The iron deficiency response in Arabidopsis thaliana requires the phosphorylated transcription factor URI

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Iron is an essential nutrient for plants, but excess iron is toxic due to its catalytic role in the formation of hydroxyl radicals. Thus, iron uptake is highly regulated and induced only under iron deficiency. The mechanisms of iron uptake in roots are well characterized, but less is known about how plants perceive iron deficiency. We show that a basic helix–loop–helix (bHLH) transcription factor Upstream Regulator of IRT1 (URI) acts as an essential part of the iron deficiency signaling pathway in Arabidopsis thaliana. The uri mutant is defective in iron-Regulated Transporter1 (IRT1) and Ferric Reduction Oxidase2 (FRO2) and their transcriptional regulators FER-like iron deficiency-induced transcription factor (FIT) and bHLH38/39/100/101 in response to iron deficiency. Chromatin immunoprecipitation followed by sequencing (ChIP-seq) reveals direct binding of URI to promoters of many iron-regulated genes, including bHLH38/39/100/101 but not FIT. While URI transcript and protein are expressed regardless of iron status, a phosphorylated form of URI only accumulates under iron deficiency. Phosphorylated URI is subject to proteasome-dependent degradation during iron resupply, and turnover of phosphorylated URI is dependent on the E3 ligase BTS. The subgroup IVc bHLH transcription factors, which have previously been shown to regulate bHLH38/39/100/101, communoprecipitate with URI mainly under Fe-deficient conditions, suggesting that it is the phosphorylated form of URI that is capable of forming heterodimers in vivo. We propose that the phosphorylated form of URI accumulates under Fe deficiency, forms heterodimers with subgroup IVc proteins, and induces transcription of bHLH38/39/100/101. These transcription factors in turn heterodimerize with FIT and drive the transcription of IRT1 and FRO2 to increase Fe uptake.

Iron deficiency | iron homeostasis | Arabidopsis | bHLH transcription factor | phosphorylation

Iron (Fe) is an essential nutrient for plants. It serves as a cofactor for more than 300 enzymes and plays an irreplaceable role in vital processes, such as respiration and photosynthesis. However, excess Fe is toxic due to reactive hydroxyl radicals generated by the Fenton reaction (1). Thus, plants tightly regulate Fe homeostasis to avoid both Fe deficiency and Fe toxicity (2).

Although Fe is abundant in most soils, it is present in aerated soils as ferric (Fe^{3+}) oxihydrates, which are practically insoluble. To overcome the low solubility, plants rely on reduction and chelation-based mechanisms to make Fe bioavailable. Arabidopsis thaliana induces a set of biochemical activities to facilitate Fe uptake. Root plasma membrane H^{+}-adenosinetriphosphatas e release protons to acidify the rhizosphere (3) and thus, increase Fe solubility in the soil. In addition, coumarin family phenolics are released into the rhizosphere to chelate and mobilize Fe^{3+} (4). Fe^{3+} is then reduced to Fe^{2+} by the membrane-bound ferric chelate reductase enzyme (5), and the resulting Fe^{2+} is then transported into root epidermal cells by Iron-Regulated Transporter1 (IRT1) (6).

In Arabidopsis, FER-like iron deficiency-induced transcription factor (FIT) plays a key role in inducing Fe uptake genes in roots in response to Fe limitation (7–9). FIT is an ortholog of a basic helix–loop–helix (bHLH) transcription factor FER that controls Fe uptake responses in tomato (10). FIT is induced by Fe deficiency and forms a heterodimer with the subgroup Ib bHLH transcription factors (bHLH38, bHLH39, bHLH100, and bHLH101) to activate the transcription of FRO2 and IRT1 during Fe deficiency (11, 12). The loss of FIT or subgroup Ib genes impairs the induction of FRO2 and IRT1 and causes Fe deficiency chlorosis (7, 13, 14). Overexpression of FIT alone does not enhance Fe deficiency responses (7), but co-overexpression of FIT with bHLH38, bHLH39, or bHLH101 constitutively activates Fe uptake genes and improves tolerance to Fe deficiency (11, 12). Similarly, FIT is required for overexpressed bHLH39 to constitutively induce FRO2 and IRT1 (15).

