min after treatment.

(D) GUS activity of Pro_{JUB1} :GUS seedlings measured by a MUG assay after treatment with 10 mM H₂O₂ for 30 min. Asterisk indicates significant difference (P < 0.05, Student's *t* test). Data are the means of three biological replicates ± sp. MU, methylumbelliferone.

toxin that induces H_2O_2 accumulation and cell death through perturbation of sphingolipid metabolism; Gechev and Hille, 2005). Collectively, these data indicate that *JUB1* expression is triggered by an intracellular rise of H_2O_2 level.

Overexpression of JUB1 Enhances Tolerance to Salt Stress

Salt stress triggers the accumulation of intracellular H_2O_2 (e.g., Chung et al., 2008). To investigate whether overexpression of *JUB1* enhances tolerance to salt stress, the effect of 150 mM NaCI on *JUB1-IOE* and *jub1-1* knockdown mutants was studied. To this end, 2-week-old *jub1-1* mutant and *JUB1-IOE* seedlings were transferred from solid MS medium to liquid MS medium containing 150 mM NaCI. In the case of *JUB1-IOE* lines, 15 μ M EST was added to induce *JUB1* expression, and, as a control, 0.15% ethanol was used. All seedlings were incubated on a shaker in a growth chamber with continuous light; after 3 d of stress, the *JUB1-IOE* plants incubated in saline medium containing EST retained fivefold higher levels of chlorophyll than the plants incubated in saline medium in the absence of EST (Figures 7A and 7B). By contrast, chlorophyll content in *jub1-1* seedlings was only half that of wild-type plants when stressed by salt (Figures 7C and 7D).

JUB1 Enhances Tolerance to H_2O_2 by Regulating Its Cellular Concentration

Similar to findings in animals, extended life span in plants has been observed to be closely related to increased tolerance to oxidative stress. In particular, several *Arabidopsis* mutants with extended longevity have been shown to exhibit superior tolerance to oxidative stress. Examples include transgenic plants overexpressing the *CBF2* and *CBF3* TFs, the *ore1*, *ore3*, and *ore9* mutants, and the very late flowering, long-living mutant *gigantea* (Kurepa et al., 1998; Woo et al., 2004; Sharabi-Schwager et al., 2010). The *gigantea* mutant exhibits enhanced tolerance to various types of oxidative stress, such as paraquat- and H_2O_2 -induced oxidative stress (Kurepa et al., 1998).

To determine whether there is a link between increased JUB1 expression and enhanced tolerance to oxidative stress, we first analyzed the effect of H_2O_2 treatment on JUB1 transgenic plants.

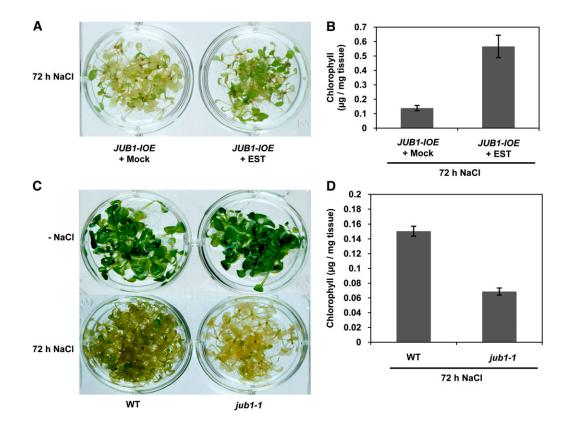


Figure 7. EST-Treated JUB1-IOE Plants Exhibit Increased Tolerance to NaCl Stress, Whereas Tolerance to NaCl Is Reduced in the jub1-1 Mutant.

(A) Two-week-old *JUB1-IOE* seedlings were transferred from solid MS medium to liquid MS medium containing 150 mM NaCl, in the absence (mock; 0.15% ethanol) or presence of 15 µM EST added to induce *JUB1* expression. Plants were incubated for 72 h. EST-treated *JUB1-IOE* seedlings are more tolerant to salt stress.

(B) Higher chlorophyll levels are retained in EST-treated JUB1-IOE seedlings after salt treatment.

(C) Seedlings of wild-type (WT) plants are less affected by 72 h salt treatment (150 mM NaCl) than those of the jub1-1 mutant.

(D) Chlorophyll content remains higher in the wild type (WT) than *jub1-1* plants after stress treatment. Data in (B) and (D) are the means of three biological replicates \pm SD.

Two-week-old *RD29A*:*JUB1* seedlings were treated with 10 mM H₂O₂. As shown in Figure 8A, seedlings subjected to treatment for 24 h retained chlorophyll more efficiently than EV-transformed control plants. Additionally, we cultured *35S*:*JUB1* and EV seedlings on sterile medium containing 0 or 10 mM H₂O₂. *35S*:*JUB1* plants remained healthier and showed less leaf bleaching in the presence of H₂O₂ than the control line (data not shown). We next tested H₂O₂ tolerance of *JUB1-IOE* lines by transferring 3-weekold seedlings from half-strength solid MS medium to fresh medium containing 15 μ M EST and 10 mM H₂O₂; control seedlings were similarly treated with H₂O₂, but EST was omitted (mock treatment: 0.15% ethanol). After 6 d, the *JUB1-IOE* plants treated with H₂O₂

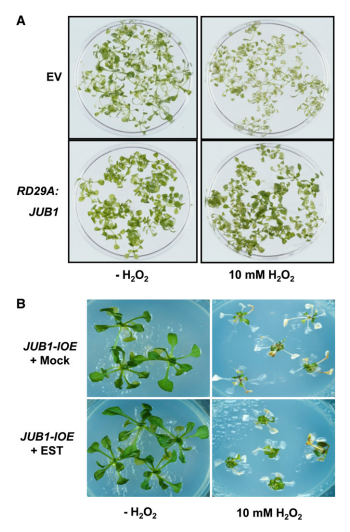


Figure 8. Overexpression of JUB1 Confers Tolerance to H₂O₂.

(A) Two-week-old *RD29A:JUB1* and EV seedlings grown on solid MS medium were transferred to liquid medium containing 10 mM H_2O_2 and incubated for 24 h. *RD29A:JUB1* seedlings remained green, whereas EV lines bleached in the presence of H_2O_2 .

(B) JUB1-IOE lines treated with 15 μ M EST survived better than plants treated with 0.15% ethanol (mock treatment) after transferring 3-weekold plants to fresh medium containing 10 mM H₂O₂ for 6 d. and EST were more vital than plants treated with H₂O₂ alone (Figure 8B). We next tested the effect of reduced JUB1 expression on H₂O₂ tolerance. Leaves detached from 35-d-old soil-grown jub1-1 and EV plants were incubated in 10 mM H₂O₂ for 5 d; EV leaves remained greener than those of the jub1-1 mutant (see Supplemental Figures 9A and 9B online). Moreover, when 12-dold seedlings were incubated in MS medium with 10 mM H₂O₂ for 6 h tissue damage was more prominent in *jub1-1* than wild-type seedlings. Accordingly, diaminobenzidine (DAB) staining revealed a higher H₂O₂ accumulation in jub1-1 than wild-type plants (Figure 9A). We confirmed a slight (\sim 14%) but significant increase in H₂O₂ concentration in 12-d-old jub1-1 seedlings compared with the wild type using an Amplex Red assay (Figure 9B). Next, we tested the effect of elevated JUB1 expression on cellular H₂O₂ level. To this end, we treated JUB1-IOE lines in the absence (control) or presence of 15 µM EST (to induce JUB1 expression) with 10 mM H₂O₂ for 6 h and observed that ESTtreated seedlings accumulated slightly less (\sim 10%) H₂O₂ than the controls (Figures 9C and 9D). Jointly these data indicate that JUB1 counteracts the cellular accumulation of H₂O₂.

A Heat Shock Protein Response Network Regulated by JUB1

To start unraveling the H_2O_2 regulatory network regulated by JUB1, we tested the expression of 187 reactive oxygen species (ROS)–responsive genes in 2-week-old seedlings of *RD29A*: *JUB1* and *jub1-1* plants before and after treatment with H_2O_2 (10 mM, 6 h) and compared results with data from wild-type seedlings. Expression analysis was performed by qRT-PCR. Genes included in the ROS expression platform are known to be induced by H_2O_2 , superoxide anion, and/or singlet oxygen (for details, see Methods) and encode TFs, heat shock proteins (HSPs), protein kinases, glutathione S-transferases (GSTs), and others (see Supplemental Data Set 2 online).

Our data revealed detectable expression of 179 ROS genes in all samples at both control and treatment conditions. We observed that expression of 36 genes was more strongly induced by H₂O₂ in *RD29A*:*JUB1* than wild-type seedlings, while expression of these genes was either not or only marginally affected by H₂O₂ treatment in jub1-1 seedlings compared with the wild type (Table 1). These genes therefore represent prime candidates for ROSresponsive genes regulated by JUB1 during H₂O₂ signaling. Genes encoding HSPs constitute the largest fraction of the genes in the regulatory network. Upon H₂O₂ treatment, 14 HSP genes were more strongly induced in RD29A:JUB1 seedlings compared with the wild type, whereas their expression was not or only marginally induced in jub1-1 plants (Table 1). In addition to HSPs, several GST genes (including GST10, 24, and 25), TFs (e.g., DREB2A, DREB2B, ANAC047, and ANAC055), and some heat shock binding proteins were also among the ROS genes induced by H₂O₂ treatment in RD29A:JUB1 transgenic plants. In accordance with the reduced induction of HSPs after stress in jub1-1 seedlings, we observed a reduced heat stress tolerance (45°C, 5 h) of jub1-1 seedlings relative to the wild type (see Supplemental Figure 10 online), whereas overexpression of JUB1 enhanced tolerance to heat stress (see Supplemental Figure 11 online).

