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
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Subset of heat-shock transcription factors required for the early response of *Arabidopsis* to excess light

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Sunlight provides energy for photosynthesis and is essential for nearly all life on earth. However, too much or too little light or rapidly fluctuating light conditions cause stress to plants. Rapid changes in the amount of light are perceived as a change in the reduced/oxidized (redox) state of photosynthetic electron transport components in chloroplasts. However, how this generates a signal that is relayed to changes in nuclear gene expression is not well understood. We modified redox state in the reference plant, *Arabidopsis thaliana*, using either excess light or low light plus the herbicide DBMIB (2,5-dibromo-3-methyl-6-isopropyl-p-benzoquinone), a well-known inhibitor of photosynthetic electron transport. Modification of redox state caused a change in expression of a common set of about 750 genes, many of which are known stress-responsive genes. Among the most highly enriched promoter elements in the induced gene set were heat-shock elements (HSEs), known motifs that change gene expression in response to high temperature in many systems. We show that HSEs from the promoter of the *ASCORBATE PEROXIDASE 2* (*APX2*) gene were necessary and sufficient for *APX2* expression in conditions of excess light, or under low light plus the herbicide. We tested *APX2* expression phenotypes in overexpression and loss-of-function mutants of 15 *Arabidopsis* A-type heat-shock transcription factors (HSFs), and identified HSFA1D, HSFA2, and HSFA3 as key factors regulating *APX2* expression in diverse stress conditions. Excess light regulates both the subcellular location of HSFA1D and its biochemical properties, making it a key early component of the excess light stress network of plants.

Photosynthesis is the process by which light is converted to chemical energy; yet, despite its critical and central role for the sustenance of life, photosynthetic performance of plants is not optimal. Under light intensities of higher than optimal growth (termed excess light or EL), light energy absorption increases linearly as light intensity increases. However, carbon assimilation becomes saturated at a certain light intensity, termed the light saturation point. Thus, in EL, most of the absorbed light energy is not used for carbon metabolism. Instead, the EL energy causes damage to proteins involved in photosynthesis, a process called photoinhibition (1–3).

Plants have evolved multiple ways to sense and respond to EL (4). Photoreceptors, such as cryptochrome, sense light directly and lead to changes in nuclear gene expression and chloroplast avoidance responses. Excess light energy can also be dissipated as heat via nonphotochemical quenching, thereby avoiding damage to the photosynthetic apparatus. In addition to inactivating photosynthesis (3), it has been reported that EL brings about accumulation of a number of metabolic intermediates in plastids that act as signals of EL stress (5–7). Finally, EL causes the accumulation of multiple reactive oxygen species, including, singlet oxygen ($^1\text{O}_2$), hydrogen peroxide (H_2O_2), and superoxide anion (O_2^-) (8) in plastids, which are neutralized by reactive oxygen species detoxifying enzymes and by the synthesis of antioxidant molecules, such as carotenoids, ascorbic acid, and tocopherol (1).

Several studies have provided evidence for a role of photosynthetic electron transport (PET) components as sensors of EL and regulators of nuclear gene expression (9, 10). A stress-response gene encoding *ASCORBATE PEROXIDASE 2* (*APX2*) is controlled by the reduced/oxidized (redox) state of the plastoquinone (PQ) pool in *Arabidopsis* (11, 12). The expression of *APX2* is significantly induced by the reduced PQ pool that is generated by either EL or low light (LL) plus DBMIB (2,5-dibromo-3-methyl-6-isopropyl-p-benzoquinone), a herbicide that blocks electron transport between the PQ pool and the cytochrome *b₆f* complex (11, 13, 14). EL-driven *APX2* expression requires H_2O_2 and functional PET (11, 13).

The molecular mechanism by which the PQ redox state triggers the induction of *APX2* is unknown, although clues to some possible mechanisms have been reported. Despite *APX2*'s fast and very strong induction upon exposure to EL (11), genetic screens have identified only constitutive *APX2* expression mutants, but no mutants affected specifically in EL-driven *APX2* induction have been found (15, 16). There is evidence linking *APX2* induction by EL to the zinc-finger transcription factor ZAT10, to the plant hormone abscisic acid, and to 3'-phosphoadenosine 5'-phosphate, a phosphonucleotide (PAP) (5, 17, 18); however, a signal transduction pathway linking gene expression in the nucleus to redox of the PQ pool remains elusive.

Here, we took advantage of reporters of EL-induced gene expression and DBMIB to identify nuclear proteins specific to a chloroplast-to-nucleus signaling process generated by signals from the PQ pool. We determine that a reduced state of the PQ pool changes the expression of about 750 nuclear genes, many of which contain heat-shock elements (HSEs). A hierarchy of interactions among a subset of heat-shock transcription factors (HSFs)—HSFA1D, HSFA2, and HSFA3—is responsible for the early gene-expression response of *Arabidopsis* to EL.

