

A Mitochondrial Complex I Defect Impairs Cold-Regulated Nuclear Gene Expression

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To study low-temperature signaling in plants, we previously screened for cold stress response mutants using bioluminescent *Arabidopsis* plants that express the firefly luciferase reporter gene driven by the stress-responsive *RD29A* promoter. Here, we report on the characterization and cloning of one mutant, *frostbite1* (*fro1*), which shows reduced luminescence induction by cold. *fro1* plants display reduced cold induction of stress-responsive genes such as *RD29A*, *KIN1*, *COR15A*, and *COR47*. *fro1* leaves have a reduced capacity for cold acclimation, appear water-soaked, leak electrolytes, and accumulate reactive oxygen species constitutively. *FRO1* was isolated through positional cloning and found to encode a protein with high similarity to the 18-kD Fe-S subunit of complex I (NADH dehydrogenase, EC 1.6.5.3) in the mitochondrial electron transfer chain. Confocal imaging shows that the *FRO1*:green fluorescent protein fusion protein is localized in mitochondria. These results suggest that cold induction of nuclear gene expression is modulated by mitochondrial function.

INTRODUCTION

Because of their sessile nature, plants frequently have to endure unfavorable environmental conditions. Consequently, plants have developed unique mechanisms to cope with environmental stresses such as chilling and freezing temperatures. Plants from temperate regions can acquire freezing tolerance after being exposed to low nonfreezing temperatures. This process is known as cold acclimation (Guy, 1990). Cold acclimation is associated with complex biochemical and physiological changes in plants, including changes in gene expression (Thomashow, 1994), leaf ultrastructure (Ristic and Ashworth, 1993), membrane lipid composition (Lynch and Steponkus, 1987; Miquel et al., 1993), enzyme activities, levels of sugars and polyamines (Levitt, 1980; Strand et al., 1997), and ion channel activities (Knight et al., 1996).

In *Arabidopsis*, a number of genes are induced during cold acclimation (Thomashow, 1999). These include *RD29A* (also known as *COR78* or *LT178*), *KIN1*, *KIN2* (*COR6.6*), *COR15A*, and *COR47* (*RD17*). The products of these genes are highly hydrophilic, but none of their functions (except those of *COR15A*) have been established. The constitutive expression of *COR15A* in *Arabidopsis* increased freezing tolerance at the chloroplast and protoplast levels (Artus et al., 1996). *COR15A* appears to function by decreasing the tendency of membranes to form the lamella-to-hexagonal II

phase, which leads to membrane damage during freezing (Steponkus et al., 1998).

DNA regulatory elements in the promoters of cold-responsive genes have been identified and named dehydration-responsive element (DRE) or C-repeat (CRT) (Baker et al., 1994; Yamaguchi-Shinozaki and Shinozaki, 1994). DRE/CRT binding proteins, *CBF1* (or *DREB1B*), *CBF2* (or *DREB1C*), and *CBF3* (or *DREB1A*) have been cloned and shown to function as transcriptional activators (Stockinger et al., 1997; Gilmour et al., 1998; Liu et al., 1998). *CBFs/DREB1s* are induced by low temperatures and are involved in the regulation of the DRE/CRT class of cold-responsive genes. Overexpression of *CBF1* resulted in the constitutive expression of the DRE/CRT class of genes and enhanced freezing tolerance (Jaglo-ottosen et al., 1998). Overexpression of a *CBF1* isolog, *DREB1A/CBF3*, also brought about expression of the DRE/CRT class of genes and increased drought tolerance as well as freezing tolerance (Liu et al., 1998). As *CBFs/DREB1s* are induced rapidly by low temperatures (Thomashow, 1999), Gilmour et al. (1998) proposed that an as yet unknown transcriptional activator may exist in an inactive form at warm temperatures but may become activated at low temperatures to turn on the transcription of *CBF/DREB1* genes.

Many plant adaptations to stresses involve mitochondria (Mackenzie and McIntosh, 1999). Mitochondria have been reported to be the main cellular organelles affected by low temperatures. Mitochondria at low temperatures exhibit changed respiration rate (Lyons and Raison, 1970), decreased cytochrome c oxidase activity (Prasad et al., 1994b), and enhanced alternative oxidase activity (Prasad et

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al., 1994b), leading to the generation of reactive oxygen species (ROS) such as superoxide and hydrogen peroxide and, consequently, oxidative stress (Rich et al., 1976; Huq and Palmer, 1978). For example, low temperatures induce oxidative stress in maize (Prasad et al., 1994a) and mitochondrial ROS accumulation (Prasad et al., 1995; De Santis et al., 1999; Gonzalez-Meler et al., 1999).

Although nuclear contributions to mitochondria have long been studied, mitochondrial effects on the nucleus have not received much attention until recently. Clearly, nuclear gene expression can be affected by the functional state of mitochondria (Parikh et al., 1987, 1989). This phenomenon is called retrograde regulation (Liao and Butow, 1993). In the yeast *Saccharomyces cerevisiae*, cells lacking mitochondrial DNA, and thus dysfunctional mitochondria, exhibit upregulation of the *CIT2* gene encoding peroxisomal citrate synthase (Liao et al., 1991). This altered nuclear gene expression caused by dysfunctional mitochondria involves two basic helix-loop-helix-leucine zipper transcription factors, Rtg1p and Rtg3p (Liao and Butow, 1993; Jia et al., 1997), and a member of the heat shock protein family, Rtg2p (Liao and Butow, 1993). In plants, our understanding of retrograde communications is limited by the lack of a suitable model system in which to study nuclear-mitochondrial interactions (Mackenzie and McIntosh, 1999).

Here, we report on the characterization and cloning of an Arabidopsis mutant, *frostbite1* (*fro1*), which has altered cold-responsive nuclear gene expression because of a defect in mitochondrial complex I. The mutant was recovered in a genetic screen based on its reduced reporter gene induction by cold stress. RNA gel blot analysis showed that *fro1* plants have lower levels of cold induction of stress-responsive genes such as *RD29A*, *KIN1*, *COR15A*, and *COR47*. *fro1* leaves appear water-soaked, which mimics wild-type leaves that have been subjected to freezing stress. The mutant leaves are constitutively leaky to ions and have reduced capacity for cold acclimation. The mutant leaves show constitutive accumulation of ROS. *FRO1* was isolated through map-based cloning. It encodes a protein with high similarity to the 18-kD Fe-S subunit of complex I (NADH dehydrogenase, EC 1.6.5.3) in the mitochondrial electron transfer chain. The *FRO1*:green fluorescent protein (GFP) fusion protein is localized in mitochondria. These results suggest that mitochondrial defects affect nuclear gene expression under low-temperature conditions, possibly through reactive oxygen messengers.

RESULTS

Identification of the *FRO1* Locus

To study stress signaling pathways in plants, we previously generated transgenic Arabidopsis plants expressing the firefly luciferase reporter gene under the control of the stress-

responsive *RD29A* promoter (Yamaguchi-Shinozaki and Shinozaki, 1993; Ishitani et al., 1997). The transgenic plants emit bioluminescence in response to low-temperature, abscisic acid, or NaCl treatment. Using these plants as the background line, mutants were isolated from an ethyl methanesulfonate-mutagenized M2 population based on altered luminescence responses under different stress conditions (Ishitani et al., 1997).

One mutant showing a lower level of luminescence under low-temperature treatment was chosen for further study. This mutant, in the *los_{cold}* category (Ishitani et al., 1997), was named *fro1* because of its translucent, water-soaked leaf phenotype, which is typically found in plants injured by chilling or freezing (Saltveit and Morris, 1990). *fro1* was backcrossed to parental *RD29A::LUC* plants (i.e., wild type). Luciferase imaging after cold treatment showed that the resulting F1 plants all behaved like the wild type (Table 1). In the selfed F2 generation, plants segregated for wild-type and mutant *RD29A::LUC* response phenotypes at an ~3:1 ratio (Table 1). These results indicate that *fro1* is caused by a single recessive nuclear mutation. *fro1* mutant plants were backcrossed to the wild type four times to remove possible unlinked mutations. All subsequent characterization was performed using the mutant that had been backcrossed.

fro1 Mutant Plants Are Defective in Cold-Regulated Gene Expression

A comparison of luminescence images from wild-type and *fro1* seedlings showed that the *fro1* mutant clearly had reduced *RD29A::LUC* expression under cold stress (Figure 1A). However, *RD29A::LUC* responses to either abscisic acid or NaCl were not substantially different between *fro1* and the wild type (Figures 1B and 1C). Quantification of the luminescence intensities revealed that cold-induced *RD29A::LUC* expression in *fro1* was only ~12% of that in the wild type, whereas abscisic acid- or NaCl-induced expression was not significantly different between the mutant and the wild type (Figure 1D).

Figure 2 shows the time course of *RD29A::LUC* expression in wild-type and *fro1* seedlings under stress treatments. Wild-type plants showed considerable *RD29A::LUC* expression in response to cold treatment for 12 h or longer. By contrast, *fro1* mutant plants did not show high levels of *RD29A::LUC* expression even after 48 or 72 h of cold treatment (Figure 2A). Abscisic acid or NaCl treatment for a few hours induced high levels of *RD29A::LUC* expression in both wild-type and *fro1* plants. The peak level of abscisic acid response was slightly higher in *fro1*, but the NaCl response was higher in the wild type (Figures 2B and 2C).