Collectively, our data indicate that JUB1 is a key regulator of plant stress responses. Under stress, JUB1 regulates expression

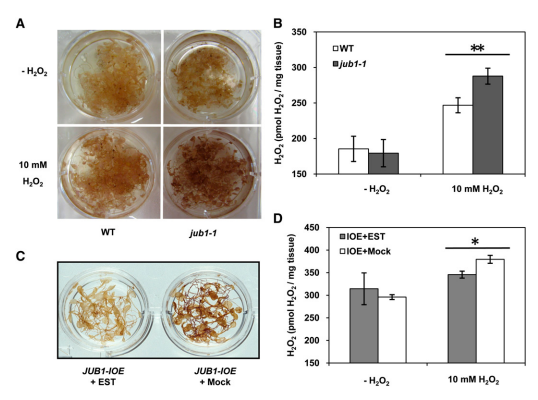


Figure 9. Overexpression of JUB1 Reduces Endogenous H₂O₂ Content, While the Opposite Is Observed in the jub1-1 Mutant.

(A) DAB staining of wild-type (WT) and *jub1-1* seedlings treated with 10 mM H_2O_2 for 6 h. Note the stronger DAB staining in *jub1-1* seedlings in the presence of H_2O_2 .

(B) Amplex Red assay. Note the higher H_2O_2 level in H_2O_2 -treated *jub1-1* seedlings compared with the wild type. Asterisks indicate significant difference (P < 0.001).

(C) DAB staining. JUB1-IOE seedlings treated with EST for 6 h accumulate less H₂O₂ than mock-treated seedlings.

(D) Amplex Red assay. Note the reduced H_2O_2 level in EST-treated *JUB1-IOE* seedlings compared with mock-treated plants (asterisk indicates significant difference; P < 0.02). Data in (B) and (D) are the means of three independent biological replicates \pm sp.

of a set of stress tolerance genes in transgenic *Arabidopsis*, thereby leading to improved stress tolerance.

JUB1 Binding Site

To identify cis-elements recognized by the JUB1 TF, we performed in vitro binding site selection using the CELD-TF fusion method (Xue, 2002, 2005). Thirty-six unique double-stranded oligonucleotides bound by JUB1-CELD fusion protein were recovered after five rounds of selection using biotin-labeled doublestranded oligonucleotides containing a 30-nucleotide random sequence and were analyzed for binding activity (Table 2). Alignment of the target sequences identified RRYGCCGT as the JUB1 consensus core binding sequence. Notably, all JUB1-selected motifs, except for pf17d127, had a fixed distance to the primer sequence flanking the 30-nucleotide random sequence, indicative of primer sequences establishing part of the JUB1 recognition site. To identify nucleotide positions required for efficient JUB1 binding, and to discover potential secondary motifs contributing to JUB1 binding, nucleotide substitution experiments were performed using the sequences of selected oligonucleotides, F15d64 and F17d127, as a starting point. This analysis demonstrated that the JUB1 binding sequence consists of two elements (Table 3). We observed at least two types of sequences that differ in their 3' parts: TGCCGT(7N)ACG and TGCCGT(7N)CCGC (N, any nucleotide). However, the second element is not essential for JUB1 binding but increases binding affinity.

DREB2A Is a Direct Target of JUB1

One of the genes induced by H_2O_2 in *RD29A*:*JUB1* seedlings that attracted our attention was *DREB2A*, a member of the AP2/ EREBP TF family whose function with respect to plant responses to various abiotic stresses, such as cold and drought, is well established (Sakuma et al., 2006a, 2006b). *DREB2A* was approximately fivefold more induced by H_2O_2 in the *RD29A*:*JUB1* overexpressor than wild-type seedlings (Table 1). However, enhanced expression of *DREB2A* in *JUB1* overexpressors was also observed in the absence of H_2O_2 treatment. We found that *DREB2A* expression was elevated by more than threefold in 35-d-old *35S*:*JUB1* plants compared with wild-type plants (Figure 10A). By contrast, reduced *DREB2A* expression (~1.7fold decrease) was observed in the *jub1-1* mutant when compared with the wild type. We also tested *DREB2A* expression in

Table 1. Genes Affected by H₂O₂ Treatment

		H ₂ O ₂ Treatment versus Control		
		RD29A:JUB1/Wild Type	jub1-1/Wild Type	
	Description	log2 FCh		
AGI		A	В	A-B
AT1G74310	HOT1_HSP101; ATP binding/ATPase/	4.69	0.99	3.70
	nucleoside-triphosphatase/nucleotide binding			
AT2G29500	17.6-kD Class I small HSP (HSP17.6B-CI)	3.57	0.05	3.52
AT4G25200	ATHSP23.6-mitochondrial small HSP23.6	3.62	0.45	3.17
AT5G12030	HSP17.6; unfolded protein binding	3.98	0.97	3.01
AT1G74590	GST TAU10 (ATGSTU10_GSTU10)	2.69	-0.22	2.90
AT1G53540	17.6-kD Class I small HSP (HSP17.6C-CI) (AA 1-156)	3.16	0.26	2.89
T1G71000	HSP binding	3.87	1.09	2.78
T2G28210	ALPHA CARBONIC ANHYDRASE2 (ACA2)	2.82	0.29	2.53
T3G09350	Armadillo/β-catenin repeat family protein	3.36	0.83	2.53
T5G51440	23.5-kD mitochondrial small HSP (HSP23.5-M)	3.54	1.03	2.51
T5G12020	17.6 KDA CLASS II HSP (HSP17.6II)	2.94	0.42	2.51
T3G12580	HSP70; ATP binding	3.43	0.92	2.51
T5G52640	ATHS83_HSP81-1_HSP81.1_HSP83ATHSP90.1; ATP	3.55	1.05	2.50
	binding/unfolded protein binding			
T3G24500	ATMBF1CMBF1C; DNA binding/transcription coactivator/TF	3.48	1.04	2.44
T5G64510	Unknown protein	3.19	0.77	2.42
T3G46230	ATHSP17.4	2.54	0.13	2.41
T1G17170	GST TAU24 (GST ATGSTU24)	2.34	-0.05	2.39
T5G48570	Peptidyl-prolyl <i>cis-trans</i> isomerase, putative/FK506 binding protein	3.26	0.91	2.35
T1G54050	17.4-kD Class III HSP (HSP17.4-CIII)	3.27	0.92	2.35
T4G12400	Stress-inducible protein, putative	3.09	0.83	2.26
T1G07160	Protein phosphatase 2C, putative/PP2C, putative	1.50	-0.70	2.20
T2G46240	ATBAG6_BAG6; calmodulin binding/protein binding	2.61	0.43	2.18
T2G38340	AP2 domain-containing TF, putative (DRE2B)	1.83	-0.34	2.10
T2G26150	HSFA2_ATHSFA2; DNA binding/TF	3.73	1.59	2.17
T5G05410	DREB2A; DNA binding/transcription activator/TF	2.28	0.15	2.14
T2G32120	HSP70T-2; ATP binding	2.47	0.34	2.13
T3G04070	ANAC047: TF	1.96	-0.14	2.13
T2G20560	DNAJ heat shock family protein	3.23	1.19	2.10
T4G37370	CYP81D8; electron carrier/heme binding/iron ion	1.93	-0.02	2.04 1.94
1400/0/0	binding/monooxygenase/oxygen binding	1.55	-0.02	1.54
T4G34410	REDOX RESPONSIVE TRANSCRIPTION	1.41	-0.53	1.94
T1017100	FACTOR1 (RRTF1)	0.71	0.01	1 00
T1G17180	GST TAU25 (ATGSTU25)	2.71	0.91	1.80
T1G16030	Hsp70b; ATP binding	2.27	0.48	1.79
T3G15500	ATNAC3_ANAC055; TF	1.51	-0.12	1.63
T3G28210	PMZ; zinc ion binding	1.66	0.06	1.60
AT4G37990	ATCAD8_CAD-B2ELICITOR-ACTIVATED GENE 3-2 (ELI3-2); aryl-alcohol dehydrogenase/mannitol	0.30	-1.29	1.58
T1014000	dehydrogenase Zing finger (C211C4 type DINC finger) femily protein	1 77	0.01	1 50
AT1G14200	Zinc finger (C3HC4-type RING finger) family protein	1.77	0.21	1.56

Genes highly induced by H_2O_2 treatment (6 h) in the *RD29A:JUB1* line but either not or marginally affected in the *jub1-1* mutant when compared to wild-type plants. A threefold change (log2 = 1.5) was selected as threshold. HSPs are indicated in bold. AGI, Arabidopsis Genome Initiative; FCh, fold change.

2-week-old JUB1-IOE seedlings shortly (~3 h) after EST treatment. DREB2A expression was almost fourfold induced in the EST-treated lines, suggesting DREB2A as a JUB1 target gene (Figure 10A). Notably, the DREB2A promoter harbors a perfect match to the full JUB1 binding sequence (TGCCGTNNNNNNACG) ~1 kb upstream of its translation start site.

To provide further evidence for regulation of *DREB2A* by JUB1, we performed luciferase-based transactivation assays in *Arabidopsis* mesophyll cell protoplasts using the \sim 1.8-kb *DREB2A* promoter (including the JUB1 binding site) fused to the firefly luciferase reporter. JUB1 significantly transactivated the 1.8-kb *DREB2A* promoter (Figure 10B).