Results

Defining Stress Conditions. Recent studies have linked the accumulation of a number of different metabolic intermediates with EL stress, leading to the proposal that the accumulation of these

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Data deposition: The Affymetrix ATH1 CEL files reported in this paper has been deposited in the ArrayExpress, www.ebi.ac.uk/arrayexpress (accession no. E-MTAB-403).

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intermediates is the initiating event for multiple stress signaling pathways (5–7). Other articles have reported that a signal originates from PET, specifically the redox state of the PQ pool (11, 19, 20). To better define the nature of the signals related to the PQ redox state, we treated plants with a number of different light stresses, as well as with LL plus the herbicide DBMIB.

First, we assessed the effectiveness of our DBMIB treatment in generating the PQ pool in a reduced state. Following incubation of leaf discs in a DBMIB (24 μ M) solution for 2 h under LL conditions, chlorophyll fluorescence and *APX2* transcript levels were determined. As expected, the DBMIB treatment increased the excitation pressure on PSII (1-qP) (21) (Fig. S1A) and *APX2* induction levels (Fig. S1B); in contrast, the 3,3,4-dichlorophenyl-1,1-dimethylurea (DCMU) treatment under LL did not induce *APX2* expression (Fig. S1B), despite its effect on increased 1-qP (Fig. S1A). These results confirmed that generation of our conditions for reducing the PQ pool with DBMIB worked as expected (11). However, both DBMIB and DCMU treatments failed to generate H₂O₂ in leaf tissue, whereas “hot” EL (hEL) treatment resulted in H₂O₂ generation (Fig. S1C).

We further explored *APX2* expression by light, high temperature, or both by exposing plant leaves to a range of EL conditions: hEL, “filtered” EL (fEL), which was achieved by using a mixture of water and ice layer (13, 22), and “cold” light (LED) (Fig. 1A). Although all EL conditions significantly increased leaf temperature, the effects of hEL on leaf temperature increases were more drastic than those of fEL and LED (Fig. S1D). All EL conditions up-regulated *APX2* expression, although *APX2* induction levels were lower in fEL and LED (Fig. 1A). In addition, it seems like *APX2* induction is positively correlated with the amount of H₂O₂ in leaves (Fig. S1E). Based on these observations, it is very likely that *APX2* expression is regulated by both light and high temperature in a synergistic way. In addition, it seems like there is a heat-independent *APX2* induction pathway. For the rest of this report, all EL experiments described were conducted using hEL and presented as EL.

EL Triggers Redox Stress and Changes the Expression of Hundreds of Genes. To understand the effect of the reduced PQ pool on global nuclear gene expression, we harvested mRNA from *Arabidopsis* leaf tissue treated with EL (for 30 min or 2 h), or LL plus DBMIB (for 30 min or 2 h), and generated expression profiles using ATH1 microarrays. We found ~1,250 genes up-regulated

and ~1,300 genes down-regulated by either treatment at either time point compared with before treatment, with 331 genes commonly up-regulated and 426 genes down-regulated by both treatments (Fig. 1B). Among the top genes up-regulated by both DBMIB and EL were genes encoding several heat-shock factors and small heat-shock proteins (some induced up to several thousand-fold), as well as *APX2*, expected from previous studies (Dataset S1) (13). Of the ~20 heat-shock factor protein family members, 10 are significantly ($P < 0.01$) up-regulated in at least one condition, and at least 30 heat-shock proteins are also significantly induced (Dataset S1). Gene Ontology term-enrichment analysis revealed nearly 20 processes overrepresented ($P < 0.01$ using the hypergeometric test) among the core up-regulated gene set, including response to heat, response to high light intensity, and response to oxidative stress, consistent with the treatments applied (Dataset S2). Inspection of promoter sequences (defined as the 500-bp upstream from the predicted ATG) from the 331 commonly up-regulated genes revealed that a high proportion of genes contained at least one of four variants of the known HSEs (e.g., GAAnnTTC) (Fig. S1F). Genes down-regulated by EL and DBMIB include many biosynthetic enzymes, cytochrome P450 enzymes, and transcription factors (Dataset S1), although the magnitude of change of down-regulated genes is smaller than for up-regulated transcripts. Gene Ontology enrichment for this gene set indicates a high overrepresentation for metabolic enzymes, especially those involved in glucosinolate biosynthesis (Dataset S2). In summary, ~750 genes are significantly altered by PQ redox state, which most notably induces expression of *APX2* and heat-shock factors, indicating a strong redox stress response by EL and DBMIB.

In addition to HSEs, we mined the predicted promoters of up-regulated genes for other previously known and novel overrepresented motifs using the ELEMENT algorithm (23) (Fig. S1F). Among genes up-regulated by 30 min of EL, we found a number of known light-regulatory elements, including the G-box (CACGTG) (24), and a GCCAC motif previously identified as overrepresented in light-induced promoters (SORLIPs) (25). We also found an ACGT motif (26), a CCGAC motif (27), and an ATCCAAT motif (28) (Fig. S1F).

