A Major Facilitator Superfamily Transporter Plays a Dual Role in Polar Auxin Transport and Drought Stress Tolerance in Arabidopsis

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INTRODUCTION

The phytohormone auxin, particularly its predominant endogenous form indole-3-acetic acid (IAA), plays a critical role in the spatial and temporal coordination of plant development. Auxin regulates a variety of unrelated processes, such as embryo, root and vascular patterning, postembryonic organogenesis, and tropisms, by directing cell division and expansion (reviewed in Woodward and Bartel, 2005). The diversity of developmental responses mediated by auxin is determined by a specific cellular signal transduction mechanism involving perception by the Transport Inhibitor Response 1/Auxin Signaling F-Box receptor proteins and interpretation by a downstream nuclear signaling pathway that ultimately mediates transcriptional developmental reprogramming (reviewed in Paciorek and Friml, 2006). Many if not all aspects of auxin action rely on its differential distribution within plant tissues manifested by local auxin maxima and minima, also referred to as auxin gradients (reviewed in Tanaka et al., 2006), with the steady-state level of auxin within a particular cell triggering the developmental output of auxin signaling.

Along with metabolism, the asymmetric auxin distribution is mainly sustained by its intercellular transport, which, uniquely among plant signaling molecules, is strictly regulated in a directional fashion termed polar auxin transport. In the shoot, auxin flows in a single direction from its primary sites of synthesis, such as the apical meristem and developing leaves, down toward the root through the stem vascular tissues (Okada et al., 1991; Rashotte et al., 2003). By contrast, two distinct antiparallel streams of auxin movement occur in the root. Shoot-derived auxin travels over the whole-root distance through the central stele downwards to the root apex, where after loading into the outer cell layers it is redirected over a short distance toward the base of the root (Mitchell and Davies, 1975; Tsurumi and Ohwaki, 1978; Rashotte et al., 2000). According to the current paradigm, the mechanistic basis for the polarized auxin cell-to-cell movement is better described by the chemiosmotic hypothesis, in which the proton gradient generated primarily by plasma membrane H+−ATPases between the neutral cytoplasm and the acidic extracellular space drives cellular auxin uptake and efflux (Rubery and Sheldrake, 1974; Raven, 1975). This model postulates the existence of plasma membrane–localized auxin influx and efflux carriers, whose coupled asymmetrical localization between adjacent cells provides the directionality of...
cellular auxin flow (Goldsmith, 1977). These specific transporters were later identified by chemical and genetic approaches using the model system Arabidopsis thaliana (reviewed in Vieten et al., 2007; Petrášek and Friml, 2009).

In the acidic apoplastic environment, a fraction of the weak acid IAA exists in its undissociated form, which can passively diffuse through the plasma membrane inside the cells, whereas the nonlipophilic and therefore less permeable proton-dissociated auxin fraction requires a carrier-mediated uptake system to enter the cell. Auxin influx is mediated by the amino acid permease-like Auxin resistant 1 (AUX1) and three closely related Like AUX1 (LAX) proteins that catalyze proton symport activities (Marchant et al., 1999; Yang et al., 2006; Swarup et al., 2008). Discovered due to the root auxin-resistant nature of the corresponding mutant, AUX1 was among the earliest polar auxin transporters identified, and its localization to a specific side of some cell types has been shown to facilitate directional auxin transport, thereby influencing a wide array of tropic and phyllotactic processes (Bennett et al., 1996; Swarup et al., 2001; Kleine-Vehn et al., 2006). In the neutral cytosolic environment, IAA exists mainly in its membrane-impermeant anionic form that requires active transport to exit the cell. Hitherto, two distinct protein families whose members possess auxin-exporting activity have been associated with cellular polar auxin efflux. The best characterized auxin efflux carriers are members of the unique and plant-specific PIN-formed (PIN) protein family. PINs possess a predicted topology similar to ion-coupled members of the Major Facilitator Superfamily (MFS) and are believed to be secondary transporters that use proton gradients as an energy source (Petrášek et al., 2006; Wisniewska et al., 2006). By contrast, some plant homologs of the human multidrug resistance/P-glycoprotein transporters belonging to the B-family of ATP binding cassette (ABC) superfamily, such as ABCB1, ABCB4, and ABCB19, have been implicated in ATP-energized auxin efflux (Geisler et al., 2005; Lin and Wang, 2005; Santella et al., 2005; Terasaka et al., 2005). Although a few cases of asymmetric subcellular localization have been reported for ABCBs (Blakeslee et al., 2007), the bias and rate of auxin transport are mainly attributable to the highly regulated polar localization of PIN transporters (Petrášek et al., 2006; Wisniewska et al., 2006).

Based on the different, sometimes opposite, phenotypes that their auxin export activities mediate, PINs and ABCs have been shown to define two distinct auxin efflux systems. Whereas PIN-mediated auxin efflux activity is required for polar embryo genesis, organogenesis, and patterning (Reinhardt et al., 2003; Biliou et al., 2005; Weijers et al., 2005), ABCBs contribute mainly to the export of auxin in vegetative tissues, directing long-distance auxin transport in mature plants (Lin and Wang, 2005; Wu et al., 2007). On the other hand, a concerted action of PINs and ABCBs has been highlighted in a number of recent studies demonstrating a role for ABCBs in providing IAA to the PIN proteins for vectorial transport (Mravec et al., 2008) or in stabilizing PINs at the plasma membrane in order to enhance specificity for auxin (Blakeslee et al., 2007; Titapiwananakun et al., 2009). Therefore, both efflux transport systems may act coordinately to generate and maintain the differential distribution of auxin.

The two largest families of transporters occurring in the plant kingdom and the only known to be ubiquitous to all classes of organisms, the ABC superfamily and the MFS, share the common feature of a particularly diverse range of substrates (reviewed in Pao et al., 1998; Rea, 2007). While numerous plant ABC transporters are known to participate in such disparate processes as polar auxin transport, xenobiotic sequestration, disease resistance, or stomatal regulation, among others, the functional significance of the vast majority of MFS transporters in such fundamental physiological processes remains unknown. MFS transporters are single-polypeptide secondary carriers capable only of transporting small solutes in response to chemiosmotic ion gradients. According to the current annotation, the Arabidopsis genome contains more than 120 genes predicted to encode members of the MFS transporter family (TransportDB, www.membranetransport.org, Ren et al., 2004; TAIR10, www.arabidopsis.org, Huala et al., 2001), which have been traditionally classified according to the extensive sequence similarities they share with MFS transporters characterized in other organisms (Pao et al., 1998). To date, the few plant MFS transporters characterized have been essentially implicated in sugar, oligopeptide, and nitrate transport (Büttner, 2007; Tsay et al., 2007). In addition, an Arabidopsis MFS member, Zinc-Induced Facilitator 1 (ZIF1), has been described as a tonoplast-localized transporter involved in zinc tolerance (Haydon and Cobbett, 2007), indicating that MFS transporters might also influence ion homeostasis in plants. We have recently shown that heterologous expression of a ZIF1 paralog, ZIF-Like 1 (ZIFL1), in Saccharomyces cerevisiae confers resistance to the synthetic auxin 2,4-dichlorophenoxyacetic acid (2,4-D) by reducing its concentration inside the yeast cell (Cabrito et al., 2009). Owing mainly to its high stability, 2,4-D has been used for decades as an exogenous source of auxin in physiological studies and mutant screens, thus allowing the discovery of polar auxin transport system components (Bennett et al., 1996).

In this study, we investigated the in vivo roles of the Arabidopsis ZIFL1 transporter using a combination of functional analyses in Arabidopsis and heterologous expression in S. cerevisiae. Our results reveal an important role for MFS transporters in modulating both polar auxin transport and drought stress tolerance. This dual function is determined by alternative splicing of the ZIFL1 gene, which allows the encoded transporter to regulate both root shootward auxin transport, by indirectly affecting cellular auxin efflux and leaf stomatal movements.

RESULTS

The ZIFL1 Promoter Is Predominantly Active in Root Tissues and Stomatal Guard Cells

To initiate the characterization of the Arabidopsis ZIFL1 gene (At5g13750), we monitored its organ- and tissue-specific expression patterns by means of reporter gene experiments. Staining of transgenic lines stably expressing β-glucuronidase (GUS) under the control of the ZIFL1 promoter revealed that this promoter is active in most plant organs (Figures 1A to 1L). In
flowers, *ZIFL1* appears to be strongly and exclusively expressed in the anther stamen filaments at all stages of floral development (Figure 1A). Interestingly, in young leaves *ZIFL1* expression was largely restricted to stomata, which exhibited a distinct GUS coloration (Figure 1B). In fact, all green mature plant organs, such as stems, siliques, and leaves, displayed exclusive staining of stomatal guard cells (Figure 1C). In light- and dark-grown seedlings, very intense *ZIFL1* promoter activity was detected at the shoot apical meristem (Figure 1D) as well as throughout the root system. Although homogenous GUS coloration was observed from the hypocotyl-root junction (Figure 1E) through all along the primary root (PR) (Figure 1F), the *ZIFL1* promoter was particularly active at the root tip (Figure 1G). In addition, while no staining could be detected in lateral root (LR) primordia (Figures 1H to 1K), *ZIFL1* was abundantly expressed in elongating LRs (Figure 1L).

We further explored the precise localization of *ZIFL1* promoter activity in root tissues using transgenic lines expressing the green fluorescent protein (GFP) under the control of the *ZIFL1* promoter (Figures 1M to 1P). In the mature portion of the PR, the GFP signal was restricted to the cortex and to a lesser extent the epidermis (Figure 1M). At the root tip, *ZIFL1* promoter activity was high in both the cortical and epidermal cell layers of the apical meristem and the transition zone, while absent from the quiescent center, the columella cells, or the LR cap (Figure 1N). A similar layer-specific pattern was observed in LRs, with *ZIFL1* expression appearing mainly restricted to the cortex at an early stage of LR elongation (Figure 1O) and spreading also to the epidermis, particularly at the tip, in older LRs (Figure 1P). By contrast, *ZIFL1* promoter activity in stomata was insufficient to allow detection of the GFP signal.

Figure 1. *ZIFL1* Promoter Activity in *Arabidopsis*.

(A) to (L) Differential interference contrast microscopy images of GUS-stained transgenic plants carrying the ProZIFL1:GUS reporter construct. *ZIFL1* promoter activity in flowers (A), a young leaf (B), guard cells (C), the shoot apical meristem (D), the hypocotyl-root junction (E), the PR (F), the PR tip (G), LR primordia at stage II (H), stage V (I), and stage VII (J), a newly emerged LR (K), and a mature LR tip (L). Bars = 1 mm in (A) and (B), 5 µm in (C), and 50 µm in (D) to (L). The arrowhead in (H) points to new cells emanating from the pericycle.