Table 2. JUB1-Selected Binding Sequences

JUB1-Selected Oligonucleotide (5'/3')		
Bio-RS-Oligo 1		
Pf15d21	TCCCAATAGGATTCGTAAAGTGCCGTGTTCcgtccgccagcgcacc	0.82
Pf15d30	AGCGAAGGATCAATTGAAGACGCCGTGATCcgtccgccagcgcacc	0.74
Pf15d25	AATTTGACGTCATATTCT AACGCCGT AGTCcgtccgccagcgcacc	0.83
Pf15d64	CAACATGAAGCTAGATGCCGTAGACcgtccgccagcgcacc	1.18
Bio-RS-Oligo 2		
Pf16d41	TCACCCCCCTTCTGAGGAACTCGGTGCCGTTCCTTTCcgtccacctgcag	0.85
Pf16d43	CCTGTGACTTTCTCGAATCATG AGTGCCGT GCTCTCTCcgtccacctgcag	0.48
Pf16d44	AGCGTATTCCCACTCCCGCTATGAGTGCCGTGCCCCCCcgtccacctgcag	0.87
Pf16d48	TGCGAACCTTGTAGTGCTCCAGGATGCCGTACACCCCcgtccacctgcag	0.84
Pf16d82	ATACTTTCCCCGAGTGTGATCGGGTGCCGTGCCCCcgtccacctgcag	0.96
Pf16d84	CCATTTCGCCTTGCTGATTGCGCGGTGCCGTGTATCTCcgtccacctgcag	0.96
Pf16d88	AAGCATTATCGTTGTTAAATACGGTGCCGTGTTCTGGCcgtccacctgcag	0.66
Pf16d98	GGGCGGGCTGGTCTCGTATTGAGATGCCGTACTTGCCcgtccacctgcag	0.92
Pf16d100	CCGATATCCTGTGAACTCAGCAAGATGCCGTCGTCCCCcgtccacctgcag	0.89
Pf16d102	TTGGTGGTGCACGTATTTGATAG GGTGCCGT GTGTTCCcgtccacctgcag	0.90
Pf16d104a	ATGTTCGGCTGGATCTATATCACGATGCCGTGCGTTGCcgtccacctgcag	1.20
Pf16d106	CCAATTCCCTTTTGCTGTTTAGTAGTGCCGTGCCCgtccacctgcag	1.01
Bio-RS-Oligo 3		
Pf17d51	GGGACTTGTATACCTGTAAGGTGCCGTACCtcatgcggtacccacgtc	0.87
Pf17d52b	GGGAGGCCTCGTGCCAACCAGTGCCGTACGtcatgcggtacccacgtc	1.07
Pf17d54	ACGTGATACACGCTCTATCAGTGCCGTGCCtcatgcggtacccacgtc	0.85
Pf17d57	AGGCCGTTAAACATACATGAGTGCCGTACGtcatgcggtacccacgtc	1.06
Pf17d58	ACACAATTGTGACGCGAAAGGTGCCGTACAtcatgcggtacccacgtc	1.02
Pf17d112	CATCGGTTTCGGCCTTGTAGGTGCCGTACCtcatgcggtacccacgtc	0.94
Pf17d114	TCCGTCCTCCGAGGATCATGATGCCGTACGtcatgcggtacccacgtc	0.77
Pf17d115	TCAGGTACAACTCTGATGCAGTGCCGTACCtcatgcggtacccacgtc	0.91
Pf17d116	CGAGCGTGGCCCAAAACACGGTGCCGTACCtcatgcggtacccacgtc	0.94
Pf17d117	TCAGCTTGGCTGGAGCTAGGATGCCGTGGGtcatgcggtacccacgtc	0.72
Pf17d118	GATCCCCCTCCTTGCCTCTAGTGCCGTACCtcatgcggtacccacgtc	0.86
Pf17d120	TTCCCAGAACCTCTAACTGGATGCCGTACCtcatgcggtacccacgtc	0.86
Pf17d121	GTACTAGATGCCGTACGtcatgcggtacccacgtc	0.84
Pf17d124	AATGTCACTGTCCCCCTACAGTGCCGTGGCtcatgcggtacccacgtc	0.91
Pf17d126	GGCTTAACCCGACAGCACAGACGCCGTGCCtcatgcggtacccacgtc	0.94
Pf17d127	TGCCCAATGCCGTGTGTAGCACGCTGCCCA	1.00
Pf17d128	AAATCCTTGTAAATCCCTAGATGCCGTACTtcatgcggtacccacgtc	0.86
Pf17d129	GGTCGCACATCTCATCATGGATGCCGTACCtcatgcggtacccacgtc	0.81
Pf17d130	ACGCACGTGTCTAGTATTGAGTGCCGTGCAtcatgcggtacccacgtc	0.99

Thirty-six JUB1-selected oligonucleotides were obtained after five rounds of in vitro DNA binding site selection. Relative binding activity (RBA) of JUB1 to oligonucleotide Pf17d127 is set to 1. Values are based on a single assay. Nucleotides in lowercase letters are from flanking primer sequences. The JUB1 core binding sequence is in bold.

We next performed an electrophoretic mobility shift assay (EMSA) to test the physical interaction of JUB1 with the *DREB2A* promoter. As shown in Figure 10C, JUB1 interacts with a 40-bp *DREB2A* promoter fragment harboring the JUB1 binding site. Finally, we used chromatin immunoprecipitation–quantitative PCR (ChIP-qPCR) to demonstrate that JUB1 binds also in vivo to the *DREB2A* promoter (Figure 10D). Thus, our data demonstrate that JUB1 is an upstream transcriptional regulator of *DREB2A*.

Metabolite Profiling Reveals Accumulation of Trehalose and Pro in *JUB1* Overexpressors

Next, we were interested to know how overexpression of JUB1 affects metabolism of the corresponding plants. Therefore, the profile of primary metabolites in rosette leaves was compared between wild-type and *JUB1* overexpression lines (*35S:JUB1* and *RD29A:JUB1*). Metabolic profiling by gas chromatography cou-

pled to mass spectrometry (GC-MS) was performed on extracts from rosette leaves of 35-d-old plants (Figure 11A) and metabolic profiles were compared (Figure 11B). A total of 51 metabolites of known chemical structure were accurately quantified in every chromatogram. These compounds mostly included amino acids, carbohydrates (sugars and sugar alcohols), and organic acids. Overall, 67% (34 out of 51) and 55% (28 out of 51), respectively, of the identified metabolites revealed a significant difference (*P < 0.05, Student's t test) when 35S:JUB1 and RD29A:JUB1 metabolite profiles were compared with the wild type (for an overview, see Figure 11C and Supplemental Data Set 3 online). Metabolite profiles of the two types of overexpression plants were highly similar to each other: of the 19 metabolites upregulated in 35S: JUB1 plants, 15 were also elevated in RD29A:JUB1 plants. Similarly, of the 15 metabolites downregulated in 35S:JUB1 plants, six were also reduced in RD29A:JUB1 plants compared with the wild

Table 3. Base Substitution or Insertion Analysis of JUB1 Binding Motifs

Synthetic Oligonucleotide Probe	JUB1 RBA	
F17d127	TGCCCAA TGCCGT GTGTAGC ACG CTGCCCA	1.00 ± 0.03
F17d127m8	TGCCCTT TGCCGT GTGTAGC ACG CTGCCCA	0.82 ± 0.05
F17d127m7	TGCCCAAAACCGTGTGTAGCACGCTGCCCA	0.46 ± 0.02
F17d127m1	TGCCCAA TGAAGT GTGTAGC ACG CTGCCCA	0.23 ± 0.01
F17d127m3	TGCCCAA TGCCAA GTGTAGC ACG CTGCCCA	0.07 ± 0.01
F17d127m9	TGCCCAA TGCCGT TTGTAGC ACG CTGCCCA	0.96 ± 0.02
F17d127m13	TGCCCAA TGCCGT GAATAGC ACG CTGCCCA	1.00 ± 0.07
F17d127m14	TGCCCAA TGCCGT GTGAAGC ACG CTGCCCA	1.02 ± 0.04
F17d127m10	TGCCCAA TGCCGT GTGTTTC ACG CTGCCCA	1.10 ± 0.02
F17d127m17	TGCCCAA TGCCGT GTGTAGA ACG CTGCCCA	1.02 ± 0.07
F17d127m4	TGCCCAA TGCCGT GTGTAGT TCG CTGCCCA	0.49 ± 0.01
F17d127m2	TGCCCAA TGCCGT GTGTAGCAAACTGCCCA	0.56 ± 0.02
F17d127m5	TGCCCAA TGCCGT GTGTAGC ACGA AGCCCA	0.92 ± 0.05
F17d127m15	TGCCCAA TGCCGT GTGTAGC ACG CTAACCA	1.05 ± 0.06
F17d127m6f	TGCCCAA TGCCGT GTGTAGC ACG CTGTTCA	1.06 ± 0.04
F17d127m11	TGCCCAA TGCCGT GTGAAATAGC ACG CTGCCCA	0.08 ± 0.01
F17d127m12	TGCCCAA TGCCGT GTGTAC AAAA C AAA CCA	0.19 ± 0.02
F17d127m18ª	TGCCCAA TGCCGT GTG AAAACCGCCAGC CA	1.01 ± 0.02
F15d64	AGCTA GATGCCGTA GACCGT CCGCCAGC GC	0.84 ± 0.02
F15d64m1	AGCTA GATGCCGTA GACCGT AAGCCAGC GC	0.45 ± 0.05
F15d64m2	AGCTA GATGCCGTA GACCGTCCAACAGCGC	0.35 ± 0.02
F15d64m3	AGCTA GATGCCGTA GACCGTCC GCATGC GC	0.59 ± 0.01
F15d64m4	AGCTA GATGCCGTA GACCGTCC GCCAAA GC	0.73 ± 0.04

Nucleotides of motif 1 and motif 2 of the JUB1 binding site are shown in bold. Mutated nucleotides are underlined. Values are means ± SD of three replicated assays. RBA, relative binding activity.