***APX2* Induction by EL or DBMIB Does Not Require New Protein Synthesis.**

About 50% of the genes induced by either EL or DBMIB were common to both conditions, but induction by EL appeared to be faster than in DBMIB. In agreement with previous reports (11, 13), *APX2* induction was seen within 5 min of exposure to EL, achieved peak levels after 1 h of exposure to EL, and then decreased (Fig. S2A). Although *APX2* mRNA levels also increased in the presence of DBMIB, the induction profile was different as the fold-change was smaller than that with EL (Fig. S2A and B). Furthermore, *APX2* mRNA levels never peaked during the 2 h of DBMIB treatment (Fig. S2B), indicating that the effects of DBMIB were slower than those of EL (11). The rapid induction kinetics of *APX2* by EL or DBMIB led us to test whether new protein synthesis is required for *APX2* induction. Leaf tissue was pretreated with 200 μ M cycloheximide (CHX) for 30 min, then exposed to EL for 2 h, and *APX2* mRNA accumulation was determined (Fig. S2A). *APX2* induction was observed in the presence of CHX for the first hour of EL treatment. After 2 h of EL, *APX2* transcript levels were higher in the presence of CHX than in the absence of CHX (Fig. S2A). One explanation for this observation is that an unidentified factor, synthesized under EL, might negatively regulate EL-driven *APX2* induction. By a pretreatment with CHX followed by 2 h of DBMIB treatment, we did not see an effect of CHX on *APX2* induction (Fig. S2B). As a positive control for the CHX treatment, we generated a transgenic line, A2P, in which *LUCIFERASE (LUC)* gene expression is under the control of the *APX2* gene promoter [*A2P_{APX2(-1507)}*] (Fig. 2A). After 2 h of EL treatment, we observed about a 15-fold change in relative luciferase units (RLU) (Fig. S2C) in the absence of CHX, whereas in the presence of CHX, there was no change in RLU, indicating that the CHX treatment effectively

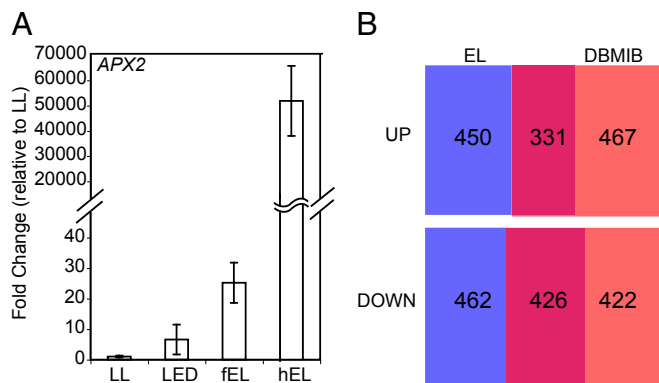


Fig. 1. EL and DBMIB change expression levels of nuclear genes including *APX2*. (A) *APX2* transcript abundance under LL, LED, fEL, and hEL. Plants were exposed to the EL treatments for 1 h. Light qualities have a total of ~1,000 μ mol photons $m^{-2} \cdot s^{-1}$. hEL was filtered using a 2-cm height column of water and ice mixture in a clear perspex tray to generate fEL. For LED treatments, only the EL exposed tissue was harvested. Average values of fold-change relative to LL were from three biological replicates. Error bars represent SD. (B) The number of genes up- and down-regulated by EL and DBMIB treatment. The areas are proportional to the number of genes in each category.

inhibited synthesis of the LUC protein (Fig. S2C). Therefore, we conclude that EL or DBMIB induced *APX2* expression in the absence of new protein synthesis.

Heat-Shock Transcription Elements Are Necessary and Sufficient for *APX2* Gene Induction by DBMIB. Previously reported genetic screens failed to identify positive regulators of *APX2* gene expression (15, 16), suggesting that these regulators may be redundantly encoded in the *Arabidopsis* genome. In addition, the rapid kinetics of the response, its independence of new protein synthesis, and the preponderance of HSEs predicted in the promoters of up-regulated genes was reminiscent of the response to heat shock in many organisms. To remove any effect of heat on *APX2* expression (29) and to investigate a heat-independent *APX2* induction mechanism, we decided to use DBMIB under LL in a temperature-controlled growth chamber. Under this condition, we were able to observe strong *APX2* induction (Fig. S2B).

To identify *cis*-acting elements responsible for *APX2* induction by DBMIB, we generated a series of transcriptional fusion constructs made with various lengths of the *APX2* promoter region fused to the *LUC* gene (Fig. 2A). Almost all transgenic lines containing the A2P construct, which encompasses the 1,507 bp upstream from the *APX2* start codon (Fig. 2A), showed an increase in LUC activity after DBMIB treatment. Several lines had greater than 10-fold LUC induction by DBMIB (Fig. 2B). The A2P3 construct, containing 346 bp upstream of the *APX2* start site (Fig. 2A), also showed strong induction by DBMIB, although the levels of LUC activity were not as high as those from the A2P construct (Fig. 2B). *LUC* in the A2P1 and A2P2 constructs, which contained 203 bp and 281 bp upstream of the *APX2* ATG, respectively (Fig. 2A), did not respond to DBMIB treatment (Fig. 2B), indicating that the 65-bp region from -282 to -346 was necessary for DBMIB-induced *APX2* induction (Fig. 2A). Deletion of this region (-282 to -346) in the A2P4 construct resulted in transgenic plants (49/50 T1 lines) with little to no LUC activity (Fig. 2A), indicating that the 65-bp region from -282 to -346 is necessary for DBMIB-induced *APX2* induction (Fig. 2B).

