Intron Retention in the 5′UTR of the Novel ZIF2 Transporter Enhances Translation to Promote Zinc Tolerance in Arabidopsis

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Abstract

Root vacuolar sequestration is one of the best-conserved plant strategies to cope with heavy metal toxicity. Here we report that zinc (Zn) tolerance in Arabidopsis requires the action of a novel Major Facilitator Superfamily (MFS) transporter. We show that ZIF2 (Zinc-Induced Facilitator 2) localises primarily at the tonoplast of root cortical cells and is a functional transporter able to mediate Zn efflux when heterologously expressed in yeast. By affecting plant tissue partitioning of the metal ion, loss of ZIF2 function exacerbates plant sensitivity to excess Zn, while its overexpression enhances Zn tolerance. The ZIF2 gene is Zn-induced and an intron retention event in its 5′UTR generates two splice variants (ZIF2.1 and ZIF2.2) encoding the same protein. Importantly, high Zn favours production of the longer ZIF2.2 transcript, which compared to ZIF2.1 confers greater Zn tolerance to transgenic plants by promoting higher root Zn immobilization. We show that the retained intron in the ZIF2 5′UTR enhances translation in a Zn-responsive manner, markedly promoting ZIF2 protein expression under excess Zn. Moreover, Zn regulation of translation driven by the ZIF2.2 5′UTR depends largely on a predicted stable stem loop immediately upstream of the start codon that is lost in the ZIF2.1 5′UTR. Collectively, our findings indicate that alternative splicing controls the levels of a Zn-responsive mRNA variant of the ZIF2 transporter to enhance plant tolerance to the metal ion.

Introduction

The transition metal zinc (Zn) is a micronutrient essential for optimal plant growth and development, serving as a catalytic co-factor or structural motif in numerous proteins required for basic cellular processes. However, its propensity to inactivate other crucial proteins also renders Zn a potentially toxic element that can adversely affect plant growth when present in excessive amounts [1]. To deal with these opposing effects and adjust to environmental fluctuations in Zn availability, plants developed a tightly controlled homeostatic network aimed at ensuring an adequate supply of the nutrient while preventing its toxic build-up at the cellular and whole-plant levels [2].

Zn is acquired from the soil as a divalent cation that once absorbed into the root epidermis moves mostly symplastically through the adjacent cell layers to the central stele. After active loading into the xylem vessels, Zn is translocated to the shoot via the transpiration stream and subsequently transferred to the phloem before allocation to aerial organs [3]. While plants adapt to zinc shortage prevalently by activating cellular Zn uptake particularly at the root-soil interface [4], cellular Zn detoxification mechanisms are primarily intended at restricting the accumulation of free cytosolic Zn²⁺, mainly through its extrusion to the apoplasm [5], chelation with specific ligands [6,7] and/or vacuolar compartmentalisation [8]. At the whole-plant level, tolerance to excess Zn is achieved through the rearrangement of its tissue partitioning via enhanced Zn sequestration in leaves [9], whereas within the root both immobilisation in the outer cell layers [10] and exclusion from the epidermis [11] contribute to limit Zn entry into the root symplasm.

All steps in plant Zn homeostasis, from initial acquisition by the root to subsequent distribution among different tissues and subcellular trafficking, require the concerted action of multiple Zn membrane transport systems. Three groups of such carrier proteins have been primarily characterised in Arabidopsis thaliana. While members of the ZIP/IRT (Zinc-regulated transporter, Iron-regulated transporter-like Protein) family have been proposed to constitute the major cellular Zn uptake system [12–14], HMAs (Heavy–metal transporting ATPases) and MTPs (Metal Tolerance Proteins), which belong to the P₁B-type ATPases and Cation Diffusion Facilitator (CDF) families, respectively, are essential for full Zn tolerance owing to their ability to drive either Zn whole-plant partitioning or organelle sequestration/exclusion [8–10,15–19]. The recent implication of two novel Arabidopsis transporters,
the fact that over 60% of the genome annotation (TAIR V10; www.arabidopsis.org), 34 MFS superfamily [25,26]. Interestingly, according to the current transporter after the ATP-Binding Cassette (ABC) Arabidopsis members, the MFS represents the second widest group of to all classes of organisms. With more than 120 predicted using chemiosmotic gradients as an energy source and ubiquitous plants is poorly understood, having been uncovered for only a dozen genes [28,29].

PCR2 (Plant Cadmium Resistance 2; [11]) and the Major Facilitator Superfamily (MFS) carrier ZIF1 (Zinc-Induced Facilitator 1; [20]), in Zn tolerance suggests that additional plant Zn transport components are still to be discovered.

On the fringe of the vast majority of plants that like A. thaliana typically possess basal Zn tolerance, a specialized flora displays naturally selected tolerance to extremely high Zn concentrations, being able to accumulate considerable Zn amounts in their aboveground parts without developing toxicity symptoms. This hyperaccumulation strategy relies primarily on efficient xylem loading and vacuolar sequestration in leaves, thus inducing a systemic Zn-deficiency status that stimulates root Zn uptake [21].

Cross-species transcriptomic profiling particularly in the hyper-accumulator Arabidopsis halleri, a close A. thaliana relative, has established that key Zn transporters, such as HMA4, MTP1, HMA3 and ZIP/IRTs, are not different but rather constitutively overexpressed in hyperaccumulators [22–24]. Considering not only differential expression between the two Arabidopsis species but also individual regulation by exogenous Zn, Becher et al. [23] provided a shortlist of 50 promising candidate genes for a role in Zn homeostasis that notably includes At2g48020 encoding a yet uncharacterised MFS carrier, which given the Arabidopsis ZIF1 role in zinc tolerance [20] raises particular interest.

MFS transporters are single-polypeptide secondary carriers using chemiosmotic gradients as an energy source and ubiquitous to all classes of organisms. With more than 120 predicted members, the MFS represents the second widest group of Arabidopsis transporters after the ATP-Binding Cassette (ABC) superfamily [25,26]. Interestingly, according to the current genome annotation (TAIR V10; www.arabidopsis.org), 34 MFS genes undergo alternative splicing, a key generator of proteome diversity and functional complexity in higher eukaryotes. Despite the fact that over 60% of the Arabidopsis multi-exon genes are currently estimated to produce more than one transcript via this mRNA processing mechanism [27], its biological significance in plants is poorly understood, having been uncovered for only a dozen genes [28,29].

Here, we report that the MFS transporter encoded by the At2g48020 gene, which we named ZIF2, sustains plant Zn tolerance in A. thaliana by mediating root vacuolar sequestration of the metal, thus preventing its translocation to the shoot. Importantly, we found that elevated Zn levels favour expression of a ZIF2 splice variant whose longer 5' untranslated region (UTR) enhances the translation efficiency of the ZIF2 transporter and hence plant Zn tolerance.

Results

The ZIF2 Promoter Is Ubiquitously Active in Arabidopsis

To initiate the characterisation of the ZIF2 gene (At2g48020), we examined its organ- and tissue-specific expression pattern by means of reporter gene experiments. Staining of transgenic Arabidopsis lines stably expressing β-glucuronidase (GUS) under the control of the ZIF2 promoter revealed that the latter is active in most plant organs (Figure 1A–H). In particular, ZIF2 was highly expressed at all stages of floral development (Figure 1A), with flower buds displaying exclusive staining of the style, stamen filaments and sepals (Figure 1B). Promoter activity was also detected in leaf mesophyll cells (Figure 1C) and strongly throughout the root system (Figure 1D–H). Although homogenous GUS coloration was observed all along the primary root (PR) (Figure 1D), the ZIF2 promoter was particularly active at the root tip (Figure 1E). We detected very intense GUS staining in lateral root (LR) primordia (Figure 1F, G), but ZIF2 was also abundantly expressed in elongating LRs (Figure 1H).