^aNucleotides shown in italics in F17d127m18 are derived from the flanking primer sequence of Pf15d64 (see Table 2).

type (Figure 11B). In general, the content of the major organic acids increased in the overexpression lines compared with the wild type. Among these, the level of glyceric acid and various tricarboxylic acid cycle intermediates, fumaric acid, malic acid, citric acid, and succinic acid, significantly increased in the 35S:JUB1 line compared with wild-type plants. Most of these components were just slightly increased in RD29A:JUB1 plants. Shikimic acid content was significantly higher in both types of overexpression plants. Our data revealed that the most drastic changes were detectable in amino acids and carbohydrates. For example, among amino acids, a significant increase was observed for Pro and 4-hydroxy-Pro levels in both overexpression plants. Polyamines and compatible osmolytes, such as Pro, are known to be involved in the plant's responses to various environmental stresses, including osmotic and salt stress, heat stress, drought, cold, and pathogen infection (e.g., Yoshiba et al., 1997; Bhatnagar-Mathur et al., 2008; Verbruggen and Hermans, 2008; Gill and Tuteja, 2010). Among the detected disaccharides, the proportion of Suc significantly increased in both overexpression lines compared with the wild type. Moreover, a significant increase in the level of trehalose was observed in both overexpression lines. Trehalose is a nonreducing disaccharide of Glc that functions as an osmoprotectant (Müller et al., 1995) and stabilizes biological structures under abiotic stress conditions in bacteria, fungi, and invertebrates (Djilianov et al., 2005). Among the sugar alcohols, the levels of maltitol, glycerol (a compatible osmolyte), and erythritol were significantly higher in both overexpression lines, and the level of myo-inositol was significantly decreased in these lines compared with the wild type. In general, it appears that *JUB1* overexpressors accumulate higher levels of trehalose, Pro, and polyols (glycerol) than wild-type plants, which is in accordance with their enhanced tolerance to abiotic stress.

Secondary Metabolite Profile of JUB1 Transgenics

Secondary metabolites confer various advantages to the plants, such as regulation of development and response to biotic and abiotic stresses. To study secondary metabolite profiles of JUB1 transgenics, liquid chromatography-mass spectrometry (LC-MS) analysis was performed on extracts from rosette leaves of 35-d-old JUB1 overexpression and jub1-1 lines. Moreover, to identify JUB1regulated genes involved in secondary metabolism, we tested by gRT-PCR the expression of 94 genes encoding enzymes and TFs that regulate the biosynthetic pathway from shikimate to phenylpropanoids, including anthocyanins, flavonols, and sinapoyl derivatives (see Supplemental Data Set 4 online). Our metabolite analysis revealed the most drastic changes for cyanidin derivatives (Figure 12). Cyanidin derivative (A11 and A9; Tohge et al., 2005) levels were significantly decreased in 35S:JUB1 plants, whereas no significant change was observed for the level of anthocyanidins in the jub1-1 mutant. Accordingly, expression of genes encoding dihydroflavonol 4-reductase (DFR; At5g42800), a key enzyme in shunting flavonols into the anthocyanins, leucoanthocyanidin dioxygenase (LDOX; At4q22880), anthocyanin glycosyltransferases (A5GT, At4g14090; A3G2"XT, At5g54060), anthocyanin acyltransferases (A5GMaT, At3g29590; A3GCouT, At1g03940), and GST (TT19, At5g17220) were significantly downregulated in JUB1 overexpression plants. Similarly, expression of anthocyanin regulatory

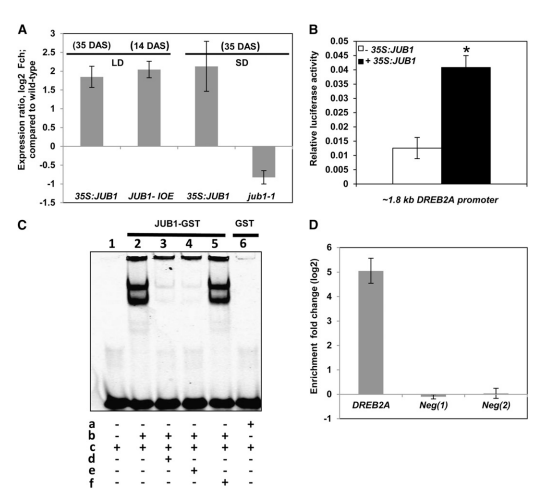


Figure 10. DREB2A Is a Direct Target of JUB1.

(A) Expression of *DREB2A* in 35S:JUB1, JUB1-IOE, and jub1-1 lines compared with the wild type. Plant ages are indicated in days after sowing (DAS). LD, long day; SD, short day. Numbers on the *y* axis indicate expression fold change (log2 basis) compared with the wild type. Data represent means \pm sD of five (LD) or three (SD) independent experiments.

(B) Transactivation of *DREB2A* expression (from its \sim 1.8-kb promoter) by JUB1 in *Arabidopsis* mesophyll cell protoplasts. The *Pro_{DREB2A}:FLuc* construct harboring the *DREB2A* promoter upstream of the firefly (*Photinus pyralis*) luciferase (FLuc) open reading frame was cotransformed with the 35S:*JUB1* plasmid (omitted in control experiments). The 35S:*RLuc* vector was used for transformation efficiency normalization. Bars indicate the sD of at least four biological replicates. The asterisk indicates significant difference to control at P < 0.05.

(C) EMSA. Purified JUB1-GST protein binds specifically to the JUB1 binding site within the *DREB2A* promoter. In vitro DNA binding reactions were performed with the 40-bp wild-type fragment of the *DREB2A* promoter containing the JUB1 motif (5'-GATGCCGTTAGAGACACG-3'). a, GST protein; b, JUB1-GST protein; c, 5'-DY682 double-stranded oligonucleotide containing the perfect JUB1 binding site; d, $100 \times$ competitor (unlabeled oligonucleotide containing perfect JUB1 binding site); e, $200 \times$ competitor (unlabeled oligonucleotide containing perfect JUB1 binding site); f, $200 \times$ mutated oligonucleotide (unlabeled with mutation in JUB1 binding site where 5'-GATGCCGTTAGAGACACG-3' was replaced by 5'-GATGCC<u>AA</u>TA-GAGACACG-3').

(D) ChIP-qPCR. Whole shoots of 35-d-old *Arabidopsis* plants expressing GFP-tagged JUB1 under the control of the CaMV 35S promoter (35S:JUB1-GFP) and wild-type plants were harvested for the ChIP experiment. qPCR was used to quantify enrichment of the *DREB2A* promoter. As negative controls, primers annealing to promoter regions of two *Arabidopsis* genes lacking a JUB1 binding site, At3g18040 (Neg 1) and At2g22180 (Neg 2), were used. Data represent means \pm sp of three independent experiments.

TFs of the MYB (*PAP1* and *PAP2*), bHLH (*TT8*), and WRKY (*TTG2*) families was also reduced in *JUB1* overexpression plants compared with the wild type. Expression of *PRODUCTION OF ANTHO-CYANIN PIGMENT2* (*PAP2*) was significantly induced in *jub1-1* knockdown plants (see Supplemental Figure 12 online). Apart from *PAP2*, expression of anthocyanin biosynthesis genes was

either not affected or only slightly reduced in *jub1-1* knockdown plants. These data indicate that JUB1, most probably in conjunction with other TFs, negatively regulates the expression of the anthocyanin biosynthesis genes. Anthocyanins represent a group of flavonoids that accumulate under conditions of various types of environmental stresses. Moreover, they accumulate

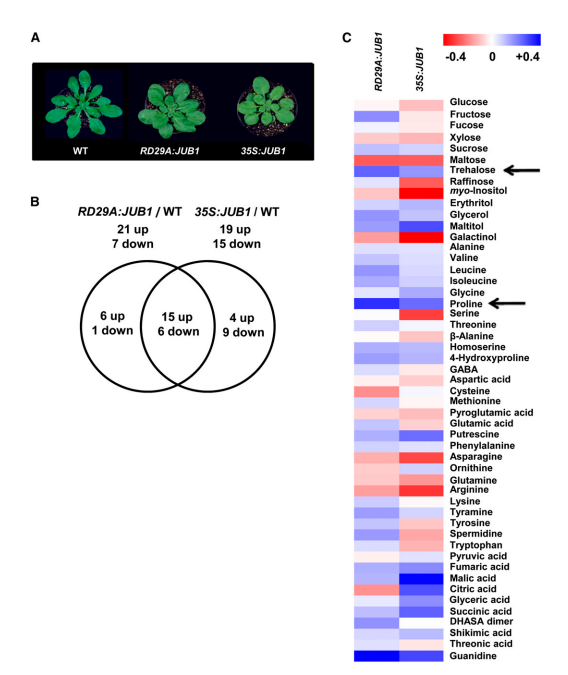


Figure 11. Primary Metabolite Profiling of JUB1 Overexpression Plants.

(A) Phenotype of 35-d-old wild-type (WT), RD29A:JUB1, and 35S:JUB1 plants subjected to metabolite profiling.

(B) Venn diagram showing an overview of metabolites that are significantly different (P < 0.05, Student's *t* test) in 35S:JUB1 and RD29A:JUB1 lines compared with the wild type.