To determine whether endogenous *RD29A* gene induction also is altered in *fro1*, RNA gel blot hybridization was performed with total RNA extracted from wild-type and *fro1* mutant seedlings treated with or without cold, abscisic acid, or osmotic stress (Figure 3). Consistent with the expression

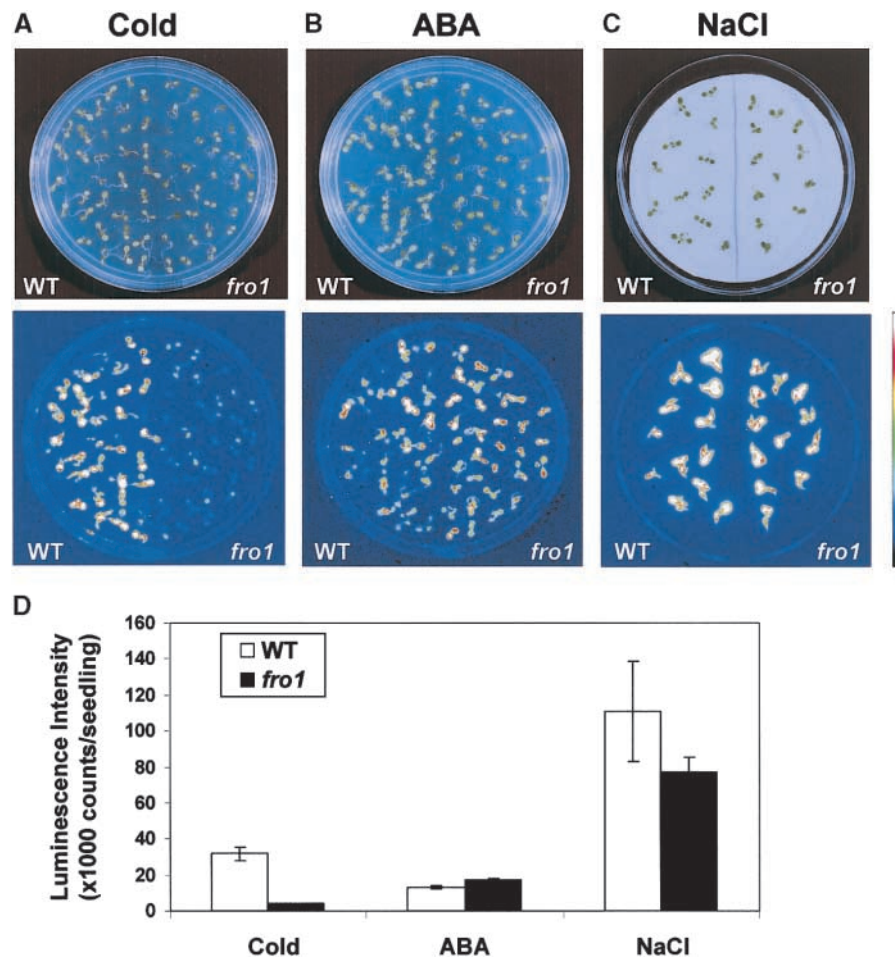
Table 1. Genetic Analysis of the *fro1* Mutant (Wild Type × *fro1*)^a

Generation	Seedlings Tested	Wild Type ^b	<i>fro1</i>	χ^2	P
F1	25	25	0		
F2	544	412	132	0.157	0.693

^aFemale × male.^bSegregation was scored by comparing luminescence intensities from each genotype on the same plate. Seedlings with intensities >5000 counts per seedling after cold stress were considered to be wild type.

of the *RD29A::LUC* transgene, lower cold induction of the endogenous *RD29A* gene was detected in *fro1* than in the wild type (Figures 3A and 3B). No significant reduction in *RD29A* expression was observed in *fro1* in response to either abscisic acid or NaCl treatment. To test if the *fro1* effect on cold regulation is specific to *RD29A*, the expression of three other cold-responsive genes, *COR15A*, *KIN1*, and *COR47*, was analyzed (Figure 3C). All three genes showed substantially reduced cold induction in *fro1*. By contrast, their expression in response to abscisic acid or NaCl was not lower in the mutant (Figure 3C).

We tested and found that the cold-induced expression of *CBF1*, *CBF2*, and *CBF3* was not lower in *fro1* (Figure 3A).

**Figure 1.** *RD29A::LUC* Luminescence Images of Wild-Type and *fro1* Seedlings.

(A) Morphology of seedlings on an agar plate and their luminescence images after treatment at 0°C for 72 h.

(B) Morphology of seedlings on an agar plate and their luminescence images after treatment with 100 μ M abscisic acid for 3 h.

(C) Morphology of seedlings on an agar plate and their luminescence images after treatment with 300 mM NaCl for 5 h.

The color scale bar at right shows the luminescence intensity from black (lowest) to white (highest).

(D) Luminescence intensities of wild-type and *fro1* seedlings after each treatment.

ABA, abscisic acid; WT, wild type.

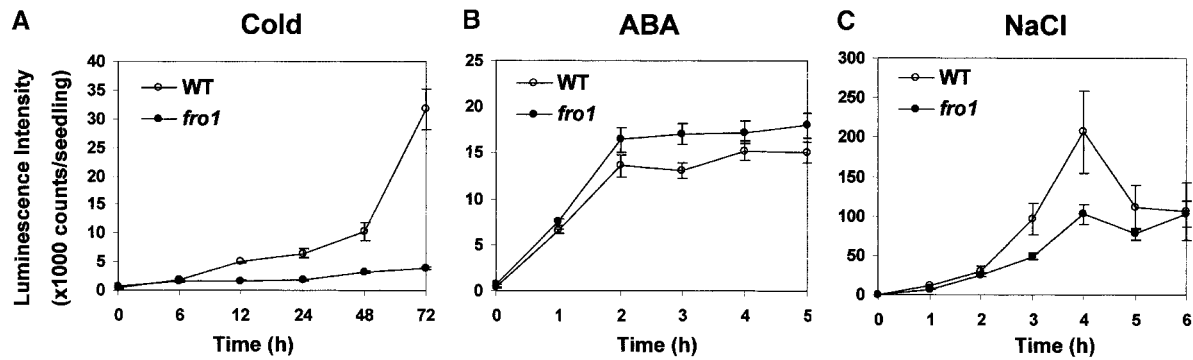


Figure 2. Time Courses of *RD29A::LUC* Expression in Wild-Type and *fro1* Seedlings in Response to Cold, Absciscic Acid, or NaCl.

(A) *RD29A::LUC* expression after low-temperature treatment at 0°C.

(B) *RD29A::LUC* expression after treatment with 100 μM abscisic acid.

(C) *RD29A::LUC* expression after treatment with 300 mM NaCl.

RD29A::LUC expression was quantified as luminescence intensity. ABA, abscisic acid; WT, wild type.

The *CBF* genes were induced early and peaked at 6 h in both the wild type and *fro1* (Figure 3A). In wild-type plants, a consistently reduced level of *CBF* induction was observed at 12 h, and the induction level recovered to some extent at 24 h. In *fro1*, *CBF* induction decreased only slightly at 12 h

(Figure 3A). At 48 h, *CBF* induction was very low in both the wild type and *fro1*. Cold induction of *RD29A*, *COR15A*, *COR47*, and *KIN1* occurred later than that of *CBF* genes, and the induction levels were lower in *fro1* throughout the time course (Figures 3A and 3C). The *fro1* mutation reduced

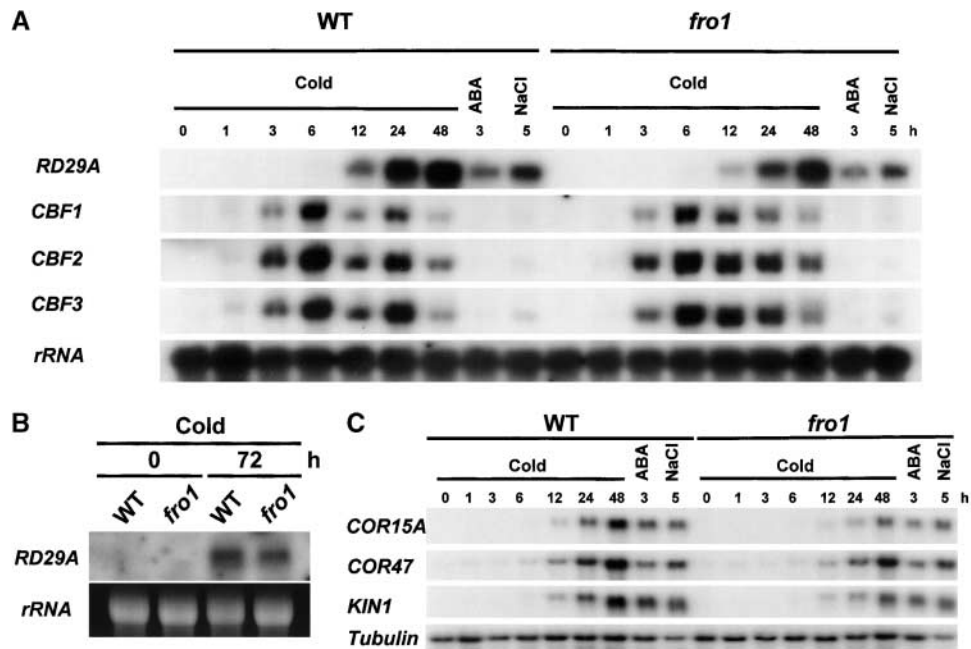


Figure 3. Gene Expression in Wild-Type and *fro1* Mutant Plants in Response to Stress Treatments.