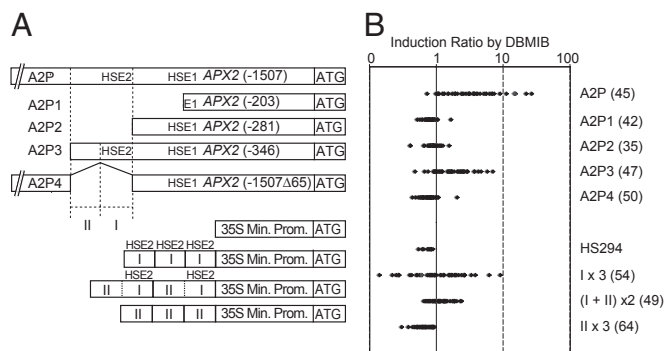


Fig. 2. HSE2 is responsible for *APX2* induction by DBMIB. (A) Constructs generated to identify *cis*-acting elements responsible for DBMIB-driven *APX2* induction. "ATG" is the translational start site for the *LUC* gene, and bars located in front of ATG represent the *APX2* promoter regions. The numbers in parentheses following construct names are the size of the *APX2* promoter regions from the translation start site of *APX2*. HSE1 and HSE2 are inverted repeat of HSE dimmers, and the numbers "1" and "2" are added for this study. The Roman numerals "I" and "II" were arbitrarily assigned to represent the portion with (I) and without (II) HSE2 in the segment from -282 to -346 . "35S Min Prom" stands for the 35S minimal promoter in the vector of pH5294. (B) LUC induction ratio by DBMIB in transgenic plants transformed by constructs in A. Each dot represents an induction ratio (after/before DBMIB treatment) after 2 h of DBMIB treatment in a single transgenic plant in its T1 generation. The numbers of T1 lines tested per construct are in parentheses.

We scanned the interval between -282 and -346 of the *APX2* promoter for known *cis*-acting elements, and found two HSEs in tail-to-tail arrangement (tTTCTgGAAg: HSE2 in Fig. 2A). These same elements have been reported to be responsible for heat-induced *APX2* expression in a tobacco leaf protoplast transient expression assay (29), and the elements are known to be the minimally required sequence for active HSF binding (30). There are also HSEs in head-to-head arrangement, another functional type (30), adjacent to the start codon (aGAAgTTCa: HSE1 in Fig. 2A); however, HSE1 does not appear to be necessary for *APX2* induction by DBMIB (Fig. 2B).

To investigate whether the 65-bp interval containing HSE2 is sufficient for DBMIB-driven *APX2* expression, we generated transgenic plants (HS294), in which *LUC* expression is under the control of the 35S cauliflower mosaic virus (35S) minimal promoter (Fig. 2A). None of the HS294-containing plants showed an increase in LUC activity after exposure to DBMIB (Fig. 2B). However, when two copies of the 65-bp interval were placed in front of the 35S minimal promoter and LUC fusion [(I + II) \times 2] (Fig. 2A), 63% (31 of 49) of T1 transgenic plants showed LUC induction by DBMIB (Fig. 2B). We further subdivided the 65-bp region into two fragments: I (29 bp) and II (36 bp) (Fig. 2A). When three copies of fragment I encompassing HSE2 (I \times 3) were attached to the 35S minimal promoter and translationally fused to *LUC* (Fig. 2A), about half (26 of 54) of the transgenic plants (T1) showed DBMIB-induced LUC induction, and the highest induction ratios in I \times 3 were greater than those of two copies of the 65-bp interval [(I+II) \times 2] (Fig. 2B). Three copies of fragment II (II \times 3), lacking HSE2 (Fig. 2A), failed to show any DBMIB-induced LUC expression in the 64 T1 plants tested (Fig. 2B). Taking these data together, we conclude that a 29-bp element containing HSE2 is both necessary and sufficient to induce gene expression after short DBMIB treatments (Fig. 2B).

Dynamics of *APX2* Induction by HSFs Under Stress Conditions. The inverted repeats of HSEs, such as HSE2, are bound by HSFs (30), thus making HSFs the primary candidates mediating *APX2* up-regulation in response to EL. In *Arabidopsis*, there are 21 HSFs (31): 15 type A, 5 type B, and 1 type C, characterized by their domain architectures. Because only type A HSFs contain transcriptional activation domains (31), and because it was reported that overexpression of *HSA2* and *HSA3* could induce *APX2* expression in the absence of stress (32–34), we focused on type A HSFs as possible regulators of *APX2* expression.