Although we still do not know how FIT transcription is increased under Fe deficiency, overexpression of bHLH39 increases FIT expression under Fe deficiency, suggesting that bHLH39 is upstream of FIT and that FIT expression is controlled in part by a feedforward regulatory loop involving bHLH39 (15). The expression of subgroup Ib bHLH genes is induced by Fe deficiency; hence, there must be upstream regulatory elements that relay the Fe deficiency signal and activate these genes. The subgroup IVc bHLH transcription factors bHLH34, bHLH104, ILR3 (bHLH105), and bHLH115 are involved in activation of the subgroup Ib genes (16–18). The loss of each subgroup IVc gene undermines the induction of subgroup Ib genes and exacerbates iron deficiency.

**Significance**

More than 2 billion people are iron deficient, because their plant-based diets are not a rich source of this essential nutrient. Despite progress in tracing how iron moves throughout the plant, we still do not fully understand how plants sense and respond to iron availability. Here, we identify an essential basic helix–loop–helix transcription factor, Upstream Regulator of IRT1 (URI), and describe its central role as a Fe-dependent switch. The phosphorylated form of URI accumulates under Fe-deficient growth conditions and activates most known Fe deficiency-induced genes. Under Fe-replete growth conditions, the phosphorylated URI undergoes proteasomal degradation dependent on the E3 ligase BTS. Our study provides a molecular mechanism for Fe-dependent regulation of Fe deficiency signaling in plants.

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Data deposition: The ChIP-seq data and the DNA microarray data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, https://www.ncbi.nlm.nih.gov/geo (accession nos. GSE137645 and GSE137201, respectively).

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Fe deficiency symptoms under low-Fe supply (16–18). Conversely, overexpression of subgroup IVc genes increases the expression of subgroup Ib genes under all Fe conditions and enhances Fe uptake. Chromatin immunoprecipitation (ChIP)-qPCR assays showed that bHLH104, ILR3, and bHLH115 bind to promoters of the subgroup Ib genes when overexpressed in Arabidopsis protoplasts (16). Transactivation assays in tobacco leaves showed that either bHLH34 or bHLH104 prompts transcription from the bHLH110 promoter (17). Subgroup IVc genes are expressed under all Fe conditions, suggesting that the regulation of their activity occurs at the protein level so as to induce the expression of subgroup Ib genes only under Fe-deficient growth conditions.

The E3 ligase BTS is implicated in the degradation of subgroup IVc bHLH transcription factors (19, 20). Presumably, the protein abundance of subgroup IVc transcription factors is maintained at a higher level in the bts mutant than in the wild type, although protein levels have not yet been examined. The increase in subgroup IVc proteins would then enhance the expression of subgroup Ib genes and constitutively activate Fe uptake genes in the bts mutant. As a result, the bts mutant is more tolerant of Fe deficiency than the wild type under Fe deficiency compared with wild-type plants. Introducing bhlh104 or bhlh115 mutant alleles into the bts background mitigated the constitutive expression of Fe uptake genes, and double mutants become less tolerant to Fe deficiency compared with the bts mutant. The bts mutation also presses the Fe toxicity observed in the bts mutant, indicating that the loss of ILR3 also prevents constitutive Fe uptake (21). Yeast 2-hybrid assays demonstrated physical interaction between BTS and bHLH104, ILR3, or bHLH115 (19).

Here, we introduce a bHLH transcription factor Upstream Regulator of IRT1 (URI) and show that URI acts as part of the bHLH104, ILR3, or bHLH115 (19) heterodimer to activate the transcription of bHLH121. The URI protein was detected and expressed only under Fe deficiency (Fig. 2F). Additionally, we tested using the GFP-tagged URI construct in yeast. Multiple lines of each construct were examined, and the results from one representative line for each construct are presented here. Both constructs restored induction of pIRT1:Luc in the uri mutant under Fe deficiency (Fig. 2A). The uri mutant grows poorly on B5 medium, and such growth impairment is completely restored by expressing either construct. Next, we examined the IRT1 protein level to verify that endogenous IRT1 is indeed being expressed. Consistent with the luciferase activity, the IRT1 protein was detected and expressed only under Fe deficiency (Fig. 2B). The map-based cloning and the complementation test confirmed that bHLH121 is the causative gene of the uri mutant phenotype.