(C) Hierarchical average linkage clustering of all detected primary metabolites. For every metabolite, the metabolic content of the wild type was considered as 1 and the metabolic content of overexpression lines was normalized to that. Metabolic ratios: red, minimum (between 0 and –0.4); blue, maximum (between 0 and + 0.4); see also Supplemental Data Set 3 online. Arrows indicate increased trehalose and Pro content in *JUB1* overexpressors compared with the wild type.

in senescing leaves preceding chlorophyll breakdown and play a photoprotective role against strong light in combination with coolness that may occur during autumn (Hoch et al., 2003; Diaz et al., 2006). The reduced level of anthocyanins in *JUB1* overexpression plants is consistent with their prolonged longevity. Of the other secondary metabolites, one of the major phenylpropanoid compounds, sinapoylmalate, was significantly induced in the *JUB1* overexpression plants (Figure 12). Similarly, transcript levels of several genes encoding enzymes of phenylpropanoid metabolism, such as *FAH1* (encodes ferulate-5-hydroxylase), *ALDH* (aldehyde dehydrogenase 2C4), and *SMT* (sinapoylglucose:malate

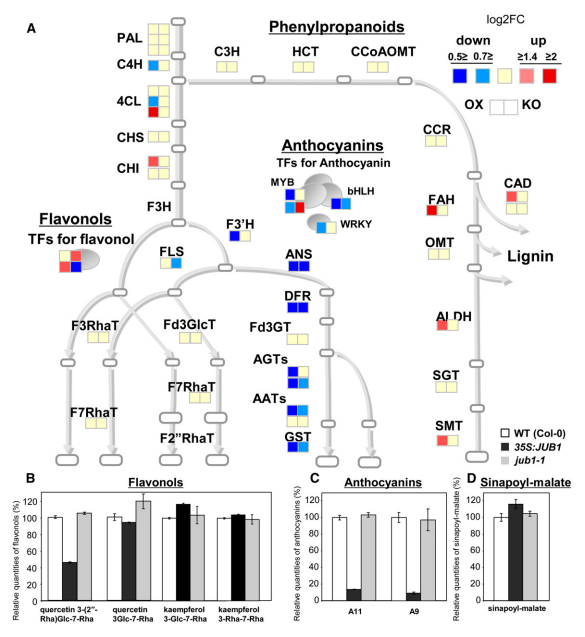


Figure 12. Gene Expression Profiling of Phenylpropanoids, Flavonols, and Anthocyanins, and Metabolite Profiling of Related Secondary Metabolites in 35S: JUB1, jub1-1, and Wild-Type Plants.

(A) Expression of 33 enzymatic genes and six TFs as measured by qRT-PCR. Intensity of fold change against wild-type (WT) expression level (log₂FC) is indicated by color. Abbreviations are given in Supplemental Table 2 online.

(B) to (D) The content of flavonols [quercetin-3-O-(2''-O-Rha)Glc-7-O-Rha, quercetin-3-O-Glc-7-O-Rha, kaempferol-3-O-Glc-7-O-Rha, and kaempferol-3-O-Rha], anthocyanins (A11 and A9; Tohge et al., 2005), and sinapoyl-malate in 35S:JUB1, jub1-1, and wild-type plants was analyzed by LC-MS. Average of two biological replicates ± sp.

sinapoyltransferase, which catalyzes the formation of sinapoylmalate from sinapoylglucose), were induced in *JUB1* overexpression plants (see Supplemental Figure 12 online).

Hormonal Adjustments in JUB1 Transgenics

Plant hormones play an important role in regulating senescence (in particular cytokinins) and the response to stress (including ABA, salicylic acid [SA], and jasmonic acid [JA]). We determined the concentrations of these hormones in rosette leaves of 43-dold 35S:JUB1, jub1-1, and wild-type plants. Significantly higher levels of isopentenyladenosine (IPA) and zeatin riboside (ZR) were detected in leaves of 35S:JUB1 transgenics compared with jub1-1 and wild-type plants (Figure 13A), while levels of zeatin (Z), dihydrozeatin (DHZ), and dihydrozeatin riboside (DHZR) were not

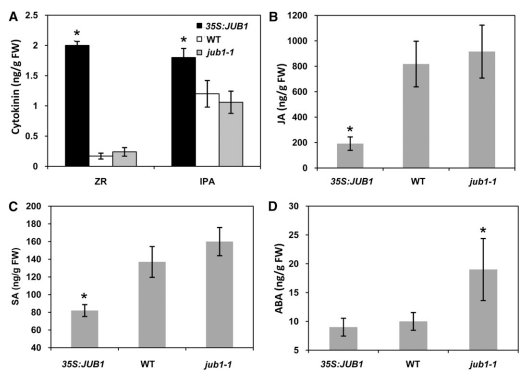


Figure 13. Hormone Contents in 35S:JUB1, jub1-1, and Wild-Type Plants.

Determination of ZR and IPA (A), JA (B), SA (C), and ABA (D) in 43-d-old 35S:JUB1, jub1-1, and wild-type plants grown at long-day conditions (16 h light/8 h dark). Values represent the means \pm SD from five independent sets of samples. Asterisks indicate significant differences compared with the wild type (WT) (P < 0.05, Student's *t* test). FW, fresh weight.

significantly altered (see Supplemental Figure 6C online). By contrast, levels of the biotic stress hormones JA and SA were significantly reduced in 35S:JUB1 lines compared with the wild type (Figures 13B and 13C), while ABA was not affected (Figure 13D). However, ABA was increased by \sim 94% in *jub1-1* mutant plants compared with the wild type (Figure 13D).

DISCUSSION

To identify novel regulators of plant senescence, we screened NAC overexpression and T-DNA insertion lines for changes in leaf senescence. We previously reported the identification of ORS1 as a TF that positively regulates senescence (Balazadeh et al., 2011). Similarly, At NAP and ORE1 (ANAC092) have been shown to act as positive regulators of senescence in Arabidopsis (Guo and Gan, 2006; Kim et al., 2009; Balazadeh et al., 2010a). Here, we discovered another member of the NAC gene family, designated JUB1, which in contrast with these previously characterized NAC factors, strongly delays senescence when overexpressed in transgenic plants and triggers precocious senescence at low expression level (in the jub1-1 mutant and artificial microRNA lines). Thus, JUB1 represents a strong negative regulator of senescence whose molecular function may differ from those of the positively acting NAC factors. Notably, the expression of all four NACs is triggered by H₂O₂, although the H₂O₂-dependent induction is slightly less pronounced for ORE1 compared with At NAP, ORS1, and JUB1 (Balazadeh et al., 2010b). Additionally, the expression of several other senescence-regulated NAC genes, such as *ANAC032*, *ATAF1*, and *ANAC102*, is triggered by H_2O_2 (Balazadeh et al., 2010b). This observation is interesting and suggests a close regulatory node connecting the accumulation of cellular H_2O_2 to the regulation of senescence and possibly bolting, a developmental process often tightly linked with the onset of leaf senescence (Levey and Wingler, 2005; Balazadeh et al., 2008a). However, a distinct role during senescence has not been reported for the other NACs so far. Recently, VNI2 (*ANAC083*; At5g13180) was found to regulate senescence by integrating ABA signaling (Yang et al., 2011); it also regulates xylem vessel specification (Yamaguchi et al., 2010). Currently, however, it remains largely unknown how these diverse cellular functions are integrated by VNI2.

Similar to observations in animals, extended longevity in plants is known to be correlated with increased tolerance to oxidative stress (Finkel and Holbrook, 2000; Muller et al., 2007). The correlation between stress tolerance and the onset of senescence and determination of life span in plants is supported by experimental evidence (Jing et al., 2003). Additionally, increased stress tolerance was also observed for late-flowering/long-living gigantea, ore1, ore3, and ore9 mutants (Kurepa et al., 1998; Woo et al., 2004). It has been suggested that aging is triggered by oxidative stress as a result of an imbalance between production and scavenging of oxygen radicals. In plants, this hypothesis is in part supported by the observation that timing of senescence is altered in mutants with a decreased level of the antioxidant L-ascorbic acid (vitamin C). The *vtc1* mutant enters senescence prematurely and is more sensitive to various oxidative stresses than the corresponding wild type (Barth et al., 2004). Compared with other ROSs, H_2O_2 has a relatively long half-life of ~ 1 ms, although its stability is influenced by the cellular pH and redox equilibrium (Reth, 2002). H_2O_2 acts as a signaling molecule that regulates plant development and adaptation to various stresses. It has been observed that a decrease of catalase (CAT2) and cytosolic ascorbate peroxidase 1 (APX1) activities during bolting time is followed by an accumulation of H_2O_2 and an enhanced expression of the senescence-associated TF *WRKY53*, suggesting H_2O_2 functions as a signal to promote senescence (Ye et al., 2000; Miao et al., 2004; Zimmermann et al., 2006).

Although the precise molecular pathways through which JUB1 regulates longevity and abiotic stress tolerance are not known at present, one possible scenario is that it does so by affecting a gene regulatory network that possibly involves DREB2A, its direct downstream target. DREB2A is an important transcription regulator acting in response to various abiotic stresses (e.g., Sakuma et al., 2006a; Kant et al., 2008). Transcriptome studies have shown that DREB2A activates a large number of abiotic stress-responsive genes involved in drought and heat stress responses (Sakuma et al., 2006b). The heat shock TF gene HsfA3 has been identified as a direct downstream target of DREB2A during heat stress (Schramm et al., 2008; Yoshida et al., 2008), and DREB2A itself is a heat shock-responsive gene (Suzuki et al., 2011). Another direct target of DREB2A is RESPONSIVE TO DESSICATION29A (RD29A; also referred to as COR78; Liu et al., 1998). Although we have not experimentally tested heat stress-dependent JUB1 expression here, global transcriptome data from Swindell (2006) identified JUB1 as a heat stressresponsive gene. Another observation of interest is that all three genes are significantly upregulated by H₂O₂ treatment (10 mM, 5 h) in Arabidopsis seedlings, although induction levels varied between the three TF genes (JUB1, ~25-fold; DREB2A, ~90fold; HsfA3, approximately threefold; see Supplemental Figure 6D online). Taken together, our data in conjunction with published reports (Schramm et al., 2008; Yoshida et al., 2008) establish an extended transcriptional cascade involving three consecutive positive regulators (JUB1-DREB2A-HsfA3), with JUB1 adopting an upstream position. Furthermore, HsfA3 has been suggested to be part of an expression amplification loop (Nishizawa-Yokoi et al., 2011) involving two additional heat shock TFs (i.e., HsfA1e and HsfA2), where HsfA1e activates HsfA2 expression (likely by direct binding to heat shock ciselements present in its promoter; Nishizawa-Yokoi et al., 2011), HsfA2 activates HsfA3 expression (Schramm et al., 2006), and HsfA3 stimulates expression of HsfA1e (Yoshida et al., 2008). HsfA2 has been shown to directly regulate the expression of APX2, which encodes a key cytosolic enzyme for the detoxification of H₂O₂ (Shigeoka et al., 2002; Nishizawa et al., 2006; Schramm et al., 2006), consistent with our observation of reduced H₂O₂ level in JUB1 overexpressors. The control network linking H₂O₂ signaling with this Hsf activation loop most likely involves additional transcriptional regulators, including MBF1c, which was shown to be required for enhanced expression of DREB2A during heat stress (Suzuki et al., 2011). Notably, MBF1c is highly responsive to abiotic stresses, including heat stress

(Suzuki et al., 2011), and is also rapidly and strongly upregulated by H_2O_2 treatment (~30-fold up already after 1 h at 10 mM H_2O_2 ; see Supplemental Figure 6E online). As we have shown here (Table 1), MBF1c is significantly more upregulated after H_2O_2 challenge in *JUB1* overexpressors than in the *jub1-1* mutant, further supporting the model of a regulatory link between JUB1 and MBF1c upstream of *DREB2A* and *HsfA3*. In accordance with this model is the observation that *MBF1c* contains a JUB1 binding site (CGCCGT) in its promoter at around 640 bp upstream of the transcription start site; however, we have not yet tested its functional relevance.

