Overexpression of the Trehalase Gene AtTRE1 Leads to Increased Drought Stress Tolerance in Arabidopsis and Is Involved in Abscisic Acid-Induced Stomatal Closure1

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Introduction of microbial trehalose biosynthesis enzymes has been reported to enhance abiotic stress resistance in plants but also resulted in undesirable traits. Here, we present an approach for engineering drought stress tolerance by modifying the endogenous trehalase activity in Arabidopsis (Arabidopsis thaliana). AtTRE1 encodes the Arabidopsis trehalase, the only enzyme known in this species to specifically hydrolyze trehalose into glucose. AtTRE1-overexpressing and Attre1 mutant lines were constructed and tested for their performance in drought stress assays. AtTRE1-overexpressing plants had decreased trehalose levels and recovered better after drought stress, whereas Attre1 mutants had elevated trehalose contents and exhibited a drought-susceptible phenotype. Leaf detachment assays showed that Attre1 mutants lose water faster than wild-type plants, whereas AtTRE1-overexpressing plants have a better water-retaining capacity. In vitro studies revealed that abscisic acid-mediated closure of stomata is impaired in Attre1 lines, whereas the AtTRE1 overexpressors are more sensitive toward abscisic acid-dependent stomatal closure. This observation is further supported by the altered leaf temperatures seen in trehalase-modified plantlets during in vivo drought stress studies. Our results show that overexpression of plant trehalase improves drought stress tolerance in Arabidopsis and that trehalase plays a role in the regulation of stomatal closure in the plant drought stress response.

Trehalose is a nonreducing sugar linking two Glc units in an α,α-1,1 configuration and is implicated in osmo-regulation and stress protection in many bacteria and fungi. The biosynthesis of trehalose in plants involves two consecutive enzymatic reactions mediated by trehalose-6-phosphate synthase (TPS) and trehalose-6-phosphate phosphatase (TPP), with trehalose-6-phosphate (T6P) as an intermediate compound (Cabib and Leloir, 1958). The synthesized trehalose can be hydrolyzed into two Glc monomers by the enzyme trehalase. Most genomes of higher plants contain elaborate trehalose biosynthesis gene families (Lunn, 2007; Avonce et al., 2010). The genome of Arabidopsis (Arabidopsis thaliana) contains 11 TPS genes (AtTPS1–AtTPS11) and 10 TPP genes (AtTPPA–AtTPP11; Leyman et al., 2001). The Arabidopsis TPS proteins carry a TPS- and a TPP-like domain and are divided into two classes based on their homology with the yeast (Saccharomyces cerevisiae) TPS (ScTps1) and TPP (ScTps2) proteins. Class I displays the highest similarity to ScTps1 and includes four proteins (AtTPS1–AtTPS4), but only AtTPS1 has demonstrated TPS activity (Blázquez et al., 1998; Van Dijck et al., 2002; Vandesteene et al., 2010). The class II proteins (AtTPS5–AtTPS11) are more similar to ScTps2, but they do not show any detectable TPS or TPP activity upon expression in yeast (Vogel et al., 2001; Ramon et al., 2009). All TPP proteins (AtTPPA–AtTPP11) contain specific phosphatase boxes (Thaller et al., 1998) but do not share homology to class I and II TPP proteins. Expression studies in yeast revealed that they all have TPP activity, and promoter-GUS-GFP lines showed that each one has a tissue-specific expression pattern in Arabidopsis (Vandesteene et al., 2012). In contrast to the large TPS and TPP gene families in Arabidopsis, trehalase is encoded by a single gene, AtTRE1 (Müller et al., 2001a; Lunn, 2007). Among the

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vascular plants, only a few desiccation-tolerant resurrection plants, such as Selaginella lepidophylla and Myrothamnus flabellifolius, accumulate substantial amounts of trehalose (Zentella et al., 1999). In most other higher plants, only trace amounts of trehalose are detected, barely exceeding 0.15 mg g\(^{-1}\) dry weight, which implies that there is insufficient trehalose for it to act as a compatible solute during environmental stress (Vogel et al., 2001; Fernandez et al., 2010).

The intermediate of trehalose biosynthesis, T6P, has emerged as an essential sugar-signaling metabolite that regulates plant metabolism and influences many aspects of plant growth and development (Schluepmann et al., 2004; Lunn et al., 2006). T6P promotes biosynthetic processes in growing tissues in response to high Suc (Delatte et al., 2011b) and induces starch synthesis in the plastids via thioredoxin-mediated activation of AGPase (Kolbe et al., 2005). In developing tissues of Arabidopsis, T6P inhibits the activity of the SNF1-related kinase1 (SnRK1; Zhang et al., 2009), which is known to repress plant growth and to promote survival under stress (Baena-González et al., 2007). T6P, SnRK1, and the sugar-regulated transcription factor bZIP11 have been postulated to be part of a regulatory network that regulates plant metabolism and in plants hosting trehalose-producing microorganisms, such as arbuscular mycorrhizal fungi, rhizobia, and the clubroot disease-inducing pathogen *Plasmodiophora brassicae* (Müller et al., 1994; Schubert and Wyss, 1995; Brodmann et al., 2002). Recently, an *Attre1* knockout line was characterized (SALK_147073C; Alonso et al., 2003) and reported to have a 2-fold increase in T6P levels when grown on sorbitol (Delatte et al., 2011a).

Many attempts have been made to enhance the stress tolerance of model plants and crops by introducing *TPS* and *TPP* genes of yeast or bacterial origin. In general, the stress tolerance of these plants increased, although trehalose levels remained low. In most cases, the plants exhibited aberrant phenotypes such as stunted roots, lance-shaped leaves, and growth retardation (Romero et al., 1997; Cortina and Culianez-Macia, 2005). The undesired abnormalities in these plants were ascribed to altered levels of T6P, perturbing the developmental processes influenced by this important signal metabolite (Eastmond et al., 2003; Schluepmann et al., 2003, 2004). Plants with improved stress tolerance but no obvious morphological defects were obtained by introducing bifunctional *TPS-TPP* constructs (Garg et al., 2002; Karim et al., 2007; Miranda et al., 2007), by placing the introduced *TPS* genes under the control of drought-inducible or tissue-specific promoters (Garg et al., 2002; Karim et al., 2007), or by overexpressing the plant endogenous *TPS1* in Arabidopsis (Avonce et al., 2004) and rice (*Oryza sativa*; Li et al., 2011). It seems unlikely that the increased drought tolerance in these transgenic lines could be ascribed to trehalose acting as an osmoprotectant, since the trehalose levels never exceeded 1 mg g\(^{-1}\) fresh weight (Garg et al., 2002; Avonce et al., 2004; Cortina and Culianez-Macia, 2005). The improved tolerance seemed instead to correlate with a higher soluble carbohydrate content and an increased photosynthetic capacity (Garg et al., 2002).