Constructs expressing each individual type A HSF under the control of the 2 \times 35S promoter were introduced into wild-type plants. We collected leaf tissue from T1 plants, and monitored *APX2* transcript levels using real-time RT-PCR (Fig. S3A). Besides *HSA2* and *HSA3* overexpression, *HSA1B*, *HSA1D*, *HSA4A*, and *HSA6B* overexpression induced *APX2* (fold-change >10) compared with transgenic control plants harboring only the binary vector or wild-type Col-0 (Fig. S3A). Because knockout mutant lines of *HSA1D*, *HSA2*, and *HSA3* were available in the same genetic background from the SALK T-DNA collection, we focused on these three HSFs. The overexpression of *HSA1D*, *HSA2*, and *HSA3* were confirmed in the next generation (Fig. S3B). There was a hierarchy of *APX2* expressions levels with *HSA1D* $>$ *HSA2* $>$ *HSA3* (Fig. 3A).

We checked whether *HSA1D*, *HSA2*, and *HSA3* genes were up-regulated in EL. *HSA1D* mRNA was detectable in LL, but its levels remained low after exposure to EL for 1 h (Fig. 3B and Fig. S3C); moreover, between 1- and 2-h treatment, no further increase was observed (Fig. 3B). mRNA levels of both *HSA2* and *HSA3* were low in LL, but dramatically increased in response to EL (Fig. 3B and Fig. S3C). A time-course of *HSA2* mRNA levels in EL revealed an induction pattern that is very similar to that of *APX2* (Fig. 3B, and Figs. S2A and S3C). In contrast, *HSA3* mRNA levels increased continuously during 2 h of EL treatment (Fig. 3B and Fig. S3C). The EL-driven induction patterns of *HSA1D*, *HSA2*, and *HSA3* were similar

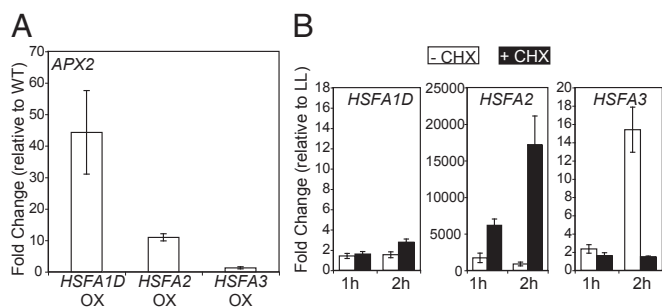


Fig. 3. Overexpression of *HSFA1D*, *HSFA2*, and *HSFA3* could induce *APX2* expression under LL. (A) Fold-changes of *APX2* under LL in *HSFA1D*-, *HSFA2*-, and *HSFA3*-overexpression lines, relative to wild-type Col-0 plants, were determined. The average and SEs were calculated from real-time RT-PCR of two or three independent T2 lines and three technical replicates. (B) Fold-changes of *HSFA1D*, *HSFA2*, and *HSFA3* by the indicated time of EL were determined using real-time RT-PCR. Fold-changes, relative to untreated LL samples, in the presence of 200 μ M CHX and in mock-treated (ethanol) samples were displayed using filled and blank bars, respectively. Two biological and four technical replicates were tested to calculate the average and SEs.

to those elicited by a combined treatment of moderate light stress and heat stress (32).

The similarity of induction kinetics in EL between *APX2* and *HSFA2* prompted us to test whether EL-driven *HSFA2* induction required new protein synthesis. We determined the induction levels of *HSFA1D*, *HSFA2*, and *HSFA3* by EL in the presence of CHX and compared induction levels to LL samples (Fig. 3B). Both *HSFA1D* and *HSFA2* were induced in the presence of CHX; they accumulated to even higher levels in CHX. In contrast, the induction of *HSFA3* was completely abolished by the CHX treatment (Fig. 3B). Thus, the induction of *HSFA1D* and *HSFA2* by EL does not require new protein synthesis, whereas new protein synthesis is required for *HSFA3* induction by EL.

We obtained homozygous T-DNA insertion lines from the SALK collection for *HSFA1D* (SALK_022404), *HSFA2* (SALK_008978), and *HSFA3* (SALK_011107). After backcrossing at least twice, we confirmed the absence of *HSFA1D*, *HSFA2*, and *HSFA3* transcripts in the T-DNA insertion mutants by semiquantitative RT-PCR (Fig. S3D). The mutants were used to test whether *HSFA1D* is required for EL-driven *HSFA2* and *HSFA3* induction, and whether *HSFA2* or both *HSFA1D* and *HSFA2* are essential for *HSFA3* induction by EL (Fig. S3 E–G). In an *hsfa1d*-null mutant, *HSFA2* and *HSFA3* mRNA accumulated in a similar manner to wild-type. Induction of *HSFA3* by EL was even faster in *hsfa1d* than wild-type (Fig. S3E). Similarly, we observed no difference from wild-type in EL-driven *HSFA3* induction levels in either the *hsfa2* or *hsfa1d hsfa2* double mutant (Fig. S3 F and G).