Our analysis presented here also identified changes in primary and secondary metabolism in *JUB1* transgenic lines. In general, our analysis revealed an accumulation of various compatible solutes, including trehalose, Pro, and various sugar alcohols (maltitol, glycerol, and erythritol) in *JUB1* overexpressors compared with wild-type plants, which may contribute to the enhanced abiotic stress tolerance of such lines. Although the role of these metabolites in senescence is not well established, examples indicate that trehalose and sugar alcohols (mannitol and inositol) delay senescence of cut flowers in some species (reviewed in van Doorn and Woltering, 2008). We also noticed a significant increase

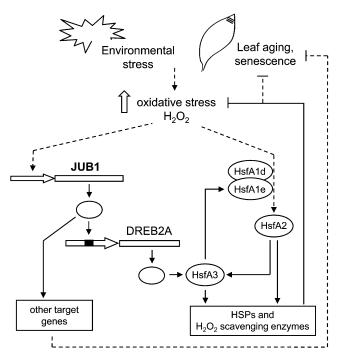


Figure 14. Model of JUB1 Action.

JUB1 is activated by H_2O_2 and during leaf senescence. JUB1 TF binds to the *DREB2A* promoter, thereby activating its expression. DREB2A positively regulates the expression of *HsfA3* (Schramm et al., 2008; Yoshida et al., 2008), thus establishing a transcriptional cascade. As suggested by Nishizawa-Yokoi et al. (2011), HsfA3 together with Hsf1A1e and HsfA2 form an expression amplification loop. HsfA3 and HsfA2 regulate the expression of *HSPs* and H_2O_2 scavenging enzymes, leading to reduced intracellular H_2O_2 levels, extended longevity, and increased stress tolerance. Increased longevity may also be regulated through other JUB1 target genes.

in Suc concentration due to JUB1 overexpression. At the level of secondary metabolites, we observed a decrease of cyanidin derivatives upon JUB1 overexpression, which was accompanied by reduced expression of anthocyanin biosynthesis genes (including DFR, LDOX, A5GT, and others) and known anthocyanin regulatory TFs (i.e., PAP1, PAP2, TTS, and TTG2). By contrast, expression of PAP2 was significantly induced in jub1-1 knockdown plants. Our data thus indicate that JUB1, possibly jointly with other TFs, has a negative effect on the expression of anthocyanin biosynthesis genes. Anthocyanins accumulate as a response to various types of environmental stresses and in senescing leaves, where they play a photoprotective role against high light stress in combination with low temperature, which occurs, for instance, in autumn (Hoch et al., 2003; Diaz et al., 2006). The reduced anthocyanin level in JUB1 overexpression plants is consistent with their prolonged longevity. Furthermore, minor flavonol glycosides, such as quercetin glycosides, were slightly decreased in leaves of JUB1 overexpressors. This may be due to the downregulated expression of F3'H (TT7) caused by the suppression of PAP1. Despite this fact, the increased levels of other effective phenolic antioxidants, like sinapoyl-malate and major glycosides of kaempferol, may contribute to the enhanced oxidative stress tolerance (Figure 12).

At the hormone level, we found higher levels of cytokinins (ZR and IPA) in *JUB1* overexpressors. Cytokinins are important regulators of senescence, and leaf senescence and the induction of SAGs can only be initiated when cytokinin levels are below a threshold (Gan and Amasino, 1995; Noodén et al., 1997). The high cytokinin level in *JUB1* overexpressors is thus in accordance with their extended lifespan. On the contrary, ABA level increased in the *jub1-1* mutant, indicating cellular stress, possibly due to disturbed H_2O_2 homeostasis in these plants.

Model for JUB1 Action

Based on the available experimental data, we propose the following model for JUB1 action (Figure 14). JUB1 transcription is activated by a rise of endogenous H_2O_2 concentration that is triggered by developmental input or environmental stress, including wounding, salinity stress, or cellulase treatment. Notably, JUB1 expression follows the cellular H₂O₂ concentration that changes with plant development, showing a peak during bolting (Zimmermann et al., 2006). The upstream TF(s) regulating H₂O₂dependent or senescence-associated JUB1 transcription remains unknown at present. JUB1 directly targets DREB2A, which functions as a positive regulator of HsfA3 and RD29A. HsfA3 itself regulates the expression of HSP genes and is part of a positive feedback loop together with HsfA1e and HsfA2 (Nishizawa-Yokoi et al., 2011), whereas the molecular function of RD29A is not known. Elevation of JUB1 expression lowers H₂O₂ concentration in plant tissues, possibly through the Hsf amplification loop, while the opposite effect (i.e., increased H₂O₂ concentration) is observed in the jub1-1 mutant, suggesting that JUB1 assists in regulating cellular H₂O₂ homeostasis. Although the precise molecular mechanism through which JUB1 regulates intracellular H₂O₂ concentration remains to be established, the current model proposes an involvement of HSPs and H₂O₂ scavenging enzymes. Notably, several GST genes (including

GST10, 24, and 25) were induced in JUB1 overexpressors after H_2O_2 treatment, while expression of these genes remained either unchanged or was slightly reduced in the *jub1-1* mutant (Table 1).

Concomitant with enhanced JUB1 expression, we observed reduced expression of many SAGs, which is in accordance with the delayed senescence observed in these lines. Notably, expression of SAGs is increased in the jub1-1 mutant, constituting JUB1 as a negative regulator of senescence. A possible explanation of the reduced expression of SAGs in JUB1 overexpressors may be derived from the observation that expression of many SAGs is enhanced by H₂O₂ (e.g., Navabpour et al., 2003; Balazadeh et al., 2010b; Genevestigator at http://www.genevestigator.com). In particular, expression of the majority of the senescence-associated NAC TFs is also triggered by H₂O₂ treatment (15 NACs in total), including At NAP, ORS1, and ORE1 (Balazadeh et al., 2010b), all of which have been shown to affect senescence positively. Thus, our model suggests that JUB1 lowers the cellular H₂O₂ level, thereby minimizing the stimulatory effect on NAC gene expression and, hence, senescence. On the contrary, reduced JUB1 expression (such as in the jub1-1 mutant) would favor the accumulation of cellular H₂O₂ that drives NAC gene expression and through this supports precocious senescence. However, there is also the additional possibility that JUB1 regulates senescence through other target genes of currently unknown molecular function (Figure 14). Future work will have to address the intricacies of the underlying regulatory network in greater detail.

METHODS

General

Standard molecular techniques were performed as described (Sambrook et al., 2001; Skirycz et al., 2006). Oligonucleotide sequences are given in Supplemental Data Set 5 online. Chemicals and reagents for GC-MS analysis were obtained from Sigma-Aldrich, Fluka, or Merck with the exception of *N*-methyl-*N*-(trimethylsilyl) trifluoroacetamide, which was obtained from Macherey-Nagel. For sequence analyses, the tools provided by the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/), MIPS (http://mips.gsf.de/), The Arabidopsis Information Resource (http://www.arabidopsis.org/), and the Plant Transcription Factor Database (http://plntfdb.bio.uni-potsdam.de/v3.0/) were used.

Plants

Seeds of Arabidopsis thaliana accession Col-0 were obtained from the Arabidopsis thaliana Resource Centre for Genomics (Institut National de la Recherche Agronomique, France; http://dbsgap.versailles.inra.fr/ publiclines/). For growth under long-day conditions, seedlings were grown in soil (Einheitserde GS90; Gebrüder Patzer) in a climate-controlled chamber with a 16-h daylength provided by fluorescent light at ${\sim}100\,\mu\text{mol}\,\text{m}^{-2}$ s⁻¹ and a day/night temperature of 20/16°C and a RH of 60/75%. After 2 weeks, seedlings were transferred to a growth chamber with a 16-h day (80 or 120 μ mol m⁻² s⁻¹) and a day/night temperature of 22/16°C and 60/75% RH. For growth under short-day conditions, the light period was reduced to 8 h. Growth in hydroponic culture, salinity treatment, and sample preparation were done as described using stage 1 plants (28 d old) (Balazadeh et al., 2010a). T-DNA insertion lines screened for extended longevity (see Supplemental Table 1 online) were obtained from the European Arabidopsis Stock Centre (http://Arabidopsis.info/). Homozygous plants were identified by PCR using the T-DNA left border primer, as well as the gene-specific primers LP and RP.

Constructs

Constructs were generated by PCR- and restriction enzyme-mediated cloning. Primer sequences are given in Supplemental Data Set 5 online. PCR-generated amplicons were checked by DNA sequence analysis (MWG). Constructs were transformed into *Arabidopsis* Col-0 via *Agrobacterium tumefaciens*-mediated transformation.