The activity of the endogenous trehalase is readily detectable in plants, and treatment with validamycin A, a trehalose analog and competitive inhibitor of trehalase, led to a 20-fold increase in trehalose in leaves of Arabidopsis plants grown on 10 mM trehalose (Müller et al., 2001a). Together, these observations indicate that the endogenous trehalase activity is high enough to restrict the accumulation of trehalose, even in transgenic plants engineered to overproduce this sugar, and that genetic down-regulation of trehalase could be a good strategy for increasing the trehalose content (Vinocur and Altman, 2005). In this work, we present a novel approach for engineering drought tolerance in Arabidopsis plants by manipulating the expression and activity of the endogenous trehalase *AtTRE1*.

**RESULTS**

**Isolation and Characterization of Arabidopsis Plants with Altered *AtTRE1* Expression**

The *AtTRE1* gene, encoding the only specific trehalase in Arabidopsis, was overexpressed in Arabidopsis plants of the Columbia-0 (Col-0) accession, and multiple homozygous 35S:*AtTRE1* lines (2.3, 3.4, 4.1, 5.2, 6.5, 9.1, and 11.1) were obtained. Seedlings of the 9.1...
and 3.4 lines show 292- and 139-fold increases in AtTRE1 expression, respectively, and were selected for further experiments based on their high trehalase transcript levels (Table I). A detailed expression study in rosette leaves of the 3.4 line revealed 52- to 178-fold increases in AtTRE1 transcripts (Supplemental Table S1). Lines from publicly available collections with transfer DNA (T-DNA) or transposon insertions in the AtTRE1 gene were screened to obtain plants that were homozygous for the insertion (Supplemental Fig. S1). One knockout line was found with no AtTRE1 transcripts detectable by quantitative reverse transcription-PCR (Table I): Attre1-1 (transposon insertion line GT_16843 from the Genetrap collection in the Landsberg erecta [Ler] accession; Sundaresan et al., 1995). The T-DNA insertion line Attre1-2 (SALK_147073C from the SALK collection in the Col-0 accession; Alonso et al., 2003) seemed to be a knockdown, since AtTRE1 transcription was decreased by a factor of 6 in seedlings (Table I) and by a factor of 7 to 14 in young rosette leaves, while it was hardly detectable in the oldest leaves (Supplemental Table S1). The Attre1-3OE line (SALK_151791 in Col-0) contains an insertion 304 bp upstream of the transcription start site and was unexpectedly found to be an overexpressing line, with AtTRE1 transcript levels that were 8 times increased in seedlings (Table I) and 3 to 11 times increased in the rosettes (Supplemental Table S1) compared with the wild type. The AtTRE1 (approximately 64-kD) protein was weakly detectable in wild-type Ler and Col-0 seedlings by immunoblotting with an anti-AtTRE1 antibody (Fig. 1A). At the same protein loading, a much stronger signal was observed in extracts from wild-type flowers and siliques, but no AtTRE1 protein was detected in wild-type rosettes (Fig. 1A). The absence of the AtTRE1 protein in wild-type rosettes is in conflict with previous immunoblots showing a faint signal of AtTRE1 in rosette leaves of bolted wild-type plants (Frison et al., 2007). The antibody used in that work was raised against two peptides of the AtTRE1 protein (residues 93–107 and 377–392), while our antibody was raised against one peptide (residues 94–107). It might thus be possible that our antibody, which recognizes a single epitope of AtTRE1 in close proximity to the transmembrane span, is less efficient in reaching its target than the antibody used by Frison et al. (2007).

There was no detectable AtTRE1 protein in any of the tissues examined from the Attre1-1 knockout, while AtTRE1 protein accumulation in the Attre1-2 knockdown was limited to a hardly visible signal in the seedlings (Fig. 1A). The 35S:AtTRE1 lines 3.4 (Fig. 1A) and 9.1 (data not shown) had high levels of AtTRE1 protein in all three tissues, indicating constitutive overexpression throughout the plant. The Attre1-3OE line showed an AtTRE1 accumulation pattern that was qualitatively similar to wild-type plants but with considerably higher levels of the AtTRE1 protein in those tissues where it was expressed. This suggests that the spatiotemporal control of AtTRE1 expression in this line is still under the control of the endogenous promoter but that the level of expression is enhanced by the presence of the T-DNA insertion within the putative promoter region. A further point to note from the immunoblot analysis is the presence of multiple immunoreactive bands in the extracts of the AtTRE1-overexpressing lines (Fig. 1A).

Trehalase activity was higher in wild-type Ler seedlings than in wild-type Col-0 plantlets and was decreased by 31% in wild-type Ler rosettes, while staying below the background of the assay in rosette leaves of the wild-type Col-0 (Table I). No trehalase activity was detected in the wild-type Col-0 (Table I). Trehalase activity in the Attre1-1 and Attre1-2 mutants (Table I). Trehalase activity in the Attre1-3OE line was 6.5-fold increased in seedlings and detectable in rosettes, albeit at low levels (Table I). Seedlings of the 35S:AtTRE1 lines displayed 109- to 119-fold higher trehalase activities, and this magnitude of activity was also seen in the rosettes (Table I). Plants engineered to constitutively express the Escherichia coli cytosolic trehalase (35S:treF) had 227-fold higher trehalase activity than wild-type Col-0 (Table I).

### Table I. Characterization of the trehalase-modified lines

<table>
<thead>
<tr>
<th>Genotype (Tissue)</th>
<th>Relative AtTRE1 Expression (Seedlings)</th>
<th>Trehalase Activity</th>
<th>Trehalase (Seedlings)</th>
<th>Trehalase (Rosettes)</th>
<th>T6P (Seedlings)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Seedlings</td>
<td>Rosettes</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>nkat mg⁻¹ protein</td>
<td>nkat mg⁻¹ protein</td>
<td>nmol g⁻¹ fresh wt</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild-type Ler</td>
<td>1.01 ± 0.2</td>
<td>0.13 ± 0</td>
<td>0.09 ± 0</td>
<td>26.4 ± 10.7</td>
<td>0.148 ± 0.03</td>
</tr>
<tr>
<td>Attre1-1</td>
<td>0.00****</td>
<td>0.00****</td>
<td>0.00 ± 0**</td>
<td>101.5 ± 14.5***</td>
<td>0.200 ± 0.03</td>
</tr>
<tr>
<td>Wild-type Col-0</td>
<td>1.04 ± 0.4</td>
<td>0.10 ± 0</td>
<td>0.00</td>
<td>22.6 ± 3.8</td>
<td>0.143 ± 0.03</td>
</tr>
<tr>
<td>Attre1-2</td>
<td>0.18 ± 0**</td>
<td>0.00***</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>35S:treF</td>
<td>9.1</td>
<td>11.07 ± 1.1******</td>
<td>11.23 ± 3.4***</td>
<td>5.5 ± 2.6***</td>
<td>0.115 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>3.4</td>
<td>12.12 ± 0.8******</td>
<td>10.71 ± 2.0***</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>35S:treF</td>
<td>23.09 ± 2.0******</td>
<td>ND</td>
<td>ND</td>
<td>0.094 ± 0</td>
</tr>
</tbody>
</table>