We further examined the precise root tissue distribution of ZIF2 promoter activity using transgenic plants expressing the green fluorescent protein (GFP) under the control of the native ZIF2 promoter (Figure 1I–L). At the PR tip, ZIF2 promoter activity was confined to the endodermal and cortical cell layers of the elongation and transition zones, while absent from the apical meristem, the quiescent centre, columella cells and the lateral root cap (Figure 1I). Similarly, in the mature portion of the PR, the GFP signal was detected in both the endodermis and the cortex (Figure 1K). A slightly different layer-specific pattern was observed in LRs, with ZIF2 expression appearing largely restricted to the cortex, particularly at the tip (Figure 1J, L). ZIF2 promoter activity in flowers and leaves was insufficient to allow detection of the GFP signal. Collectively, the above results suggest that ZIF2 exerts a prominent role in the root system, particularly in the PR.

High Zn Induces ZIF2 Expression and Promotes an Intron Retention Event in Its 5'UTR

According to the current genome annotation (TAIR V10), the Arabidopsis ZIF2 gene contains 17 exons (Figure 2A) and generates two distinct mRNAs that differ solely in their 5'UTRs. In order to verify the accuracy of these predictions and examine the tissue-specific distribution of the ZIF2 transcripts, we performed RNA Ligase-Mediated-5' Rapid Amplification of cDNA Ends (RLM-5' RACE) in different vegetative and reproductive tissues, i.e. in seedlings, adult leaves and flowers. Given the high ZIF2 promoter activity detected throughout the root system (see Figure 1) and the differential expression of the ZIF2 orthologue reported in the Zn hyperaccumulator A. halleri [23], we also performed RLM-5' RACE in roots of seedlings exposed to Zn toxicity. As shown in Figure 2B, we were able to identify nine alternative 5'UTRs and named the corresponding transcripts ZIF2.1 (At2g48020.1 according to TAIR V10) to ZIF2.9. A common start codon is predicted for all ZIF2 transcripts, and we verified the absence of an in-frame ATG or alternate start codon.
in the nine 5’UTRs as well as of any alterations in the coding sequence. Hence, all ZIF2 transcripts are predicted to encode the same 53-kDa 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 [25]. Distribution of the ZIF2 transcripts was found to exhibit tissue specificity (Figure 2B). Interestingly, the ZIF2.1 and ZIF2.2 splice variants, which result from retention of a 139-nt intron in the 5’UTR, appeared to be predominantly expressed in roots challenged with high Zn concentrations, hinting at a role for this intron retention event in roots exposed to excess Zn.

We then examined the tissue-specific distribution of the ZIF2.1 and ZIF2.2 splice variants by RT-PCR (Figure 3A, B). Consistent with the promoter activity profile shown in Figure 1, RT-qPCR analysis revealed that ZIF2 is expressed throughout plant development, particularly in roots and flowers of mature plants, and to a lesser extent in young seedlings, while lower transcript levels were detected in other aboveground tissues such as developing siliques, stems or leaves (Figure 3B). By contrast, the

Figure 1. The ZIF2 promoter is active in most Arabidopsis tissues. (A–H) Differential interference contrast microscopy images of GUS-stained transgenic plants carrying the ProZIF2:GFP-GUS reporter construct. ZIF2 promoter activity in floral buds (A), a mature flower (B), a young leaf (C), the primary root (D), the primary root tip (E), a lateral root primordium at stage V (F) and stage VII (G) and a mature lateral root (H). Scale bars, 1 mm (A), 200 μm (B, C), or 50 μm (D–H). (I–L) Confocal laser scanning microscopy images of transgenic root tissues carrying the ProZIF2:GFP-GUS reporter construct. ZIF2 promoter activity in the primary root (I, K) and a mature lateral root (J, L). ep, epidermis; c, cortex; en, endodermis; cc, central cylinder. Cell walls were stained with propidium iodide. The GFP and propidium iodide signals are visualized by green and red coloration, respectively. Scale bars, 25 μm (I, J), or 10 μm (K, L).

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and ZIF2.2 transcripts were detected at very low levels in all tissues analysed, except in roots of mature plants where the expression level of the two splice variants combined was found to represent more than half of total ZIF2 expression. ZIF2.1 and ZIF2.2 expression was also markedly induced during leaf senescence (Figure 3A, B).

We next asked whether ZIF2 expression would be responsive to exogenous Zn in A. thaliana roots. Both total ZIF2 expression and the equivalent amounts of the ZIF2.1 and ZIF2.2 splice variants detected in root tissues under control Zn conditions (30 μM) appeared unchanged by Zn deficiency (Figure 3C, D). However, concomitant with a significant induction of ZIF2 expression, a clear shift in the ZIF2 splicing pattern occurred after at least 48 h of challenge with high Zn (Figure 3C, D; Figure S1). Indeed, root ZIF2.2 expression was markedly up-regulated in a concentration-dependent manner following exposure to Zn excess, even as slight

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**Figure 2. The Arabidopsis ZIF2 gene encodes multiple transcripts.** (A) Exon/intron organization of the ZIF2 gene and T-DNA insertion site in the zif2-1 mutant. Boxes and lines between boxes denote exons and introns, respectively. The triangle depicts the site of the T-DNA insertion. F1, F1’, F2, F3, R1, R2, and R3 indicate the location of the primers used to detect ZIF2 expression in Figures 3A, 3C, 7A, S1 and S4. Scale bar, 200 bp. (B) Structure of the alternative ZIF2 transcripts. The coding sequence (CDS) is shown in black and the UTRs in white. Grey boxes indicate portions of the 5'-UTR overlapping with the retained intron. RACE1 to RACE6 indicate the location of the gene-specific primers used in the 5’ RLM-RACE experiment. Occurrence of each ZIF2 transcript was determined according to its detection (number of corresponding 5’ RLM-RACE clones) or not (-) by RLM-5’ RACE in the indicated tissue. Scale bar, 200 nt.

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as 100 μM, while a considerable increase in ZIF2.1 steady-state levels was also observed but of lower magnitude and only under toxic Zn excess (250 μM). Thus, both total expression of the ZIF2 gene and alternative splicing of its precursor mRNA (pre-mRNA) are regulated by the plant’s Zn status, with high levels of the metal ion promoting retention of the 5’UTR intron.