(A) and (C) RNA gel blot hybridization with total RNA (20 μg) from wild-type and *fro1* mutant seedlings treated with low temperature (0°C) for the indicated times, abscisic acid (100 μM) for 3 h, or NaCl (300 mM) for 5 h. Gene probes used for RNA gel blot hybridization are indicated at left.

(B) RNA gel blot hybridization with total RNA (20 μg) from seedlings treated with low temperature (0°C) for either 0 or 72 h.

25S rRNA and tubulin were used as loading controls. ABA, abscisic acid; WT, wild type.

the cold induction of *COR15A* and *KIN1* more than that of *RD29A* and *COR47*. The decrease in cold induction of the endogenous *RD29A* (Figures 3A and 3B) does not seem to be as great as that of the *RD29A::LUC* transgene (Figure 2A). This is probably because the transgene had <700 bp of the *RD29A* promoter region (Ishitani et al., 1997). In comparison, the endogenous *RD29A* gene may contain more regulatory elements in its promoter, introns, or untranslated regions and thus may be subjected to more complex regulation.

***fro1* Mutant Leaves Are Translucent and Resemble Wild-Type Leaves That Have Been Subjected to Freezing**

Some of the well-known freezing injuries include water-soaked phenotypes in leaves and reduced plant growth (Saltveit and Morris, 1990). Some chilling-sensitive plants also appear water-soaked in response to chilling damage (Saltveit and Morris, 1990). Under normal growth conditions, leaves of *fro1* mutant plants appear water-soaked and translucent, with dark green color (Figure 4D). This dramatic water-soaked leaf appearance is similar to that of wild-type leaves that have been frozen (Figures 4A to 4C), although

there may not be a mechanistic connection, because the latter is a physical process brought about by ice formation. The translucent leaf phenotype of *fro1* was typically found in rosette leaves and sometimes in cauline leaves as well (data not shown).

A comparison of leaf cross-sections did not reveal gross structural differences between *fro1* and the wild type (Figure 5). However, less turgid and irregularly shaped cells were found frequently in *fro1* leaves (Figures 5B and 5D) compared with wild-type leaves (Figures 5A and 5C). Chloroplasts (Figure 5F) and mitochondria (Figure 5H) in *fro1* plants did not appear different from those of the wild type (Figures 5E and 5G). Interestingly, cell walls in *fro1* were substantially thinner than those in the wild type (Figures 5E to 5H).

***fro1* Leaves Are Constitutively Leaky to Cellular Electrolytes and Are Impaired in Cold Acclimation**

In wild-type leaves that have been frozen, the integrity of membranes is compromised. The electrolyte leakage test (Sukumara and Weiser, 1972; Ristic and Ashworth, 1993) is considered a good indicator of cell membrane integrity. Because the water-soaked appearance of *fro1* leaves resembles that of wild-type leaves that have been frozen, we

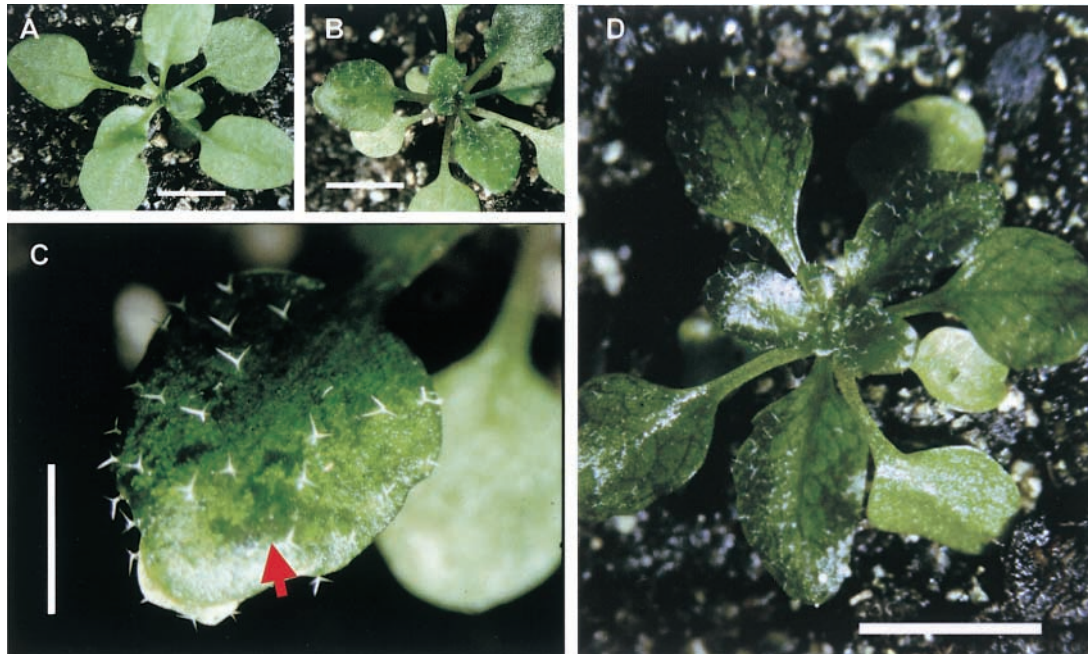


Figure 4. Water-Soaked Leaf Phenotype of *fro1* Mutant Plants.

- (A) Two-week-old wild-type plant grown at 22°C. Bar = 5 mm.
 (B) Two-week-old wild-type plant after 4 h of freezing treatment at –10°C followed by 24 h of incubation at 4°C. Bar = 5 mm.
 (C) Frozen wild-type leaf magnified from (B). The arrow indicates a water-soaked region of the leaf. Bar = 2 mm.
 (D) Three-week-old *fro1* mutant plant grown at 22°C. Bar = 5 mm.

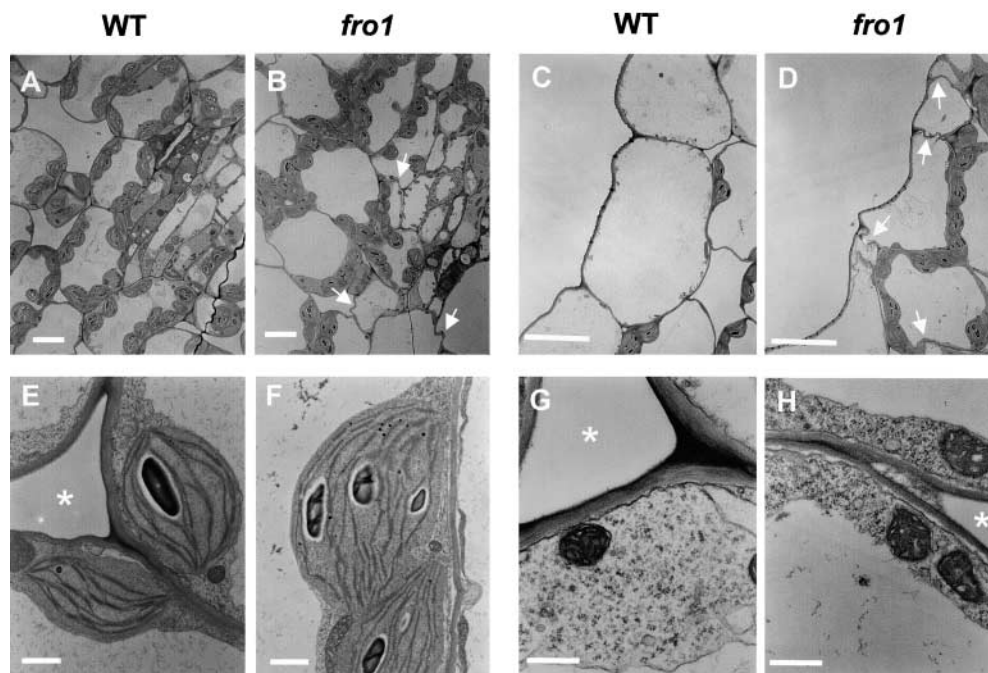


Figure 5. Comparison of Ultrastructure between Wild-Type and *fro1* Mutant Plants.

(A), (C), (E), and (G) Wild-type leaves. Chloroplasts (E) and mitochondria (G) are shown.

(B), (D), (F), and (H) *fro1* mutant leaves. Chloroplasts (F) and mitochondria (H) are shown.

White arrows indicate the irregular cell shape in *fro1*. Asterisks indicate intercellular spaces. WT, wild type. Bars = 20 μm for (A) and (B), 10 μm for (C) and (D), 1 μm for (E) and (F), and 0.5 μm for (G) and (H).

tested whether *fro1* leaves are more leaky to electrolytes. Even without any cold treatment, *fro1* mutant leaves showed a higher level of electrolyte leakage, as indicated by higher relative conductivity (Figure 6A). After 3 days of incubation at 4°C, electrolyte leakage in detached leaves of the wild type and *fro1* was measured. The results indicated that the extent of electrolyte leakage did not change much in either the wild type or *fro1*. After 1 day of exposure to -1°C, *fro1* leaves showed a large increase in electrolyte leakage, whereas wild-type leaves showed a relatively slight increase (Figure 6A). These results indicate that the membrane integrity of *fro1* is impaired and that *fro1* mutant seedlings are constitutively leaky and hypersensitive to freezing.