(M) to (P) Confocal laser scanning microscopy images of transgenic root tissues carrying the ProZIFL1:GFP reporter construct. *ZIFL1* promoter activity in the PR (M), the PR tip (N), and a young (O) or mature (P) LR tip. Cell walls were stained with iodide propidium. The GFP and iodide propidium signals are visualized by green and red coloration, respectively. Bars = 50 µm.
Taken together, these results indicate that ZIFL1 may exert a prominent role in roots, particularly at the PR and LR tips, as well as in stomatal guard cells.

The ZIFL1 Gene Encodes Two Splice Isoforms, ZIFL1.1 and ZIFL1.3, Exhibiting Distinct Tissue and Subcellular Distribution

According to the current genome annotation (TAIR10; www.arabidopsis.org), the Arabidopsis ZIFL1 gene contains 17 exons (Figure 2A) and generates three distinct transcripts (Figure 2B). The first, ZIFL1.1 (At5g13750.1), corresponds to the full-length transcript resulting from constitutive splicing of the precursor mRNA (pre-mRNA), while the second, ZIFL1.2 (At5g13750.2), likely arises from selection of an alternative transcription start site in the second intron. The predicted start codons of these two alternative transcripts are in frame; hence, their coding sequences are identical downstream of the ZIFL1.2 ATG. Finally, the ZIFL1.3 (At5g13750.3) splice variant derives from selection of an alternative 3’ splice site in the fourteenth intron, with the second of two contiguous AGs being recognized (see Supplemental Figure 1 online). The ZIFL1.1 and ZIFL1.3 transcripts are therefore identical, except for the absence of two nucleotides in ZIFL1.3 that leads to the inclusion of a premature stop codon in the 15th exon in ZIFL1.3. The ZIFL1.1 transcript is predicted to encode the full-size transporter, displaying the typical MFS transporter signature motif that includes two transmembrane domains, each consisting of six membrane-spanning segments delimiting a central hydrophilic loop (Figure 2C). On the other hand, the ZIFL1.2 and ZIFL1.3 transcripts encode putative transporters that retain the characteristic MFS structure but lack the two first N-terminal or the two last C-terminal membrane-spanning segments, respectively.

In order to verify the accuracy of these predictions and examine the tissue-specific distribution of the ZIFL1 transcripts, we first undertook an RT-PCR approach. Given that the ZIFL1.1 and ZIFL1.3 transcripts differ by just two nucleotides, only primers concomitantly amplifying both transcripts could be designed, while an independent forward primer was used to detect ZIFL1.2. Consistent with the promoter activity profile shown in Figure 1, RT-PCR analysis revealed that ZIFL1 is expressed throughout plant development (Figure 2D), with the ZIFL1 mRNAs being most abundant in young seedlings and root tissues, whereas very low transcript levels were detected in flowers and green tissues. Globally, the ZIFL1.1/ZIFL1.3 and ZIFL1.2 expression patterns were comparable except that the ZIFL1.1/ZIFL1.3 transcripts appeared to be markedly more abundant in senescent leaves and expression of ZIFL1.2 was slightly more pronounced in flowers.

Intriguingly, during this analysis, we were able to easily clone the full-length ZIFL1.1 and ZIFL1.2 transcripts using cDNA obtained from roots, but various attempts to isolate the third transcript from this particular tissue revealed unsuccessful. This prompted us to investigate further the endogenous tissue distribution of the ZIFL1 transcripts by means of a sequencing approach. To this end, we PCR amplified and sequenced a ZIFL1 fragment, including the 2-nucleotide deletion present only in the ZIFL1.3 transcript, independently from root and leaf cDNA. As shown in Supplemental Figure 1 online, when using the F4/R3 primer pair, which concomitantly amplifies the three transcripts, only the sequence corresponding to ZIFL1.1/ZIFL1.2 could be detected in the ZIFL1 fragment obtained from root tissues, whereas the ZIFL1.3 transcript was also detectable, albeit at lower levels, in leaves. When using the F1/R3 primer pair, which specifically amplifies the ZIFL1.1 and ZIFL1.3 splice variants, we were able to identify only the ZIFL1.1 sequence in the root-derived fragment, whereas a mixture of rather equivalent amounts of ZIFL1.1 and ZIFL1.3 was found in the fragment derived from leaf tissues. These sequencing results thus revealed clear tissue specificity of the ZIFL1 transcripts, with roots exclusively expressing ZIFL1.1/ZIFL1.2, whereas in leaves the ZIFL1.3 splice variant also represented a significant fraction of the ZIFL1 transcript pool.

To allow inference of the in vivo function(s) of the ZIFL1 transporter, we isolated two mutant alleles carrying T-DNA insertions in the ZIFL1 gene. The first allele (SALK_030680) was designated zifl1-1 in accordance with Hayden and Cobbett (2007), whereas the second was identified from the GABI-Kat collection (GABI_052H08) and named zifl1-2. Sequence analysis of the genomic DNA/T-DNA junctions determined that both insertions are located in the 11th exon of ZIFL1 (Figure 2A). RT-PCR analysis of ZIFL1 expression in zifl1-1 and zifl1-2 homozygous seedlings using primers annealing upstream of the insertion sites revealed transcript levels comparable to wild-type plants, but no expression was detected when primers flanking or annealing downstream of the T-DNA segments were used (Figure 2E). This indicates that the two mutant alleles produce a truncated version of the three ZIFL1 transcripts that lacks the sequence corresponding to the entire second transmembrane domain and are thus unlikely to encode functional membrane transporters (Shin et al., 2004). These results strongly suggest that both zifl1-1 and zifl1-2 are true loss-of-function mutants.

Because the subcellular localization of the three alternative isoforms was expected to provide important clues in the functional analysis of the ZIFL1 gene, we next generated C-terminal yellow fluorescent protein (YFP) fusions of each isoform and independently cloned them under the control of the 35S promoter. Surprisingly, transient expression of these constructs in Arabidopsis protoplasts suggested that the ZIFL1.1-YFP fusion protein localizes to the tonoplast, whereas ZIFL1.3-YFP appears to be targeted to the plasma membrane (Figures 3A to 3C). By contrast, no fluorescence could be detected in any of the transfection assays performed with the ZIFL1.2-YFP construct. To further verify the subcellular localization of the ZIFL1.1 and ZIFL1.3 isoforms, we generated stable transgenic Arabidopsis zifl1 mutant lines carrying either ZIFL1.1 or ZIFL1.3 fused at the C-terminal end with the GFP sequence and placed under the control of the native ZIFL1 promoter. Confocal microscopy analysis of root apices expressing ZIFL1.1-GFP again indicated tonoplast localization for ZIFL1.1 in both the cortical and epidermal cell layers (Figures 3D to 3F), whereas no fluorescence could be detected with this fusion protein in stomatal guard cells. In agreement with the results from our promoter-GFP reporter gene experiments, we were also unable to visualize...
transgene-derived fluorescence in guard cells of plants expressing the ZIFL1.3-GFP fusion protein. Interestingly, in these transgenic plants, no ZIFL1.3-GFP signal was detected in root apices, despite detailed analysis of a large number of independent transformants and consistent with the absence of ZIFL1.3 transcript detected in wild-type root tissues (see Supplemental Figure 1 online). In order to unequivocally ascertain the subcellular localization of the ZIFL1.1 and ZIFL1.3 splice forms, we subsequently performed colocalization experiments in tobacco (Nicotiana tabacum) leaf epidermal cells using specific tonoplast and plasma membrane markers (Nelson et al., 2007). As shown in Figures 3G to 3V, the ZIFL1.1-GFP fusion protein colocalized with the tonoplast marker, whereas the ZIFL1.3-GFP fusion protein matched the distribution of the plasma membrane marker. Collectively, these findings indicate that the full-length ZIFL1.1 and the truncated ZIFL1.3 transporters are localized at the tonoplast and the plasma membrane, respectively.

Loss of ZIFL1 Function Causes Hypersensitivity to Drought Stress and Auxin-Related Defects

To investigate the biological role(s) of the Arabidopsis ZIFL1 transporter, we performed a detailed phenotypical analysis of the zifl1-1 and zifl1-2 mutants. Under optimal growth conditions, zifl1 mutant adult plants appeared indistinguishable from the corresponding wild type (Columbia-0 [Col-0]), as illustrated in Figure 4A (control conditions), showing also normal flowering time and fertility (data not shown). However, we consistently observed that zifl1 mutant plants were more sensitive to lack of regular watering, as demonstrated by their faster wilting under limited water supply when compared with wild-type plants (Figure 4A, drought stress conditions). This effect was quantified using a detached leaf assay and, as depicted in Figure 4B, both the zifl1-1 and zifl1-2 mutants clearly exhibited enhanced transpiration rates, losing between 40 and 50% of their fresh weight, versus only ~20% for wild-type plants, upon 3 h of water stress. Similar results were obtained when entire plants (rosette stage)
Figure 3. Subcellular Localization of the ZIFL1 Isoforms.

(A) to (C) Confocal laser scanning microscopy images of *Arabidopsis* wild-type mesophyll protoplasts transiently expressing either YFP alone (A) or the ZIFL1.1·YFP (B) or ZIFL1.3·YFP (C) fusion proteins under the control of the 35S promoter. Arrowheads point to the YFP signal on the inner side of the chloroplasts and the nucleus. The YFP and chloroplast autofluorescence signals are visualized by green and red coloration, respectively. Bars = 10 µm.

(D) to (F) Confocal laser scanning microscopy images of an *Arabidopsis* zif1-2 mutant root tip (D) and a mature epidermal (E) or cortical (F) root cell stably expressing the ZIFL1.1·GFP fusion protein under the control of the endogenous ZIFL1 promoter. The GFP and iodide propidium signals are visualized by green and red coloration, respectively. The arrowhead points to a transvacuolar strand. Bars = 50 µm in (D) and 5 µm in (E) and (F).

(G) to (V) Confocal laser scanning microscopy images of tobacco leaf epidermal cells transiently coexpressing the ZIFL1.1·GFP ([G] and [K]) or the ZIFL1.3·GFP ([O] and [S]) fusion proteins with the tonoplast marker γ-TIP-mCherry ([H] and [P]) or the plasma membrane marker PIP2A-mCherry ([L] and [T]) under the control of the 35S promoter. Merged images of whole-cell ([I], [M], [Q], and [U]) or close-up ([J], [N], [R], and [V]) views are shown. Arrowheads point to transvacuolar strands, and asterisks indicate fluorescence signals approaching the nucleus only on the side facing the exterior of the cell. The GFP and mCherry signals are visualized by green and red coloration, respectively. Bars = 20 µm.
Figure 4. Drought-Related Phenotypes of ZIFL1 Loss-of-Function Mutants.

(A) Representative images of 5-week-old wild-type (Col-0) and zifl1 mutant (zifl1-1 and zifl1-2) plants grown under normal water supply (control conditions) or 7 d after terminating irrigation (drought stress). Bars = 1 cm.