Relative AtTRE1 expression, trehalase-specific activity, trehalase, and T6P in seedlings and rosettes of the trehalase-modified lines are significantly different from the wild type (Ler and Col-0) as indicated: *P < 0.075, **P < 0.05, ***P < 0.01, ****P < 0.001, *****P < 0.0001 (Student’s t test). Values shown are means ± SD (n = 3; each replicate represents a pool of at least 12 seedlings or 10 rosettes). NA, Not applicable; ND, not determined.
Growth of the transgenic and wild-type seedlings was evaluated on one-half-strength Murashige and Skoog (0.5× MS) medium in the presence or absence of 25 mM trehalose. In the presence of this concentration of trehalose, the wild-type seedlings had stunted root growth and failed to develop rosette leaves (Fig. 1B), confirming the previous observations of Ramon et al. (2007). An even more pronounced growth inhibition was observed in \textit{Attre1-1} and \textit{Attre1-2} seedlings, while all of the \textit{AtTRE1} overexpressors were highly tolerant to the externally supplied trehalose. The roots and rosette leaves of the \textit{35S::AtTRE1} lines 9.1 and 3.4 were noticeably larger than those of the \textit{Attre1-3OE} seedlings, suggesting a positive correlation between enhanced growth on trehalose-containing medium and trehalase activity. However, the \textit{35S::treF} line, having the highest trehalase activity of all the lines tested, grew less well than the \textit{35S::AtTRE1} seedlings. The apoplastic \textit{AtTRE1} enzyme presumably has direct access to the external trehalose in the medium and so will readily hydrolyze it to Glc. With the extracellular hydrolysis of trehalose, plants might also avoid taking up trehalose into the cells, where it has the potential to perturb their metabolism and signaling processes. In contrast, the cytosolic \textit{treF} trehalase will only be able to hydrolyze trehalose that has been taken up into the cell. The intracellular trehalose can still perturb trehalose-sensitive processes, potentially counteracting the positive effects of the increased Glc supply.

Trehalose levels were determined in the different mutants. The \textit{Attre1-1} seedlings accumulated up to 101.5 nmol trehalose g\(^{-1}\) fresh weight, which was almost four times more than the levels found in wild-type \textit{Ler} (26.4 nmol g\(^{-1}\) fresh weight; Table I). Trehalose contents in seedlings of the strong \textit{35S::AtTRE1} line 9.1 and \textit{Attre1-3OE} ranged from 5.5 to 9.3 nmol trehalose g\(^{-1}\) fresh weight and were both significantly lower than the 22.6 nmol trehalose g\(^{-1}\) fresh weight observed in wild-type \textit{Col-0}. The \textit{Attre1-1} seedlings accumulated 0.20 nmol T6P g\(^{-1}\) fresh weight, which was somewhat higher than the 0.15 nmol T6P g\(^{-1}\) fresh weight seen in wild-type \textit{Ler} (Table I). The trehalase-overexpressing lines showed a tendency to have slightly lower levels of T6P than wild-type Col-0.

**Figure 1.** Characterization of \textit{AtTRE1}-modified lines. A, Immunoblot (IB) of \textit{AtTRE1} accumulation (approximately 64 kD) in protein extracts from seedlings, rosettes, flowers, and siliques and the corresponding Coomassie Brilliant Blue (CBB) stain showing the Rubisco large subunit as a constitutive control. B, Growth of trehalase mutants and wild-type (Wt) seedlings on 0.5× MS medium with and without 25 mM trehalose.
seedingls, but the differences were not statistically significant.

**ATRE1-Overexpressing Lines Are More Tolerant to Severe Drought Stress**

To test the effect of *ATRE1* on drought stress tolerance, our transgenic lines were tested in a severe drought stress experiment on soil-grown plants, as described in “Materials and Methods.” When the plants were 2 weeks old, water was gradually restricted in the same manner for all plants over a period of 20 d, and then plant were subsequently rewatered (Supplemental Fig. S2). After a period of 48 h, photographs and fresh weights were taken to examine the recovery of the plants, which is the ability to survive and rehydrate upon rewatering. Dry weights were assayed to study the growth performance of the plants during the drought stress period. As previously shown in the work of Meyre et al. (2001), we observed a difference in drought stress tolerance between the accessions *Ler* and Col-0. Col-0 plants acclimate to drought by increasing their root-to-shoot ratio and increasing their water use efficiency, while *Ler* plants exhibit more of a drought-escape strategy by flowering early and reallocating nutrients from the rosette to the flowers, which leads to a rapid yellowing of rosette leaves.

In the Col-0 background, two wild-type plants could survive out of five individuals, while four of the *Attre1-3OE* plants and three of the 35S::*ATRE1*-overexpressing plants survived out of five (Fig. 2A; Table II). The 35S::*treF* and *Attre1-2* plants were hypersensitive toward drought stress, since none of the individuals survived the drought treatment. The aboveground organs (rosette and inflorescence) of the surviving *ATRE1* overexpressors had almost twice the fresh weight of the surviving wild-type Col-0 control plants (Table II), which reflects the enhanced ability of the *ATRE1* overexpressors to recover after drought stress. Rosette dry weights of the *Attre1-2* mutant were decreased, while inflorescence dry weights were increased, suggesting that this mutant invested more energy in flowering than wild-type Col-0 plants (Table II). No differences in dry weights were observed between the wild-type Col-0 and the *ATRE1* overexpressors *Attre1-3OE* and 3.4. The rosette dry weights of the overexpressor 9.1 were higher than the wild-type Col-0, indicating that this 35S::*ATRE1* line has an increased capacity to grow during dry conditions. Interestingly, the phenotypes in rosette growth of the *Attre1-2* mutant and the *ATRE1* overexpressors are reversed under well-watered conditions (data not shown). The 35S::*treF* line grew worse than the wild-type Col-0 during drought (Table II) and nonstressed conditions (data not shown).

All plants of the *Ler* accession survived the drought stress treatment; rosette leaves were mostly senescent, while the inflorescences were fully developed (Fig. 2A). Dry weights of the rosettes (and flower stem) were slightly increased in the *Attre1-1* mutant compared with wild-type *Ler*, although not at a significant level (Table II). This tendency is more pronounced under well-watered conditions (data not shown). *Attre1-1* plants formed significantly fewer leaves during the recovery period, most of which were cauline leaves on secondary inflorescences (Table II). Interestingly, the *Attre1-1* line displayed decreased survival rates compared with wild-type *Ler* when slowing down the drying rate of the soil (data not shown).