ZIF2 Encodes a Functional Tonoplast-Localised Transporter

To determine the subcellular localisation of the protein encoded by the ZIF2 transcripts, we generated C-terminal yellow fluorescent protein (YFP) or GFP fusions of each cDNA under the control of the 35S promoter. Confocal microscopy analysis of the YFP signal upon transient or stable expression in Arabidopsis protoplasts (Figure 4A–C) or root apices (Figure 4D–F), respectively, suggested that ZIF2 localises to the vacuolar membrane. In co-localisation experiments in tobacco leaf epidermal cells using specific tonoplast and plasma-membrane RFP markers [30], the ZIF2.2-GFP fusion protein co-localised exclusively with the tonoplast marker, whereas it did not match the distribution of the plasma membrane marker (Figure 4G–R), confirming that the ZIF2 transporter is primarily targeted to the tonoplast of plant cells.
The transport properties of the plant ZIF2 carrier were then explored by means of heterologous expression in Saccharomyces cerevisiae. Correct expression of the GFP-ZIF2 fusion protein in yeast was confirmed by immunoblotting (Figure S2A) and fluorescence microscopy (Figure S2B). To determine whether the plant transporter is able to influence Zn delivery to yeast cells, the GFP-ZIF2 fusion was introduced into the \( \text{Dzrt1zrt2} \) and \( \text{Dzrc1cot1} \) mutant strains, which are sensitive to low and high Zn concentrations, respectively [31,32]. As seen in Figures 5 and S2C, yeast cells expressing GFP-ZIF2 displayed no significant growth difference with respect to cells transformed with the empty vector in normal nutrition medium (2.5 \( \mu \text{M Zn}^{2+} \)). Due to the lack of the Zn importers Zrt1 and Zrt2, the \( \text{Dzrt1zrt2} \) mutant exhibited as expected [32] a severe growth retardation under Zn-limiting conditions that was further exacerbated by GFP-ZIF2 expression. Conversely, GFP-ZIF2 was able to markedly alleviate the growth defects induced at high Zn concentrations by loss of the vacuolar transporters Zrc1 and Cot1 [31]. Similar although less pronounced trends were observed when ZIF2 was expressed in the wild-type background (Figure 5 and Figure S2C). To correlate these effects with an enhanced Zn release capacity of yeast cells, we measured the total Zn concentration of the different strains grown under various Zn conditions (Table 1). While intracellular Zn accumulation was similar for all cell types under normal Zn supply, expression of the plant carrier significantly reduced the intracellular Zn concentration in all tested backgrounds when compared with cells carrying the empty vector under both low and high Zn supplies. These findings further confirm that the ZIF2 transporter is able to mediate Zn extrusion from yeast cells.

By contrast, expression of the plant transporter was unable to rescue the growth defects induced at high Co concentrations by loss of the Zrc1 and Cot1 transporters [31], nor was it able to influence yeast response to several other plant beneficial ions (Figure S3A). ZIF2, also known as ERD6 (Early Response to Dehydration)-like 7, belongs to the Monosaccharide-like Transporter subfamily of MFS transporters and indeed one of its closest \( S. \text{cerevisiae} \) homologues is the inositol transporter Itr1 [33]. Nevertheless, GFP-ZIF2 was unable to rescue the \( \text{Ditr1} \) deletion mutant’s deficient growth under limiting \( \text{nno-} \text{inositol} \) concentrations (Figure S3B). Conversely, expression of GFP-ZIF2 conferred enhanced yeast resistance to acetate, malate and citrate (Figure S3C), indicating that besides Zn, ZIF2 is also able to modulate weak acid sensitivity in yeast.

Loss of ZIF2 Function Causes Zn Hypersensitivity in Arabidopsis

To uncover the \( \text{in vivo} \) role of the ZIF2 transporter, we isolated an \( A. \text{thaliana} \) mutant allele (SALK_037804), harbouring a T-DNA insertion in the tenth intron of the \( \text{ZIF2} \) gene, which we named...
zif2-1 (see Figure 2A). RT-PCR analysis of ZIF2 expression in zif2-1 homozygous seedlings using primers annealing upstream of the insertion site revealed transcript levels comparable to wild-type plants, but no expression was detected when primers flanking or annealing downstream of the T-DNA segment were used (Figure S4). This indicated that the mutant allele produces a truncated version of the ZIF2 transcripts that lacks nearly the entire sequence corresponding to the second transmembrane domain and is thus unlikely to encode a functional membrane transporter [34], strongly suggesting that zif2-1 is a true loss-of-function mutant.

When grown in vitro under optimal conditions, zif2-1 mutant seedlings appeared morphologically indistinguishable from the corresponding wild type (Col-0), showing normal shoot growth, chlorophyll content and root system development (Figure 6A, Table S1). In the presence of excessive Zn amounts, Arabidopsis seedlings develop toxicity symptoms, typically including shoot growth retardation, leaf chlorosis and inhibition of PR elongation [1]. These three Zn toxicity hallmarks were visibly exacerbated in zif2-1 when compared to the wild type following exposure to a wide Zn toxicity range, though to a lesser extent than in the previously characterised zif1-2 mutant (Figure 6). By contrast, no significant differences between zif2-1 and wild-type seedlings could be observed at the PR elongation level under conditions of Zn deficiency, even when assayed on media with low cation content [6] (Figure S5A). In contrast to the zif1-2 mutant [7,20], no zif2-1 phenotype was observed under conditions of Fe depletion (Figure S5B) or in the presence of excessive amounts of Cd or Ni (Figure S6). At least at the tested concentrations, sensitivity of the mutant to excess of other essential, beneficial or rhizotoxic cations (Figure S6) or to myo-inositol (Figure S7) was also unaltered, as was sensitivity to sucrose and glucose, whose protective effect on Zn toxicity was unchanged by the zif2-1 mutation (Figure S8). Taken together, these data suggest that ZIF2 acts specifically as a Zn detoxifier in Arabidopsis.

To exclude the possibility that the observed phenotype results from disruption of another gene, we transformed the zif2-1 mutant with a genomic fragment spanning the entire ZIF2 gene and including the same promoter sequence used in the reporter gene experiments. These complementation lines exhibited complete restoration of wild-type sensitivity to excess Zn (Figure S9), thus confirming that the ZIF2 transporter participates in basal Zn tolerance in Arabidopsis.

Overexpression of the ZIF2.2 Splice Variant with Longer 5'UTR Confers Higher Plant Zn Tolerance

To gain insight into the physiological relevance of the ZIF2 5'UTR intron retention event promoted in roots by high Zn exposure, we selected three transgenic Arabidopsis lines independently expressing either the ZIF2.1 or ZIF2.2 cDNA under the control of the 35S promoter in the wild-type background that similarly displayed an approximately five-fold increase in expression of either splice variant when compared to wild-type plants (Figure 7A, 7B). Importantly, our RT-PCR analyses also showed that ZIF2.1 levels in the ZIF2.2-overexpressing lines were similar to those detected in the wild type, indicating that the 5’UTR intron is not spliced from the transgenic ZIF2.2 cDNA. No obvious phenotypical alterations were observed between wild-type and ZIF2-overexpressing seedlings grown under control Zn conditions (Figure 7C, Table S1). However, ZIF2.1 or ZIF2.2 overexpression substantially attenuated the detrimental effects induced by excessive Zn amounts over a broad range of Zn supplies.

Figure 5. The Arabidopsis ZIF2 transporter confers Zn resistance to S. cerevisiae. Comparison of the growth curves of non-adapted wild-type, Δzrt1zrt2 and Δzrc1cot1 mutant yeast cells, harbouring either the cloning vector pGREG576 or the GFP-ZIF2-encoding pGREG576_ZIF2 plasmid, in Zn-free liquid medium supplemented with 0.062 μM (wild-type and Δzrt1zrt2 strains), 2.5 or 20 μM (all strains), and 100, 250 or 500 μM (wild-type and Δzrc1cot1 strains) ZnSO4. Results are representative of three independent experiments (a replicate is shown in Figure S2C).