(B) Water loss rates of rosette leaves detached from 5-week-old irrigated plants of the wild type (Col-0), the zifl1 mutants (zifl1-1 and zifl1-2), and two independent genomic complementation lines (zifl1-1comp and zifl1-2comp). Results are representative of three independent experiments, and values represent means ± SD (n = 4).

(C) Stomatal apertures of rosette leaves detached from 5-week-old irrigated plants of the wild type (Col-0) and the zifl1 mutants (zifl1-1 and zifl1-2) after 3 h of light, dark, or ABA (3 μM) treatment. Values represent the mean of four independent experiments ± SD. Asterisks denote statistically significant differences between zifl1 mutants and the wild type (*P < 0.05, **P < 0.01, and ***P < 0.001; Student’s t test).
were employed instead of detached leaves (Table 1), indicating that the ZIFL1 transporter is involved in the regulation of drought stress tolerance in Arabidopsis.

Guard cells play a key role in optimizing plant CO₂ uptake and concomitant water loss at the leaf interface by precisely controlling stomatal apertures in response to physiological and environmental stimuli (reviewed in Araújo et al., 2011). Together with the specific ZIFL1 promoter activity, we detected in guard cells (Figure 1C), the drought stress hypersensitivity of the zifl1 mutants prompted us to investigate whether ZIFL1 disruption affects stomatal movements. Indeed, microscopy measurements of stomatal apertures revealed that the stomatal pore was significantly larger in the zifl1 mutants than in the wild type (Figure 4C). This effect was relatively modest under light but markedly more pronounced in the dark or upon treatment with the stress hormone abscisic acid (ABA), two major effectors triggering stomatal closure (Mittelheuser and Van Steveninck, 1969; Thimann and Satler, 1979). Nevertheless, zifl1 mutant stomata retain full sensitivity to both dark- and ABA-mediated stomatal closure, indicating that loss of ZIFL1 function does not affect the corresponding signaling transduction pathways. These results indicate that the ZIFL1 transporter is required for efficient stomatal closure.

Given the high ZIFL1 expression in root tissues (Figures 1, 2D, and 3D) and the ZIFL1 heterologous expression results we obtained previously in yeast with the 2,4-D herbicide and the naturally occurring auxin IAA (Cabrito et al., 2009), we next sought to evaluate the PR response of the zifl1 mutants to exogenous application of these two auxins. Although PR growth was unaffected under control conditions, zifl1-1 and zifl1-2 mutant PR elongation exhibited hypersensitivity to the inhibitory effect of 2,4-D, as well as significantly enhanced sensitivity to inhibition by IAA, at least at low concentrations of the natural auxin (Figures 5A and 5B, Table 1). To determine whether the identified phenotypes were restricted to root elongation, we also examined the effect of auxin application on hypocotyl elongation of dark-grown seedlings. Again, no phenotype was observed under control conditions, but both 2,4-D and IAA inhibited hypocotyl elongation to a greater extent in zifl1-1 and zifl1-2 mutants than in the wild type (see Supplemental Figure 2A online). Remarkably, such an exacerbated sensitivity was not detected when zifl1 mutant seedlings were challenged with other naturally occurring (e.g., indole-3-butyric acid [IBA]) or synthetic (e.g., 1-naphthaleneacetic acid [1-NAA]) auxinic compounds (see Supplemental Figure 3 online), strongly suggesting that the ZIFL1 transporter confers resistance specifically to 2,4-D and IAA.

A detailed characterization of the root system architecture of seedlings grown under control conditions revealed that zifl1 mutant and wild-type roots produce a similar number of total LR structures, including LR primordia and emerged LRs. However, the proportion of emerged LRs was reduced by ~25% in zifl1 mutant seedlings, and these emerged LRs were significantly shorter than those of the wild type (Figures 5A and 5C). This indicates that the zifl1 mutants are not defective in LR initiation processes but rather in LR emergence and/or postemergence elongation. Furthermore, we quantitatively assessed root tip reorientation after gravistimulation and found that both zifl1-1 and zifl1-2 exhibit defective root gravitropic bending relative to wild-type seedlings (P < 0.001, Student’s t test; Figure 5D). As no differences were observed in linear PR elongation, this phenotype is likely to result from an intrinsic defect in the gravitropic response. Therefore, the zifl1 mutants are impaired in two developmental processes typically triggered by auxin. In fact, this phytohormone provides a critical stimulatory signal during the two steps of LR development, the initiation and emergence phases (Wu et al., 2007; Swarup et al., 2008), eliciting also differential growth rates in response to tropisms, in particular to root gravitropism (Friml et al., 2002).

The striking phenotypical similarity displayed by the zifl1-1 and zifl1-2 mutant alleles strongly suggested that the observed defects result from loss of ZIFL1 function. To exclude the possibility of the effect of a mutation in another gene, we transformed both mutants with a genomic fragment spanning the

### Table 1. Complementation of zifl1 Mutant Phenotypes by the ZIFL1.1 and ZIFL1.3 Isoforms

<table>
<thead>
<tr>
<th>Genotype</th>
<th>PR Elongation (cm)</th>
<th>Water Loss (%) FW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
<td>0.025 μM 2,4-D</td>
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<tr>
<td>Col-0</td>
<td>4.56 ± 0.46</td>
<td>2.93 ± 0.33</td>
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<tr>
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<tr>
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<tr>
<td>zifl1-2/ProZIFL1-ZIFL1.3-GFP</td>
<td>4.42 ± 0.39</td>
<td>2.39 ± 0.41 (1.7e⁻³)</td>
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Numbers between parentheses indicate the P values (wild type versus zifl1 mutants or ZIFL1 transgenic lines) obtained by Student’s t test.
aEffect of 2,4-D and IAA on PR elongation of 12-d-old seedlings of the wild-type (Col-0), the zifl1 mutants (zifl1-1 and zifl1-2), and two independent mutant lines complemented with either the ProZIFL1-ZIFL1.1-GFP or ProZIFL1-ZIFL1.3-GFP construct. Results are representative of three independent experiments, and values represent means ± SD (n = 16).
bWater loss rates of 5-week-old irrigated plants of the wild type (Col-0), the zifl1 mutants (zifl1-1 and zifl1-2), and two independent mutant lines complemented with either ProZIFL1-ZIFL1.1-GFP or ProZIFL1-ZIFL1.3-GFP. Results are representative of three independent experiments, and values represent means ± SD (n = 4). FW, fresh weight.
Figure 5. Auxin-Related Phenotypes of ZIFL1 Loss-of-Function Mutants.
entire ZIFL1 gene, which included the same promoter sequence used in the reporter gene experiments. The water loss rates of the corresponding transgenic complementation lines were similar to those of wild-type plants (Figure 4B). In addition, zif1 mutants complementation lines exhibited complete restoration of PR and hypocotyl elongation wild-type sensitivity to 2,4-D and IAA (Figure 5B; see Supplemental Figure 2A online). Finally, genomic complementation of the zif1 mutants fully suppressed the LR formation (Figure 5C) and gravitropic bending defects (Figure 5D). We have thus confirmed that disruption of the ZIFL1 gene is responsible for the identified zif1 mutant phenotypes. Moreover, the fact that we observed full rescue of all phenotypes strongly suggests that the full-length ZIFL1 promoter is comprised in the selected sequence.

ZIFL1.1 Functions in Auxin-Related Processes, Whereas ZIFL1.3 Confers Drought Stress Tolerance

In light of the above results, we next wanted to examine the contribution of the three alternative ZIFL1 isoforms to the phenotypes established for the zif1 mutants. To address this issue, we first generated Arabidopsis transgenic lines independently expressing each ZIFL1 transcript under the control of the 3SS promoter in the wild-type background for phenotypical characterization.

Regarding the auxin-related phenotypes, we found that overexpression of ZIFL1.1 confers increased resistance to the inhibitory effects of exogenously applied 2,4-D and IAA, both at the level of PR (Figure 6A) and hypocotyl (see Supplemental Figure 2B online) elongation. Figure 6B shows that, although the number of total LR structures produced by the ZIFL1.1-overexpressing lines was not significantly different from the wild type, the ratio between LR primordia and emerged LRs was markedly unwedged to emerged LRs, whose length was increased by almost 30% in these plants. Overexpression of ZIFL1.1 also conferred enhanced gravity bending ability to the transgenic roots (P < 0.001, Student t test; Figure 6C). Therefore, the auxin-related phenotypes of ZIFL1.1-overexpressing lines were strikingly opposite to those discovered for the zif1 mutants. By contrast, overexpression of ZIFL1.3 had no effect on any of these auxin-related processes. However, while no change in the transpiration potential of ZIFL1.1-overexpressing leaves was detected, leaves from ZIFL1.3-overexpressing plants exhibited significantly lower transpiration rates than those of the wild type (Figure 6D). Consistent with this and contrary to the zif1 mutants, ZIFL1.3-overexpressing plants noticeably closed their stomata more efficiently than wild-type or ZIFL1.1-overexpressing plants, irrespective of the condition tested (Figure 6E). Finally, the ZIFL1.2-overexpressing lines, which were included in all the phenotypical assays performed, did not exhibit any evident phenotype (Figures 6A to 6D; see Supplemental Figure 2B online), indicating that this ZIFL1-specific transcript has no function, at least in the parameters tested.

To better ascertain ZIFL1.1 and ZIFL1.3 functional specificity, we then assessed the ability of these isoforms to suppress the zif1 mutant defects by phenotyping the transgenic lines expressing either the ZIFL1.1-GFP or the ZIFL1.3-GFP fusion protein under the control of the ZIFL1 promoter in the zif1-1 and zif1-2 mutant backgrounds. As depicted in Table 1, complementation with ZIFL1.1-GFP fully abolished the PR elongation hypersensitivity to exogenous auxins of the zif1 mutants but had no effect on their water loss rates. Conversely, ZIFL1.3-GFP fully complemented the drought-related phenotype but not the auxin-related defects. In addition to demonstrating that the GFP fusion proteins are functionally active, these results, together with those from the overexpression studies, provide evidence that the two alternative ZIFL1 splice isoforms have distinct biological functions, with ZIFL1.1 modulating root auxin-related processes and ZIFL1.3 regulating drought stress tolerance.

ZIFL1.1 Influences Cellular Auxin Efflux in Yeast and Arabidopsis

The opposite alterations conferred by loss and gain of ZIFL1.1 function in auxin-related processes led us to hypothesize that this isoform regulates auxin transport in the Arabidopsis root. In a previous study, we reported that heterologous expression of the ZIFL1.3 isoform confers enhanced 2,4-D and IAA resistance in S. cerevisiae, most likely sustained by an increased efflux from the yeast cell, at least in the case of 2,4-D (Cabrito et al., 2009). We therefore decided to reexamine the auxin transport activity of ZIFL1.3 in parallel to that of ZIFL1.1 in the yeast Δtpo1 mutant, which lacks an MFS transporter involved in 2,4-D and IAA resistance (Teixeira and Sá-Correia, 2002). Correct expression of the GFP-ZIFL1.1 and GFP-ZIFL1.3 fusion proteins was confirmed by immunoblotting (see Supplemental Figure 4A online).