To understand the role of URI in IRT1 induction, we asked if URI expression is induced by Fe deficiency and where URI is expressed. Publicly available microarray data and our own analysis revealed that URI gene expression is not responsive to Fe availability, and URI is broadly expressed in all organs, including all layers of roots (SI Appendix, Fig. S1). We monitored the localization of the URI protein using the pURI:URI complementation lines. The GFP signal was observed in all root cell layers, consistent with the gene expression data (Fig. 2C), and suggested nuclear localization of the URI protein. The fluorescence intensity was similar between Fe-sufficient and Fe-deficient roots. We generated transgenic lines that express a β-glucuronidase (GUS) reporter gene regulated by the URI promoter (pURI:GUS) and examined the tissue distribution. The histochemical assay verified that URI is ubiquitously expressed in both shoots and roots, and the expression level is unchanged by Fe conditions (Fig. 2C).

Global Gene Expression Analysis Showed Deregression of Fe Deficiency-Induced Genes in the uri Mutant. Previously published gene expression data from Fe-sufficient and Fe-deficient wild-type roots were compiled to create a list of genes that show changes in gene expression between the 2 Fe conditions (Dataset S1). We used a 1.5-fold difference as a cutoff, and we identified 188 genes that are induced under −Fe conditions and 36 genes that are repressed under −Fe conditions. Next, we explored how loss of URI affected the expression of these Fe-regulated genes (Fig. 3A). Among the 188 Fe deficiency-induced genes, 42 genes (e.g., IREG3, ZIF1) were normally regulated in uri. Seven genes (e.g., WRKY22, NAC2) were hyperinduced in uri under −Fe conditions, suggesting that URI could function as a repressor, either directly or indirectly, of these genes. Meanwhile, 139 genes were not induced (or induced to a lesser extent) in uri compared with the wild type under −Fe conditions. This included transcription factors (e.g., FIT, bHLH39, bHLH101, PYE, MYB10, MYB72), genes involved in coumarin biosynthesis and release (e.g., 4CL1, F6H1, SHH, CYP82C4, ABCG37, BGLU42), Fe transport- and mobilization-related genes (e.g., IRT1, IRT2, FRO3, OPT3, NAS4), and Fe homeostasis genes (e.g., IMA1, BTS, BSL1, BSL2). All of these genes were not properly induced in uri under −Fe conditions, indicating that URI is directly or indirectly involved in activation of most of the known Fe deficiency-induced genes.

URI Encodes the bHLH Transcription Factor bHLH121. Using map-based cloning, we identified the URI gene as At3g19860, encoding the bHLH121 transcription factor bHLH121. The uri mutation is a C to T nucleotide substitution in the fourth exon that introduces a stop codon at amino acid position 103 (Q103*). The uri mutation is recessive, suggesting that the truncated protein is unlikely to be functional. Because no transfer (T)-DNA insertion alleles that disrupted URI function were found, we sought to confirm the identification of URI via complementation. We introduced a GFP-tagged URI construct into the uri mutant. In addition to the URI genomic DNA construct, including its native promoter (pURI:URI), a separate construct that has the URI coding sequence fused to the cauliflower mosaic virus (CaMV) 35S promoter (35S:URI) was tested.
Ferric chelate reductase activity. Plants were grown on half-strength B5− (Soil; expression. (graphed after luminescence imaging (Light). Plants were grown for 3 wk in erase substrate was supplied before imaging. The same plants were photo-

URI Directly Regulates the Majority of Fe-Responsive Genes. Knowing that URI is a bHLH transcription factor and that many of the Fe deficiency-inducement genes are not properly expressed in uri, we asked which genes are direct targets of URI using ChIP-seq analysis. We used the complementation line (pURI:URI) and prepared chromatin from Fe-sufficient and Fe-deficient roots. Chromatin fragments were communoprecipitated with an anti-GFP antibody, and sequencing libraries were created. One set of fragmented chromatin was incubated with immunoglobulin G, and the resulting library was used as a negative control during the peak analysis. We identified 2,366 binding regions and mapped them to the Arabidopsis genome (Dataset S2). A total of 1,623 potential target genes were identified (Fig. 4D). Of the 188 genes that are induced and the 36 genes that are repressed under −Fe conditions, 71 and 22, respectively, were direct targets of URI. Thus, URI directly controls ∼50% of Fe-regulated genes (Fig. 4C).