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Table 1. Effect of GFP-ZIF2 expression on the zinc content of yeast cells.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>External Zn(^{2+}) concentration ((\mu)M)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>0.062</td>
</tr>
<tr>
<td></td>
<td>2.5</td>
</tr>
<tr>
<td></td>
<td>500</td>
</tr>
<tr>
<td>WT+pGREG576</td>
<td>2.38±0.36</td>
</tr>
<tr>
<td>WT+GFP-ZIF2</td>
<td>2.78±0.18</td>
</tr>
<tr>
<td>zrt1zrt2+pGREG576</td>
<td>0.48±0.47</td>
</tr>
<tr>
<td>zrt1zrt2+GFP-ZIF2</td>
<td>1.25±0.34</td>
</tr>
<tr>
<td>zrc1cot1+pGREG576</td>
<td>ND</td>
</tr>
<tr>
<td>zrc1cot1+GFP-ZIF2</td>
<td>ND</td>
</tr>
</tbody>
</table>

Total Zn concentration (pg/cell) of wild-type, zrt1zrt2 and zrc1cot1 mutant yeast cells, harbouring either the cloning vector pGREG576 or the pGREG576_ZIF2 plasmid, grown in Zn-free liquid medium supplemented with 0.062, 2.5 or 500 \(\mu\)M ZnSO\(_4\) (means ± SD, n=4). Statistically significant differences from the respective empty-vector control are indicated with the P values (in parentheses) obtained by Student's t-test. ND, not detectable.

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We next sought to understand the different functional impact of the two ZIF2 splice variants on Zn tolerance and whole-plant partitioning in Arabidopsis. Given that ZIF2.1- and ZIF2.2-overexpressing seedlings displayed very similar steady-state levels of the respective ZIF2 transcripts (see Figure 7A, 7B), which encode the same transporter (see Figure 2B), the differences in phenotype magnitude are unlikely to result from distinct mRNA levels and/or protein stability/activity, thus pointing to a translational regulatory mechanism.

To assess the ZIF2 protein levels produced by each ZIF2 splice variant in planta, we took advantage of our ZIF2-YFP overexpression lines (see Figure 4E, F), providing a setting in which the abundance of the ZIF2 transporter can be accurately linked to the amount of each splice variant through quantification of the corresponding YFP signal and transgenic levels. Two transgenic Arabidopsis lines independently expressing either ZIF2.1-YFP or ZIF2.2-YFP under the control of the 35S promoter in the wild-type background were selected. Both the ZIF2.1-YFP and the ZIF2.2-YFP transgenic lines exhibited increased resistance to excess Zn when compared to the wild type (Figure S12), demonstrating the functionality of the ZIF2-YFP fusion proteins. Moreover, as with our other set of overexpression plants (see Figures 7 and S10), both ZIF2.2-YFP transgenic lines were significantly more tolerant to Zn toxicity than the two lines expressing ZIF2.1-YFP (Figure S12).


The ZIF2.2 Splice Variant Produces More ZIF2 Protein In Planta than ZIF2.1

We next sought to understand the different functional impact of the two ZIF2 splice variants on Zn tolerance and whole-plant partitioning in Arabidopsis. The ZIF2.2 function influences Zn tissue partitioning in Arabidopsis, whereas zif2-1 mutant roots concentrated significantly less Zn than those of the wild type. The opposite Zn partitioning trend was observed in ZIF2-overexpressing plants, which accumulate respectively less and more Zn in their shoot and root tissues than wild-type plants. This tendency, while also observed under control conditions, was substantially more pronounced at the two highest Zn concentrations (Figure 8). As the total amount of Zn quantified at the whole seedling level was globally comparable in all genotypes (Figure S11), the Zn shoot-to-root ratio was enhanced and reduced by ZIF2 loss of function and overexpression, respectively. These results indicate that ZIF2 function alters Zn partitioning between roots and shoots by driving root Zn immobilization and therefore shoot Zn exclusion, particularly under conditions of Zn excess. Importantly, the effects of ZIF2 overexpression on Zn distribution were significantly more pronounced in plants ectopically expressing the ZIF2 transcript with the longer 5'UTR, ZIF2.2, than in the ZIF2.1 transgenics (Figure 8), in clear agreement with the physiological data obtained for both types of lines (see Figures 7 and S10).

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The ZIF2 5'UTR Intron Determines Translation Efficiency in a Zn-Responsive Manner

To investigate whether the ZIF2.1 and ZIF2.2 5'UTRs are sufficient to determine different translation efficiencies, the effect of each splice variant’s 5’UTR on the expression of a reporter gene (LUC) was examined by means of an in vivo assay using isolated Arabidopsis protoplasts [35,36]. The ZIF2.1_5’UTR-LUC and ZIF2.2_5’UTR-LUC constructs were expressed under the control of...
Zn when compared with LUC activity was detected upon transfection with the 35S promoter, and LUC activity was normalized to the GUS reporter construct. This structure is lost in the ZIF2.1 splice variant as the intron removes roughly half of the sequence participating in the structure formation. To determine whether the predicted 5’UTR secondary structure element plays a role in translational regulation of the two alternatively-spliced mRNAs, we performed two successive rounds of site-directed mutagenesis on the ZIF2.2_5’UTR-LUC construct to generate the ZIF2.2_5’UTR-ΔM-LUC (where CTCA was mutated to AGTC) and ZIF2.2_5’UTR-R-LUC (where TGAG was mutated to GACT) constructs, in which the secondary structure was destabilized and restored, respectively. (Figure 10B). In the absence of Zn, a two-fold increase in LUC activity was detected upon transfection with any of the three ZIF2.2_5’UTR-LUC constructs when compared with the ZIF2.1_5’UTR-LUC transcript (Figure 10C). However, destabilization of the secondary structure strikingly reduced the pronounced inductive effect that Zn exerts on ZIF2.2_5’UTR-LUC translation, while restoration of the structure fully rescued translational induction by the metal ion (Figure 10C). The results from this last experiment indicate that the mechanism(s) underlying Zn regulation of translation driven by the ZIF2 5’UTR depend, at least to a large extent, on an RNA secondary structure that is present when the 5’UTR intron is retained.

### Discussion

Since the identification of ZIPs as the first plant Zn transporters [37], several carriers from well-established metal transporter families have been linked to plant Zn homeostasis. The few plant MFS carriers characterised so far have been mainly implicated in sugar, nitrate, oligopeptide and phosphate transport [38–40]. Here we report that besides ZIF1 [20], another MFS carrier contributes to Arabidopsis Zn tolerance, hinting at a broader role for this class of transporters in plant heavy-metal homeostasis.

Conclusive evidence for a ZIF2 role in basal Zn tolerance stems from our functional analysis of Arabidopsis ZIF2 loss-of-function and overexpression lines, exhibiting respectively enhanced sensitivity and resistance to Zn toxicity. The ZIF2 tonoplast localisation detected in planta is consistent with two previous proteomic studies of the Arabidopsis vacuole [41,42] and strongly suggests that ZIF2 is involved in vacuolar Zn compartmentalisation. Indeed, plant vacuoles play an essential role during metal detoxification and sequester up to 80% of cellular Zn [43]. In agreement with our physiological data, Zn concentration determination in seedlings exposed to high Zn shows that the shoot-to-root ratio of the zif2-1 mutant is unaltered to the shoot, while Zn
Arabidopsis ZIF2 Transporter Mediates Zn Tolerance
Figure 7. The ZIF2.1 and ZIF2.2 splicing variants contribute differentially to Zn tolerance in Arabidopsis. (A) RT-PCR profile of ZIF2.1 and ZIF2.2 expression in roots of 7-d-old seedlings of the wild type (Col-0) and ZIF2.1 or ZIF2.2 overexpression lines. The location of the F1 and R1 primers used is shown in Figure 2A. Expression of the UBQ10 gene was used as a loading control. Results are representative of three independent experiments. (B) Real-time RT-PCR analysis of total ZIF2 mRNA expression as well as of ZIF2.1 and ZIF2.2 transcript levels in roots of 7-d-old seedlings of the wild type (Col-0) and ZIF2.1 or ZIF2.2 overexpression lines using UBQ10 as a reference gene. Results are from two independent experiments and values represent means ± SE (n = 4). Different letters indicate statistically significant differences between genotypes (P < 0.05; Student’s t-test).