Electrolyte leakage at subzero temperatures was investigated with detached leaves from both cold-acclimated and nonacclimated plants (Figure 6B). Without cold acclimation, *fro1* leaves showed higher electrolyte leakage over the wild type at all temperatures tested. In nonacclimated plants, the LT_{50} values were -2.8 and -3.5°C for *fro1* and the wild type, respectively. After 2 days of cold acclimation at 4°C, the wild type showed a large increase in freezing tolerance. In comparison, *fro1* showed a much smaller increase in freezing tolerance under the same conditions (Figure 6B). The LT_{50} values for cold-acclimated *fro1* and wild-type

plants were -4.2 and -6.5°C, respectively. These results confirm that *fro1* is more sensitive to freezing and, additionally, is impaired in cold acclimation.

fro1 Shows Constitutive Accumulation of ROS

Prasad et al. (1994a) observed an accumulation of hydrogen peroxide when chilling-sensitive maize was exposed to cold conditions (either 14 or 4°C). ROS have been known to cause damage to cellular membranes (Kagan, 1988). It is possible that the membrane leakiness in *fro1* is a consequence of oxidative damage. Therefore, the presence of ROS in wild-type and *fro1* plants was tested with nitroblue tetrazolium (NBT) staining for superoxide and 3,3'-diaminobenzidine (DAB) staining for hydrogen peroxide. Without stress treatment, wild-type leaves did not show significant NBT staining, suggesting a lack of superoxide (Figure 7A). However, *fro1* leaves showed intense NBT staining without any stress treatment, indicating that a high level of superoxide was present in the mutant (Figure 7B).

After cold treatment, superoxide was detected in both the wild type and *fro1*, but the level in *fro1* was much higher than that in the wild type (Figures 7C and 7D). Similarly, DAB

staining showed that without cold treatment, *fro1* but not the wild type accumulated hydrogen peroxide (Figures 7E and 7F). After cold treatment, wild-type leaves showed slight DAB staining, whereas *fro1* exhibited very substantial staining (Figures 7G and 7H). In control treatments with superoxide dismutase for NBT staining or with ascorbic acid for DAB staining, no staining was detected in the wild type or *fro1*, suggesting that the staining was ROS specific. These results show that the *fro1* mutation causes constitutive accumulation of ROS.

Growth Retardation and Germination Sensitivity to Osmotic Stress in *fro1*

On Murashige and Skoog (1962) (MS) agar medium supplemented with 3% Suc, *fro1* seeds germinated ~1 day later than wild-type seeds. After germination, *fro1* plants also grew more slowly than the wild type. First leaf appearance after germination in *fro1* took 1.7 more days than in the wild type, and inflorescence and first flower in *fro1* also appeared with a significant delay (Figure 8A). *fro1* mutant plants were smaller in size than wild-type plants (Figures 8B and 8C). Nevertheless, *fro1* plants eventually reached heights similar to those of wild-type plants (Figures 8D and 8E), even though *fro1* leaves remained smaller than wild-type leaves. The green color and vigor of *fro1* mutant plants lasted longer than those of wild-type plants. Whereas the wild type started to dry at 7 weeks after imbibition, *fro1* was still green (Figure 8D) at that time and only began to dry at >8 weeks after imbibition.

During mutant handling, we noticed that *fro1* on MS agar medium supplemented with 3% Suc showed a slightly lower germination rate. When Suc was removed from the medium, the germination rates of the wild type and *fro1* were almost

the same (data not shown). Because of this observation, we compared *fro1* and wild-type seed germination in response to various concentrations of sugars or salt (Figure 9). To avoid potential complications caused by MS agar medium, filter papers were used for the germination test. On control filter papers soaked with water, maximal germination was achieved for both *fro1* and the wild type on day 4 after imbibition. Thus, germination rate was scored on day 4 after imbibition. Under our conditions, Suc or mannitol did not affect wild-type germination (Figures 9A and 9B). However, *fro1* germination was affected by all of these sugars in a concentration-dependent manner (Figures 9A and 9B), suggesting that the reduced germination rate in *fro1* probably was the result of an osmotic effect.

Glc at high concentrations reduced wild-type germination, but the effect was much more dramatic on *fro1* (Figure 9C). NaCl inhibited both wild-type and *fro1* germination, but the inhibition on *fro1* was slightly more pronounced (Figure 9D). These observations suggest that *fro1* seed germination is more sensitive to inhibition by osmotic stress. We further investigated the osmotic effect on *fro1* vegetative growth by monitoring root elongation. Seven-day-old wild-type and *fro1* seedlings were transferred onto media containing different concentrations of mannitol. Root growth was measured 4 days later. No difference in root growth was observed between *fro1* and the wild type (data not shown).

Isolation of the *FRO1* Gene

To isolate the *FRO1* gene, a positional cloning strategy was used. *fro1* mutant plants in the C24 background were crossed to wild-type plants of the Columbia ecotype. F1 plants from the cross were selfed, and the resulting F2 seeds were collected. A total of 781 *fro1* mutants from the

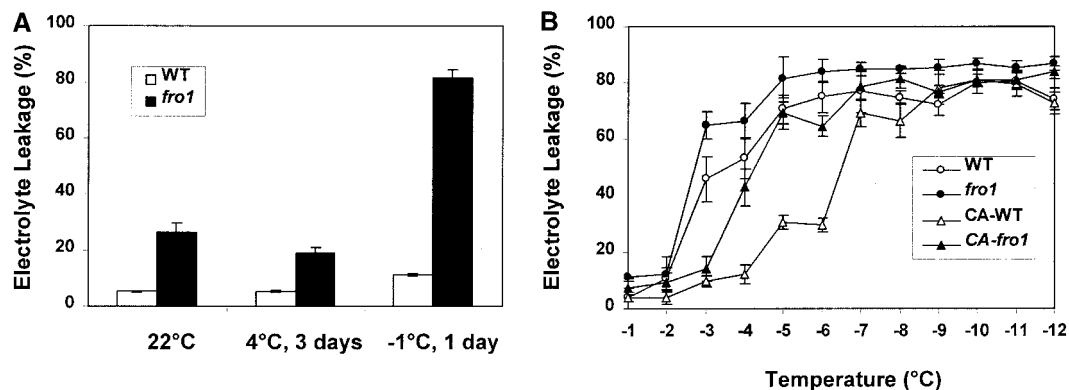


Figure 6. Constitutive Leakiness and Cold Acclimation Defect in *fro1* Plants.

(A) Electrolyte leakage in wild-type and *fro1* leaves from whole plants without stress or with treatment at 4°C for 3 days or at -1°C for 1 day. (B) Electrolyte leakage at different temperatures. For cold acclimation, seedlings were incubated under light at 4°C for 2 days before the test. CA-*fro1*, cold-acclimated *fro1*; CA-WT, cold-acclimated wild type; WT, wild type.

F2 population were selected on the basis of its translucent, water-soaked phenotype and used for mapping with simple sequence length polymorphism (SSLP) (Bell and Ecker, 1994) and cleaved amplified polymorphic sequence (Konieczny and Ausubel, 1993) markers. Preliminary mapping results indicated that *FRO1* was located at the bottom of chromosome 5. Because molecular markers polymorphic between C24 and Columbia were scarce, new SSLP makers were developed based on simple sequence repeats identified from Arabidopsis genomic sequences within this region.

Further mapping with these markers delimited *FRO1* to a short region at the lower arm of chromosome 5, south of marker K8K14-C1 (Figure 10A). Because *FRO1* showed a very tight linkage to marker K9I9-3, *FRO1* very likely was on transformation-competent artificial chromosome clone

K9I9. Therefore, clone K9I9 was introduced into *fro1* mutant plants via *Agrobacterium tumefaciens*-mediated in planta transformation (Liu et al., 1999). *RD29A::LUC* imaging and morphological observations showed that *fro1* mutant plants transformed with K9I9 all behaved like the wild type. These results demonstrate that *FRO1* is contained within transformation-competent artificial chromosome clone K9I9.

Because K9I9 had been sequenced and annotated, several candidate genes were amplified from *fro1* and wild-type plants and sequenced. After sequence analysis, a mutation was found in a hypothetical open reading frame, K9I9.16 (K9I9.10 by Kazusa DNA Research Institute, Japan, at <http://www.kazusa.or.jp>), which spans from 37,239 to 38,522 bp of K9I9. The mutation is a G-to-A change at an intron-exon junction at nucleotide 396 from the ATG start codon. This

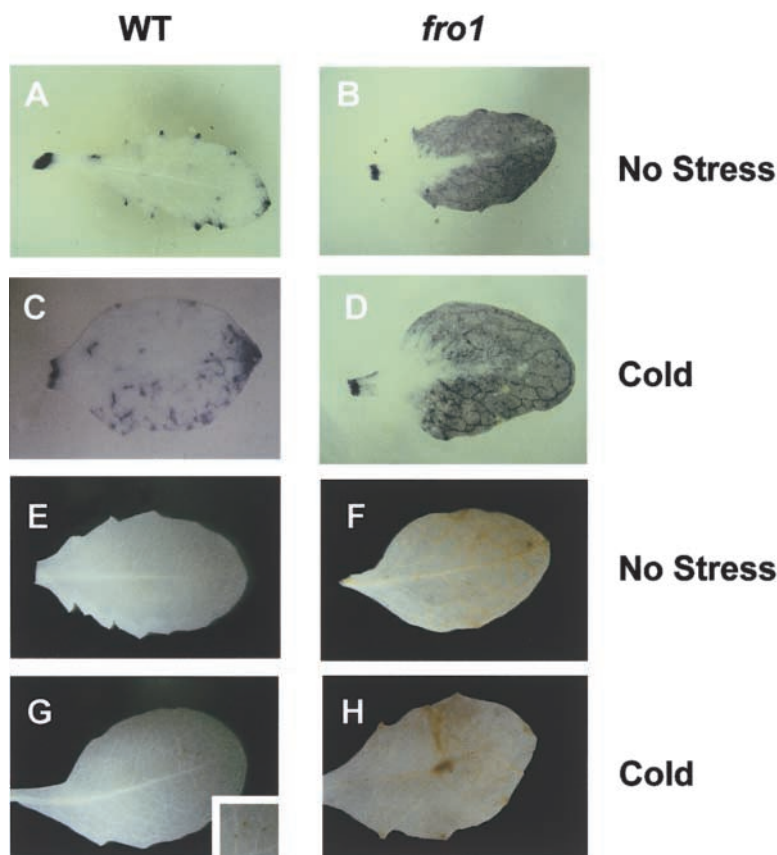


Figure 7. Detection of ROS in *fro1* Leaves.