Figure 5. (continued).

(A) Representative images of 12-d-old wild-type (Col-0) and zif1 mutant (zif1-1 and zif1-2) seedlings grown on control medium or medium supplemented with 0.025 μM 2,4-D or 0.05 μM IAA.

(B) Effect of 2,4-D (left panel) and IAA (right panel) on PR elongation of 12-d-old seedlings of the wild type (Col-0), the zif1 mutants (zif1-1 and zif1-2), and two independent genomic complementation lines (zif1-1comp and zif1-2comp). Results are representative of three independent experiments, and values represent means ± sd (n = 16).

(C) LR phenotype of 12-d-old seedlings of the wild type (Col-0), the zif1 mutants (zif1-1 and zif1-2), and two independent genomic complementation lines (zif1-1comp and zif1-2comp). Total number of LRs (top panel), frequency of LR primordia (LRP) and emerged LRs (ELR) (middle panel), and LR length (bottom panel) are presented. Results are representative of three independent experiments, and bars represent means ± sd (n = 8).

(D) Root gravitropic response of 7-d-old seedlings of the wild type (Col-0), the zif1 mutants (zif1-1 and zif1-2), and two independent genomic complementation lines (zif1-1comp and zif1-2comp) after 24 h of 90° gravistimulation. The length of each bar represents the frequency of seedlings showing the direction of root tip curvature within the corresponding 30° sector. Results are representative of three independent experiments (n ≥ 30). Asterisks denote statistically significant differences between zif1 mutants and the wild type (∗P < 0.01 and ∗∗ P < 0.001; Student’s t test).
Figure 6. Phenotypes of Transgenic Arabidopsis Lines Overexpressing Individual ZIFL1 Isoforms.

(A) Effect of 2,4-D (left panel) and IAA (right panel) on PR elongation of 12-d-old seedlings of the wild type (Col-0) and ZIFL1-overexpressing lines (ZIFL1.1OX, ZIFL1.2OX, and ZIFL1.3OX). Results are representative of three independent experiments, and values represent means ± SD (n = 16).
Subcellular localization studies showed that both plant isoforms are targeted to the yeast plasma membrane (see Supplemental Figure 4B online), in agreement with the localization observed in planta for the ZIFL1.3 but not for the ZIFL1.1 isoform (Figure 3). The latter observation is not so surprising, as plant proteins, particularly transporters, may not always be targeted properly when expressed in yeast (Bassham and Raikhel, 2000).

We took advantage of ZIFL1.1 mislocalization, as plasma membrane targeting is a prerequisite to study ZIFL1 transport properties in the Δtpo1 background (Cabrito et al., 2009). As seen in Supplemental Figure 4C online, Δtpo1 mutant cells expressing GFP-ZIFL1.1 or GFP-ZIFL1.3 similarly exhibited higher final biomass and a reduced lag phase in the presence of 2,4-D or IAA. This enhanced resistance was correlated with a significant reduction in the accumulation of [14C]2,4-D or [14C]IAA in the LR phenotype of 12-d-old seedlings of the wild type (Col-0) and or IAA. This enhanced resistance was correlated with a significant reduction in the accumulation of [14C]2,4-D or [14C]IAA in nonadapted yeast Δtpo1 cells suddenly exposed to either radiolabeled auxin (Table 2). Compared with yeast cells harboring the empty vector, net auxin accumulation was reduced by ∼50% in cells expressing either of the two plant isoforms, indicating that ZIFL1.1 and ZIFL1.3 mediate similar auxin transport activity, at least in S. cerevisiae. In agreement with previous reports in yeast heterologous systems, Figure 7A shows that the IAA accumulation rate was reduced by ∼30% when cells were challenged with the polar auxin efflux inhibitor naphthoylphthalamic acid (NPA), probably owing to the inhibitory effect this chemical exerts on endogenous yeast transporters (Geisler et al., 2005; Yang and Murphy, 2009). However, this negative effect was not further intensified by expression of either plant isoform, indicating that the IAA transport activity mediated by ZIFL1.1 and ZIFL1.3 is not inhibited by NPA in S. cerevisiae.

We next investigated whether the ZIFL1.1 and ZIFL1.3 transporters also influence cellular auxin levels in Arabidopsis using a radiolabeled auxin accumulation assay in excised root tips. The results presented in Table 2 show that the saturable accumulation of [14C]2,4-D or [14C]IAA in nonadapted yeast Δtpo1 cells suddenly exposed to either radiolabeled auxin (Table 2). Compared with yeast cells harboring the empty vector, net auxin accumulation was reduced by ∼50% in cells expressing either of the two plant isoforms, indicating that ZIFL1.1 and ZIFL1.3 mediate similar auxin transport activity, at least in S. cerevisiae. In agreement with previous reports in yeast heterologous systems, Figure 7A shows that the IAA accumulation rate was reduced by ∼30% when cells were challenged with the polar auxin efflux inhibitor naphthoylphthalamic acid (NPA), probably owing to the inhibitory effect this chemical exerts on endogenous yeast transporters (Geisler et al., 2005; Yang and Murphy, 2009). However, this negative effect was not further intensified by expression of either plant isoform, indicating that the IAA transport activity mediated by ZIFL1.1 and ZIFL1.3 is not inhibited by NPA in S. cerevisiae.

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Given the encouraging results described above, we next aimed at determining whether the ZIFL1.1 transporter plays a role in the polar transport of IAA in roots. To this end, we first examined the effect of different chemical inhibitors of polar auxin transport on the elongation of zif11 mutant and ZIFL1.1-overexpressing PRs (Table 3). While both root types responded to the auxin influx inhibitor naphthoxyacetic acid (NOA) as the wild type, mutant and ZIFL1.1-overexpressing lines exhibited enhanced and reduced resistance, respectively, to the inhibitory effect that the auxin efflux inhibitors NPA and triiodobenzoic acid (TIBA) exert on PR elongation (Chen et al., 1998; Luschnig et al., 1998). These data reinforce the notion that ZIFL1.1 influences exclusively cellular auxin efflux, hinting at its involvement in polar auxin transport. By contrast, ZIFL1.3-overexpressing lines behaved as the wild type in this assay, further supporting the finding that the ZIFL1.3 truncated isoform plays no role in auxin-related processes.

To validate our hypothesis at the whole-plant level, we indirectly visualized auxin distribution by histochemical staining of wild-type and zif11-mutant root tips carrying the well-described ProDR5:GUS auxin-responsive reporter construct (Ulmasov et al., 1997; Delker et al., 2010). Under control conditions, DRS activity in both types of roots was confined to a small collection of cells comprising the quiescent center and the stem cell niche (Figure 7C; see Supplemental Figure 5 online), as previously reported (Sabatini et al., 1999). Localized application of IAA at either the hypocotyl-root junction or the extreme root tip of wild-type and zif11 mutant roots similarly led to a substantially more robust staining in these particular cell types. In addition, in the wild-type background, GUS staining further extends into the

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**Figure 6.** (continued).

**(B)** LR phenotype of 12-d-old seedlings of the wild type (Col-0) and ZIFL1.1-overexpressing lines (ZIFL1.1OX, ZIFL1.2OX and ZIFL1.3OX). Total number of LRs (top panel), frequency of LR primordia (LPR) and emerged LRs (ELR) (middle panel), and LR length (bottom panel) are presented. Results are representative of three independent experiments and bars represent means ± so (n = 8).

**(C)** Root gravitropic response of 7-d-old seedlings of the wild type (Col-0) and ZIFL1.1-overexpressing lines (ZIFL1.1OX, ZIFL1.2OX, and ZIFL1.3OX) after 6 h of 90° gravistimulation. The length of each bar represents the frequency of seedlings showing the direction of root tip curvature within the corresponding 30° sector. Results are representative of three independent experiments (n ≥ 30).

**(D)** Water loss rates of rosette leaves detached from 5-week-old irrigated plants of the wild type (Col-0) and ZIFL1.1-overexpressing lines (ZIFL1.1OX, ZIFL1.2OX, and ZIFL1.3OX). Results are representative of three independent experiments and values represent means ± so (n = 4).

**(E)** Stomatal apertures of rosette leaves detached from 5-week-old irrigated plants of the wild type (Col-0) and ZIFL1.1-overexpressing lines (ZIFL1.1OX and ZIFL1.3OX) after 3 h of light, dark, or ABA (3 μM) treatment. Values represent the mean of four independent experiments ± so. Asterisks denote statistically significant differences between ZIFL1-overexpressing lines and the wild type (**P < 0.01 and ***P < 0.001; Student’s t test).
distal elongation zone, with a clear gap in the region spanning from the quiescent center to the transition zone. By contrast, when IAA was applied at the hypocotyl-root junction of zif1 mutant roots, staining barely spread into the apical meristem, with little if any staining evident in the transition or distal elongation zones. The DR5 activity pattern of the mutant roots, staining barely spread into the apical meristem, remaining also unaffected compared with the wild type, in agreement with previous reports (Chen et al., 1998; Müller et al., 1998). As seen in Figure 7D, root-replacing IAA by benzoic acid as a weak acid diffusion control (Chen et al., 1998; Müller et al., 1998), we included the Arabidopsis Δtpo1 mutant, a null allele in the Col-0 ecotype (Luschnig et al., 1998), in this analysis. The entire assay was also recapitulated from the quiescent center to the transition zone. By contrast, impairing rootward transport rates to a similar extent as in the wild type, as already reported for the eir1-4 mutant (Rashotte et al., 2000). Despite being slightly less pronounced in the zif1-1 than in the eir1-4 mutant background, the opposite effect was observed in ZIF1.1-overexpressing roots.

Together with the results from the PR elongation assay in presence of NPA (Table 3), the data presented in Figure 7D demonstrate that ZIF1.1-mediated IAA transport activity is NPA sensitive in plant. In combination with its NPA insensitivity in yeast (Figure 7A), this further suggests that the tonoplast ZIF1.1 transporter does not function as an auxin transporter itself, but rather favors shootward auxin flows driven by typical plasma membrane IAA efflux carriers whose IAA export activity is inhibited by NPA in both plant and heterologous systems (Noh et al., 2001; Murphy et al., 2002; Petrášek et al., 2006; Yang and Murphy, 2009).