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Figure 8. The ZIF2 transporter affects root-shoot partitioning in Arabidopsis. Zn concentration, expressed on a dry weight (DW) basis, in the shoot (upper panel) and root (middle panel), as well as the corresponding shoot/root ratio (lower panel), of 21-d-old seedlings of the wild type (Col-0), the ZIF2.1-OX1 and ZIF2.2-overexpressing lines (ZIF2.1OX2 and ZIF2.2OX1) grown on control medium (30 μM Zn²⁺) or under excess Zn supply (250 μM Zn²⁺). Scale bar, 1 cm. (D) Effect of Zn toxicity on shoot biomass (upper panel), chlorophyll content (middle panel) and PR elongation (lower panel) of wild-type (Col-0) and ZIF2.1- or ZIF2.2-overexpressing (ZIF2.1OX2 or ZIF2.2OX1) seedlings. Results are representative of three independent experiments and values represent means ± SD (n = 4 for shoot biomass/chlorophyll content and n = 16 for PR elongation). Different letters indicate statistically significant differences between genotypes (P < 0.05; Student’s t-test).

doi:10.1371/journal.pgen.1004375.g008

is preferentially retained in ZIF2-overexpressing roots, demonstrating that ZIF2 function influences Zn root-to-shoot translocation under Zn excess. Given that ZIF2 is primarily expressed in the root endodermis and cortex, we propose that the encoded transporter protects from Zn toxicity by promoting vacuolar immobilization of the metal ion in these two cell layers, thus restricting symplastic movement towards the stele and subsequent xylem loading and translocation to the shoot. Richard et al. [44] showed that low root-driven Zn translocation rates to the shoot contribute to higher Zn tolerance, as reported for the Arabidopsis HMA3, MTP3, MTP1 and ZIF1 [10,19,20,45]. The partial expression pattern overlap among these vacuolar transporters and ZIF2 suggests either some degree of functional redundancy or a concerted action to regulate root Zn symplastic movement. Putative roles of the ZIF2 transporter in other plant organs where substantial ZIF2 expression was detected, such as in flowers, cannot be excluded as already shown for instance for the HMA2 and HMA4 promoters. Similarly to ZIF2, HMA2 and HMA4 promoter activity is high in developing anthers, and a double hma2hma4 mutant exhibits a male-sterile phenotype [17].

The identification of a transporter’s physiological substrate(s) can be instrumental in elucidating the molecular mechanisms governing its function, but remains an extremely challenging task. Importantly, our yeast heterologous expression and in planta phenotypical data point to a ZIF2 effect restricted to Zn, similarly to MTP1 and MTP3 that exhibit relatively high Zn selectivity [10,45] but in contrast to HMA or ZIPs that often display broad metal selectivity contributing to plant metal homeostasis crosstalk [19,46]. More puzzling are our preliminary findings indicating that, despite belonging to the Monosaccharide-like subfamily, activity of the ZIF2 transporter does not seem to influence sugar sensitivity in planta nor in yeast. Instead, heterologous expression studies show that ZIF2 is able to mediate Zn efflux in yeast cells without requiring additional plant-specific factors, as already reported for HMA2, HMA4, MTP1 and PCR2 [5,11,16]. By contrast, the transport activity of ZIF1 is unable to complement a Zn hypersensitive yeast mutant, thus indicating that the mode of action of the two MFS carriers in mediating Zn tolerance is strikingly different. Haydon et al. [7] identified the substrate of ZIF1 as nicotianamine, a low molecular mass chelator with high affinity for a range of transition metals, and proposed that ZIF1, by affecting nicotianamine vacuolar partitioning, specifically promotes Zn vacuolar sequestration via Zn²⁺/H⁺ antiporters. As no plasma-membrane Zn exporter has been identified so far in S. cerevisiae [47], it seems reasonable to exclude ZIF2 activation of endogenous transporters catalysing Zn efflux. On the other hand, as MFS transporters are known to transport small solutes in response to chemiosmotic gradients and as ZIF2 affects weak acid delivery in yeast, it is tempting to speculate that ZIF2 indirectly mediates Zn vacuolar sequestration by transporting a small chelator or organic acid or by influencing the tonoplastic proton gradient. Indeed, plant cells are known to take advantage of proton gradients to actively sequester ions inside the vacuole [48] and the Arabidopsis H⁷⁺-ATPase (V-ATPase), which energizes transport across the tonoplast, is required for full Zn tolerance [49].

Our results also indicate that the Zn metal ion positively regulates ZIF2 expression at multiple levels. Firstly, root ZIF2
Figure 9. The ZIF2.1 and ZIF2.2 5' UTRs exert differential effects on translation. (A) Representative confocal laser scanning microscopy images of root tips of transgenic Arabidopsis 7d-old seedlings expressing either the ZIF2.1-YFP or ZIF2.2-YFP fusions under the control of the 35S promoter. Detection settings for YFP visualization were identical for all genotypes. The YFP signal is visualized by green coloration. Scale bars, 50 μm. (B) Transgene transcript and protein levels in Arabidopsis ZIF2.1-YFP and ZIF2.2-YFP overexpression lines. Quantification of ZIF2-YFP transcript (upper panel) and corresponding fusion protein (lower panel) levels in roots of 7-d old ZIF2-YFP overexpression seedlings grown under control conditions. Relative ZIF2-YFP gene expression was quantified by real-time RT-PCR using UBQ10 as a reference gene (means ± SD, n = 4). The fluorescence intensity of the YFP signal in root tips (means ± SD, n = 12) was normalized to ZIF2-YFP transcript levels to determine the relative abundance of the ZIF2-YFP protein. Asterisks denote statistically significant differences from the ZIF2.1-YFP OX1 line (***P ≤ 0.001; Student’s t-test). (C) Quantification of ZIF2-YFP protein levels in Arabidopsis ZIF2.1-YFP and ZIF2.2-YFP overexpression lines. Western blot analysis of ZIF2-YFP protein levels in 5-d old seedlings from the wild type (Col-0) and ZIF2.1-YFP OX or ZIF2.2-YFP OX transgenic lines. The blot image is representative of two independent experiments and values represent means ± SD (n = 4). Different letters indicate statistically significant differences (P ≤ 0.01) between Zn concentrations for each construct and P values for the comparison of the two constructs under each condition are shown (Student’s t-test). doi:10.1371/journal.pgen.1004375.g009
transcript levels are substantially induced following exposure to elevated Zn supplies, corroborating an essential role in Zn detoxification, as already inferred for ZIF1 [20] or MTP3 [10]. Secondly, splicing of the ZIF2 pre-mRNA is Zn-regulated, with high levels of the metal favouring retention of the 5' UTR intron and hence expression of the longer ZIF2.2 splice variant. The splicing factors regulating this intron retention event remain unknown. In animal systems, splice site selection is known to be primarily regulated by the highly conserved serine/arginine-rich (SR) and heterogeneous nuclear ribonucleoprotein particle (hnRNPs) protein families, which bind specific exonic or intronic sequences in the pre-mRNA. While these splicing factors have been implicated in the modulation of alternative splicing also in plants, their direct endogenous targets and the mechanisms controlling differential splice site usage have not been elucidated [50,51].