(A) and (B) NBT staining for superoxide in unstressed leaves of wild-type (A) and *fro1* (B) plants.

(C) and (D) NBT staining for superoxide in cold-treated (4°C for 2 days) leaves of wild-type (C) and *fro1* (D) plants. Staining is shown as dark blue.

(E) and (F) DAB staining for hydrogen peroxide in unstressed leaves of wild-type (E) and *fro1* (F) plants.

(G) and (H) DAB staining for hydrogen peroxide of cold-treated (4°C for 2 days) leaves of wild-type (G) and *fro1* (H) plants. Staining is shown as dark yellow. Dark yellow spots representing DAB staining in the wild type are shown in the inset in (G).

WT, wild type.

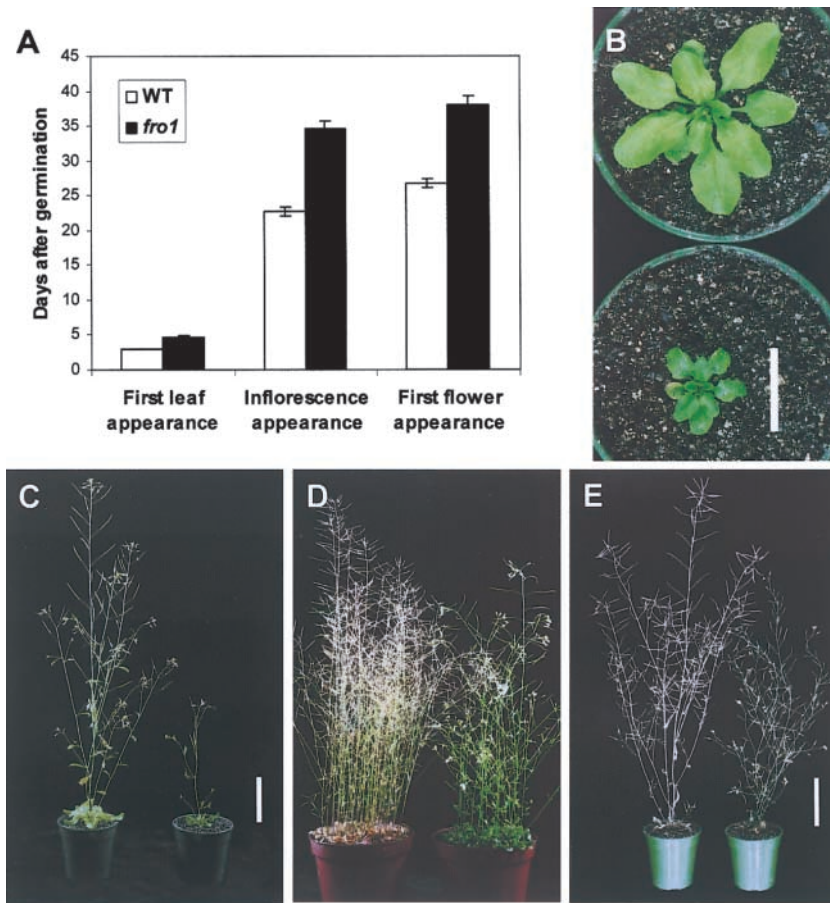


Figure 8. Difference in Growth and Development Rates between Wild-Type and *fro1* Mutant Plants.

(A) Difference in organ appearance after germination ($n = 10$).
 (B) Twenty-day-old wild-type (top) and *fro1* (bottom) plants. Bar = 2 cm.
 (C) Thirty-day-old wild-type (left) and *fro1* (right) plants. Bar = 5 cm.
 (D) Seven-week-old wild-type (left) and *fro1* (right) plants.
 (E) Eight-week-old wild-type (left) and *fro1* (right) plants. Bar = 5 cm.
 WT, wild type.

mutation is predicted to cause missplicing, which would create a premature stop codon 13 bp downstream from the mutation. This candidate gene (i.e., K9I9.16), along with a 1.6-kb sequence 5' upstream of the translation start codon and 260 bp of the 3' untranslated region, was cloned into a binary vector and introduced into *fro1* mutant plants. Twelve independent transgenic lines were obtained, and all grew like the wild type (Figure 10B). In the T2 generation, luciferase imaging analysis of cold-treated seedlings revealed the segregation of wild-type and *fro1* phenotypes (Figures 10C and 10D). All seedlings showing the wild-type *RD29A::LUC* phenotype were found to have the K9I9.16 transgene based on their hygromycin resistance. These results prove that K9I9.16 is the *FRO1* gene.

***FRO1* Encodes the NADH Dehydrogenase Subunit of Mitochondrial Respiratory Chain Complex I**

FRO1 cDNA was isolated by reverse transcriptase-mediated PCR. A comparison between the cDNA and genomic sequences revealed four introns and five exons in the *FRO1* gene. This experimentally deduced gene structure is the same as the computer-annotated version generated by the Arabidopsis Genome Initiative (Palm et al., 2000). GenBank searches found that *FRO1* has high amino acid sequence similarities to the 18-kD Fe-S subunit of mitochondrial respiratory chain complex I (NADH dehydrogenase) from diverse organisms. For example, *FRO1* shows 46 to 47% amino acid identity and 63 to 64% similarity to its human and bovine

orthologs, respectively. Because the bovine protein has been characterized biochemically and its sequence confirmed by amino acid sequencing (Walker et al., 1992), an alignment is shown between FRO1 and the bovine protein (Figures 11A and 11B). The calculated molecular mass of FRO1 is 17 kD, similar to that of the Fe-S subunit of complex I (18-kD). Hydropathy analysis revealed that FRO1 is highly hydrophilic, with no potential transmembrane domains (data not shown).

RNA gel blot analysis showed that *FRO1* expression was constitutive and not regulated substantially by cold, abscisic acid, or NaCl treatment, except for a slight upregulation after 1 h of cold treatment (Figure 11C). Interestingly, no *FRO1* transcript could be detected in *fro1* mutant plants, suggesting instability of the misspliced mutant mRNA. This lack of *FRO1* transcript suggests that *fro1* might be a null mutation (Figure 11C).

Like the other known 18-kD Fe-S subunits of complex I, a putative mitochondrial targeting signal peptide was found in FRO1, with a cleavage point between residues 31 and 32, as analyzed by the MitoProt program at <http://www.mips.biochem.mpg.de/cgi-bin/proj/medgen/mitofilter> (Claros and Vincens, 1996). To confirm FRO1 subcellular localization, FRO1 was fused at its C terminus in frame with the GFP reporter under the control of the double 35S promoter. The

construct was introduced into wild-type C24 plants via *Agrobacterium*-mediated transformation. Stable transgenic plants were obtained and used for FRO1:GFP localization with a confocal microscope.

A clear particulate pattern of GFP expression was observed in the FRO1:GFP transgenic *Arabidopsis* roots (Figure 12). The size of the particles is consistent with that expected of mitochondria. The pattern of GFP subcellular localization in *FRO1:GFP* transgenic *Arabidopsis* (Figure 12A) is identical to that of β -ATPase:GFP, which is known to be localized in mitochondria (Figure 12D) (Logan and Leaver, 2000). These results strongly suggest that FRO1 is localized in mitochondria. The subcellular localization of FRO1 did not change under cold treatment (Figures 12B and 12C).