**Trehalase Is Strongly Expressed in the Stomata**

To investigate the spatiotemporal expression patterns of the *AtTRE1* gene, promoter *TRE1-GUS-GFP*
and to a lesser extent in the developing seeds (Fig. 3J). was due to GUS expression in the placenta and funiculus (Fig. 3, H and I). The abundant GUS signal in the carpels 0.01 (Student’s test; n = 5). NA, Not applicable (see text); ND, not determined. The number of surviving plants out of five individuals, fresh weights of the surviving plants, dry weights of all the plants, and the number of newly appeared leaves in Ler plants are listed. Values are averages ± SD (n = 2–5). Significant differences are as indicated: *P < 0.075, **P < 0.05, ***P < 0.01 (Student’s t test; n = 5). NA, Not applicable (see text); ND, not determined.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Surviving Plants</th>
<th>Rosette Fresh Weight</th>
<th>Rosette Dry Weight</th>
<th>Flower Stem Fresh Weight</th>
<th>Flower Stem Dry Weight</th>
<th>Newly Appeared Leaves</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type Ler</td>
<td>5</td>
<td>NA</td>
<td>0.016 ± 0</td>
<td>0.110 ± 0.01</td>
<td>ND</td>
<td>4.2 ± 1.1</td>
</tr>
<tr>
<td>Attre1-1</td>
<td>5</td>
<td>NA</td>
<td>0.019 ± 0.01</td>
<td>0.136 ± 0.04</td>
<td>ND</td>
<td>1.6 ± 0.5***</td>
</tr>
<tr>
<td>Wild-type Col-0</td>
<td>2</td>
<td>0.110 ± 0.01</td>
<td>0.055 ± 0.01</td>
<td>0.030 ± 0.01</td>
<td>0.033 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>Attre1-2</td>
<td>0</td>
<td>NA</td>
<td>0.044 ± 0.01*</td>
<td>NA</td>
<td>0.049 ± 0.01***</td>
<td></td>
</tr>
<tr>
<td>Attre1-3OE</td>
<td>4</td>
<td>0.183 ± 0.07</td>
<td>0.050 ± 0.01</td>
<td>0.055 ± 0.03</td>
<td>0.038 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>9.1</td>
<td>3</td>
<td>0.193 ± 0.04</td>
<td>0.077 ± 0.01**</td>
<td>0.083 ± 0.01</td>
<td>0.027 ± 0.01</td>
<td></td>
</tr>
<tr>
<td>3.4</td>
<td>3</td>
<td>0.211 ± 0.01</td>
<td>0.051 ± 0.02</td>
<td>0.093 ± 0.06</td>
<td>0.031 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>35S::treF</td>
<td>0</td>
<td>NA</td>
<td>0.038 ± 0.01***</td>
<td>NA</td>
<td>0.029 ± 0.01</td>
<td></td>
</tr>
</tbody>
</table>

lines were constructed and subjected to GUS staining assays. During germination, GUS expression was mainly detected in the radicle at the early stages of testa rupture and root protrusion but shifted toward the hypocotyl and cotyledons with the onset of hypocotyl stretching (Fig. 3A). In 5-d-old seedlings, GUS staining was seen in the hypocotyl and shoot apical meristem and was very concentrated in the cotyledons, particularly in the stomatal guard cells (Fig. 3B). Very remarkable here was the sharp boundary between the basal part of the hypocotyl, with a diffuse GUS signal, and the apical part of the hypocotyl, with GUS expression limited to stomata. This observation might be explained by the fact that, at close proximity to the root, stomatal complexes are organized in a pattern reminiscent of the root epidermis pattern, which is clearly different from the epidermally specialized stomatal structures in the upper part of the plant (Berger et al., 1998). By the time the seedlings reached 10 d old, no GUS activity was apparent in the hypocotyl or cotyledons, but GUS remained strongly expressed in the cotyledons, but GUS expression was limited to stomata. This observation might be related with any differences in stomatal responses upon treatment of detached leaves with 20 μM ABA in the light. In the absence of ABA, the Attre1-1 leaves had significantly larger stomatal apertures than wild-type Ler, but the Attre1-2 knockdown did not show any difference from the corresponding wild-type Col-0 controls (Fig. 4A). However, both the knockout and knockdown showed striking insensitivity to ABA, with no difference in stomatal aperture after 2 h of exposure to ABA, in marked contrast to the wild-type Ler and Col-0 plants, whose stomatal apertures were 43% and 20% lower, respectively, after ABA treatment. The Attre1-3OE and 35S::Attre1 9.1 overexpressing lines showed the opposite behavior to the knockout and knockdown, with stomatal apertures smaller than the wild type in the absence of ABA and decreasing even further than the wild type after ABA treatment. The other trehalase-overexpressing lines, 35S::Attre1 3.4 and 35S::treF, had similar stomatal apertures to wild-type Col-0 in the absence of ABA but showed more pronounced stomatal closure than the control plants after ABA treatment. These results suggest that increased trehalase activity enhances the sensitivity of the guard cells to exogenous ABA and in some cases Attre1-modified lines limit their transpirational water losses in response to water stress by excising the rosette leaves and measuring water loss over time. Water loss was significantly greater in the Attre1-1 and Attre1-2 mutants than in the corresponding wild-type Ler and Col-0 plants, whereas the Attre1-overexpressing lines Attre1-3OE and 9.1 showed lower water loss than the wild-type control (Fig. 2B). The 35S::treF line was indistinguishable from wild-type Col-0 up to about 90 min but had a slightly slower rate of water loss thereafter. These results support the idea that Attre1 regulates transpiration. ABA is a key regulator of stomatal closure, and several reports have found evidence of connections between trehalase metabolism and ABA signaling (Avonce et al., 2004; Gómez et al., 2010; Vandesteene et al., 2012). Therefore, we investigated whether the altered phenotype of the trehalase mutants under drought stress was correlated with any differences in stomatal responses with water restriction, followed by a recovery period of 48 h upon rewatering. ABA in the absence of ABA but showed more pronounced stomatal closure than the control plants after ABA treatment. These results suggest that increased trehalase activity enhances the sensitivity of the guard cells to exogenous ABA and in some cases...

Attre1 Affects Transpiration and Stomatal Movements in Vitro

Given the specific Attre1 expression in stomatal guard cells, we investigated to what extent the...
restricts stomatal opening even in the absence of external ABA. Moreover, AITRE1 appears to be essential for ABA-induced closure of guard cells, since both the Attre1-1 and Attre1-2 mutants were unable to close their stomata in response to ABA addition. Inhibition of seed germination is another classical response to ABA treatment. Interestingly, neither the AITRE1 mutants nor the treF overexpressor seemed to interfere with the ABA responsiveness in seeds, since seed germination in the presence of ABA was similarly affected in both the transgenic and wild-type lines (data not shown).