Noticeably, our data indicate that the 5' UTR intron is not spliced from the ZIF2.2 transgene in our ZIF2.2-overexpressing lines, while splicing of this intron occurs at considerable rates from the ZIF2.2vU/C transcript transiently expressed in protoplasts. On the other hand, the inductive effect that Zn exerts on the intron retention event in wild-type roots is lost in the single-cell system. These apparent discrepancies could be due to the distinct tissue sources (roots versus leaves) and/or to differences in transcription rates, which are well known to affect alternative splicing [52], between the different expression contexts. The molecular mechanisms underlying the regulation of alternative splicing in plants are poorly understood. Despite numerous reports that abiotic stress markedly affects plant mRNA splicing, virtually nothing is known about the effects of metal ions. Interestingly, in humans Zn has been shown to modulate alternative splicing of the hypoxia-inducible factor HIF1 [53] and the pro-apoptotic Bim [54] genes.

Finally, and given that information on the functional relevance of alternative splicing in plant systems is also surprisingly scarce, a key finding of this study is that two ZIF2 splice forms differentially contribute to Zn detoxification. In fact, our two sets of overexpression data indicate not only that ZIF2.2 confers greater Zn tolerance than ZIF2.1, but also that the different functional impact of the two mRNAs is not due to differences in their steady state levels. As the two alternative transcripts encode the same transporter, it is also highly improbable that differences in protein stability or activity account for the observed variations in resistance to Zn toxicity. Instead, we show that the stronger Zn tolerance phenotype conferred by ZIF2.2 results from its enhanced translation efficiency. Interestingly, Zn significantly promotes translation of both ZIF2 splice variants, but this effect is dramatically more pronounced for ZIF2.2. Furthermore, our results indicate that the ZIF2 5'UTR alone is sufficient for this translational control. Indeed, 5'UTR sequence and structural features have been proposed to play an essential role in translational regulation in plants [55,56], as described for other eukaryotes. It is interesting to note that expression of an Arabidopsis tonoplast Zn/Mg transporter, AtMHX, was shown to be repressed at the translational level through the inhibitory effect of an upstream open-reading-frame present in the 5'UTR of the corresponding gene [57,58]. Importantly, we were able to demonstrate that the mechanism(s) underlying Zn-responsive ZIF2.2 translation depend largely on a predicted stable stem loop structure that is lost when the ZIF2 5'UTR intron is spliced out. From the publicly available databases, the 5'UTR intron retention event in ZIF2 appears to be fairly specific to this Arabidopsis gene, as no significant sequence homology is found with other genes in A. thaliana nor in other plant species except with the ZIF2 orthologue in Arabidopsis lyrata. Similarly, the identified stable stem loop appears to be a unique feature of A. thaliana, as no common mRNA secondary structure could be predicted in the 5'UTR of plant ZIF2 orthologues.

Selective recruitment of certain physiologically relevant mRNAs under stress conditions that trigger a global repression of the initiation step of translation has been widely reported in metazoans. Such a mechanism is beginning to emerge as a key process in plant adaptation to environmental stresses, interestingly for instance during sub-lethal cadmium poisoning [59]. In particular, secondary structures in the 5'UTRs have been implicated in cap-independent mRNA translation in maize under heat and other stress conditions [60,61] as well as in Arabidopsis under heat stress [55]. The findings presented here indicate that alternative splicing controls the levels of a Zn stress responsive mRNA variant of the ZIF2 transporter to enhance plant tolerance to the metal ion.

Materials and Methods

Plant Materials and Growth Conditions

The Arabidopsis thaliana (L.) Heynh., ecotype Colombia (Col-0), was used in all experiments. Seeds of the T-DNA insertion mutants zif2-1 (SALK_037804) and zif1-2 (SALK_011408; [20]) were obtained from the Nottingham Arabidopsis Stock Centre (NASC) (Nottingham, UK). The exact zif2-1 T-DNA insertion site was confirmed using gene-specific primers (Table S2) and primers annealing at the T-DNA border, which also allowed PCR-based genotyping to identify homozygous lines. Plant transformation was achieved by the floral-dip method [62] using Agrobacterium tumefaciens strain EHA105. Seeds were surface-sterilized and sown on Murashige and Skoog [63] medium solidified with 0.8% agar, stratified for 3 d, placed in a growth chamber and transferred to soil after 2–3 weeks. Plants were cultivated under long-day conditions (16-h light, 22°C/8-h dark, 18°C; 60% RH).

Gene Expression Analyses

RLM-5'RACE was performed using the FirstChoice RLM-RACE kit (Ambion) on total RNA extracted from 21-d old seedlings, adult leaves, flowers or roots exposed to high Zn according to the manufacturer’s instructions and using various gene-specific primers (Table S2). Subsequently, PCR products were purified, cloned into the p-Gem-T Easy vector (Promega) and sequenced. For native ZIF2 promoter reporter gene experiments, a fragment including the 1233 bp immediately upstream of the start codon was PCR-amplified (Table S2) from genomic DNA and inserted via the SacI/SacII restriction sites into the pKGWFS7 plasmid [64]. After agroinfiltration of this ProZIF2:6FGFUS construct into wild-type plants, eight independent transformants all showing similar tissue-specific GUS expression patterns were recovered. Histochemical staining of GUS activity was performed as described by Sundaresan et al. [65]. RT-PCR analyses were conducted as previously described [66] using primers designed to detect ZIF2, ZIF1, ZIP1, ROC10 (CYCLOPHILIN) and UBQ10 (UBQ10/UBQ10/10) expression (Table S2). The results shown are representative of three independent experiments. Real-time RT-PCR was performed using specific primers (Table S2) on a CFX384 Touch Real-Time PCR Detection System (Bio-Rad) using the Absolute SYBR GREEN ROX mix (Thermoscientific) according to the manufacturer’s instructions. For each condition tested, two RNA extractions from different biological samples and two reverse transcription reactions for each biological repeat were performed. Data were processed using Q-Genie [67] that took the respective primer efficiency into consideration.
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**A**

Structure

**B**

ZIF2.1 5'UTR
MFE -114 kcal/mol

Start codon

ZIF2.2 5'UTR
MFE -139.2 kcal/mol

Intron start

**C**

![Graph showing relative LUC/GUS expression](image)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Relative LUC/GUS</th>
<th>p-value</th>
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<tr>
<td>0 μM Zn²⁺</td>
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<td>100 μM Zn²⁺</td>
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Legend:
- □ 0 μM Zn²⁺
- ■ 100 μM Zn²⁺

P-values:
- P = 2.90e-3
- P = 5.80e-3
Figure 10. An RNA secondary structure element sustains Zn regulation of translation driven by the ZIF2.2 5' UTR. (A) Organization of the 5' end of the ZIF2 gene. The orange and blue rectangles represent the intronic and coding sequences, respectively, and the region forming a stable RNA structure is indicated as a black line. (B) Secondary structure predictions for the ZIF2.1 and ZIF2.2 5'UTRs flanked by 200 bp of the coding sequence. A potentially stable secondary structure element is enclosed in an oval. Four consecutive bases in the destabilized structure element or stable RNA structure is indicated as a black line. (B) Secondary structure predictions for the genomic DNA and inserted into the promoterless version of the gene and including the 1233-bp promoter via the PacI restriction sites into the YFP- or Pro35S:ZIF2.2-YFP. These YFP constructs were also transfected into Arabidopsis protoplasts (see below). Transient co-expression of the GFP constructs with the tonoplast marker γ-Tonoplast Intrinsic Protein (TIP-mCherry) or the plasma membrane marker Plasma membrane Intrinsic Protein 2A (PIP2A-mCherry) [30] and the pBIN-NA construct [68] in leaf abaxial epidermal cells of Nicotiana tabacum was performed via the agroinfiltration procedure described by Voornet et al. [69] using A. tumefaciens strain GV3101.