DISCUSSION

In this study, we identified the *fro1* mutation, characterized *fro1* mutant phenotypes, cloned the *FRO1* gene, and analyzed its expression and the subcellular localization of its gene product. *fro1* mutant plants are impaired in cold regulation not only of the *RD29A::LUC* transgene but also of sev-

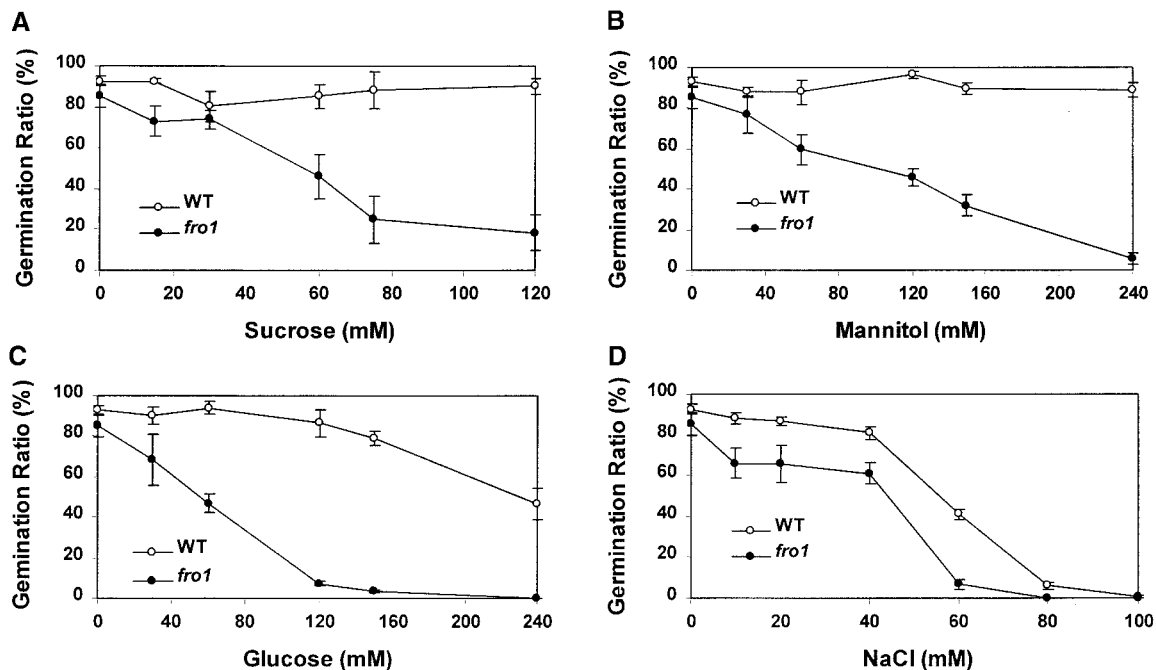


Figure 9. Effect of Osmotic Stress on Seed Germination in *fro1* and the Wild Type.

Germination ratio of the wild type and *fro1* on filter papers saturated with different concentrations of Suc (A), mannitol (B), Glc (C), and NaCl (D). A clear appearance of the radicle was considered as germination, which was scored on day 4 after incubation at room temperature. WT, wild type.

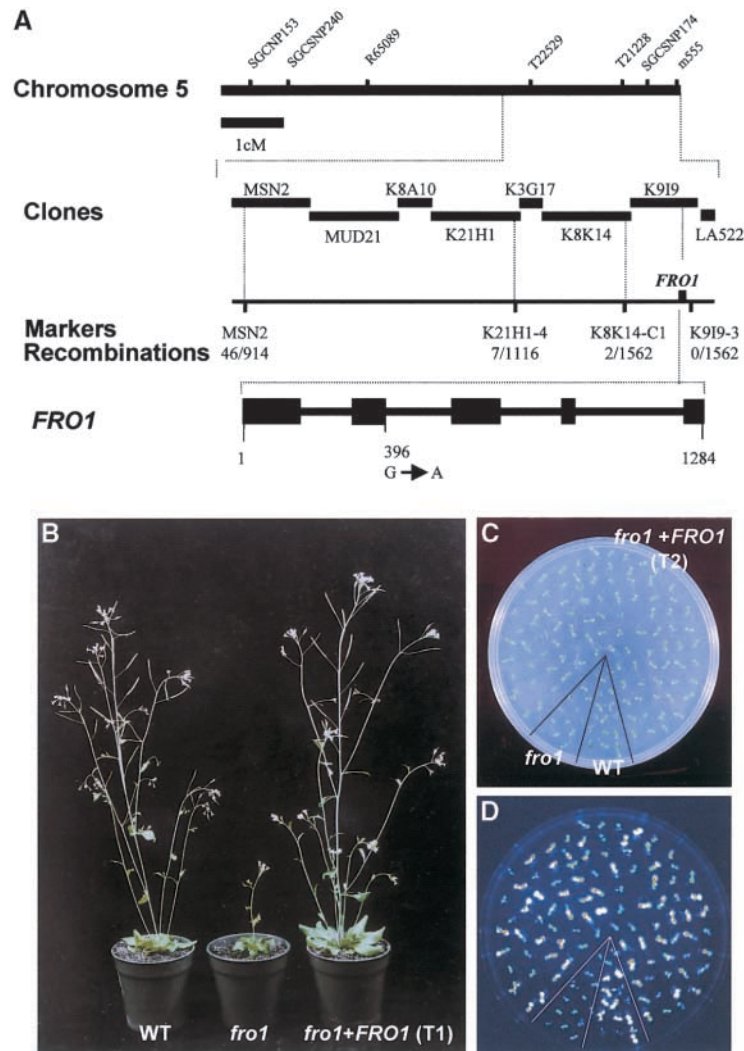


Figure 10. Map-Based Cloning of *FRO1* and Molecular Complementation of *fro1* Mutants.

(A) Markers are SSLP markers except for K8K14-C1, which is a cleaved amplified polymorphic sequence marker. The number of recombinant chromosomes/number of total chromosomes examined at each locus is indicated. The *FRO1* gene structure was obtained by comparing its cDNA sequence with the genomic sequence. Black boxes indicate exons, and solid lines between boxes indicate introns. The position of the *fro1* mutation is indicated. cM, centimorgan.

(B) Morphological comparison between wild type (WT), *fro1*, and *fro1* transformed with *FRO1* (*fro1*+*FRO1*; T1 generation) at the same developmental stage (5 weeks old).

(C) and **(D)** Morphology **(C)** and luminescence **(D)** of wild-type (WT) and *fro1* seedlings of a segregating T2 population from *fro1* transformed with *FRO1* (*fro1*+*FRO1*).

eral endogenous cold-responsive genes, including *RD29A*, *COR47*, *COR15A*, and *KIN1*. The fact that *fro1* reduces the cold induction of *RD29A*, *COR47*, *COR15A*, and *KIN1* but not the *CBF* genes is not surprising, considering that the *sfr6* mutation was reported to impair the induction of DRE/CRT-type genes but not the induction of *CBF* genes (Knight et al., 1999). It is possible that *fro1* may affect factors required for the proper function of CBF transcriptional activa-

tors. The significance, if any, of the increased induction of the *CBF* genes at 12 h after cold treatment is unclear at present.

fro1 plants show a translucent, water-soaked appearance, which may be caused by membrane leakiness. *fro1* mutant plants are defective in one of the components of complex I of the electron transfer chain in mitochondria. This molecular lesion leads to the accumulation of ROS such as superoxide and hydrogen peroxide, which may be

the intermediary signals that alter the cold induction of nuclear genes. Thus, the *fro1* mutant provides a novel example of the retrograde regulation of cold-responsive nuclear gene expression by functionally compromised mitochondria.

Plant mitochondria have the same basic electron transfer system as animal cells. The system consists of the following complexes: complex I, NADH dehydrogenase; complex II, succinate dehydrogenase; complex III, cytochrome *bc*₁; complex IV, cytochrome *c* oxidase. Complex I, the first enzyme, transfers electrons from NADH to ubiquinone. In plants, complex I consists of >35 polypeptides (Leterme and Boutry, 1993), some of which are encoded in mitochondrial DNA. In addition to the common complexes, plant mitochondria have four additional NAD(P)H dehydrogenases (Rasmusson et al., 1998) and an alternative oxidase (Vanlerberghe and McIntosh, 1997). These membrane-bound additional NAD(P)H dehydrogenases and alternative oxidase bypass complex I and complexes III and IV, respectively. As a result, a lower proton electrochemical gradient across the inner membrane would be established if the two alternative electron pathways were used. Nevertheless, the electron transfer system still would be functional. Thus, a defect in complex I in plants is not lethal, as has been shown in other complex I mutant plants (e.g., NMS1; Sabar et al., 2000).

The lower cold induction of the endogenous *RD29A* in *fro1* was not as dramatic as that of the *RD29A::LUC* transgene. This probably is the result of sequence differences between the endogenous *RD29A* gene and the transgene. In the *hos1*

mutant, the endogenous *RD29A* gene also is not affected as much as the *RD29A::LUC* transgene (Ishitani et al., 1998). These observations suggest that there might be additional regulatory elements in the endogenous *RD29A* gene promoter, intron, or untranslated regions, which might be enough to confer some cold responsiveness or transcript stability. In this regard, it is interesting that *COR15A* and *KIN1* were affected more dramatically by *fro1* than the endogenous *RD29A* and *COR47* (Figure 3). Perhaps the additional regulatory elements are not present in *COR15A* and *KIN1*.

As a result of the defect in complex I, *fro1* mutant plants accumulate ROS constitutively. Damage by ROS is dosage dependent. A very high dosage of ROS may cause hypersensitive cell death (Alvarez et al., 1998). However, *fro1* mutant plants can complete their life cycle and set a normal number of seeds. Thus, the endogenous ROS level in *fro1* is not high enough to cause cell death.