### Table 2. Accumulation of Radiolabeled Auxins in Yeast and Arabidopsis

<table>
<thead>
<tr>
<th>Genotype</th>
<th>2,4-D</th>
<th>IAA</th>
<th>NAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yeast&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Δtpo1 + pGREG576</td>
<td>13.97 ± 1.29</td>
<td>28.70 ± 2.08</td>
<td>ND</td>
</tr>
<tr>
<td>Δtpo1 + GFP-ZIFL1.1</td>
<td>7.26 ± 0.89 (1.2e^-5)</td>
<td>12.96 ± 2.36 (2.1e^-7)</td>
<td>ND</td>
</tr>
<tr>
<td>Δtpo1 + GFP-ZIFL1.3</td>
<td>7.40 ± 0.64 (6.7e^-7)</td>
<td>15.03 ± 1.70 (2.4e^-7)</td>
<td>ND</td>
</tr>
<tr>
<td>Arabidopsis&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Col-0</td>
<td>100.00 ± 11.64</td>
<td>100.00 ± 11.99</td>
<td>100.00 ± 14.38</td>
</tr>
<tr>
<td>zif1-1</td>
<td>141.70 ± 10.40 (3.8e^-1)</td>
<td>120.52 ± 9.26 (1.6e^-6)</td>
<td>103.94 ± 14.91 (0.22)</td>
</tr>
<tr>
<td>zif1-2</td>
<td>134.88 ± 19.89 (6.5e^-6)</td>
<td>121.83 ± 12.41 (1.4e^-5)</td>
<td>106.19 ± 16.16 (0.13)</td>
</tr>
<tr>
<td>ZIFL1.1OX</td>
<td>77.38 ± 8.43 (1.2e^-7)</td>
<td>85.71 ± 9.19 (1.3e^-4)</td>
<td>99.53 ± 9.87 (0.45)</td>
</tr>
<tr>
<td>ZIFL1.3OX</td>
<td>96.74 ± 16.00 (0.27)</td>
<td>95.24 ± 11.74 (0.12)</td>
<td>106.98 ± 17.67 (0.11)</td>
</tr>
</tbody>
</table>

Numbers between parentheses indicate the P values (yeast Δtpo1 mutant versus ZIF1.1 or ZIF1.3-expressing Δtpo1 lines, or Arabidopsis wild type versus zif1 mutants or ZIF1.1-overexpressing lines) obtained by Student’s t test. ND, not determined.

<sup>a</sup>Accumulation ratio (intra/extracellular) of [14C]2,4-D or [14C]IAA in nonadapted yeast Δtpo1 mutant cells harboring either the cloning vector pGREG576 or the pGREG576/ZIFL1.1 or pGREG576/ZIFL1.3 plasmids. Results are from three independent experiments, and values represent means ± sd (n = 6).

<sup>b</sup>Relative [14C]2,4-D, [14C]IAA, or [14C]NAA accumulation in root tips from 5-d-old seedlings of the wild type (Col-0), the zif1 mutants (zif1-1 and zif1-2), and the ZIF1.1.OX or ZIF1.3.OX transgenic lines. Data are expressed relative to wild-type accumulation. Results are from three independent experiments, and values represent means ± sd (n = 12).
Figure 7. ZIFL1.1 IAA Transport Activity in Yeast and Arabidopsis.

(A) Effect of the auxin polar efflux inhibitor NPA on [14C]IAA accumulation in nonadapted yeast Δtpo1 mutant cells harboring either the cloning vector pGREG576 or the pGREG576_ZIFL1.1 or pGREG576_ZIFL1.3 plasmids. Values represent the mean of four independent experiments ± sd.

(B) IAA efflux activity of root tips from 5-d-old wild-type (Col-0), zif1-1-mutant, and ZIFL1.1-overexpressing (ZIFL1.1OX) seedlings. Efflux is calculated as a percentage of the initial [14C]IAA loading. Results are representative of three independent experiments and bars represent means ± sd (n = 18).

(C) Representative differential interference contrast microscopy images of PRs from 5-d-old wild-type (Col-0) and zif1-1-mutant (zif1-1-1 and zif1-1-2) seedlings expressing the auxin-responsive ProDR5:GUS construct. Seedlings were either not treated (Control) or treated with 1 µM IAA for 3 h at the hypocotyl-root junction (Rootward IAA) or at the root tip (Shootward IAA). Bars = 25 µm.

(D) Rootward and shootward IAA transport in PRs from 5-d-old seedlings of the wild type (Col-0), the eir1-4 and zif1-1-mutants, and the ZIFL1.1OX transgenic line in the absence (-NPA) or presence (+NPA) of 10 µM NPA. Values represent the mean of three independent experiments ± sd. For each
As discussed elsewhere (Petrásek and Friml, 2009; Löfke et al., 2013), carrier-mediated auxin transport can be altered by regulating a given transporter’s abundance, subcellular trafficking, or activity levels. Since an experimental design to measure PIN activity is still lacking, we focused on examining the distribution and steady-state levels of PIN2 at the plasma membrane by immunofluorescence labeling of native PIN2 at the PR tip (Figure 8A). As expected, no PIN2 signal was detected in the eir1-4 mutant background (Luschnig et al., 1998), whereas proper polar localization of PIN2 at the plasma membrane of wild-type root tips (i.e., shootward and rootward orientation in cells of the epidermis/LR cap and cortex, respectively) (Müller et al., 1998; Boonsirichai et al., 2003; Rahman et al., 2007), was observed. This asymmetric subcellular distribution was unaltered upon prolonged IAA treatment, consistent with previous reports that the polarity of PIN2 localization is not auxin responsive (Peer et al., 2004; Rahman et al., 2007). Importantly, even under IAA challenge, mislocalization of the auxin efflux carrier was not observed in either ZIFL1.1 loss-of-function or overexpression lines, indicating that ZIFL1.1 function does not interfere with PIN2 polar targeting. However, in the presence of 0.1 µM IAA, PIN2 fluorescence levels appeared to be altered in the ZIFL1.1 loss-of-function and transgenic lines. We therefore assessed PIN2 abundance by quantifying the corresponding immunofluorescence signal at the plasma membrane of root tip epidermal cells (Figure 8B). Under control conditions, PIN2 incidence at the cell surface was not altered by ZIFL1.1 function, as illustrated by the equivalent PIN2 levels detected in wild-type, zif1-1 mutant, and ZIFL1.1-overexpressing root tips. In seedlings grown in the presence of IAA, a substantial reduction in PIN2 plasma membrane abundance was detected in the wild type, consistent with PIN2 protein degradation following prolonged IAA treatments (Sieberer et al., 2000; Vieten et al., 2005; Abas et al., 2006; Kleine-Vehn et al., 2008a; Baster et al., 2013).

Table 3. Effect of Polar Auxin Transport Inhibitors on Wild-Type, zif1-mutant, and ZIFL1-Overexpressing Seedlings

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Relative PR elongation (%)&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Control</th>
<th>10 µM NOA</th>
<th>10 µM NPA</th>
<th>1 µM TIBA</th>
</tr>
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<tbody>
<tr>
<td>Col-0</td>
<td>100.00 ± 5.89</td>
<td>76.06 ± 6.27</td>
<td>38.09 ± 9.84</td>
<td>75.67 ± 9.91</td>
<td></td>
</tr>
<tr>
<td>zif1-1</td>
<td>100.00 ± 4.76</td>
<td>74.28 ± 8.18 (0.26)</td>
<td>59.00 ± 6.57 (6.0e&lt;sup&gt;-8&lt;/sup&gt;)</td>
<td>90.19 ± 9.82 (1.7e&lt;sup&gt;-4&lt;/sup&gt;)</td>
<td></td>
</tr>
<tr>
<td>zif1-2</td>
<td>100.00 ± 6.79</td>
<td>73.02 ± 6.36 (0.10)</td>
<td>55.02 ± 9.98 (2.6e&lt;sup&gt;-5&lt;/sup&gt;)</td>
<td>87.65 ± 4.82 (1.1e&lt;sup&gt;-4&lt;/sup&gt;)</td>
<td></td>
</tr>
<tr>
<td>ZIFL1.1OX</td>
<td>100.00 ± 3.60</td>
<td>73.40 ± 6.97 (0.15)</td>
<td>24.08 ± 3.60 (7.5e&lt;sup&gt;-6&lt;/sup&gt;)</td>
<td>57.15 ± 7.55 (1.3e&lt;sup&gt;-4&lt;/sup&gt;)</td>
<td></td>
</tr>
<tr>
<td>ZIFL1.3OX</td>
<td>100.00 ± 6.06</td>
<td>74.53 ± 6.90 (0.27)</td>
<td>38.04 ± 6.43 (0.49)</td>
<td>78.49 ± 8.63 (0.21)</td>
<td></td>
</tr>
</tbody>
</table>

Numbers between parentheses indicate the P values (the wild type versus zif1 mutants or ZIFL1-overexpressing lines) obtained by Student’s t test.<sup>a</sup>

Notably, we found that under IAA challenge, PIN2 stability at the plasma membrane of root tip epidermal cells was significantly decreased and enhanced by ZIFL1.1 loss of function and overexpression, respectively. These results are in clear agreement with the gathered physiological data (Figures 5 and 6) and indicate that, in the context of a stronger polar IAA stream, activity of the ZIFL1.1 transporter influences the steady-state levels of PIN2 at the plasma membrane, further supporting the notion that ZIFL1.1 acts as a positive regulator of shootward auxin transport.

ZIFL1 Catalyzes Potassium and Proton-Coupled Transport Activities

In a first attempt to identify the physiological substrate(s) of the two plant ZIFL1 splice isoforms, we further examined the response of Δtpo1 mutant yeast cells expressing either GFP-ZIFL1.1 or GFP-ZIFL1.3 to various additional compounds. As seen in Supplemental Figure 4D online, expression of either GFP-ZIFL1.1 or GFP-ZIFL1.3 similarly conferred enhanced yeast resistance to two weak acids, malate and acetate, as well as to the metal ions Al<sup>3+</sup> and Tl<sup>3+</sup>, as already reported for the latter in the case of the ZIFL1.3 isoform (Cabrito et al., 2009). Hence, ZIFL1 is also able to modulate ion and weak acid sensitivity in yeast. Interestingly, we also found that expression of either ZIFL1 splice variant dramatically increases yeast sensitivity to the metal ion Cs<sup>+</sup> (see Supplemental Figure 4D online). Cesium possesses similar chemical properties to potassium, and physiological studies in plants have demonstrated that Cs<sup>+</sup> competes with K<sup>+</sup> influx by entering root cells through at least some of the K<sup>+</sup> uptake systems (White and Broadley, 2000; Qi et al., 2008). We therefore decided to investigate whether the ZIFL1 transporter can also influence K<sup>+</sup> delivery to the yeast cell by evaluating the capacity of the two genotype, the percentage of root IAA transport inhibition by NPA is indicated above the corresponding bars (P < 0.01 for all genotypes in both polarities, except for shootward transport in ZIFL1.1OX, where P < 0.001). Asterisks denote statistically significant differences between zif1 mutants or the ZIFL1.1OX line and the wild type (**P < 0.01, *** P < 0.001; Student’s t test).
ZIFL1 isoforms to rescue the deficient growth of the Δqr2 deletion mutant under limiting potassium concentrations (Vargas et al., 2007). Figure 9A shows that both ZIFL1.1-GFP and ZIFL1.3-GFP were able to markedly alleviate the pronounced growth defect induced by loss of Qdr2 at low K⁺ concentrations. This finding strongly suggests that both the Arabidopsis ZIFL1.1 and ZIFL1.3 isoforms possess potassium transport activity.