Homozygous null-mutants were exposed to EL and the levels of *APX2* expression were quantified. Although both *hsfa1d* and *hsfa2* mutant plants exhibited a reduction of *APX2* transcript accumulation in response to EL, (Fig. 4A, Left), there was still a significant increase in *APX2* RNA after EL treatment. Moreover, after 1 h of EL, *APX2* induction in an *hsfa3*-null mutant was indistinguishable from wild-type.

To better understand the individual contributions of *HSFA1D*, *HSFA2*, and *HSFA3* to *APX2* expression, we exposed single- and higher-order mutants of *hsfa1d*, *hsfa2*, and *hsfa3* to three environmental stresses: EL, heat (40 °C), or DBMIB (Fig. 4A). In EL, the *hsfa1d hsfa2 hsfa3* triple-mutant had drastically lower *APX2* induction levels compared with all other genotypes (Fig. 4A). The role of *HSFA1D*, *HSFA2*, and *HSFA3* in EL-driven *APX2* expression was also confirmed by the near elimination of LUC activity in the triple-mutant compared with wild-type after 2 h of EL treatment (Fig. S3H).

In EL, the contribution of *HSFA1D* for *APX2* expression is greater than *HSFA2*, which is greater than *HSFA3* (Fig. 4A, Left). Although overall induction levels caused by heat were lower than those caused by EL, *HSFA1D* was also the major contributor to heat-induced *APX2* expression (Fig. 4A, Center). Under DBMIB treatment, *HSFA1D* again played the major role in *APX2* expression and the contribution of *HSFA2* and *HSFA3* for DBMIB-induced *APX2* expression was marginal (Fig. 4A, Right). In summary, *HSFA1D* is the major regulator of *APX2* induction by multiple stresses, whereas the relative contribution of *HSFA2* and *HSFA3* to *APX2* gene expression varied with the stress.

The contributions of the three HSFs to growth and development were assessed by comparing the triple-mutant to wild-type Col-0 in LL vs. EL (Fig. S4). Under a 12-h LL/12-h dark cycle, 3-wk-old *hsfa1d hsfa2 hsfa3* plants were smaller than wild-type (Fig. S4A, LL) and flowered early (Fig. S4B). After 2 d of EL 12-h/Dark 12-h, we observed that emergent leaves of *hsfa1d hsfa2 hsfa3* were yellow; however, over time, older leaves appeared to recover (Fig. S4A, EL).

Posttranslational Regulation of *HSFA1D* by EL. *HSFA1D* transcription did not appear to be regulated by EL. Thus, we tested whether this protein is regulated posttranscriptionally in response to EL. The subcellular localization of *HSFA1D* under LL and EL was determined using transgenic lines expressing a *HSFA1D*-YFP-HA fusion protein in the $A2P_{APX2(-1507)/hsfa1d hsfa2 hsfa3}$ background. We isolated 12 transgenic lines harboring the *HSFA1D*-YFP-HA construct (Fig. S5A). Each line had various amounts of the fusion protein, and the basal LUC activities under LL were relatively proportional to the amount of the fusion protein (Fig. S5A and B). After exposure to 1 h of EL, all transgenic lines showed increased *APX2* expression as measured by LUC activities (EL1h/LL) (Fig. S5C) but untransformed *hsfa1d hsfa2 hsfa3* plants did not. These data indicate that the *HSFA1D*-YFP-HA fusion protein is biologically functional for EL-driven *APX2* induction, and that *HSFA1D* could functionally replace *HSFA2* and *HSFA3* in *APX2* induction under EL.

In transgenic line 24, in which basal LUC activity was similar to that in wild-type, *HSFA1D*-YFP-HA localized mainly in the cytoplasm (Fig. 5A–C). In contrast, in transgenic line 14, which overexpressed the fusion protein and had the highest basal LUC activity in LL (Fig. S5A and B), *HSFA1D*-YFP-HA was