One well-known form of cellular damage caused by ROS is lipid peroxidation in cellular membranes (Kagan, 1988). Phospholipid hydroperoxides, a product of lipid peroxidation, form clusters that can function as channels. As a result, membrane permeability to electrolytes increases (Kagan, 1988). Therefore, the translucent and water-soaked *fro1* leaf phenotype may be attributed to increased membrane permeability as a result of lipid peroxidation by ROS. This idea is supported by the electrolyte leakage test, which revealed high relative electric conductivity in *fro1* (Figure 6). This damaged membrane integrity may result in the irregular

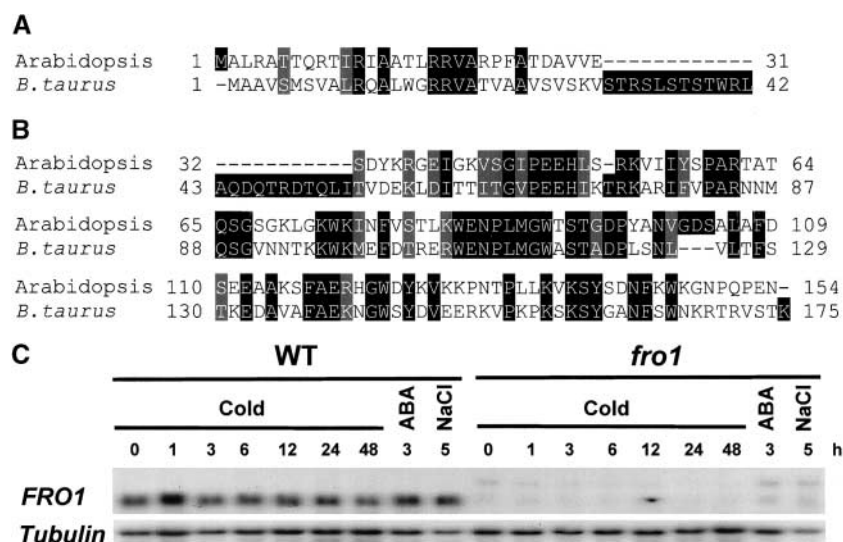


Figure 11. Amino Acid Alignment between *FRO1* and Its Homolog from *Bos taurus*.

(A) and (B) Alignment between predicted mitochondrial targeting sequences (A) and mature proteins after the targeting sequences are cleaved (B). Amino acid sequence alignment was performed with ClustalW (<http://dot.imgen.bcm.tmc.edu:9331/multialign/Options/clustalw.html>). Identical amino acids are highlighted in black, and conservative substitutions are highlighted in gray.

(C) *FRO1* expression in wild-type and *fro1* mutant plants. Seedlings were treated with low temperature (0°C) for the indicated times, abscisic acid (100 μM) for 3 h, and NaCl (300 mM) for 5 h. Tubulin was used as a loading control. ABA, abscisic acid; WT, wild type.

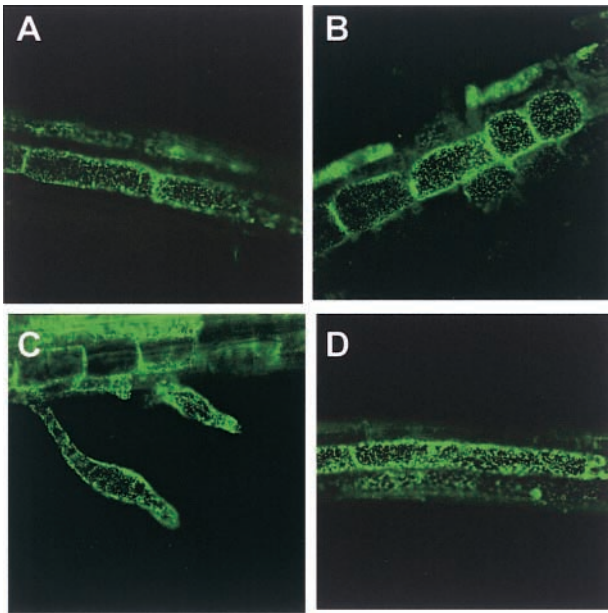


Figure 12. Subcellular Localization of FRO1:GFP.

(A) to (C) Green fluorescence from root tissues of *Arabidopsis* plants transformed with FRO1:GFP was detected using a confocal microscope. Before confocal imaging, seedlings were subjected to the following treatments: no stress (A), cold stress at 0°C for 12 h (B), or cold stress at 0°C for 24 h (C).

(D) Green fluorescence from root tissue of mitochondria-targeted *Arabidopsis* β -ATPase:GFP (Logan and Leaver, 2000) is shown as a positive control.

cells found in *fro1* leaves (Figure 5). Also, the hypersensitivity to osmotic stress seen in *fro1* during seed germination may be caused by its impaired membrane integrity.

Complex I impairments in plants have been reported in the maize NCS2 mutant (Marienfeld and Newton, 1994) and tobacco CMS (cytoplasmic male sterile) I and II mutants (Gutierrez et al., 1997). Interestingly, like *fro1*, the tobacco mutants CMS I and II also showed a slow growth phenotype (Gutierrez et al., 1997), and the maize NCS2 mutant displayed a moderate defect in growth (Coe, 1983; Newton and Coe, 1986). *fro1* mutant plants, however, are not sterile, as the NCS2 and CMS I/II mutants are. The slow growth is likely a consequence of reduced metabolic activity in *fro1*. Reduced metabolic activity in the mutant also may be responsible for its apparent delayed senescence and its thinner cell wall. Increased oxidative stress has been correlated with accelerated senescence (Finkel and Holbrook, 2000). Because *fro1* has increased levels of ROS, it is an interesting and novel example of the negative correlation between senescence and oxidative stress.

It has been reported that hydrogen peroxide activates ANP1, a mitogen-activated protein kinase kinase (Kovtun et al., 2000). In the *Arabidopsis* protoplast system, RD29A was

not induced by exogenous hydrogen peroxide (Kovtun et al., 2000). Thus, it is possible that the signaling of DRE/CRT genes may not be mediated by the ANP1 pathway. Consistent with this notion, none of the cold-responsive genes tested in this study showed constitutive expression, even though *fro1* accumulated high levels of ROS (Figures 3 and 7). Nevertheless, ROS accumulation in *fro1* may be responsible for the defect in cold-responsive gene expression.

Cold regulation of gene expression is known to involve calcium signaling (Monroy and Dhindsa, 1995; Knight et al., 1996). Oxidative stress has been shown to affect cytoplasmic calcium signaling (Price et al., 1994). Therefore, it is possible that *fro1* alters calcium signaling under low temperatures through ROS. Recently, using C2C12 skeletal myocytes that are defective in mitochondrial DNA or are stressed with respiratory chain inhibitors, Biswas et al. (1999) showed increased basal cytosol Ca^{2+} levels and altered nuclear gene expression. Exactly how the ROS in *fro1* alters cold-activated calcium signaling is not known. Because the ROS could trigger calcium signaling without cold treatment, it may desensitize cells toward the cold-induced calcium signal, thus making the mutant plants less responsive to cold in terms of gene expression.

METHODS

Plant Materials and Growth Conditions

Transgenic *Arabidopsis thaliana* (ecotype C24) plants expressing the RD29A::LUC transgene (referred to as wild-type herein) were mutagenized by ethyl methanesulfonate to generate M2 seeds. One-week-old M2 seedlings on 0.6% agar plates containing 3% Suc and Murashige and Skoog (1962) (MS) salts (JRH Biosciences, Lenexa, KS) were screened for altered luciferase expression in response to low temperature, abscisic acid, or osmotic stress with a video imaging system composed of a charge-coupled device camera (CCD-512SB; Princeton Instruments, Trenton, NJ), a controller (Princeton Instruments), and a computer with WinView image-processing software (Princeton Instruments). For luminescence image analysis of seedlings, surface-sterilized seeds were plated on MS agar (0.6%) plates supplemented with 3% Suc and placed at room temperature ($22 \pm 1^\circ\text{C}$) under continuous light after 2 to 3 days of cold stratification. When appropriate, seedlings were transferred to soil pots and allowed to grow in a growth chamber with cycles of 16 h of light at 22°C and 8 h of dark at 18°C .

Stress Treatments

Stress was applied to 1-week-old wild-type and mutant seedlings grown on the same MS agar plate. For cold treatment, the plates were placed at 0°C in the dark for the designated times. For abscisic acid treatment, 100 μM abscisic acid [($\pm$)-*cis,trans*-abscisic acid; Sigma, St. Louis, MO] dissolved in sterile water was sprayed uniformly on the leaves of the seedlings. Abscisic acid-treated plates were kept at room temperature ($22 \pm 1^\circ\text{C}$) under cool-white light for the

designated times. For NaCl treatment, seedlings were transferred to filter paper saturated with 300 mM NaCl in MS solution. The seedlings then were incubated under light at room temperature for the designated times. For luminescence imaging, 1 mM luciferin was sprayed evenly onto seedling leaves at the end of each treatment. The luciferin-sprayed plates were kept in the dark for 5 min for chlorophyll fluorescence to decay. Luminescence images were made with a charge-coupled device camera system. Detailed procedures on imaging and screening have been described previously (Xiong et al., 1999; Lee et al., 2002).

RNA Analysis

Nine-day-old seedlings grown on MS agar plates were used for RNA analysis. After stress treatments, total RNA was extracted and analyzed as described previously (Liu and Zhu, 1997). The *RD29A*-specific probe was from the 3' noncoding region (Liu and Zhu, 1997). The *CBF1*, *CBF2*, and *CBF3* (Gilmour et al., 1998) probes were obtained by amplifying a gene-specific region with the following primers: *CBF1*-F (5'-CGATAGTCGTTCCATTTTGT-3') and *CBF1*-R (5'-TTGCTAGATTCGAGACGAGCC-3'); *CBF2*-F (5'-TTCGATTTTAT-TTCCATTTTGG-3') and *CBF2*-R (5'-CCAAACGTCCTTGAGTCT-TGAT-3'); and *CBF3*-F (5'-GAGGAGCCACGTAGAGGCC-3') and *CBF3*-R (5'-TAAAACTCAGATTATTTCAT-3'). *COR15A* and *COR47* cDNAs (Gilmour et al., 1992; Lin and Thomashow, 1992) were provided by M.F. Thomashow (Michigan State University, East Lansing, MI).