Prompted by an earlier microarray study reporting ZIFL1 induction upon plant exposure to Cs⁺ (Hampton et al., 2004), we next investigated whether ZIFL1 function affects PR elongation in response to both K⁺ and Cs⁺ (Figure 9B). Like the loss-of-function mutants, ZIFL1.1-overexpressing plants responded as the wild type to the inhibitory effect induced by K⁺. However, in clear contrast with the hypersensitivity conferred by the zif1 mutations, the ZIFL1.1 transgenic line displayed decreased

Figure 8. PIN2 Immunolocalization in Arabidopsis zif1-1 Mutant and ZIFL1.1-Overexpressing Root Tips.

(A) Representative confocal laser scanning microscopy images of the PIN2 signal in root tips from 5-d-old wild-type (Col-0), eir1-4 and zifl1-1 mutant, and ZIFL1.1-overexpressing seedlings treated or not for 2 d with 0.1 μM IAA. Detection settings for staining visualization were identical for all genotypes. Arrowheads indicate the polarity of PIN2 localization. c, cortex; e, epidermis; l, LR cap. Signal intensities are coded blue (low) to yellow (high) corresponding to increasing intensity levels. Bars = 10 μm.

(B) Quantification of the PIN2 signal at the plasma membrane in root tip epidermal cells from seedlings of the wild type (Col-0), the eir1-4 and zifl1-1 mutants, and the ZIFL1.1OX transgenic line. Average fluorescence (pixel) intensity values represent the mean of three independent experiments ± SD (n > 24). Asterisks denote statistically significant differences from the wild type under each condition (*P < 0.05 and *** P < 0.001; Student’s t test).
tolerance to inhibitory concentrations of Cs⁺ (Figure 9B). This led us to examine LR formation in the presence of Cs⁺. As shown in Figure 9C, cesium induced the production of LR structures to a similar extent in wild-type, *zi1fl1* mutant, and *ZIFL1.1* overexpression lines. Furthermore, the ratio between LR primordia and emerged LRs was not significantly altered by either the mutations or enhanced *ZIFL1.1* levels, and the positive effect exerted by Cs⁺ on LR elongation was also similar in all genotypes (Figure 9C). Therefore, the effects of loss and gain of *ZIFL1* function on LR emergence are suppressed in presence of...
Cs\(^+\), which along with the effects of its own toxicity is perceived by root cells as a potassium deficiency (Hampton et al., 2004). These results thus indicate that ZIFL1.1 may play a role in Cs\(^+\) and/or K\(^+\) homeostasis in plant root cells.

As MFS transporters are believed to function mainly as proton motive force–driven secondary transporters, catalyzing uniport, symport, or antiport activities (Pao et al., 1998), we analyzed the proton dependence of ZIFL1.1 and ZIFL1.3 activity in yeast. The intracellular pH regulation is essentially sustained in S. cerevisiae by the action of the plasma membrane H\(^+\)-ATPase Pma1 (Serrano, 1978). As the activity of this proton pump and the passive proton influx through the yeast plasma membrane can be estimated by monitoring the pH of the external medium, we compared the acidification curves of \(\Delta qdr2\) cells expressing the two plant transporters under growth-limiting potassium conditions. Low potassium levels were previously shown to lead to a reduced rate of extracellular medium acidification by yeast cells lacking \(\Delta qdr2\) when compared with the wild-type strain (Vargas et al., 2007). Importantly, expression of either the GFP-ZIFL1.1 or GFP-ZIFL1.3 fusion protein was found to increase the rate of H\(^+\) efflux in \(\Delta qdr2\) cells grown in medium with low potassium (Figure 9D), demonstrating that the two Arabidopsis transporters influence proton transport across the plasma membrane of yeast cells challenged with potassium deprivation.

Finally, to investigate whether the ZIFL1.1 isoform is also involved in proton transport activity in Arabidopsis, we assessed vascular acidification in root tips using an acidification marker, the dye acridine orange. As seen in Figure 9E, acidification of the central vacuoles was notably enhanced in \(zifl-1\) mutant roots, while ZIFL1.1-overexpressing seedlings exhibited a considerable reduction in root vascular acidification when compared with the wild type. These results strongly suggest that ZIFL1.1 mediates proton efflux from the vacuolar compartment in Arabidopsis root tip cells.

**DISCUSSION**

**Alternative Splicing Determines a Dual Function for the ZIFL1 Transporter**

This work reveals an important role for an MFS transporter in modulating both polar auxin transport and stomatal movements. In fact, a key finding of this study is that alternative splicing determines a dual function for the *Arabidopsis* ZIFL1 transporter, with the full-length ZIFL1.1 and the truncated ZIFL1.3 proteins regulating polar auxin transport and drought tolerance, respectively, as demonstrated by the exclusive ability of the corresponding GFP-tagged isoforms to rescue two distinct phenotypes displayed by \(zifl1\) insertion mutants. Proof of such a double function also stems from our overexpression studies, which further show that while both the ZIFL1.1 and ZIFL1.3 splice variants code for functional membrane transporters, the remaining alternative transcript, ZIFL1.2, may either not be translated or encode a transporter with no function, at least in the biological processes analyzed.

Alternative splicing, which generates multiple transcripts from the same pre-mRNA, is an important generator of proteomic diversity and functional complexity in higher eukaryotes. Despite the fact that over 60% of the *Arabidopsis* multiexon genes are currently estimated to undergo alternative splicing (Marquez et al., 2012), information on the functional significance of this key posttranscriptional regulatory mechanism in plants is surprisingly scarce, having been uncovered for only about a dozen genes (Carvalho et al., 2012). We report here a compelling example of the functional impact of alternative splicing, which by dictating the subcellular and tissue distribution of two splice forms allows the same gene to fulfill strikingly different biological roles in Arabidopsis.

The ZIFL1.3 transcript arises from selection of an alternative 3′ splice site, causing a frame-shift mutation that results in the production of an MFS protein lacking the last C-terminal membrane-spanning domains. We found that this splicing event directs dual targeting of the ZIFL1 transporter, with the full-length ZIFL1.1 isoform localizing to the tonoplast, while the truncated ZIFL1.3 isoform is localized to the plasma membrane. On the other hand, the two splice forms possess strikingly similar activities when heterologously expressed in yeast, suggesting that the region absent from the ZIFL1.3 protein influences mainly its subcellular localization in planta but not its substrate specificity or transport activity, at least when sequestered from the plant system. The only other report of plant splice variants clearly fulfilling distinct functions is that of the *Arabidopsis* SR45 splicing factor, with SR45.1 playing a role in flower petal development and SR45.2 being required for normal root growth (Zhang and Mount, 2009). However, while both SR45 transcripts are ubiquitously expressed in plant tissues, we were able to detect the endogenous ZIFL1.3 transcript only in leaf and not in root tissues. Moreover, the ZIFL1.3-YFP fusion is undetectable in root tissues, whereas the ZIFL1.1-YFP fusion is observed at high levels in root cell vacuolar membranes. This suggests that tissue-specific factors differentially affect ZIFL1 pre-mRNA processing and/or splice variant stability in leaves and roots. Identification of the precise molecular mechanisms controlling the tissue distribution of the ZIFL1 transcripts is beyond the scope of this study but represents an exciting avenue of future work.

Unfortunately, although the GFP-tagged ZIFL1.3 isoform is functionally active, we were unable to visualize it in leaf stomatal guard cells where significant ZIFL1 promoter activity is detected. Most transporters or channels are present at very low copy numbers in this particular cell type and can hardly be visualized using their own promoter (Nagy et al., 2009; Meyer et al., 2010). Nevertheless, the unique ability of the ZIFL1.3 isoform to reduce transpiration rates by regulating stomatal movements, which is in agreement with the description of ZIFL1 as a signature gene in the transcriptional response of three different *Arabidopsis* ecotypes exposed to elevated ambient CO\(_2\) levels (Li et al., 2006), along with the subcellular localization observed in protoplasts and tobacco cells strongly support the notion that this isoform indeed resides at the guard cell plasma membrane. The restricted ZIFL1.3 functionality also confirms the absence of a link between the root and drought phenotypes. In this respect, ZIFL1 function resembles that of the *Arabidopsis* multidrug resistance–associated protein ABCC5, which confers partial drought insensitivity upon water stress by regulating stomatal apertures,
but also controls PR elongation and branching (Gaedeke et al., 2001; Suh et al., 2007). In a subsequent study, Nagy et al. (2009) elegantly demonstrated that guard cell-targeted expression of ABCG5 partially relieves the defective stomatal responses of the corresponding null mutant without rescuing its root phenotype. By contrast, the ability of the Arabidopsis vacuolar H+-pyrophosphatase AVP1 to withstand drought stress clearly correlates with its positive effect on root development through the facilitation of auxin fluxes (Li et al., 2005; Park et al., 2005).

**ZIFL1.1 Participates in Auxin Efflux during Shootward Transport in the Root**

Importantly, by directly measuring root IAA transport rates in both polarities and by indirectly visualizing auxin maxima in the root tip, we demonstrate that the full-length ZIFL1.1 transporter participates in shootward auxin redistribution from the apex to the elongation zone in the Arabidopsis root. Accordingly, many if not all aspects of the pleiotropic phenotypes caused by loss and gain of ZIFL1.1 function are consistent with impaired shootward auxin transport, which is actually sufficient to sustain both PR elongation and gravitropic bending (Rashotte et al., 2000). A role in shootward auxin transport is also in agreement with the particularly strong ZIFL1.1 expression detected in the epidermal cell layer of the root apical meristem and the transition zone, where shootward localization of the auxin carriers AUX1 and PIN2 has been demonstrated to control shootward auxin movement and hence root gravitropism (Swarup et al., 2005; Wisniewska et al., 2006). Interestingly, the ZIFL1.1 transporter is also highly expressed in root meristematic cortical cells, where rootward localization of PIN2 is required for an optimal gravitropic response (Rahman et al., 2010). More puzzling is the finding that ZIFL1.1 function is specifically required at the emergence phase of LR formation: first, because the auxin shootward stream in the PR promotes LR initiation rather than emergence (Wu et al., 2007), and second because the ZIFL1 promoter is strongly activated in the LR but only after emergence. A recent study showed that increased activity of the auxin influx carrier LAX3 in cortical and epidermal cells directly overlaying new LR primordia promotes LR emergence, likely by reinforcing the auxin-dependent localized induction of a subset of cell wall-remodeling enzymes that are crucial for cell wall separation at this step (Swarup et al., 2008). In addition, the effect that ZIFL1.1 exerts on LR elongation may result mainly from the auxin transport it mediates in the LR itself, as previously shown for the auxin efflux carrier ABCB1 (Wu et al., 2007).