We also evaluated which genes showed differential binding depending on Fe conditions; 63 target genes showed an increase in URI binding, and 169 target genes showed decreased URI binding under −Fe conditions (Fig. 4D and Dataset S3). Among the 63 target genes with increased URI binding, 21 target genes were induced and 1 target gene was repressed under −Fe conditions (SI Appendix, Table S1). Of these Fe deficiency-induced genes, 19 genes are misexpressed in the uri mutant, indicating that URI acts as a principal activator for these genes under −Fe conditions. This included the Fe deficiency-induced transcription factors bHLH39, bHLH101, MYB10, MYB72, and PYE; Fe transport-related genes IRT1, FRO3, OPT3, NRAMP4, and NAS4; and Fe homeostasis genes IMA1, B5S, B5S1, ORG1, and CGLD27. One gene, VTE2, is repressed under −Fe conditions and showed increased URI binding, indicating URI acts as a repressor. Of the 169 target genes with decreased URI binding, 1 target gene was repressed and 3 target genes were induced under −Fe conditions. FER3 is repressed under −Fe conditions and showed decreased URI binding. This implies that URI binding contributes positively toward the expression of FER3 under +Fe conditions. Three other target genes (4CL1, GIK, KMD2) are induced under −Fe conditions, suggesting that URI acts as a repressor of these target genes. Noticeably, only a few target genes showing decreased URI binding are Fe regulated. This implies that Fe deficiency signaling is primarily due to induction of Fe deficiency responses.

We performed gene ontology (GO) enrichment analysis with URI target genes and asked in what other biological processes URI plays a role. Fe homeostasis-related terms were the most overrepresented constituents and included the Fe deficiency-induced genes bHLH38, bHLH101, IAM2, IRP3, and IRP6 (25).
These genes were missing from the list of Fe-responsive genes, because they are omitted from the ATH1 microarray platform. In addition to Fe homeostasis genes, we observed enrichment in GO terms associated with response to oxidative stress, regulation of circadian rhythm, defense response to bacterium, and lateral root formation. Other overrepresented GO terms included response to light stimulus, response to various hormones, phosphate starvation, water deprivation, and cold. This implicates URI in the regulation of various biological processes in addition to its principal role in activating most known Fe-regulated genes under −Fe conditions. We note that URI has been suggested to play a role in the potassium starvation response by binding directly to the HAK5 promoter and up-regulating its expression under low-K conditions (26). However, under our ChIP-seq conditions, HAK5 was not identified as a URI direct target.

**ChIP-qPCR Confirmed URI Binding to the Promoters of Subgroup Ib Genes.** To verify the ChIP-seq results, ChIP-qPCR assays were performed with the subgroup Ib transcription factors bHLH38, bHLH39, bHLH100, and bHLH101. ChIP-seq did not detect FIT as a direct target of URI and was included here as a negative control. Chromatin was prepared from the roots of the complementation line (pURI:URI) as well as the wild type to gauge the background level of anti-GFP antibody. Primer pairs were designed to amplify the E box or the G-box elements in the promoters covering peaks identified from ChIP-seq (SI Appendix, Fig. S2A). One primer pair residing within the coding region was included as a negative control. The promoter regions of subgroup Ib genes showed significant enrichment in Fe-deficient roots, demonstrating that the binding of URI to these targets increases under −Fe conditions (SI Appendix, Fig. S2B). The coding regions of subgroup Ib genes were not enriched in either Fe condition, consistent with the result from the ChIP-seq analysis, where no reads mapped to the coding region (Fig. 4B). The FIT gene was not enriched either in the promoter region or in the coding region, agreeing with the ChIP-seq results. ChIP-PCR confirmed that URI directly binds to the promoters of subgroup Ib genes and that the binding is increased under −Fe conditions.

**Ectopically Expressed Subgroup Ib bHLH Transcription Factors Restored the Induction of IRT1 in the uri Mutant.** As subgroup Ib bHLH transcription factors are direct targets of URI, we asked if ectopic expression of these target genes would rescue the uri phenotype. In order to override the requirement for URI, we expressed each of the subgroup Ib bHLH genes under the control of constitutive 35S promoter in the uri mutant; 35S promoter-driven FIT and PYE (35S:FIT and 35S:PYE) were also expressed in the uri mutant to explore the architecture of the regulatory network. FIT is not a direct target of URI; PYE is a direct target but not involved in the activation of IRT1.