For 35S:JUB1, the JUB1 open reading frame was amplified by PCR from Arabidopsis Col-0 leaf cDNA and inserted into pUni/V5-His-TOPO (Invitrogen). The cDNA was cloned via added Pmel-PacI sites into a modified pGreen0229-35S plant transformation vector (Skirycz et al., 2006). For RD29A:JUB1, the RD29A promoter (1 kb upstream of translation start site) was amplified from Arabidopsis (Col-0) genomic DNA, cloned into pCR2.1 (Invitrogen), and then transferred via BamHI and Ncol sites into pCAMBIA1305.1-hygromycin, giving rise to plasmid RD29A: pCAMBIA. The JUB1 coding region was amplified by PCR from leaf cDNA using primers JUB1-forward and JUB1-reverse and cloned downstream of the RD29A promoter in plasmid RD29A:pCAMBIA via primer-added Ncol and Pmll sites. For 35S:JUB1-GFP, the full-length JUB1 open reading frame was amplified without its stop codon. The PCR product was cloned into the pENTR/D-TOPO vector using the pENTR Directional TOPO cloning kit (Invitrogen). The sequence-verified entry clone was then transferred to the pK7FWG2 vector (Ghent University) by LR recombination (Invitrogen). For JUB1-IOE, the JUB1 coding region was amplified by PCR from Arabidopsis leaf cDNA using primers JUB1-IOE-fwd and JUB1-IOE-rev, inserted into pBluescript SK⁺, and then cloned via Xhol and Spel sites into the pER8 vector (Zuo et al., 2000). For the Pro, JUB1: GUS fusion, an \sim 1.8-kb 5' genomic fragment upstream of the translation initiation codon was amplified by PCR from Arabidopsis Col-0 genomic DNA, inserted into plasmid pGEM-T Easy (Promega), and fused via HindIII (present in pGEM-T Easy) and Ncol restriction sites to the GUS reporter gene in pCAMBIA1305.1-hygromycin (CAMBIA). For JUB1-amiRNA, the Web MicroRNA Designer platform (http://wmd2.weigelworld.org/cgi-bin/ mirnatools.pl?page=1) was used to design amiRNA sequences (21mers). For ProDREB2A:FLuc, the ~1.8-kb DREB2A promoter containing the JUB1 binding site was amplified by PCR from Arabidopsis genomic DNA and inserted into the pENTR/D-TOPO vector (Invitrogen). The sequence-verified promoter was then transferred to the p2GWL7.0 vector harboring the firefly (Photinus pyralis) luciferase (FLuc) coding region (Licausi et al., 2011) by LR recombination (Invitrogen).

Expression Profiling by qRT-PCR

Total RNA extraction, synthesis of cDNA, and gRT-PCR were performed as described (Caldana et al., 2007; Balazadeh et al., 2008b). Expression analysis platforms contained primer pairs for 168 SAGs (Parlitz et al., 2011), 179 ROS-responsive genes, and 94 phenolic secondary metabolite biosynthetic genes. Genes included in the SAG platform are highly upregulated during natural senescence in wild-type Arabidopsis plants (Buchanan-Wollaston et al., 2005; van der Graaff et al., 2006; Balazadeh et al., 2008b). ROS-responsive genes were extracted from the literature (Gechev et al., 2004, 2005; Davletova et al., 2005a, 2005b; Gadjev et al., 2006) and in-house experiments. Primers for the metabolite platform were designed for the biosynthetic genes from primary metabolism to flavonoid production; the platform also included primers for other phenolic secondary metabolite biosynthetic genes and TFs that control anthocyanin biosynthesis. Genes included in the qRT-PCR platforms, including primer sequences, are given in Supplemental Data Set 5 online. Primers were designed using QuantPrime (Arvidsson et al., 2008). PCR reactions were run on an ABI PRISM 7900HT sequence detection system (Applied Biosystems Applera), and amplification products were visualized using SYBR Green (Applied Biosystems). ACTIN2 served as reference gene; primers were Actin2-F (5'-TCCCTCAGCACATTCCAGCAGAT-3') and Actin2-R (5'-AACGATTCCTGGACCTGCCTCATC-3').

DNA Binding Site Selection

In vitro binding site selection was performed using the CELD system with the pTacJUB1-LCELD6XHis construct, employing three biotin-labeled double-stranded oligonucleotides (i.e., Bio-RS-Oligo 1, RS-Oligo 2, and Bio-RS-Oligo 3), which contained 30-nucleotide random sequences that differed in flanking primer sequence (Xue, 2005). JUB1-selected oligonucleotides were cloned and sequenced. The DNA binding activity of JUB1-CELD was measured using methylumbelliferyl β -D-cellobioside as substrate (Xue, 2002). DNA binding assays with a biotin-labeled singlestranded oligonucleotide or a biotin-labeled double-stranded oligonucleotide without a target binding site were used as controls.

Transactivation Assays

Arabidopsis mesophyll cell protoplasts were prepared according to the protocol of Sheen (2002). The construct containing the \sim 1.8-kb DREB2A promoter fragment in front of the FLuc coding region (ProDREB2A:FLuc) was cotransformed in the presence or absence of the 35S:JUB1 plasmid. The 35S:RLuc vector (Licausi et al., 2011) was used for normalization against transformation efficiency. Firefly luciferase and Renilla luciferase (RLuc) were assayed using the Dual Luciferase Reporter Assay System (Promega). Six micrograms of DNA were used for transient transformation of protoplasts according to Yoo et al. (2007); 16 h after incubation, protoplasts were lysed by adding 400 μL of passive lysis buffer, and the resulting suspension was briefly vortexed. Forty microliters of luciferase assay buffer was added to the same volume of crude extract and FLuc activity was measured. Forty microliters of Stop and Glow buffer was then added and Renilla chemiluminescence was measured. Relative light units were determined in a GloMax 20/20 luminometer (Promega) using a 10-s measurement. Data were collected as ratio (FLuc activity:RLuc activity). Protoplasts transformed with only the promoter-FLuc, 35S:RLuc reporter plasmid (no TF), were analyzed as background controls.

EMSA

JUB1-GST fusion protein was purified from Escherichia coli expression strain BL21 Star (DE3) pRARE, which was generated by transforming the pRARE plasmid isolated from Rosetta (DE3) pRARE cells (Merck) into E. coli BL21 Star (DE3) (Invitrogen). Protein expression was induced in a 100-mL expression culture using 1 mM isopropyl thio-β-D-galactoside, and cells were harvested 4 h after induction at 30°C. Cells were sonicated in lysis buffer (20 mM sodium phosphate buffer, pH 7.3, 150 mM NaCl, 1 mM EDTA, 1 mM DTT, and 1 mM phenylmethanesulfonyl fluoride). Supernatant of centrifuged sample was used for purification using a 1-mL GSTrap HP column (GE Healthcare) coupled to the Äkta-Purifier FPLC system (GE Healthcare). Aliquots of the flow-through fractions were analyzed by SDS-PAGE and Coomassie Brilliant Blue staining. Onemilliliter elution fractions containing the purified JUB1-GST fusion protein were pooled and dialyzed against PBS buffer (20 mM Na-phosphate, pH 7.4, and 150 mM NaCl). Protein concentration was determined by the Bradford assay (Bradford, 1976). 5'-DY682-labeled DNA fragments were ordered from MWG. Sequences of labeled DNA fragments, unlabeled competitors, and mutated fragments are given in Supplemental Data Set 5 online. Annealing was performed by heating the primers to 100°C followed by slow cooling to room temperature. The binding reaction was performed at room temperature for 20 min as described in the Odyssey Infrared EMSA kit instruction manual. DNA-protein complexes were separated on 6% retardation gel, while DY682 signal was detected using the Odyssey Infrared Imaging System from LI-COR Biosciences.

In Vivo Binding of JUB1 to the DREB2A Promoter

To investigate in vivo binding of JUB1 to its DNA binding site in the *DREB2A* promoter, we used ChIP-qPCR, using whole shoots from long day–grown,

35-d-old Arabidopsis plants expressing GFP-tagged JUB1 protein from the CaMV 35S promoter (35S:JUB1-GFP). Wild-type plants were used as negative control. For the ChIP, we followed a protocol previously described by Kaufmann et al. (2010) employing anti-GFP antibody to immunoprecipitate protein-DNA complexes. The ChIP experiment was run in three independent replications. qPCR was used to test binding of JUB1 to its binding site within the DREB2A promoter; the primers flanked the JUB1 binding site. As a negative control, we used primers annealing to promoter regions of two other Arabidopsis genes (At3g18040 and At2g22180) lacking a JUB1 binding site. Primer sequences are given in Supplemental Data Set 5 online. We analyzed ChIP-qPCR data relative to input, as this includes normalization for both background levels and input chromatin going into the ChIP. The amount of genomic DNA coprecipitated by GFP antibody (ChIP signal) was calculated in comparison to the total input DNA used for each immunoprecipitation in the following way: cycle threshold $(C_T) = C_T(ChIP) - C_T(Input)$. To calculate fold enrichment, normalized ChIP signals were compared between 35S:JUB1-GFP and wild-type plants, where the ChIP signal is given as the fold increase in signal relative to the background signal.

H₂O₂ Measurements

The H₂O₂ staining agent, DAB (D5637, Sigma-Aldrich), was dissolved in water and adjusted to pH 3.8 with KOH. To avoid auto-oxidation, the DAB solution was freshly prepared (Fryer et al., 2002). Whole seedlings were infiltrated under vacuum with 0.5 mg mL⁻¹ DAB staining solution and further incubated for 12 h in medium; chlorophyll was removed by incubating seedlings in 90% ethanol at 70°C for 10 min. H₂O₂ was visualized as brown color due to DAB polymerization.