The opposite alterations in auxin inhibition of PR elongation caused by loss and gain of ZIFL1.1 function are consistent with a role in root cellular auxin efflux. Indeed, mutations in a transporter influencing auxin influx would be expected to confer reduced auxin sensitivity, in agreement with the impaired ability of the corresponding mutant to accumulate inhibitory levels of auxin, as shown for the AUX1 influx carrier (Bennett et al., 1996). By contrast, nearly all auxin influx carrier mutants overaccumulate auxin at the root tip and are consequently more sensitive to exogenous auxins (Chen et al., 1998; Luschnig et al., 1998; Lin and Wang, 2005; Terasaka et al., 2005). Accordingly, loss of ZIFL1 function enhances and reduces auxin accumulation and efflux activity of excised root segments. The correlation between these phenotypes and root transport activity is not only qualitative but also quantitative, as both aspects are more pronounced in the presence of the synthetic auxin 2,4-D than of the naturally occurring IAA. These data were complemented by assessing the sensitivity of the different genotypes to polar auxin influx and efflux inhibitors. Importantly, only the auxin efflux inhibitors NPA and TIBA induce a differential PR response, with zfl1 mutants and ZIFL1.1 overexpression lines displaying respectively enhanced resistance and sensitivity to these compounds, as shown previously for numerous auxin efflux mutants (Chen et al., 1998; Luschnig et al., 1998; Lin and Wang, 2005). An apparent discrepancy with the proposed model regards the results obtained using the two synthetic auxins 2,4-D and 1-NAA, which have been employed experimentally to distinguish between IAA import and efflux activities, respectively (Delbarre et al., 1996; Yamamoto and Yamamoto, 1998). Rather unexpectedly, in PR elongation and root accumulation assays, no effect of loss of ZIFL1 function was found for 1-NAA, while hypersensitivity was observed for 2,4-D. However, Luschnig et al. (1998) also reported the absence of a discernible effect of 1-NAA on PR elongation of the eir1-4 mutant. On the other hand, a role in auxin influx should result in enhanced resistance, not sensitivity, of the zfl1 mutants to 2,4-D. Furthermore, while it is known that 2,4-D is poorly transported in a polar fashion, it can also constitute a weak substrate of key auxin efflux carriers among both ABCBs and PINs (Yang and Murphy, 2009). As this auxin significantly induces ZIFL1 expression (Cabrito et al., 2009), it is conceivable that the plant takes advantage of the ZIFL1.1 role in auxin transport to reallocate this function to xenobiotic resistance.

**The Tonoplastic ZIFL1.1 Transporter Assists in Polar Auxin Transport**

Several lines of evidence indicate that ZIFL1.1 modulates auxin transport indirectly. First, a tonoplast-localized transporter cannot directly catalyze auxin efflux at the plasma membrane, and though IAA conjugates and IBA provide a readily available pool of free IAA upon hydrolysis in the endoplasmic reticulum lumen or β-oxidation in the peroxisomes, respectively (reviewed in Woodward and Bartel, 2005), storage or compartmentalization of such compounds in the vacuole has not been reported. Secondly, zfl1 mutants exhibit only subtle changes in root gravitropism and PR elongation sensitivity to exogenous auxins, which are attributable to minor alterations in shootward auxin transport at the root tip. Moreover, apart from PR branching, zfl1 mutants fail to display morphological alterations under normal conditions, where the large majority of the described auxin efflux mutants exhibit specific defects in PR and/or gravitropic growth (Chen et al., 1998; Luschnig et al., 1998; Noh et al., 2001; Lin and Wang, 2005; Terasaka et al., 2005). Last but not least, ZIFL1.1 activity is NPA sensitive in planta but NPA insensitive in yeast. Our heterologous expression studies show that ZIFL1.1 is able to mediate IAA transport in yeast cells without requiring additional plant-specific factors. Indeed, transport activity of plant auxin carriers can be demonstrated in various heterologous systems (Geisler et al., 2005; Yang et al., 2006; Blakeslee et al., 2007; Yang and Murphy, 2009), but the
activation of other endogenous transporters catalyzing auxin efflux cannot always be ruled out. We found that NPA inhibition of yeast IAA transport activity is unaffected by expression of the ZIFL1.1 transporter, while in planta NPA treatment fails to reduce shootward transport in the zif1-1 mutant to the same extent as in the wild type. The latter observation, which is even more evident in the eir1-4/pin2 mutant background (this study; Rashotte et al., 2000), suggests that, by reducing shootward auxin transport rates, absence of the carrier attenuates sensitivity to the auxin efflux inhibitor. Indeed, in contrast with ZIFL1.1, PIN2-mediated IAA export activity in yeast and other heterologous systems is clearly NPA sensitive (Paciorek et al., 2006; Blakeslee et al., 2007; Yang and Murphy, 2009).

We further propose that fine-tuning of polar IAA efflux by ZIFL1.1 relies on a protective effect on PIN2 plasma membrane stability under conditions of high IAA flow that normally trigger PIN2 degradation. Recent studies have established auxin as the main regulator of its own asymmetric distribution (reviewed in Löfke et al., 2013). Dynamic polar sorting of PINs at the plasma membrane is sustained by repeated steps of endocytotic internalization and recycling back to the plasma membrane via exocytosis (Geldner et al., 2001; Dhonukshe et al., 2007). This constitutive cycling of PIN proteins controls not only their subcellular localization, but also their plasma membrane abundance (Kleine-Vehn et al., 2008b). While physiological IAA concentrations inhibit the internalization step of PIN2 cycling, promoting its stability at the plasma membrane (Paciorek et al., 2005), extended treatments with high IAA concentrations trigger PIN2 protein degradation through lytic vacuolar targeting and proteasomal activity, thus reducing its plasma membrane incidence (Sieberer et al., 2000; Kleine-Vehn et al., 2008a; Laxmi et al., 2008). The combination of these auxin antagonistic effects on intracellular trafficking and proteasome-mediated degradation allows the positional control of PIN2 activity sustaining root gravitropism (Abas et al., 2006; Baster et al., 2013).

As the vacuolar ZIFL1.1 transporter does not appear to affect PIN2 polar distribution, it is tempting to speculate that ZIFL1.1 activity modulates PIN2 steady-state levels at the plasma membrane by interfering with its vacuolar targeting and/or degradation, as already reported for the Modulator of PIN regulators (Malenica et al., 2007). At present, we cannot exclude the possibility of a more general effect on auxin efflux, as PIN2-mediated shootward auxin transport has been shown to be supported for instance by the action of ABCB4 (Terasaka et al., 2005; Lewis et al., 2007). On the other hand, ZIFL1.1 activity could also influence PIN2 activity levels, namely, through changes in plasma membrane electrochemical potential. In fact, our yeast extracellular and plant vacuolar acidification assays indicate that ZIFL1.1 may facilitate polar auxin transport by enhancing the release of protons from the vacuole, thus increasing H+ availability for plasma membrane ATPases and the proton-driving force for cellular auxin transport. Interestingly, AVP1 has been shown to promote auxin-mediated organ development by influencing apoplastic acidification through its action on the abundance and activity of the plasma membrane P-ATPase and by directly affecting trafficking or stability of the PIN1 carrier (Li et al., 2005).

Potassium as the Physiological Substrate of the ZIFL1 Transporter

The identification of the physiological substrate of a membrane transporter is of prime importance in dissecting the precise molecular mechanisms underlying its function. Our yeast experiments show that the effect of ZIFL1 expression is not restricted to auxins, influencing also the response to other weak acids and a few ions. Interestingly, ZIFL1 markedly increases yeast sensitivity to Cs+. Results obtained by Hampton et al. (2004) indicate that the main effect that cesium induces in plants along with its own toxicity is potassium starvation. In support of this notion, complementation experiments with the Δqdr2 yeast mutant indicate high-affinity K+ transport activity for both ZIFL1 splice isoforms. Indeed, potassium fluxes, in particular those mediated by K+ channels, have been previously shown to modulate auxin-induced cell elongation growth (Christian et al., 2006; Fuchs et al., 2006). Moreover, deletion of the Arabidopsis plasma membrane K+ transporter Tiny Root Hair 1 (TRH1) leads to auxin-related phenotypes similar to those reported here for the zif1 mutants (Vicente-Agullo et al., 2004). This unexpected function of the potassium carrier was suggested to rely on its ability to generate ionic and electric gradients that would favor auxin efflux through specific transporters (Vicente-Agullo et al., 2004), as proposed here for the ZIFL1.1 transporter. Interestingly, Rigos et al. (2013) recently demonstrated that TRH1 activity is required for proper polar localization of the PIN1 carrier in the root. Cesium suppression of the LR emergence and elongation effects along with the altered PR elongation in response to cesium caused by altered ZIFL1.1 levels provide a first link between the potassium and auxin transport activities of this MFS carrier. The Arabidopsis Nitrate Transporter NRT1.1 has been recently proposed to repress LR growth at low nitrate availability by promoting shootward auxin transport out of these roots, thus connecting nutrient sensing and auxin-dependent developmental adaptation (Krouk et al., 2010). Interestingly, ZIFL1 expression appears to be highly repressed in the nrt1.1 knockout background (Muñoz et al., 2004).

Finally, the notion of K+ as the physiological substrate of the ZIFL1 MFS transporter is also consistent with ZIFL1.3’s function in the regulation of stomatal movements, since potassium fluxes are well-established regulators of guard cell turgor in response to endogenous and environmental cues (Ward et al., 2009). Nagy et al. (2009) showed that ABC5C possesses high-affinity myo-inositol hexakisphosphate transport activity, thus linking this substrate to a pleiotropic function in PR elongation, seed mineral cation and phosphorus status regulation, stomatal aperture control, and drought stress tolerance. This study provides compelling evidence that the two splice isoforms of the ZIFL1 MFS carrier share K+ transport activity likely to influence membrane proton gradients and that their distinct biological roles stem solely from their different subcellular and tissue distribution.

METHODS

Plant Materials and Growth Conditions

The Arabidopsis thaliana Col-0 ecotype was used as the wild type in all experiments. Seeds of the T-DNA insertion mutants zif1-1 (SALK_030680) and zif1-2 (GABI_052H08) were obtained from Nottingham
Arabidopsis Stock Centre. The exact T-DNA insertion sites were confirmed using gene-specific primers (see Supplemental Table 2 online) and primers annealing at the T-DNA borders, which also allowed PCR-based genotyping to identify homologous lines.

Plant transformation was achieved by the floral dip method (Clough and Bent, 1998) using Agrobacterium tumefaciens strain EHA105. Highly similar results were obtained following extensive characterization of all transgenic lines generated for each construct, and representative results for one line are presented.

Seeds were surface sterilized, sown on Murashige and Skoog (1962) medium solidified with 0.8% agar, stratified at 4°C for 3 d in the dark, and placed in a growth chamber where they were transferred to soil after 2 to 3 weeks. All phenotypical assays were performed in a climate-controlled growth cabinet. Plants were cultivated either under 16-h (long-day) or 8-h (short-day) photoperiod (80 μmol m−2 s−1 white light) conditions at 23°C (light period)/18°C (dark period) and 60% RH. Hormones, antibiotics, and other compounds were purchased from Sigma-Aldrich.