First, the induction of the pIRT1:LUC reporter gene was examined in transgenic uri mutants carrying each construct. The uri mutant and transgenic uri mutants with either 35S:FIT or 35S:PYE did not induce pIRT1:LUC under Fe deficiency (Fig. 5A). In contrast, transgenic uri mutants with 35S promoter-driven subgroup Ib bHLH genes restored the luciferase induction under −Fe conditions. This demonstrates that the loss of URI is compensated by ectopically expressed subgroup Ib bHLH proteins. The defective root growth under low-Fe supply and the growth impairment on the B5 medium were also rescued. This confirms that subgroup Ib genes can reconstitute the downstream signaling cascade in the absence of URI. Importantly, these transgenic uri mutants induced the luciferase gene not only under −Fe conditions but also, in +Fe conditions.

![Fig. 2. URI identification. (A) Complementation of uri with GFP-tagged URI constructs. Luminescence imaging (LUC) of 7-d-old plants grown on Fe-sufficient (+) or Fe-deficient (−) media. The luciferase substrate was supplied before imaging. The same plants were photographed after luminescence imaging (Light). Plants were grown on B5 medium for 17 d and photographed. (B) IRT1 protein accumulation. Plants were grown on half-strength B5 medium for 14 d and transferred to +Fe or −Fe conditions for 3 d. Total protein was extracted from roots. The abundance of IRT1 was monitored by anti-IRT1 antibody. Tubulin (Tub) was used as a loading control and monitored using an antitubulin antibody. (C) Confocal fluorescence microscopy of roots from...](www.pnas.org/doi/10.1073/pnas.1916892116)
Expression level refers to the expression under Fe deficiency. Consistent with the reporter activity, the uri mutant and transgenic uri mutants with either 35S:FIT or 35S:PYE did not accumulate IRT1 protein regardless of Fe conditions (Fig. 5B). The uri transgenic lines with ectopically expressed subgroup Ib genes restore the accumulation of IRT1 protein in Fe-deficient roots. IRT1 protein was also detected in Fe-sufficient roots, which agrees with the constitutive luciferase activity.

Phosphorylated URI Protein Accumulates under Fe Deficiency. As the gene expression and the protein abundance of URI are not changed by Fe conditions, yet we see differential binding to target gene promoters, we asked if URI protein is modified in response to Fe availability. We first detected URI proteins in complementation lines pURI:URI and 35S:URI using an anti-GFP antibody. Both transgenic lines produced URI-GFP protein bands in both +Fe and −Fe roots with similar intensity. Noticeably, the blot revealed an extra band in −Fe roots that migrates slower than the expected size of the URI-GFP protein (Fig. 6A). The mobility shift toward a higher molecular weight often indicates a covalent addition of functional groups. As phosphorylation is the most common mechanism among posttranslational modifications (27), we examined if the mobility shift in URI-GFP protein is caused by phosphorylation. We treated protein extracts with the lambda phosphatase (λPP) and tested if the removal of phosphoryl groups eliminates the slower migrating band. The λPP is a manganese-dependent protein phosphatase, and its activity is inhibited by zinc ions (28). When the protein extract was incubated with manganese, the higher-molecular weight band was still detected when incubated with zinc ions. This demonstrates that the mobility shift in URI is caused by phosphorylation. Therefore, a phosphorylated form of URI accumulates under Fe deficiency.

URI is predicted to have multiple phosphorylation sites according to PhosPhAt 4.0 (29). Using URI protein purified from Fe-deficient plants and tandem mass spectrometry, we mapped 10 phosphoserine and 1 phosphothreonine residues in our analysis (Fig. 6C). Single nucleotide polymorphisms (SNP) information from the 1001 Genomes project (30) revealed Arabidopsis ecotypes with nonsynonymous mutations at 2 of these positions, S205 and S227, suggesting that phosphorylation of these residues may not be essential for URI function (Fig. 6C).

Phosphorylated URI Is Subject to Proteasome-Dependent Degradation under +Fe Conditions. To better understand the posttranslational modification of the URI protein, we investigated Fe-dependent changes in more detail using the complementation line (pURI:URI). The seedlings were grown on B5 medium for 14 d and transferred to Fe-sufficient or Fe-deficient growth conditions for 3 d. Roots were sampled each day, and total protein was extracted for western blot analysis. Consistent with the prior result, a single band matching to the expected size of the URI-GFP fusion protein was detected in all conditions at a constant intensity (Fig. 6D). The phosphorylated URI-GFP band appeared in Fe-deficient roots as early as day 1. To evaluate the Fe status of root samples, IRT1 protein was examined using an IRT1 antibody. The IRT1 band was detected in Fe-deficient roots after day 1, coinciding with the appearance of the phosphorylated URI.