Subcellular Localisation Studies

To generate ZIF2 protein fusions with the YFP and GFP reporters, each ZIF2 transcript (5’UTR/coding sequence except the stop codon) was PCR-amplified (Table S2) using root cDNA as a template, and independently inserted under the control of the 35S promoter via the Xhol/PacI restriction sites into the YFP- or GFP-tagged versions of the pBA002 vector. Two transgenic lines displaying a strong fluorescence signal in the root were recovered upon transformation of wild-type plants with either Pro35S::ZIF2.1-YFP or Pro35S::ZIF2.2-YFP. These YFP constructs were also transfected into Arabidopsis protoplasts (see below). Transient co-expression of the GFP constructs with the tonoplast marker γ-Tonoplast Intrinsic Protein (TIP-mCherry) or the plasma membrane marker Plasma membrane Intrinsic Protein 2A (PIP2A-mCherry) [30] and the pBIN-NA construct [68] in leaf abaxial epidermal cells of Nicotiana tabacum was performed via the agroinfiltration procedure described by Voornet et al. [69] using A. tumefaciens strain GV3101.

Yeast Manipulations

The S. cerevisiae mutant strains Act1ΔZIF2 [32] and Act1Δact1 [31], along with the corresponding wild-type strain DY1457, were used in the Zn experiments. All other studies were performed in the parental strain BY4741 and the derived deletion mutant Act1 [33]. Cloning of the Arabidopsis ZIF2 full-length coding sequence into the pBRE5376 vector [70]; Euroscarf collection) and expression analysis of the corresponding GFP-ZIF2 fusion protein by fluorescence microscopy and western blotting were performed as described previously [71]. Growth curve and spot assays were conducted in MMB-U liquid or agarized medium [71] supplemented with the indicated compound (Sigma-Aldrich) at the desired concentration. Results presented are representative of at least three independent experiments.

Plant Phenotypical Assays

All assays were performed in a climate-controlled growth cabinet under long-day conditions. After 5 d of vertically-oriented growth on control medium (30 μM Zn), seedlings were transferred to fresh medium containing the indicated compound (Sigma-Aldrich) at the specified concentration. PR elongation, along with shoot biomass and chlorophyll content [72] were evaluated after an additional 1 and 3 weeks of growth, respectively. Results are representative of at least three independent experiments.

Genomic Complementation and Overexpression Analyses

For genomic complementation, a 4450-bp fragment encompassing the entire ZIF2 gene and including the 1233-bp promoter sequence described above was PCR-amplified (Table S2) from genomic DNA and inserted into the promoterless version of pBA002 via the NolI/Xhol restriction sites. Representative results for three lines recovered upon introduction of the construct into ZIF2-1 mutant plants are shown. ZIF2 overexpression constructs were generated as described for the YFP and GFP plasmids, except that the corresponding fragments were inserted into the pBA002 background via the Xhol/IvI restriction sites. After agroinfiltration of wild-type plants, three transgenic lines independently overexpressing each ZIF2 transcript were selected.

Protoplast Transient Expression Assays

Arabidopsis protoplasts were generated as described by Yoo et al. [35] and transfected by polyethylene glycol transformation [73]. To generate the Pro35S::ZIF2.5’UTR fusion constructs with the LUC reporter, the ZIF2.1 and ZIF2.2 5’UTRs were PCR-amplified (Table S2) and independently cloned into the Pro35S:LUC vector [35] via the SalI/XhoI restriction sites. Site-directed mutagenesis of the Pro35S::ZIF2.2 5’UTR-LUC construct was performed using gene-specific primers (Table S2) and the NZYMutagenesis kit (NZYTech) according to the manufacturer's instructions. Protoplast transient expression assays along with GUS and LUC activity measurements and RNA protoplast extraction were performed as described previously [74]. Results shown are representative of at least three independent experiments.

Microscopy and Western Blot Analysis

Differential interference contrast and confocal images were taken with a DM LB2 microscope (Leica) and an LSM 510 laser scanning microscope equipped with a Meta detector (Zeiss), respectively. Excitation/detection wavelengths used to detect fluorescence were 488/500–530 nm for GFP, 514/535–590 nm for YFP, 543/565–615 nm for mCherry, 534/560 nm for propidium iodide and 458/560 nm for autofluorescence. For YFP signal quantification, fluorescence detection parameters (laser intensity, offset, gain and pinhole settings) were set so the fluorescence signal emitted by ZIF2.2-YFPOX1 root tips was just below the saturation threshold. Twelve micrographs per genotype were captured in the median section of roots from 5-d-old seedlings imaged in 50 mM Phosphate Buffer using identical confocal settings to allow comparison between genotypes. Post-imaging, average fluorescence intensity within the whole region (ca. 20 μm²) spanning the extreme root apex to the first elongating cells or the zone immediately above the first elongating cells was recorded. Results are representative of three independent experiments.

Total protein was extracted from 5-d-old seedlings using 2X Laemmli Extraction Buffer [75] and equal amounts of extracts were resolved on a 8% SDS/polyacrylamide gel before proteins were transferred to a PVDF membrane (Immobilon-P, Millipore), which was incubated with anti-GFP primary antibody (Roche;
1:500 dilution) and then with anti-mouse peroxidase-conjugated secondary antibody (Amersham Pharmacia; 1:20000 dilution), before membrane-associated peroxidase activity was revealed by ECL.

Zinc Measurements
To measure plant zinc concentration, pooled shoot and root tissues from 3-week old seedlings grown on 30, 125 or 250 μM Zn were processed as previously described [66]. The total Zn concentration of yeast cells was measured as described by Arrivault et al. [10]. The zinc concentration of the digests was quantified using the Atomic Emission Spectrometry — Inductively Coupled Plasma — Optical Emission System (Perkin-Elmer Optima 2100 DV) at the Laboratório de Análises, Instituto Superior Técnico (Lisbon, Portugal) according to method 3120B described by Eaton et al. [76]. Zinc standards for analytical calibration were from Merck KGaA. Four independent samples were processed per genotype.

Software
Real-time RT-PCR results were analysed using the CFX Manager 3.0 software (Bio-Rad) and the Q-Gene application [67]. Microscopy images and scanned images of root assays and western blots were processed using LSM 510 software (Zeiss) and ImageJ (http://rsbweb.nih.gov/ij/), respectively. The entire 5’UTR (plus 200 nt of coding sequence) alternative structures and their minimum free energies (MFEs) were calculated using RNAFold [77]. The structures presented in Figure 10 are the centroid representations calculated with default parameters.