To investigate whether AITRE1 expression is induced by ABA, wild-type Col-0 seedlings grown in axenic liquid cultures were incubated for 60 min in the

Figure 3. Histochemical localization of GUS activity in pTRE1::GUS-GFP lines during germination at the stages of root protrusion and hypocotyl stretching (A), in 5-d-old seedlings (B), in 10-d-old seedlings (C and D), in 5-week-old bolted plants (E), in the mature, second real leaf from a 5-week-old plant (F), in flower buds (G), in the flower stem, in cauline leaves, and in flower buds on secondary inflorescences (H), in the inflorescence (I), and in a developing seed at early maturation stage (J).

Figure 4. Leaf stomatal features. A, Stomatal pore sizes without ABA (black bars) and with 20 μM ABA (white bars). Stomatal apertures of the trehalase-modified lines within each treatment are significantly different from the wild type (Wt Ler and Wt Col-0) at P < 0.01 (*) and P < 0.0001 (**) by Student’s t test. Values represent averages ± SE (n = 80). B, Relative AITRE1 expression in wild-type seedlings treated for 60 min with 10 μM ABA is significantly increased compared with the solvent control (0 μM ABA) at P < 0.05 (*) by Student’s t test. Values represent averages ± SD (n = 3; each replicate represents a pool of at least 20 seedlings). C, Stomatal index of the trehalase-modified lines is significantly different from the wild-type index (Wt Ler and Wt Col-0) at P < 0.05 (*), P < 0.001 (**), and P < 0.0001 (***) by Student’s t test on data following the arcsine transformation. Values are means ± SE (n = 60).
presence of 10 μM ABA (diluted from a stock solution in methanol) or an equivalent amount of methanol as a solvent control. AtTRE1 expression was almost 2-fold higher in the ABA-treated seedlings than the controls (Fig. 4B), suggesting that AtTRE1 expression is both induced by ABA and modulates downstream responses to ABA.

Altering Trehalase Activity Affects the Stomatal Index and the Leaf Temperature of Water-Stressed Plants

Stomatal conductance is a major factor determining the rate of water loss via transpiration and is dependent on both the number of stomata and the extent to which they are opened. The stomatal index of the trehalase overexpressors was determined to assess whether changes in the numbers of stomata could account for their enhanced capacity to recover after drought stress. The Attre1-3OE, 35S::AtTRE1 9.1, and 35S::treF plants had a 9% to 15% lower stomatal index than the corresponding wild-type Col-0 plants (Fig. 4C) and favor a decreased water loss through the stomata upon drought stress. The Attre1-2 knockout showed a small but significant increase of 7% in stomatal index compared with wild-type Col-0, whereas the Attre1-1 mutant had the same value as the wild-type Ler control. The reciprocal changes in stomatal index resulting from overexpression or loss of trehalase activity in the Col-0 background suggest that trehalose metabolism could influence epidermal cell division and differentiation into guard cells. This is supported by GUS staining observations using 3-week-old, plate-grown seedlings of the promoter TRE1::GUS-GFP lines, where AtTRE1 expression was clearly visible in developing stomatal guard cells (Supplemental Fig. S3).

Thermal imaging is a powerful in vivo technique for detecting and analyzing mutants with altered rates of transpiration due to differences in stomatal density and/or stomatal aperture. Plants with lower stomatal conductance have lower rates of transpiration, leading to higher leaf temperatures, and vice versa for seedlings with a higher stomatal conductance. Seedlings of the various lines were grown under well-watered conditions for 8 d and then subjected to drought for 4 d before analyzing their leaf surface temperature by infrared imaging (Fig. 5A). Leaves of the Attre1-1 knockout mutant were 0.44°C colder than wild-type Ler (Fig. 5B), which is entirely consistent with the increased stomatal aperture (Fig. 4A) and greater water loss (Fig. 2B) observed in detached leaves from the mutant. In contrast, drought-stressed Attre1-2 seedlings displayed slightly higher leaf temperatures than the corresponding wild-type Col-0 plants. Assuming that the transpiration rate is dependent on the leaf size (von Caemmerer and Farquhar, 1981), the leaf area of the Attre1-2 line showed an increase of 25% compared with the wild-type Col-0 and could be held responsible for the lower transpiration rate in this mutant. Moreover, infrared assays with adult plants suggested that severe rather than mild

Figure 5. Transpiration of trehalase-modified seedlings restricted in water for 4 d. A, False-color infrared images of the drought-stressed plantlets. B, Normalized leaf temperatures of the transgenic seedlings as calculated from the quantification of three infrared images (approximately 3,000 square pixels) as shown in A are significantly different from wild-type (Wt) Ler and Col-0 at P < 0.0001 (*, Z score). Values are weighted means ± SD.
drought stress might be necessary to overrule the effect of an increased growth rate in Attre1-2 plants, leading to a stomatal response similar to that of the Attre1-1 knockout (data not shown). All of the trehalase-overexpressing lines had higher leaf temperatures than wild-type Col-0, with the temperature difference ranging from 0.29°C to 0.50°C. This is consistent with the lower stomatal aperture and stomatal index observed in these plants (Fig. 4, A and C) as well as their increased ABA sensitivity, leading to lower rates of transpiration when the plants are drought stressed. Altogether, these data support the hypothesis that the trehalase plays a significant role in controlling stomatal conductance and that overexpression of either the endogenous trehalase (AtTRE1) or a heterologous form (E. coli treF) reduces water loss during drought stress in Arabidopsis plants.

DISCUSSION

Overexpression of AtTRE1 Improves Drought Tolerance in Arabidopsis

Trehalose functions as a stress protectant and osmoprotectant in a wide variety of organisms (Elbein et al., 2003). Engineering trehalose metabolism in plants by introducing microbial TPS and TPP genes has been reported to improve plant survival and growth under abiotic stress conditions (Garg et al., 2002; Avonce et al., 2004; Miranda et al., 2007). Although 4-fold increases in trehalose content were obtained in Arabidopsis and up to 10-fold increases in rice, the absolute amounts of trehalose remained rather low and seemed unlikely to be making much contribution to osmoregulation or stress protection. The limited accumulation of trehalose was ascribed to the high endogenous trehalase activity in the plants. T6P, the intermediate of the trehalose biosynthesis, is a recognized key signaling molecule involved in regulating many metabolic and developmental processes in plants (Schluepmann et al., 2004; Kolbe et al., 2005; Lunn et al., 2006; Zhang et al., 2009) and has been proposed to play a role in the acquisition of abiotic stress tolerance in insects and fungi (Mizoguchi et al., 1996; Cheong et al., 2003). Therefore, it has been suggested that the improved drought tolerance of plants expressing microbial forms of TPS and/or TPP could be linked to changes in the level of T6P, leading to downstream effects such as the accumulation of soluble carbohydrates (Garg et al., 2002; Schluepmann et al., 2003). Nevertheless, these proposals remain largely speculative, and we cannot rule out the possibility that trehalase has a signaling or regulatory function as well.