This means that URI acts as a switch and determines when to initiate the Fe deficiency signaling cascade. Bypassing such control resulted in constitutive activation of Fe uptake activities and negatively impacted plant growth (SI Appendix, Fig. S3). Transgenic uri mutants constantly expressing subgroup Ib genes displayed inhibited root growth under Fe-sufficient conditions (Fig. 5C).
at a consistent level during Fe resupply, whereas the intensity of the phosphorylated URI-GFP band diminished as early as 6 h after transfer and became undetectable 12 h after Fe resupply (Fig. 6E). The IRT1 band diminished as early as 12 h and disappeared by 24 h. This demonstrated that the accumulation of the phosphorylated URI is Fe deficiency specific and that the removal of phosphorylated URI preceded IRT1 degradation. We asked if the disappearance of the phosphorylated URI-GFP band is due to protein degradation. Seedlings were incubated with MG132, a 26S proteasome inhibitor, during Fe resupply. When the proteasome inhibitor is present, phosphorylated URI persisted for at least 24 h (Fig. 6E). The level of nonphosphorylated URI remained unchanged. IRT1 protein, which is also subjected to proteasome-dependent degradation, persisted longer compared with the dimethyl sulfoxide (DMSO) control.

We then asked if BTS is involved in URI degradation. BTS is an Fe binding E3 ligase and interacts with subgroup IVc bHLH transcription factors. As yeast 2-hybrid failed to show a direct interaction between BTS and URI (19), we took an alternative approach and introduced pURI:URI into the bts-3 mutant. We examined the URI protein in the bts-3 mutant and compared the phosphorylation of URI between +Fe and −Fe conditions. Unlike the wild type, phosphorylated URI accumulated in bts-3 under +Fe conditions (Fig. 6F). This suggested that URI degradation is dependent on BTS and that BTS might regulate URI as it does subgroup IVc bHLH transcription factors (20).

URI Interacts with ILR3 and bHLH115 More Frequently under −Fe Conditions. Subgroup IVc bHLH transcription factors have been shown to share common binding targets with URI, such as the genes encoding group Ib bHLH transcription factors and PYE. The E3 ligase BTS physically interacts with 3 of the group IVc proteins (bHLH104, ILR3, and bHLH115) and mediates 26S proteasome-dependent degradation of at least 2 of these transcription factors (20). BTS is required for the degradation of phosphorylated URI under +Fe conditions. Therefore, we hypothesized that URI and group IVc bHLH transcription factors form heterodimers, bind to common target promoters, and become substrates for the E3 ligase BTS. Indeed, genome-wide yeast 2-hybrid screening with transcription factors from Arabidopsis indicated an interaction between URI and bHLH34 as well as bHLH104 (31). The orthologous transcription factors MdbHLH104...
and MdbHLH121 from apple were shown to interact in yeast 2-hybrid and pull-down assays with proteins expressed in *Escherichia coli* (32).

To test if URI interacts with the other subgroup IVc bHLH transcription factors, we conducted a pull-down assay using in vitro-translated ILR3 and bHLH115 proteins that were fused to a Halo tag. The complementation line (*pIRT1:Luc*) luminescence imaging (LUC) of 7-d-old plants grown on +Fe or −Fe media. The luciferase substrate was supplied before imaging. The same plants were photographed after luminescence imaging (Light). Plants were grown on B5 medium for 17 d and photographed. (B) IRT1 protein accumulation. Plants were grown on half-strength B5 medium for 14 d and transferred to +Fe or −Fe conditions for 3 d. Total protein was extracted from roots. The abundance of IRT1 was monitored with an anti-IRT1 antibody. Tubulin (Tub) was used as a loading control and monitored using an anti-tubulin antibody. WT, wild type.

**Fig. 5.** Ectopic expression of subgroup Ib bHLH transcription factors restores IRT1 induction. (A) Ectopic expression of subgroup Ib genes, FIT, and PYE in the *uri* mutant carrying *pIRT1:Luc*. Luminescence imaging (LUC) of 7-d-old plants grown on +Fe or −Fe media. The luciferase substrate was supplied before imaging. The same plants were photographed after luminescence imaging (Light). Plants were grown on B5 medium for 17 d and photographed. (B) IRT1 protein accumulation. Plants were grown on half-strength B5 medium for 14 d and transferred to +Fe or −Fe conditions for 3 d. Total protein was extracted from roots. The abundance of IRT1 was monitored with an anti-IRT1 antibody. Tubulin (Tub) was used as a loading control and monitored using an anti-tubulin antibody. WT, wild type.