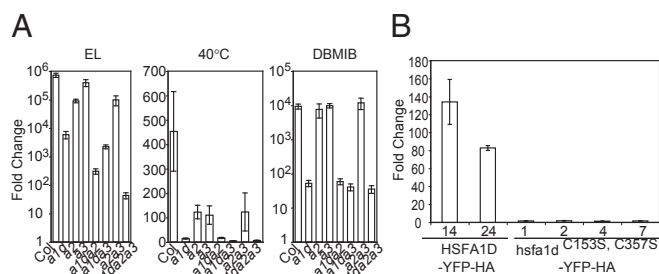


Fig. 4. *APX2* induction by EL is mediated by HSFs and their posttranslational modification. (A) Fold-changes of the *APX2* gene by EL, heat shock (40 °C), or DBMIB in the single- (*a1d*, *a2*, *a3*), double- (*a1da2*, *a1da3*, and *a2a3*), and triple- (*a1da2a3*) knockout mutants of *HSFA1D*, *HSFA2*, and *HSFA3*. Real-time RT-PCR was used to determine the fold changes relative to before the treatment. Leaf tissue was treated with EL for 1 h and with heat shock or DBMIB for 2 h. Error bars indicate SE of two biological and two technical replicates. The y axes of EL and DBMIB are on log₁₀ scales. (B) Real-time RT-PCR determined fold changes of *APX2* by 1 h of EL exposure in transgenic plants expressing *HSFA1D*-YFP-HA and *hsfa1d*^{C153S, C357S}-YFP-HA under *HSFA1D* native promoter. Both constructs were introduced to *hsfa1d hsfa2 hsfa3*. Two biological and two technical replicates were used to calculate the average and SEs.

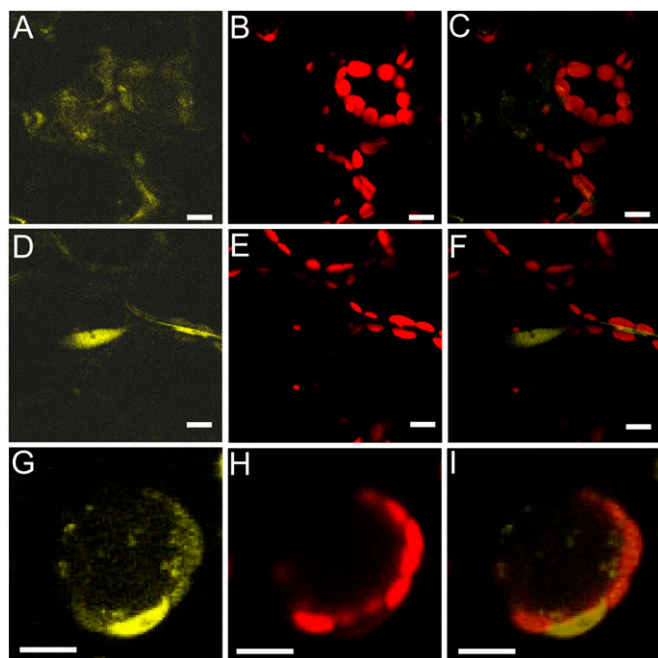


Fig. 5. Translocation of HSF1D from cytoplasm to the nucleus under EL. (A–C) Cytoplasmic localization of yellow fluorescence signals from the HSF1D-YFP-HA fusion proteins in a LL condition. (D–F) After 30 min of EL treatment, the fluorescence signals from HSF1D-YFP-HA were detected in the nucleus (D–F), and the fluorescence signals from *hsf1d*^{C153S, C357S}-YFP-HA (G–I) localized in cytoplasm and in the nucleus, with much higher signals in the nucleus (G–I). The merged images (C, F, and I) were generated by overlaying yellow fluorescence signal images (A, D, and G) and chlorophyll fluorescence images (B, E, and H) together. (Scale bars, 10 μ m.)

localized both in the cytoplasm and the nucleus (Fig. S5D). After 30 min of EL treatment, signals were observed exclusively in the nucleus in both lines (Fig. 5 D–F and Fig. S5E). Taken together, these data show that nuclear accumulation of HSF1D is correlated with EL-driven *APX2* expression. In addition to nuclear localization, activation of HSF1D might be required to fully induce *APX2* gene expression under EL, because LUC activity increased disproportionately in line 14 (Fig. S5C).

In mammals, disulfide bond formation between three HSF monomers is important for its activity as a transcriptional regulator, presumably because of stabilization of the trimer (35, 36). To understand whether this function is conserved in *Arabidopsis*, we tested the effects of disulfide bonding in EL-driven *APX2* expression. HSF1D contains two cysteine residues at amino acid positions 153 and 357. These two cysteine residues were mutated to serines and then a construct expressing the mutant protein, *hsf1d*^{C153S, C357S}-YFP-HA (under the control of the *HSF1D* promoter) was introduced to *hsf1d hsf2 hsf3*. Four transgenic lines were isolated and their expression levels were determined. Protein levels of *hsf1d*^{C153S, C357S}-YFP-HA were comparable to those of HSF1D-YFP-HA lines 14 and 24 (Fig. S5F); however, *APX2* induction levels by EL in *hsf1d*^{C153S, C357S}-YFP-HA lines were significantly lower than HSF1D-YFP-HA (Fig. 4B and Fig. S5G), indicating that disulfide bonding is critical for transcriptional activity of HSF1D for EL-driven *APX2* expression. Of note, *Arabidopsis hsf1d*^{C153S, C357S}-YFP-HA was still able to localize to the nucleus (Fig. 5 G–I), in contrast to the cysteine-to-serine mutant form of mammalian HSFs, which remain in the cytoplasm (35). Taken together, these data show that nuclear translocation of HSF1D is coupled to its transcriptional activation in EL. These two processes then control the degree of *APX2* induction.