Gene Expression and Sequencing Analyses
RT-PCR analyses were performed as previously described (Remy et al., 2012) using primers designed to detect ZIFL1.1, ROC10, and UBO10 expression (see Supplemental Table 2 online). For native ZIFL1 promoter reporter gene experiments, a fragment including the 1778 bp immediately upstream of the first start codon was PCR amplified (see Supplemental Table 2 online) from genomic DNA and inserted via the SacI-Sacl restriction sites into the pKGWFS7 plasmid (Karimi et al., 2002). After agroinfiltration of the resulting ProZIFL1::GUS:GFP construct into wild-type plants, 13 independent transformants were recovered. For synthetic DR5 promoter reporter gene experiments, wild-type and zif1 mutant plants were transformed with the ProDR5::GUS construct inserted in the pEarleyGate301 vector (Ulmasov et al., 1997; Earley et al., 2006; Delker et al., 2010), and six independent transformants per genotype were isolated. Histochemical staining of GUS activity was performed as described by Sundaresan et al. (1995).

Sequencing analyses were performed on a 3130x/Genetic Analyzer using the BigDye Terminator v1.1 cycle sequencing kit according to the manufacturer’s instructions (Applied Biosystems).

Phenotypical Assays
Water loss assays were performed on irrigated nonbolting plants by weighing either detached entire rosette or four rosette leaves at various time intervals at room temperature (~22°C). Stomatal closure assays were performed as described by Bright et al. (2006), with more than 60 apertures recorded per treatment. PR elongation assays and LR parameter evaluation were performed as previously described (Remy et al., 2012). LR primordial stages were scored under the microscope and defined according to Malamy and Benfey (1997). For hypocotyl elongation assays, seeds were sown directly on control or auxin-containing medium plates set in the vertical position and stratified for 5 d in the dark. For root tip reorientation assays, 5-d-old seedlings grown vertically on control medium were transferred to fresh medium and allowed to grow for an additional 2 d. Plates were then rotated 90°C anticlockwise and kept in the dark before the angle of root tips from the vertical plane was scored. Stomatal apertures, PR and hypocotyl elongation, and LR length and root deviation angles were measured on scanned images using the ImageJ software (http://rsbweb.nih.gov/ij/).

Subcellular Localization Studies
To generate the ZIFL1-YFP and ZIFL1-GFP fusions, each ZIFL1 transcript was PCR amplified (see Supplemental Table 2 online) using root (ZIFL1.1 and ZIFL1.2) or leaf (ZIFL1.3) cDNA as a template and independently inserted via the XhoI-Pacl restriction sites into the YFP- or GFP-tagged versions of the pBA002 vector (Kost et al., 1998). Arabidopsis protoplasts were generated as described by Yoo et al. (2007), transfected with the YFP constructs by polyethylene glycol transformation (Abel and Theologis, 1994), and analyzed by confocal microscopy. Transient coexpression of the GFP constructs with the tonoplasma marker γ-Tonoplast Intrinsic Protein (TIP)-mCherry or the plasma membrane marker Plasma Membrane Intrinsic Protein 2A (PPIP2A)-mCherry (Nelson et al., 2007) and the pBIN-NA construct (Silhavy et al., 2002) in leaf abaxial epidermal cells of tobacco (Nicotiana tabacum) was performed via the agroinfiltration procedure described by Voinnet et al. (2003) using Agrobacterium strain GV3101.

For generation of stable transgenic lines, the 35S promoter in the pBA002-ZIFL1.1-GFP and pBA002-ZIFL1.3-GFP plasmids was replaced with the ZIFL1 promoter. For each construct, six transgenic lines were recovered upon transformation of zif1-1 or zif1-2 mutant plants.

Genomic Complementation and Overexpression Analyses
For genomic complementation, a 5599-bp fragment encompassing the entire ZIFL1 gene and including the 1778-bp promoter sequence described below was PCR amplified (see Supplemental Table 2 online) from genomic DNA and inserted into the promoterless version of the pBA002 vector via the HindIII-XbaI restriction sites. The corresponding construct was introduced into zif1-1 and zif1-2 mutant plants, with three complementation lines being recovered for each mutant allele. ZIFL1 overexpression constructs were obtained as described for the YFP plasmids, except that the corresponding fragments were inserted via the XhoI-Aacl restriction sites into the pBA002 background. After agroinfiltration of wild-type plants, three transgenic lines independently overexpressing each ZIFL1 transcript were selected.

Yeast Experiments
The parental Saccharomyces cerevisiae strain BY4741 (MATα, his3Δ1, leu2Δ0, met15Δ0, ura3Δ0) and the derived deletion mutants BY4741Δtop1 (MATα, his3Δ1, leu2Δ0, met15Δ0, ura3Δ0, YLL028w::kanMX4) and BY4741Δtop2Δ (MATα, his3Δ1, leu2Δ0, met15Δ0, ura3Δ0, YLL121W::kanMX4) were used in this study. The Arabidopsis ZIFL1.1 and ZIFL1.3 coding sequences were cloned into the pREG576 vector (Jansen et al., 2005), and expression of the corresponding GFP fusion proteins was tested by immunoblotting and fluorescence microscopy analyses as described by Cabrito et al. (2009). Strains and vector were acquired from the Euroscarf collection. Susceptibility and spot assays were performed as described previously (Cabrito et al., 2009) in minimal growth MMB-U liquid and agarized medium, respectively. To test the chemical stress inducers, 2,4-D, IAA, CsCl, Al3+(SO4)2−, TICl−, l-malic acid, and acetic acid stock solutions (pH was adjusted to 4 for the weak acids) were added at the specified concentration. Growth under K+ limitation was evaluated on agarized amonium phosphate basal medium (KNA) supplemented with 0.2 or 2 mM KCl as previously described (Vargas et al., 2007). Auxin accumulation assays were performed as described by Cabrito et al. (2009), using 0.3 mM/0.5 mM of cold 2,4-D/IAA (Sigma-Aldrich) and 0.5 μM/0.5 μM [3H]labeled 2,4-D/IAA (American Radio-labeled Chemicals) in the absence or presence of NPA (10 μM). To compare the in vivo active export of protons, the external medium pH was monitored as in Vargas et al. (2007), using liquid KNA medium supplemented with 0.5 mM KCl.

Auxin Transport Assays
Radiolabeled auxin accumulation was assayed in root-tip segments as described by Ito and Gray (2006), except that radiolabeled auxins were
used at a final concentration of 4.5 μM [14C]IAA (50 mCi/ml), 7.5 μM [14C]2,4-D (50 mCi/ml), or 0.3 μM [14C]NAA (10 mCi/ml) (American Radiolabeled Chemicals). To examine [14C]IAA efflux, the same protocol was applied, except that after the 1-h incubation period, root tips were rinsed with three changes of uptake buffer and further incubated in this buffer for 1 h before harvesting.

Rootward and shootward root auxin transport was assayed as described by Lewis and Muday (2009). After treatment with either 100 nM [3H]IAA or [3H]benzoic acid (American Radiolabeled Chemicals) applied as agar droplets for 13 h at the hypocotyl-root junction (rootward) or for 5 h at the root tip (shootward), 5-mm segments from the root apex (rootward) or spanning from 2 to 7 mm from the root tip (shootward) were excised.

Immunofluorescence Protein Localization Studies

For PIN2 immunofluorescence localization assays, 3-d-old seedlings grown vertically on control medium were transferred to fresh medium supplemented or not with 0.1 μM IAA and allowed to grow for an additional 2 d. Whole-mount immunolocalization in Arabidopsis roots was performed as described previously (Sauer et al., 2006). Rabbit anti-PIN2 primary antibody was used at a dilution of 1:1000 and fluorochrome-conjugated anti-rabbit-CY3 secondary antibody (Dianova) was diluted 1:600.

Microscopy

Differential interference contrast and confocal images were taken with a DM LB2 microscope (Leica) and an LSM 510 or 710 laser scanning microscope equipped with a Meta detector (Zeiss), respectively. For cell wall staining, roots were incubated in 10 μg/ml iodide propidium for 10 min. Acidine orange staining was performed as described by Hirano et al. (2011). Fluorescence detection parameters (laser intensity, offset, gain, and pinhole settings) were set so that the fluorescence signal emitted by zifl-1 mutant root tips was just below the saturation threshold. All micrographs were acquired using identical fluorescence parameters to allow comparison with wild-type and ziflF1,1-overexpressing root tips. Postimaging, average fluorescence intensity of seven epidermal cells within a single file located 600 μm from the root apex was recorded. For PIN2 immunofluorescence signal quantification, at least seven root tip micrographs per genotype per condition were captured using identical confocal settings. Postimaging, average fluorescence intensity of the PIN2 signal at the plasma membrane was recorded within each single epidermal cell file. Excitation wavelengths used to detect fluorescence were 488 nm for GFP and acidine orange, 514 nm for YFP, 543 nm for iodide propidium and mCherry, 458 nm for autofluorescence, and 561 nm for CY3. Emitted fluorescence was monitored at detection wavelengths longer than 560 nm for iodide propidium and autofluorescence, between 565 and 615 nm for mCherry, between 535 and 590 nm for YFP and acidine orange, between 500 and 550 nm for GFP, and between 569 and 649 nm for CY3.

Accession Numbers

Arabidopsis Genome Initiative locus identifiers for the genes mentioned in this article are as follows: ZIFL1 (At5g13750), ROC1 (At4g38740), UBQ10 (At4g05320), PIN2 (At5g57090), γ-TIP (At2g36830), and PIP2A (At3g53420). T-DNA insertion mutants are as follows: zifl-1 (SALK_030680), zifl-2 (GABI_052H06), and eir1-4 (SALK_089114).

Supplemental Data

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

Supplemental Figure 1. Sequencing Analysis of the ZIFL1 Transcript Tissue Distribution.

Supplemental Figure 2. Auxin-Related Hypocotyl Phenotype of ZIFL1 Loss-of-Function Mutants and ZIFL1-Overexpressing Lines.

Supplemental Figure 3. Effect of IBA and 1-NAA on the zifl1-1 Mutant.

Supplemental Figure 4. Heterologous Expression of ZIFL1.1 and ZIFL1.3 in Yeast.

Supplemental Figure 5. Auxin-Responsive ProDR5:GUS Expression in Wild-Type and zifl1 Mutant Roots.

Supplemental Table 1. Benzoic Acid Diffusion in Roots of the Wild Type, eir1-4, and zifl1-1 Mutants and ZIFL1.1-Overexpressing Lines.

Supplemental Table 2. Sequences of Primers Used.

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AUTHOR CONTRIBUTIONS

E.R., P.B., M.C.T., J.F., I.S.-C., and P.D. designed the research. E.R., R.A.B., M.C.T., J.F., I.S.-C., and P.D. analyzed the data. The article was written by E.R. and P.D. with contributions from T.R.C., M.C.T., and I.S.-C.

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