**Discussion**

In our screen for plants with altered expression of the Fe transporter IRT1, we identified *URI*, an essential gene encoding a bHLH transcription factor that plays a key role in the Fe regulatory cascade. The 159 bHLH transcription factors in *Arabidopsis* have been grouped into 26 subfamilies (33–37), and URI belongs to subgroup IVb along with bHLH11 and PYE (bHLH47). PYE is a transcriptional repressor that attenuates the expression of Fe deficiency-induced genes in roots, such as *NAS4* and *ZIF1*, that are needed for the internal mobilization of Fe and its transport to the shoot (19). bHLH11 was recently also reported to act as a negative regulator of Fe homeostasis (38), implicating all 3 members of subgroup IVb in the Fe deficiency response. URI orthologs can be found throughout the plant kingdom, suggesting conservation of the signaling cascade to which URI belongs.

Up until now, there were 2 sets of Fe-regulated genes described in the literature for *Arabidopsis*: FIT-dependent and FIT-independent genes (7–9, 13, 39). Our discovery that URI regulates both sets of genes places URI early in the Fe regulatory cascade. For example, among FIT-independent genes are the subgroup Ib bHLH1 transcription factors that are direct targets of URI, and they regulate the expression of FIT and FIT-independent genes. URI controls ∼50% of Fe-regulated genes directly binding to their promoters (Fig. 4C), and loss of URI perturbs the expression of ∼73% of Fe-regulated genes (Fig. 3B). PYE is a direct target of URI, placing PYE direct targets under indirect control of URI. *BTS* and *BTS1* (40, 41) are also direct targets of URI, and once again, this places the proteins that they control via targeted degradation indirectly under the control of URI (Fig. 7).

One of the newest additions to the Fe regulatory cascade is an 8-member family of I RM O N MAN peptides, which are thought to be phloem-mobile signals that act to control Fe uptake through an as yet unknown pathway (42). These are strongly Fe regulated, and we show here that *IMA1* (FEP3) (43) and *IMA2* (FEP2) (43) are also direct targets of URI.

Although more than a dozen transcription factors have now been placed into the Fe regulatory network, up until our discovery of URI, only FIT had been shown to be essential for growth under Fe deficiency (7–9). In addition to both being essential transcription factors required for growth under Fe deficiency, FIT and URI share other characteristics. FIT and URI form heterodimers with multiple members of partially redundant groups of transcription factors: FIT with members of the subgroup Ib transcription factors (11, 12) and URI with members of...
the subgroup IVc transcription factors. URI and FIT are both posttranslationally modified via phosphorylation (44), and finally, both URI and FIT are subject to proteasome-mediated degradation (45, 46). URI differs from FIT in being ubiquitously expressed throughout the plant regardless of Fe status, whereas FIT is expressed mainly in the epidermal layer in the roots of Fe-deficient plants (7). Surprisingly, we do not know which transcription factor(s) directly up-regulate FIT expression under Fe deficiency. Overexpression of the group Ib proteins is sufficient to turn on IRT1, suggesting that it may also turn on FIT, perhaps as a heterodimer with FIT.

Pulling together data from both our study and the literature, there is evidence for interaction of URI with all 4 of the subgroup IVc transcription factors. We were able to coimmunoprecipitate (co-IP) 2 subgroup IVc transcription factors, ILR3 and bHLH115, using roots from plants expressing URI-GFP. We have also shown that URI can interact with these 2 transcription factors in an in vitro pull-down assay. BioGrid reports 16 interactors for URI from a high-throughput yeast 2-hybrid analysis (31), including the other 2 members of the subgroup IVc proteins: bHLH34 and bHLH104. The A. thaliana subgroup IVc transcription factors have been reported to form homodimers as well as heterodimers with each other. Similar results have been reported for the subgroups IVb and IVc proteins from apple, where MdbHLH104 was shown to interact with MdbHLH105, MdbHLH115, MdbHLH11, and MdURI (32).