To address this possibility, we investigated the effects of altering trehalase activity in Arabidopsis plants. As expected, loss of the endogenous trehalase in the Attre1-1 mutant led to a significant increase in the amount of trehalose, whereas lines that overexpressed AtTRE1 had lower trehalose contents (Table I).

It is striking, however, that the trehalose accumulation in the Attre1-1 mutant was no more than 4-fold greater compared with wild-type plants. Similar results were obtained in Nicotiana tabacum plants transformed with a double gene construct to overexpress both a TPS and a TPP of microbial origin, which accumulated only 3-fold higher trehalose levels, even when the endogenous trehalase was inhibited by treatment with 1 mM validamycin A (Goddijn et al., 1997). These data indicate that endogenously synthesized trehalose might be broken down by another glycoside hydrolase, such as the broad-specificity acid α-glucosidase reported in some legume species (García et al., 2005). Alternatively, trehalose might be secreted out of the plant, perhaps from the roots into the soil, providing nutrients for beneficial bacteria and fungi (Fernandez et al., 2012). A further possibility is that trehalose exerts strong feedback inhibition on its own biosynthesis above a certain threshold.

There were no statistically significant changes in the levels of T6P when altering the trehalase activity in Arabidopsis plants (Table I), so any phenotypic differences seem most likely to be linked to the altered trehalose content. Curiously, decreasing the level of trehalose by overexpression of trehalase led to improved survival and growth of the plants under drought stress conditions, while the Attre1-1 knockout and Attre1-2 knockdown with elevated trehalose levels appeared to be more sensitive to drought stress. In one sense, these results were counterintuitive, given the commonly held view that trehalose functions as a compatible solute and stress protectant in many organisms. However, the low absolute concentrations of trehalose found in either wild-type or genetically engineered plants almost certainly rule out a role for this sugar as a quantitatively important compatible solute unless its distribution is highly localized in specific cell types. Unfortunately, currently available techniques for measuring trehalose are not sensitive enough to determine its distribution between different cells in plant tissues.

Since genetic modification of the trehalose biosynthesis pathway in plants led to differences in growth (Schluepmann et al., 2004; Lunn et al., 2006), it is important to consider whether the altered drought stress performances of the trehalase-modified lines were due to specific drought responses rather than differences in growth. This is surely the case for the AtTRE1 overexpressors, since the reduced growth seen in nonstress conditions (data not shown) was not reflected but was reversed during drought stress (Table II). The 35S::treF-overexpressing line grew slower than the wild-type Col-0 during normal (data not shown) and dry conditions (Table II) but was more sensitive toward drought. Although the Attre1-1 knockout grew better in well-watered conditions (data not shown), it is hard to conclude whether the slightly, but not significantly, increased growth rate affected the overall performance of this mutant during drought stress (Table II). The same indecisiveness applies for the Attre1-2 line. This knockdown displayed increased growth rates during
well-watered conditions (data not shown) and mild drought stress (seedlings; Fig. 5A) but invested less energy in rosette growth and more energy in flowering than the wild-type Col-0 during severe drought stress (adult plants; Table II).

To investigate the function of trehalase during drought stress, it was important to know where and when trehalase is expressed in Arabidopsis plants. Analysis of the pTRE1::GUS-GFP promoter-reporter lines revealed strong expression in sink organs such as young rosette leaves, flower buds, and ripening siliques (Fig. 3, E, F, and I). This expression pattern is similar to that of the Arabidopsis TPS1 gene (van Dijken et al., 2004), suggesting that there may be factors in common regulating the expression of these two genes. T6P is proposed to link the growth of developing tissues to the carbohydrate status of the plant via regulation of the SnRK1 protein kinase (Schluepmann et al., 2003; Lunn et al., 2006; Zhang et al., 2009). In such tissues, trehalose could be an unwanted, perhaps deleterious, by-product of T6P signaling, and so the presence of high trehalase activity would help to prevent its accumulation. The AtTRE1 gene is also highly expressed during seed germination, budding, flowering, and seed maturation (Fig. 3, A and G–J). Whatever the functions of AtTRE1 are during these developmental stages, they do not appear to be essential, because the Attre1-1 knockout and Attre1-2 knockdown germinate, flower, and set seed under benign controlled growth conditions. However, the marked phenotypes of the AtTRE1-overexpressing and knockout lines under drought stress conditions (Fig. 2A) strongly indicate a role for trehalase and trehalose metabolism in abiotic stress tolerance.

Trehalase Affects the Stomatal Index and ABA-Induced Stomatal Closure

The pTRE1::GUS-GFP promoter-reporter lines showed strong expression of AtTRE1 in stomatal guard cells (Fig. 3, D and F). Stomatal closure can be studied in wild-type plants by an ABA treatment of detached leaves, floated on a KCl-containing buffer in the light (Fig. 4A). Expression of the Arabidopsis AtTRE1 gene is essential for this response, as the stomata in leaves from the Attre1-1 knockout and Attre1-2 knockdown showed no effect of ABA treatment on stomatal aperture (Fig. 4A). In contrast, the stomata in leaves from the AtTRE1 overexpressors were found to be hypersensitive to ABA (Fig. 4A). The higher leaf temperature of drought-stressed AtTRE1 overexpressors is indicative of a lower stomatal conductance and transpiration rate and could be explained by the increased stomatal sensitivity to endogenous high ABA levels, which would be expected under abiotic stress conditions. Interestingly, the Attps1-12 mutant, lower in TPS activity and T6P content than wild-type plants, was also found to have a smaller stomatal aperture (Gómez et al., 2010). Although trehalose was not measured, the decreased rates of T6P synthesis in the mutant suggest that it probably has lower levels of trehalose, like the trehalase overexpressors, which also showed reduced stomatal aperture. Thus, potential changes in the level of trehalose might be a contributing factor to the stomatal phenotypes of the Attps1-12 mutant, the Attre1 mutants, and the AtTRE1 overexpressors. Taken together, these data indicate that AtTRE1 and trehalose metabolism fulfill an important role in the control of water loss through the stomata upon drought stress in Arabidopsis plants.

Potassium ions, together with both inorganic (e.g. chloride) and organic (e.g. malate) anions, account for the majority of osmotic changes associated with stomatal opening and closing, but other solutes can also be important under certain conditions. When the levels of light are high enough, Suc accumulation plays a dominant role in raising the osmotic potential of the guard cells driving stomatal opening. The precise mechanism for modulating sugar levels during stomatal movement is not fully understood, but it may involve the transport of Suc into and out of the guard cells as well as the synthesis, redistribution, and catabolism of Suc within the guard cells (Outlaw and Manchester, 1979; MacRobbie, 1998). At present, we have insufficient information to assess whether perturbing trehalase activity and trehalose metabolism in the guard cells affects all types of stomatal movements or if the effects are specifically linked to sugar-related mechanisms of stomatal opening and closing.