Probe for *KIN1* (Kurkela and Franck, 1990) was a 0.4-kb *EcoRI* fragment of the Arabidopsis EST clone YAP368T7. As a loading control, 25S rRNA, actin2, and β -tubulin gene were amplified by PCR with the following primer pairs: 25S rRNA-F (5'-GGGATTACCCGCTGAGTTTA-3') and 25S rRNA-R (5'-CGTCTCCACAAGCGTATCAA-3'); Actin-F (5'-TGTCGCCATCCAAGCTGTTCTCT-3') and Actin-R (5'-CCATCGGGTAATTCATAGTTCTTCTCG-3'); and Tubulin-F (5'-CGTGGATCACAGCAATACAGAGCC-3') and Tubulin-R (5'-CCTCCTGCACTTCCACTTCGTCTCT-3').

Microscopic Analysis

Leaves from 3-week-old wild-type and *fro1* mutant plants were used to compare their ultrastructure at the transmission electron microscopy facility at the University of Arizona. Fully expanded leaves of wild-type and *fro1* plants were fixed in 0.1 M phosphate buffer, pH 7.4, 4% formaldehyde, and 1% glutaraldehyde, postfixed in 1.0% osmium tetroxide, dehydrated in an ethanol series, and embedded in Epon araldite. Sections were observed with a transmission electron microscope (JEOL 100CXII). All procedures were performed according to standard protocols (Hayat, 2000).

For confocal microscopy, *FRO1:GFP* transgenic seedlings selected on MS agar medium supplemented with 25 mg/L hygromycin were mounted on glass slides, and green fluorescence images were made using a Bio-Rad MRC-1024 confocal laser scanning microscope with a 488-nm excitation laser and a 522/DF35 emission filter.

Electrolyte Leakage Test

The electrolyte leakage test was performed to compare membrane integrity and cold acclimation capability between wild-type and *fro1* plants. To investigate potential relationships between the water-

soaked leaf phenotype in *fro1* and membrane leakage, ~3-week-old plants of the wild type and *fro1* were treated at 4°C for 3 days or at -1°C for 24 h. With 22°C-grown plants of the wild type and *fro1* as controls, several rosette leaves from either treated or untreated plants were detached and transferred to tubes with 25 mL of deionized water. The conductivity of the solution was measured after shaking overnight at room temperature.

To evaluate cold acclimation, one fully developed rosette leaf was detached from ~3-week-old plants and placed immediately into a test tube containing 100 μ L of deionized water with only the petiole submerged in water. The tubes were placed in a refrigerated circulator with the temperature preset at 0°C. After 1 h of incubation, ice chips were added to provide ice nuclei in the tubes. The circulator was programmed so that the bath temperature decreased step-wise to -12°C at a rate of 1°C every 30 min. The tubes were removed upon reaching the designated temperatures and were placed on ice immediately to allow gradual thawing. After complete thawing, the leaflet and the solution in the tube were transferred to another tube containing 25 mL of deionized water followed by overnight shaking at room temperature.

After conductivity measurement of the samples, the tubes with the leaflets were autoclaved. After cooling to room temperature, the conductivity of the solution was measured again. The percentage of electrolyte leakage was calculated as the percentage of the conductivity before autoclaving divided by that after autoclaving.

Detection of Reactive Oxygen Species

For superoxide detection, leaves detached from ~3-week-old plants were vacuum-infiltrated with 0.1 mg/mL nitroblue tetrazolium in 25 mM Hepes buffer, pH 7.6. In a control treatment, 10 mM $MnCl_2$ and 10 units/mL superoxide dismutase were added to the buffer. Samples were incubated at room temperature in the dark for 2 h. For hydrogen peroxide staining, leaves were vacuum-infiltrated with 0.1 mg/mL 3,3'-diaminobenzidine in 50 mM Tris-acetate buffer, pH 5.0. As a control, ascorbic acid at a final concentration of 10 mM was added to the staining medium. Samples were incubated for 24 h at room temperature in the dark. To remove chlorophylls, the stained samples were transferred to 80% ethanol and incubated at 70°C for 10 min. For cold treatment, plants were placed at 4°C for 2 days before staining.

Germination Test

Surface-sterilized seeds were incubated at 4°C for 4 to 5 days to achieve germination uniformity. Then, seeds were planted on filter paper (on a plate) saturated with solutions containing Suc, Glc, mannitol, or NaCl at various concentrations. The plates then were incubated at room temperature under light to allow germination. Germination was scored at day 4. A clear appearance of the radicle was considered as germination.

Positional Cloning

For genetic mapping of the *fro1* mutation, *fro1* was crossed with the wild type in ecotype Columbia with the *glabrous1* mutation. The resulting F1 plants were allowed to self, and F2 seeds were collected. Homozygous *fro1* mutations in the segregated F2 population were selected based on their morphological phenotypes. Mapping of the

mutation was performed using simple sequence length polymorphism (Bell and Ecker, 1994) or cleaved amplified polymorphic sequence (Konieczny and Ausubel, 1993) markers. Primers for simple sequence length polymorphism markers were as follows: MSN2-F (5'-ACGTAAACGAGTCGCCACGT-3') and MSN2-R (5'-GTGAGG-AGTTTGGTATAGCT-3'); K21H1-4F (5'-AACCAAGAGAACCTTGT-TT-3') and K21H1-4R (5'-GATTGGGATTTCTTCCTCAT-3'); and K919-3F (5'-TTTGATAACTAATTAAGGGGAAA-3') and K919-3R (5'-AGCCATAAAACAGCAATCA-3'). Primers for cleaved amplified polymorphic sequence makers were K8K14-C1F (5'-AACTAGCA-CCTGCAAATAGTATT-3') and K8K14-C1R (5'-CTTCTTCTTCTT-AAATAGCTCGAAA-3'). The resulting PCR products were digested with HhaI and resolved in a 1% agarose gel.

Plasmid Construction and Plant Transformation

The K919 transformation-competent artificial chromosome clone was obtained from ABRC (Columbus, OH) and was used for in planta transformation of *fro1* plants. For *fro1* single gene complementation, a genomic DNA fragment of *FRO1* from 1622 bp upstream of the start codon to 260 bp downstream of the stop codon was amplified by LA Taq polymerase (Takara Shuzo, Shiga, Japan) using K919 transformation-competent artificial chromosome DNA as a template with the following primers: K919.10FXb (5'-AAATATCTAGAATAT-ACAGAAAGATTGATGTTT-3') and K919.10RH3 (5'-ATAATAAGC-TTCTCTCTTTCATAATCCAATCAC-3'). The resulting 3178-bp fragment was T-A cloned into the pBluescript II SK- EcoRV site and then subcloned into pCAMBIA1200 between the XbaI and HindIII sites, resulting in pCAMFRO1g23.

To make the *FRO1*:GFP construct, *FRO1* cDNA was isolated by reverse transcriptase-mediated PCR. Total RNA prepared by Trizol (Life Technologies, Rockville, MD) was used to synthesize the first strand using SuperScript II (Life Technologies) with a poly-T primer, T7PtxbaI (5'-GCTCTAGATAATACGACTCACTATAGGGTTTTTTTTT-3'). First-strand cDNA diluted 20 times was used for amplification of the *FRO1* open reading frame using the following primer pair K919.10C1F (5'-AATTTTCCAGATTTCTCTAATTGACGATGG-3') and K919.10C1R (5'-GGAAGAAGGTGTAACATCAGTTTCTGG-3'). The amplified fragment was subcloned into the pBluescript II SK EcoRV site, resulting in pBS2SK-T-*FRO1*c5.

Using pBS2SK-T-*FRO1*c5 as a template, the *FRO1* coding region was amplified with LA Taq polymerase to make the GFP fusion construct. The primers used for the amplification were *FRO1*cNcoIF (5'-CTAATTGACCATGGCGCTTCGTGCTACTACTC-3') and *FRO1*cNcoIR (5'-TAAGAAGGTGCCATGGCGTTTTCTGGTTG-3'). The resulting fragment was subcloned into the pBluescript II SK EcoRV site, resulting in pBS2SK-T-*FRO1*cNcoI4. The *FRO1* coding region then was isolated from pBS2SK-T-*FRO1*cNcoI4 with NcoI treatment. The NcoI fragment was cloned into the pAVA393 NcoI site (von Arnim et al., 1998), resulting in pAVA393-*FRO1*c5NcoI4. Insert direction was checked by PCR using the 35S promoter-specific primer pAVA35S-F (5'-CTCCACTGACGTAAGGGATGAC-3') and the *FRO1*cNcoIR primer. The HindIII and BglII fragments of pAVA393-*FRO1*c5NcoI4 were cloned into the HindIII and BglII sites of pCAMBIA1390, resulting in pCAMBIA1390-*FRO1*c5NcoI4-27.

All binary vectors for plant transformation were transferred to *Agrobacterium tumefaciens* GV3101 (pMP90) by electroporation at 1250 V with capacitance of 25 μ F and resistance of 400 Ω . After appropriate antibiotic selection and PCR confirmation, selected *Agrobacterium* was grown at 28°C in Luria-Bertani medium (1% [w/v]

bacto-tryptone, 0.5% [w/v] bacto-yeast extract, and 1% [w/v] NaCl, pH 7.0) or YEB (5% [w/v] meat extract, 1% [w/v] bacto-yeast extract, 5% [w/v] peptone, 5% [w/v] Suc, and 10 mM MgSO₄, pH 7.4) overnight and then used for in planta floral vacuum infiltration.

Accession Numbers

The accession numbers for the *FRO1* orthologs mentioned in this article are NP_002486.1 (human) and Q02375 (*Bos taurus*).

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