Supporting Information
Figure S1 Expression kinetics of the ZIF2.1 and ZIF2.2 splice variants under zinc toxicity. RT-PCR profile of ZIF2.1 and ZIF2.2 expression in roots of 7-d old wild-type (Col-0) seedlings challenged for 0, 24, 48 or 96 h with 250 μM Zn. The location of the F1 and R1 primers used is shown in Figure 2A. Expression of the UBQ10 gene was used as a loading control. Results are representative of two independent experiments.

Figure S2 Heterologous expression of the Arabidopsis ZIF2 transporter in yeast. (A) Western blot analysis of the GFP-ZIF2 fusion protein in wild-type yeast cells harbouring either the cloning vector pGREG576 or the GFP-ZIF2-encoding pGREG576_ZIF2 plasmid, in liquid medium supplemented with 3.5 mM CoSO4, 1 mM NiSO4, 1 mM CuCl2, 40 mM FeSO4 or 25 mM MnCl2. Results are representative of at least two independent experiments. (B–C) Susceptibility to myo-inositol (B) and weak acids (C) of wild-type or Δzrt1 mutant yeast cells, harbouring either the cloning vector pGREG576 or the GFP-ZIF2-encoding pGREG576_ZIF2 plasmid, determined by spotting dilution series of cell suspensions (1, 1.5, and 1:10). (PDF)

Figure S4 ZIF2 expression in the Arabidopsis zif2-1 mutant. RT-PCR analysis of ZIF2 expression in 14-d old wild-type (Col-0) and mutant (zif2-1) seedlings. The location of the F1’, R1, F2, R2, F3 and R3 primers used is shown in Figure 2A. Expression of the UBQ10 gene was used as a loading control. Results are representative of three independent experiments.

Figure S5 Phenotype of the Arabidopsis zif2-1 mutant under zinc or iron deficiency. Effect of Zn deficiency under different MS strengths (A) and of ferrozine-induced Fe deficiency (B) on PR elongation of wild-type (Col-0) and zif2-1 or zif2-2 mutant seedlings. Results are representative of two independent experiments and values represent means ± SD (n = 16). Asterisks denote statistically significant differences from the wild type under each condition (**P<0.01; Student’s t-test).

Figure S6 Supplemental ion response characterisation of the Arabidopsis zif2-1 mutant. Effect of excess of essential/beneficial cations and of rhizotoxic cations on shoot biomass (upper panel), chlorophyll content (middle panel) and PR elongation (lower panel) of wild-type (Col-0) and mutant (zif2-1) seedlings. Results are representative of two independent experiments and values represent means ± SD (n = 16). No statistical differences between mutant and wild type were detected under each condition (P>0.05; Student’s t-test).

Figure S7 Phenotypical characterisation of the Arabidopsis zif2-1 mutant in the presence of different concentrations of myo-inositol. Effect of myo-inositol on PR elongation of wild-type (Col-0) and mutant (zif2-1) seedlings. Results are representative of two independent experiments and values represent means ± SD (n = 16). No statistical differences between mutant and wild type were detected under each condition (P>0.05; Student’s t-test).

Figure S8 Phenotypical characterisation of the Arabidopsis zif2-1 mutant in the presence of both zinc and sugar. Effect of sucrose or glucose combined with Zn on shoot biomass (upper panel), chlorophyll content (middle panel) and PR elongation (lower panel) of wild-type (Col-0) and mutant (zif2-1) seedlings. Results are representative of two independent experiments and values represent means ± SD (n = 8 for shoot biomass/chlorophyll content and n = 16 for PR elongation). Asterisks denote statistically significant differences from the wild type under each condition (**P<0.01, ***P<0.001; Student’s t-test).

Figure S9 Genomic complementation of the Arabidopsis zif2-1 mutant phenotype. Effect of Zn toxicity on shoot biomass (upper panel), chlorophyll content (middle panel) and PR elongation (lower panel) of seedlings of the wild type (Col-0), the zif2-1 mutant and three independent genomic complementation lines, zif2-3comp1-3. Results are representative of three independent
experiments and values represent means ± SD (n=8 for shoot biomass/chlorophyll content and n=16 for PR elongation). Asterisks denote statistically significant differences from the wild type under each condition (*P<0.05, **P<0.01, ***P<0.001; Student’s t-test).

(PDF)

**Figure S10** Phenotype of *Arabidopsis* ZIF2.1 and ZIF2.2 overexpression lines. Effect of Zn toxicity on shoot biomass (upper panel), chlorophyll content (middle panel) and PR elongation of wild-type (Col-0) and ZIF2 overexpression lines. Effect of Zn toxicity on shoot biomass (upper panel) or two (lower panel) independent experiments and average fluorescence (pixel) intensity is shown (means ± SD, n=12). Asterisks denote statistically significant differences from the ZIF2.1-YFPox1 line (***P<0.001; Student’s t-test).

(PDF)

**Figure S11** Whole-seedling zinc concentration of *Arabidopsis* wild-type, zif2-1 mutant and ZIF2 overexpression lines. Zn concentration, expressed on a dry weight (DW) basis, in 21-d old seedlings of the wild type (Col-0), the zif2-1 mutant and ZIF2 overexpression lines (ZIF2.1OX2 and ZIF2.2OX1) grown on control medium (30 μM Zn+) or under excess Zn supply (125 or 250 μM Zn+). Bars represent means ± SD (n=4). No statistical differences between genotypes were detected under each condition (P>0.05; Student’s t-test).

(PDF)

**Figure S12** Phenotype of *Arabidopsis* ZIF2.1-YFP and ZIF2.2-YFP overexpression lines. Effect of Zn toxicity on PR elongation of wild type (Col-0) and ZIF2.1-YFP or ZIF2.2-YFP overexpression lines (ZIF2.1-YFPox1-2 or ZIF2.2-YFPox1-2) seedlings. Results are representative of two independent experiments and values represent means ± SD (n=16). Different letters indicate statistically significant differences between genotypes under each condition (P<0.05; Student’s t-test).

(PDF)

**Figure S13** Quantification of ZIP2-YFP protein levels in *Arabidopsis* ZIP2-YFP overexpression lines. Quantification of the YFP signal by confocal laser scanning microscopy in primary root tip (upper panel) and primary root (lower panel) cells from 5-d old seedlings of the ZIP2.1-YFPox and ZIP2.2-YFPox transgenic lines. In each panel, detection settings for YFP visualization were identical for all genotypes. Results are representative of three (upper panel) or two (lower panel) independent experiments and average fluorescence (pixel) intensity is shown (means ± SD, n=12). Asterisks denote statistically significant differences from the ZIP2.1-YFPox1 line (***P<0.001; Student’s t-test).

(PDF)

**Table S1** Growth parameters of *Arabidopsis* ZIP2 loss-of-function and overexpression lines under control conditions.

(PDF)

**Table S2** Sequences of primers used.

(PDF)

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We thank D. Eide for the yeast AZT1202 and AZT1101 strains, N.-H. Chua for the pBAM002 vectors, M. Zeidler for the pBIN-NA plasmid, M. Baito for technical assistance and R. Carvalho, V. Nunes, E. Baena-González, C. Martinho, A. Confraria and P. Crozet for support and helpful discussions.

**Author Contributions**

Conceived and designed the experiments: ER MCT AA ISC PD. Performed the experiments: ER TRC RAB MAMH. Analyzed the data: ER TRC MCT AA ISC PD. Contributed reagents/materials/analysis tools: ISC PD. Wrote the paper: ER PD.
22. Fankhauser C, Casal JJ (2004) Phenotypic characterization of a photomorpho-