Loss of AtTRE1 expression in the Attre1-2 knockdown led to an increased stomatal index, while overexpression of trehalase in the same Col-0 background generally decreased the stomatal index (Fig. 4C). These differences in stomatal number probably contribute to changes in stomatal conductance in the mutants, which are indicated by the differential rates of water loss from detached leaves (Fig. 2B) and the differences in leaf surface temperature (Fig. 5A) shown by the various lines. Stomatal number is determined by the rates of epidermal cell division and mechanisms triggering differentiation into guard cells and suppression of differentiation in the surrounding epidermal cells. Histochemical assays with pTRE1::GUS-GFP seedlings showed an intense AtTRE1 expression in the developing stomatal guard cells, suggesting that trehalase could play a role in guard cell differentiation. TPS1, the only known enzyme that synthesizes T6P in Arabidopsis (Vandesteene et al., 2012), is reported to interact with the cell cycle-dependent kinase CDKA1 that likely acts in the stomatal pathway, with tubulin, and with the kinesin KCA1 (Bergmann and Sack, 2007; Geelen et al., 2007), implicating trehalose metabolism in control of the cell cycle. However, the physiological significance of this connection and the role of AtTRE1, if any, are unclear.

Arabidopsis Plants That Overexpress the E. coli treF Are More Sensitive to Drought Stress

Experiments with detached leaves showed that the Attre1-3treF and 35S::AtTRE1 9.1 plants, which overexpress
the endogenous apoplastic trehalase (AtTRE1), had smaller stomatal apertures than wild-type Col-0 leaves in the absence of ABA (Fig. 4C) and lower rates of water loss (Fig. 2B). In contrast, the 35S::treF plants, engineered to overexpress the heterologous cytosolic trehalase (treF) from E. coli, had essentially the same stomatal aperture as wild-type Col-0 plants in the absence of ABA and only slightly lower rates of water loss, despite having even higher trehalase activity. This suggests that the endogenous AtTRE1, a cytosolic enzyme, might be another factor contributing to the appearance of multiple AtTRE1 forms on the immunoblots. Phosphorylation of the AtTRE1 protein seems most likely to occur while the protein is being synthesized or in transit within the cell, as there appears to be little phosphorylation of proteins once they are secreted into the apoplast (Durek et al., 2010).

AtTRE1 Is Required for ABA-Induced Stomatal Closure

The degree of stomatal closure observed in the different AtTRE1 overexpressors does not seem to be dependent on the amount of AtTRE1 protein accumulated in the leaves. For example, the Attre1::3OE line had less AtTRE1 than the 35S::AtTRE1 lines but seemed highly sensitive toward ABA (Figs. 1A and 4A). This finding suggests that the endogenous AtTRE1 promoter in Attre1::3OE might be activated upon ABA detection. This was confirmed by the observed increase in AtTRE1 expression of wild-type seedlings upon ABA addition (Fig. 4B), which also agrees with results from microarray analyses of Arabidopsis plants treated with ABA (Zimmermann et al., 2004). Analysis of the AtTRE1 promoter region revealed a DNA-binding site for MYB4 (Supplemental Fig. S4), a transcription factor acting in response to environmental stresses (Chen et al., 2002). Moreover, a W-box promoter motif for MYB102 and WRKY transcription factors (Supplemental Fig. S4), known to be involved in ABA signaling upon dehydration and osmotic stress (Dennemann and Sheen, 2003; Ren et al., 2010), was also detected in the AtTRE1 promoter (ATHENA; O’Connor et al., 2005). Moreover, since MYB4 and MYB102 are both members from the R2R3-type MYB family, known to control the identity and fate of many plant cells (Stracke et al., 2001), these transcription factors could possibly induce AtTRE1 expression during developmental processes such as guard cell differentiation (Supplemental Fig. S3).

Interestingly, Arabidopsis microarrays also showed that AtTRE1 expression is light dependent, with a strong increase in AtTRE1 transcripts during extended night (Zimmermann et al., 2004), which indicates that AtTRE1 expression might be induced in the dark when stomatal closure is required. Frison et al. (2007) suggested that the additional higher molecular mass bands of AtTRE1 seen on immunoblots may represent posttranslationally modified forms of the Arabidopsis trehalase (Fig. 1A). To reach its apoplastic location, the AtTRE1 protein is likely to be transported via the endomembrane system and then secreted by vesicular exocytosis; therefore, it may be subject to glycosylation at one or more sites (Frison et al., 2007). Within the AtTRE1 protein, four residues (Ser-71, Thr-128, Ser-463, and Ser-530) and one hotspot (residues 191–213) are predicted with high confidence to be phosphorylated, and a further 38 residues have potential to be phosphorylated (Heazlewood et al., 2008). Moreover, Ser-195, which is part of the phosphorylation hotspot, falls within the RXXS/T motif, which is a characteristic target site for the SnRK2.2 and SnRK2.3 kinases (Furihata et al., 2006). This suggests that phosphorylation by this class of protein kinases might be another factor contributing to the appearance of multiple AtTRE1 forms on the immunoblots. Phosphorylation of the AtTRE1 protein seems most likely to occur while the protein is being synthesized or in transit within the cell, as there appears to be little phosphorylation of proteins once they are secreted into the apoplast (Durek et al., 2010).

CONCLUSION

The results presented here unequivocally demonstrate the importance of trehalase in stomatal function in Arabidopsis and provide a number of important leads for investigating the transcriptional and post-translational regulation of trehalase expression and activity in guard cells. Furthermore, we have shown that manipulating the expression of the endogenous plant trehalase can have a beneficial effect on the plant’s response to water deficiency and that this is a promising strategy to engineer improvements in drought stress tolerance in crop species.