Discussion

Our studies demonstrate a role for a subset of HSFs in rapid changes in nuclear gene expression in response to a redox-generated plastid stress signal. We have shown a role for three *Arabidopsis* HSFs—HSF1D, HSF2 and HSF3—regulating *APX2* expression in response to a change in redox generated by either EL or DBMIB, although HSF6B may also have a minor role. HSF1D plays a dominant role in the rapid response of plants to EL stress. In EL, preexisting HSF1D accumulates in the nucleus, where it likely becomes activated by changes in disulfide bonding. Although no new protein synthesis is required for deployment of this rapid response, the response is likely to be sustained through the transcriptional activation of genes for *HSFs* and other transcription factors.

Multiple intracellular signaling pathways have been shown to play a role in the genomic response of plants to stress. Remarkably, these stress signals are almost always generated in chloroplasts, a highly active metabolic compartment where oxygen is generated during photosynthesis. Recent reports have identified several metabolic intermediates or degradation products whose accumulation signals changes in nuclear gene expression during EL stress, including: β -cyclocitral, a breakdown product of the carotenoid β -carotene (6); MEcPP, a precursor of isoprenoids (7); and PAP (5). Although all of these stresses were created after EL treatments of plants, it is not clear at this time how to compare conditions from laboratory to laboratory. For example, nuclear gene expression in response to β -cyclocitral accumulation shares about 80% common genes with genes whose expression is altered by the $^1\text{O}_2$ signaling pathway (conditions in which *APX2* may play no role). Similarly, signaling by MEcPP may be condition-specific. Contrary to the results of Xiao et al. (7), our global expression studies indicated that *HPL1* gene expression is down-regulated (Dataset S1), rather than up-regulated. No obvious candidates exist for identification of the activator of *APX2* gene expression in response to nuclear accumulation of PAP except for ZAT10, which is also up-regulated in PAP overaccumulators, such as *altered expression of apx2 8* (5). In the case of PAP, nuclear localization of the phosphonucleotide can regulate the activity of exoribonuclease, thereby directly altering RNA pools; however, this is likely to indirectly affect *APX2* and as yet it is unclear whether this pathway intersects with the HSF response.

Although there appears to be a number of signals that initiate EL stress signaling pathways, the transcription factors that rapidly regulate expression of hundreds of nuclear genes are mostly unknown. Previous studies reported that during heat stress, HSF2 binds to the HSE2 element involved in *APX2* induction by EL (29). In addition to heat, we showed that *APX2* was induced by the reduced PQ pool in the absence of heat stress. *APX2* induction by LED (Fig. 1A), fEL (Fig. 1A), and LL plus DBMIB (Figs. S1B and S2B) reflects that there are at least two pathways for *APX2* expression: heat-dependent and heat-independent pathways.

It is also possible that these conditions generate a shared secondary signal that is transduced to the nucleus to regulate *APX2* gene expression. A good candidate for this molecule is H_2O_2 , which is generated during both heat and EL stress conditions (8, 37) (Fig. S1 C and E). However, the relationship between reduced PQ pool and H_2O_2 generation needs additional investigation because we could not—nor would we expect to—observe significant differences in the amount of H_2O_2 by our DBMIB treatment (Fig. S1C), although the DBMIB treatment could induce *APX2* expression (Figs. S1B and S2B).

A model of the signaling network would predict that in conditions of fluctuating light intensity or prolonged high-intensity light, the PQ pool becomes reduced, and generates unidentified signaling molecules in chloroplasts (Fig. S6). The extent to which the different chloroplast signals (PAP, β -cyclocitral, MEcPP) function in this HSF-mediated *APX2* expression in parallel or independent pathways remains to be investigated. Indeed, given that there are substantial differences in the relative induction of

APX2 under different experimental conditions (Fig. 1A), it is possible that multiple pathways function in its regulation. The unidentified molecule in this PQ-mediated pathway possibly facilitates HSF1D movement to the nucleus. Disulfide bond formation among HSF1Ds activates the HSF transcription complex to regulate genes such as *APX2* and *HSFA2*. *HSFA2* together with HSF1D further increases *APX2* transcription. If the EL condition is prolonged, newly synthesized unknown protein factors may induce expression of *HSFA3* and other transcription factors, which in turn play a role in induction or repression of hundreds of nuclear genes involved in growth and metabolism. This transcription program may then protect the plant from numerous environmental stresses.

In conclusion, the identification of the gene networks involved in plants' responses to stress is an area of considerable interest to plant biotechnologists. Our nascent studies identifying elements within the common set of EL and DBMIB up-regulated promoters indicate a convergence of plant responses to light and other abiotic stresses, including cold, dehydration, and heat at the level of transcriptional regulation, and may help to decipher this complex genetic program.

Materials and Methods

See [Dataset S3](#) for a list of primers used in this study.

Plant Growth Conditions and Stress Treatments. *Arabidopsis* plants were grown on a Linsmaier and Skoog medium (Caisson Laboratories) in growth chambers, or SALK Institute greenhouse number 3 (22 °C) for 3 to ~4 wk

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