The Arabidopsis subgroup IVc proteins have been shown to be degraded by the E3 ligase BTS. Our observation that phosphorylated URI accumulates in the bts-3 mutant under +Fe conditions (Fig. 6F) suggests that degradation of phosphorylated URI is also dependent on BTS. Because URI controls both FIT-dependent and FIT-independent genes, the removal of phosphorylated URI under +Fe conditions will turn off the transcription of many Fe deficiency-induced genes. A role for BTS under Fe sufficiency agrees with previous results from our laboratory showing that, in the bts-3 mutant, many Fe deficiency-induced genes are expressed even under Fe-sufficient growth conditions, and as a result, this mutant accumulates high levels of Fe (40).
observations in rice, where the BTS orthologs, HRZ1/2, are crucial for repressing iron deficiency responses and protecting cells from iron toxicity in the presence of excess iron (47).

If URI is working as a heterodimer with the subgroup IVc proteins as we suggest, we would expect URI and the subgroup IVc transcription factors to share direct targets. Although we have used ChIP-seq to generate a comprehensive list of URI direct targets in roots of Fe-sufficient and Fe-deficient plants, only a limited set of potential subgroup IVc direct target genes has been examined. Based on transient expression in protoplasts followed by ChIP-qPCR, bHLH104, ILR3, and bHLH115 have been shown to bind to the promoters of the 4 subgroup Ib transcription factor genes (bHLH38/39/100/101) as well as PYE (16). In addition, using chromatin prepared from Fe-deficient roots and ChIP-qPCR, ILR3 has been shown to bind to E boxes in the promoters of bHLH39, FER1, FER3, FER4, VTL2, NEET, and NAS4 (48). All of these genes are URI direct targets. Of course, binding to the same promoter regions does not prove that URI and the subgroup IVc proteins bind as a heterodimer. The transient expression and overexpression data available for various subgroup IVc proteins suggest that, under some conditions, the IVc proteins are certainly capable of binding in the absence of URI. For example, transient expression of GFP constructs driven by each of the subgroup Ib promoters (bHLH38/39/100/101) in Nicotiana benthamiana could be induced by coexpression of bHLH34 or bHLH104 (17).

Putting together our work on URI with work from many different laboratories, we propose a model (Fig. 7) in which phosphorylated URI is allowed to accumulate when plants become Fe deficient and interacts with subgroup IVc bHLH transcription factors. These heterodimers bind to the promoters of many Fe deficiency-induced genes, including subgroup Ib bHLH transcription factor genes, PYE, BTS, and BTSL. Subgroup Ib transcription factors increase the expression of FIT, and form heterodimers with FIT to activate the transcription of IRT1 and FRO2. FIT-dependent gene expression increases iron uptake in roots. PYE negatively regulates genes that are involved in Fe mobilization and increases iron availability in plant cells. The transcription of E3 ubiquitin ligase BTS and BTSL is up-regulated by Fe deficiency, but the protein activity is promoted by Fe binding. After Fe becomes available, E3 ubiquitin ligases (orange) interact with the transcription factor complexes, leading to their proteasomal degradation. This Fe-dependent degradation of URI and FIT via BTS family members provides the mechanism to turn off Fe deficiency signaling and prevent Fe overload. The difference in tissue distribution determines which E3 ligase is used. BTSL and FIT are root specific; BTS and URI are expressed throughout the plant.
Gene Expression Analysis. Details of RNA extraction, microarray analysis, and qRT-PCR are described in SI Appendix, SI Materials and Methods. The primer sequences are listed in SI Appendix, Table S2. Microarray data are available with the accession number GSE137201 in the Gene Expression Omnibus.

Protein Analysis. Details of protein extraction, phosphatase treatment, immunoblot analysis, in vitro protein expression, and a pull-down assay are described in SI Appendix, SI Materials and Methods.

ChIP-seq and ChIP-qPCR. ChIP is described in detail in SI Appendix, SI Materials and Methods. The primer sequences for ChIP-qPCR are listed in SI Appendix, Table S2. ChIP-seq data are available with the accession number GSE137645 in the Gene Expression Omnibus.

Commmunprecipitation, Phosphomapping, and Mass Spectrometry. The complementation line (pUR1:UR1) was used to purify UR1 proteins. Details of immunoprecipitation, phosphomapping, and mass spectrometry are described in SI Appendix, SI Materials and Methods.

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