MATERIALS AND METHODS

Plant Material

To obtain 35S::AtTRE1 overexpressors, the AtTRE1 coding sequence (Awg24040) with a 2.5-kb hemagglutinin-tagged sequence was amplified using TRE1-RgIII and TRe1-Smal primers (for primer sequences, see Supplemental Table S1). This PCR product was cloned as a Smal-Smal restriction fragment into a modified pCB302 minibinary expression vector (Hwang and Sheen, 2001). Arabidopsis (Arabidopsis thaliana) Col-0 plants were transformed by floral dip with Agrobacterium tumefaciens C58C1 carrying the expression vector. Seven independent homozygous lines were selected, which contained single insertions based on their 3:1 (resistant:susceptible) segregation patterns on glufosinate ammonium (50 mg L−1). The Escherichia coli treF gene was amplified with treF-F and treF-R primers (Supplemental Table S1). The obtained reaction product was inserted as an EcoRI restriction fragment into the pGreen binary vector (Hellens et al., 2000) between the cauliflower mosaic virus 35S promoter and terminator. A. tumefaciens-transformed Col-0 plants were selected by BASTA spraying. From the T2 plant 35S::treF 10.3, 138 out of 138 progeny were resistant to BASTA. The pTRE1::GUS-GFP lines were created by cloning a region of approximately 2 kb upstream of the AtTRE1 transcription start site plus the 5′-untranslated region with pTRE1-atTF primers in the pHGF55 vector (Karimi et al., 2007), using the Gateway technology (Invitrogen) according to the manufacturer’s instructions. A.
Attre1-2 protein (1:1000; GenScript) and as secondary antibody a horseradish peroxidase. The gene-specific primers TRE1-LP3 and TRE1-RP3. The sample loading control, and proteins from the other gel were analyzed by immunoblotting. For detection of the AtTRE1 protein, we used as primary antibody a horseradish peroxidase (Dialab). Protein concentrations were determined according to Bradford (1976). For each sample, aliquots containing 20 μg of protein were analyzed by SDS-PAGE on a horizontal 12.5 % polyacrylamide gel with a stacking gel of 4 % and a running gel of 12.5 % acrylamide. The gel was stained with Coomassie Brilliant Blue. The proteins were transferred to a nitrocellulose membrane and visualized with amido black. The membranes were incubated with primary antibodies, rinsed with TBS-T (20 mM Tris-HCl, pH 7.5, 0.5 % Tween-20) and then incubated with horseradish peroxidase-conjugated goat anti-rabbit or anti mouse antibodies (1:10 000; Santa Cruz) and visualized with enhanced chemiluminescence (Amersham). The resulting blots were scanned and analyzed by the ImageJ software. The stomatal index was determined from the same photographs taken and plant fresh and dry weights were assessed after a recovery period of 48 h.

Trehalase Activity
The following samples were harvested and frozen in liquid nitrogen: (1) at least 20 10-d-old seedlings on 0.5 % MS medium; (2) 10 4-week-old rosettes from soil-grown plants; and (3) a pool of flowers/silicages originating from at least 10 different 6-week-old plants grown in soil. The frozen tissues were ground to a fine powder with a mortar and pestle. Total RNA isolation, complementary DNA synthesis, and quantitative PCR were performed as described by Vandesteene et al. (2010). Primer sequences of AtTRE1 (TRE1-fw and TRE1-rv) and UBQ10 (UBQ10-fw and UBQ10-rv) are listed in Supplemental Table S2.

Immunoblot Analysis
The following samples were harvested and frozen in liquid nitrogen: (1) at least 20 10-d-old seedlings on 0.5 % MS medium; (2) 10 4-week-old rosettes from soil-grown plants; and (3) a pool of flowers/silicages originating from at least 10 different 6-week-old plants grown in soil. The frozen tissues were ground to a fine powder with a mortar and pestle. Total RNA isolation, complementary DNA synthesis, and quantitative PCR were performed as described by Vandesteene et al. (2010). Primer sequences of AtTRE1 (TRE1-fw and TRE1-rv) and UBQ10 (UBQ10-fw and UBQ10-rv) are listed in Supplemental Table S2.

Metabolite Measurements
Metabolite measurements were executed with three replicates of 12 seedlings each harvested 11 d after germination and grown at 22°C under continuous light (110 μE m⁻² s⁻¹). Plant material was immediately frozen in liquid nitrogen and homogenized with a mortar and pestle. Aliquots (15-20 mg) of the frozen tissue powder were extracted with chloroform-methanol as described by Lunn et al. (2006). Trehalose was quantified fluorometrically as described by Mollo et al. (2011). T6P and other phosphorylated compounds were determined by liquid chromatography coupled to liquid tandem mass spectrometry as described by Lunn et al. (2006).

Drought Assay
Drought tolerance tests were performed in individual pots filled with a mixture of soil (20 g) and vermiculite (8 g). For each transgenic line and wild-type control, five plants were grown for 2 weeks in controlled, well-watered conditions (relative soil water content [RWCsoil] of 80%). During the following 20 d, the amount of water was gradually and equally reduced in the pots until the symptoms of drought stress were strongly pronounced, which occurred at an RWCsoil of 22%. To achieve an identical drying-out process among the different plants, pot weights were monitored and synchronized on a daily basis. At the end of the drought stress period, plants were rewatered. Photographs were taken and plant fresh and dry weights were assessed after a recovery period of 48 h.

Histochemical and Histological Analysis
For GUS staining assays, in vitro-grown seedlings and bolted plants were sampled 2, 5, 10, and 35 d after germination, while flowers of soil-grown plants were harvested from 6-week-old plants. GUS staining was performed according to Beeckman and Engler (1994). For the staining of rosette leaves, vacuum infiltration was applied for 2 h. After the staining step, plant material was cleared in 85% lactic acid (Sigma-Aldrich) and mounted on glass microscope slides. Samples were examined with a M165C binocular microscope (Leica) and by differential interference contrast microscopy (BX51; Olympus).

Leaf Detachment, Stomatal Aperture, and Stomatal Counts
For stomatal aperture bioassays and leaf detachment studies, plants were grown in well-watered conditions. The first fully expanded leaves from 3-week-old plants were detached and used in both assays. For determining the stomatal aperture, leaves were floated in a stomata-opening buffer (10 mM MES-KOH, pH 6.15, 10 μM KCl) for 2 h under continuous light. To test the response to ABA, 20 and 0 μM (control) ABA was added to the buffer, and leaves were incubated 2 h more. Peels of abaxial epidermis were mounted on a microscope slide with double-sided sticky tape, moistened with a few drops of assay solution, and covered with a slip. The microscopy slides were examined within 30 min after preparation with a BX51 light microscope (Olympus). Photographs of epidermal sections were taken from three separate leaves and from 10 different areas per leaf. Stomatal apertures were measured using ImageJ software. The stomatal index was determined from the same photographs used for measuring the stomatal aperture. The stomatal index (% of a given leaf area was calculated as the ratio of the total number of stomatal guard cells to the total number of epidermal cells (sum of guard cells and pavement cells) multiplied by 100.

Thermal Imaging
Seeds were grown in individual pots filled with a mixture of soil (20 g) and vermiculite (8 g) under humid conditions (RWCsoil of 80%) for 8 d and
subsequently subjected to drought by not watering for 4 d (RWC_{soil} of approximately 70%). The 12-d-old plantlets were photographed in the same figures. We thank Gustavo Godesblat (Gernt University) for his expertise with stomata assays. We thank Nathalie Wuysts (Université Catholique de Louvain) for developing software to analyze infrared images (FlirImage/Processor for Image). Received November 21, 2012; accepted January 19, 2013; published January 22, 2013.

LITERATURE CITED