Quantitative measurement of H_2O_2 production was performed using the Amplex Red hydrogen peroxide/peroxidase assay kit (Molecular Probes) following the manufacturer's instructions. Briefly, samples were ground in liquid nitrogen, and 30 mg of ground frozen tissue from each sample was placed in an Eppendorf tube and kept frozen. Four hundred milliliters of 20 mM sodium phosphate buffer, pH 6.5, was immediately added into the tube and mixed. The extraction was centrifuged at 10,000g for 10 min at 4°C, and the supernatant was used for the assay. Measurements were performed at excitation and emission wavelengths of 560 and 590 nm, respectively, using a 96-well LS55 luminescence spectrometer (PerkinElmer). H_2O_2 levels are given in pmol/mg frozen tissue.

Primary Metabolite Profiling by GC-MS

To perform this study, we grew all plants alongside each other under carefully controlled conditions. Metabolite extraction, derivatization, and relative metabolite levels were determined using an established GC-MS protocol as described previously (Roessner et al., 2001; Lisec et al., 2006). Metabolites were identified in comparison to database entries of authentic standards (Kopka et al., 2005; Schauer et al., 2005).

Secondary Metabolite Profiling by LC-MS

Secondary metabolite analysis by LC-MS was performed as described by Tohge and Fernie (2010). All data were processed using Xcalibur 2.1 software (Thermo Fisher Scientific). The obtained data matrix was normalized using an internal standard (Isovitexin; CAS 29702-25-8). Metabolites were identified and annotated based on comparisons with data in our previous publications (Tohge et al., 2005, 2007; Hirai et al., 2007; Yonekura-Sakakibara et al., 2008), metabolite databases (reviewed in Tohge and Fernie, 2009), and standard compounds (Yonekura-Sakakibara et al., 2008; Nakabayashi et al., 2009).

Hormone Analyses

The extraction and analysis of ABA, SA, JA, and the cytokinins Z, ZR, DHZ, DHZR, and IPA were performed as described previously (Abreu and

Munné-Bosch, 2009), except that internal deuterated standards and ultraperformance liquid chromatography-MS/MS (instead of HPLC-MS/ MS) were used for the analysis. Briefly, 100 mg of leaf samples was ground in liquid nitrogen and extracted with 1.5 mL methanol using sonication. After centrifugation, the supernatant was collected and the pellet was reextracted with isopropanol:glacial acetic acid (99:1) to fully extract cytokinins. The two supernatants were dried completely under a nitrogen stream and redissolved in 150 µL methanol. Then, supernatants were combined, filtered through a 0.22-µm polytetrafluoroethylene filter (Waters), and injected into the LC-MS/MS system. MS/MS analyses were performed on an API 3000 triple quadrupole mass spectrometer (PE Sciex). All analyses were performed using the Turbo Ion Spray source in negative ion mode for ABA, SA, and JA and in positive ion mode for cytokinins. Internal standards (deuterium-labeled hormone analogs, including d₄-SA, d₆-ABA, d₅-JA, d₆-IPA, d₅-Z, and d₅-ZR purchased from OlChemIm) were added to each sample immediately after grinding, thus allowing the calculation of specific recovery rates for each compound. Quantification by MS/MS using the Multiple Reaction Monitoring method was performed as described by Abreu and Munné-Bosch (2009). Multiple Reaction Monitoring acquisition was done monitoring the following transitions: ABA, 263/153; SA, 137/93; JA, 209/59; IPA, 336/204; Z, 220/136; ZR, 352/220; DHZ, 222/136; DHZR, 354/222.2; d₆-ABA, 269/ 159; d₄-SA, 141/97.2; d₅-JA, 241/64; d₆-IPA, 342/210; d₅-Z, 225/137; and d_5 -ZR, 357/225. The declustering potential and collision energy were optimized for each compound.

Treatments

For EST induction, 12-d-old seedlings were incubated in liquid MS medium containing 15 µM EST (control treatment: 0.15% ethanol). The seedlings were kept on a rotary shaker for 30 min or 2, 6, or 24 h, harvested, and, after removal of the roots, immediately frozen in liquid nitrogen. JUB1 expression level was determined by qRT-PCR. For EST induction on plates, MS medium was supplemented with 10 µM EST (control: 0.1% ethanol). Cellulase treatment was performed as described (Rentel et al., 2004). Briefly, 12-d-old Col-0 seedlings grown on agar plates were transferred to liquid medium supplied with 0.1% cellulase R-10 (Onozuka R-10; Yakult). Seedlings were incubated for 5 h on a rotary shaker at continuous light, and expression of JUB1 was determined by qRT-PCR. For histochemical GUS assays, ProJUB1:GUS seedlings were treated with 0.5% cellulase R-10 for 3 h. For methyl viologen (MV) treatment, 12 d-old Col-0 seedlings grown on agar plates were transferred to liquid medium, supplied with 10 µM MV (Sigma-Aldrich). Seedlings were incubated for 5 h on a rotary shaker, and expression of JUB1 was determined by qRT-PCR. For histochemical GUS assays, ProJUB1:GUS seedlings were treated with 50 µM MV for 3 h. For salt treatment, seedlings grown on MS medium were transferred to liquid MS medium containing 150 or 200 mM NaCl, followed by incubation for the indicated times; alternatively, seeds were sown on MS medium containing 100 mM NaCl.

Microscopy

Distribution of JUB1-GFP fusion protein was analyzed by confocal fluorescence microscopy using an Eclipse E600 microscope (Nikon).

Other Methods

Histochemical GUS assays was performed as described by Plesch et al. (2001). Fluorometric determination of GUS activity was done using 4-MUG (Sigma-Aldrich) as substrate (Jefferson et al., 1987). Chlorophyll content was determined using a SPAD analyzer (N-tester; Hydro Agri). Alternatively, frozen *Arabidopsis* leaves were ground in liquid nitrogen, resuspended in 1 mL of 96% (v/v) ethanol, and homogenized for 1 min.

The soluble fraction, which contains chlorophyll, was separated by centrifugation (5 min, 13,000 rpm), and the amount of chlorophyll in the extract was determined spectrophotometrically at 650 nm. Ion leakage in the first six leaves was determined as described (Guo and Gan, 2006).

Statistical Analyses

Unless otherwise specified, statistical analyses were performed using Student's *t* test embedded in Microsoft Excel. Only the return of P < 0.05 was designated as statistically significant.

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the following accession numbers: *ACTIN2* (At3g18780), *ALDH* (At3g24503), At *NAP* (At1g69490), *DREB2A* (At5g05410), *FAH1* (At4g36220), *JUB1* (At2g43000), *MBF1c* (At3g24500), *ORE1* (At5g39610), *ORS1* (At3g29035), *SMT* (At2g22990), *TT7* (At5g07990), and *VNI2* (At5g13180). Additional accession numbers are given in Supplemental Tables 1 and 2 online and Supplemental Data Sets 1, 2, 4, and 5 online.

Supplemental Data

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

Supplemental Figure 1. Early Leaf Senescence in *JUB1-amiRNA* Lines.

Supplemental Figure 2. Localization of JUB1-GFP Fusion Protein in Guard Cells of Transgenic *Arabidopsis*.

Supplemental Figure 3. Phenotypic and Physiological Analyses of Transgenic Plants Overexpressing *JUB1* under Control of Stress-Inducible Promoter *RD29A*.

Supplemental Figure 4. Confirmation of T-DNA Insertion in *jub1-1 M*utant.

Supplemental Figure 5. Dark-Induced Senescence Is Affected in Detached Leaves of *JUB1* Overexpressors.

Supplemental Figure 6. Gene Expression and Hormone Levels in *Arabidopsis* Plants Subjected to Different Treatments.

Supplemental Figure 7. H₂O₂ Content and *JUB1* Expression Level in Individual Leaves of 35-d-Old Wild-Type Plants.

Supplemental Figure 8. Effect of NaCl, Cellulase, and Methyl Viologen on *JUB1* Expression.

Supplemental Figure 9. Effect of H_2O_2 on Detached Leaves of the *jub1-1* Mutant.

Supplemental Figure 10. The *jub1-1* Mutant Exhibits Decreased Heat Stress Tolerance Compared with the Wild Type.

Supplemental Figure 11. Enhanced Heat Stress Tolerance of *JUB1* Overexpressor.

Supplemental Figure 12. Expression of Secondary Metabolite-Associated Genes in 35S:*JUB1, jub1-1,* and Wild-Type Plants, Determined by qRT-PCR.

Supplemental Table 1. NAC Transcription Factor T-DNA Insertion Lines Included in the Screen for Extended Longevity.

Supplemental Table 2. Abbreviations of Enzyme Names.

Supplemental Data Set 1. Expression of 168 Senescence-Associated Genes in 47-d-Old 35S:JUB1, jub1-1, and Wild-Type Plants.

Supplemental Data Set 2. Expression of 179 ROS Genes in 2-Week-Old, H₂O₂-Treated (10 mM, 6 h) and Nontreated *RD29A:JUB1, jub1-1*, and Wild-type Plants.

Supplemental Data Set 3. Metabolite Composition of Rosette Leaves of 35-d-Old Wild-type, 35S:JUB1, and RD29A:JUB1 Plants.

Supplemental Data Set 4. Expression of 94 Secondary Metabolite-Associated Genes in 35-d-Old 35S:JUB1, jub1-1, and Wild-Type Plants.

Supplemental Data Set 5. Oligonucleotide Sequences.

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AUTHOR CONTRIBUTIONS

B.M.-R. and S.B. designed the research and supervised the group. A.D.A., A.W., H.S., P.G., and S.B. performed the research. M.-I.Z. generated the 35S:JUB1 lines. H.D. produced recombinant JUB1-GST protein. M.A.A.-F. and S.M.-B. performed the hormone analysis. C.A., T.T., and A.R.F. did the metabolite profiling. G.-P.X. performed the CELD experiment. K.K. and S.B. performed the ChIP experiment. B.M.-R. and S.B. wrote the article with contributions from the other authors.

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