RESEARCH PAPER

Alkaline stress and iron deficiency regulate iron uptake and riboflavin synthesis gene expression differently in root and leaf tissue: implications for iron deficiency chlorosis

En-Jung Hsieh and Brian M. Waters*

Department of Agronomy and Horticulture, University of Nebraska-Lincoln, Lincoln, NE 68583-0915, USA

* Correspondence: bwaters2@unl.edu

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Abstract

Iron (Fe) is an essential mineral that has low solubility in alkaline soils, where its deficiency results in chlorosis. Whether low Fe supply and alkaline pH stress are equivalent is unclear, as they have not been treated as separate variables in molecular physiological studies. Additionally, molecular responses to these stresses have not been studied in leaf and root tissues simultaneously. We tested how plants with the Strategy I Fe uptake system respond to Fe deficiency at mildly acidic and alkaline pH by measuring root ferric chelate reductase (FCR) activity and expression of selected Fe uptake genes and riboflavin synthesis genes. Alkaline pH increased cucumber (Cucumis sativus L.) root FCR activity at full Fe supply, but alkaline stress abolished FCR response to low Fe supply. Alkaline pH or low Fe supply resulted in increased expression of Fe uptake genes, but riboflavin synthesis genes responded to Fe deficiency but not alkalinity. Iron deficiency increased expression of some common genes in roots and leaves, but alkaline stress blocked up-regulation of these genes in Fe-deficient leaves. In roots of the melon (Cucumis melo L.) fefe mutant, in which Fe uptake responses are blocked upstream of Fe uptake genes, alkaline stress or Fe deficiency up-regulation of certain Fe uptake and riboflavin synthesis genes was inhibited, indicating a central role for the FeFe protein. These results suggest a model implicating shoot-to-root signaling of Fe status to induce Fe uptake gene expression in roots.

Key words: Bicarbonate, cucumber, fefe mutant, iron deficiency chlorosis, iron uptake, melon, shoot-to-root signaling.

Introduction

Iron (Fe) is an important micronutrient that plays crucial roles in plant growth, development, and reproduction (Walker and Waters, 2011; Kobayashi and Nishizawa, 2012; Vigani et al., 2013). Iron uptake into roots of graminaceous plant species (known as Strategy II) is characterized by production and secretion of phytosiderophores that chelate Fe(III) for uptake (Kobayashi and Nishizawa, 2012). Iron uptake by non-graminaceous angiosperm species (known as Strategy I) is characterized by rhizosphere acidification by H⁺-ATPase proteins, reduction of Fe(III) to Fe(II) by ferric chelate reductase (FCR) proteins, and uptake of Fe(II) by iron transporter proteins (Kobayashi and Nishizawa, 2012). The activity of these Fe uptake proteins is up-regulated in Fe-deficient roots.

Many molecular components of the Strategy I Fe uptake system have been identified. FERRIC REDUCTION OXIDASE 2 (FRO2) encodes the primary root surface FCR in Arabidopsis (Robinson et al., 1999), and the corresponding genes in cucumber (Cucumis sativus L.) and melon (Cucumis melo L.) are called FROI (Waters et al., 2007, 2014). Iron (II) transporter genes include IRT1 (Eide et al., 1996; Varotto
et al., 2002; Mukherjee et al., 2006) and NRAMP1 (Curie et al., 2000; Thomine et al., 2000; Cailléat et al., 2010). AtFRO2, AtIRT1, AtNRAMP1, and numerous other genes are up-regulated under Fe deficiency by the basic helix-loop-helix (bHLH) transcription factor AtFIT (Colangelo and Guerinot, 2004; Jakoby et al., 2004; Yuan et al., 2005), a homolog of the tomato FER protein (Ling et al., 2002; Colangelo and Guerinot, 2004). AtFIT gene expression is typically up-regulated by Fe deficiency (Colangelo and Guerinot, 2004; Jakoby et al., 2004; Lucena et al., 2006). A group of four closely related Arabidopsis bHLH genes, AtbHLH38, AtbHLH39, AtbHLH100, and AtbHLH101, are classified in clade Ib of the bHLH superfamily (Wang et al., 2007). The AtFIT protein regulates expression of its target genes as a heterodimeric complex of AtFIT and a clade Ib bHLH protein (Colangelo and Guerinot, 2004; Yuan et al., 2008; Wang et al., 2013). The melon C940-fe mutant (fefe) (Nugent and Bhella, 1988, Nugent, 1994) is similar to the fit mutant, in that it does not up-regulate FCR activity and rhizosphere acidification under Fe deficiency, and is chlorotic (Nugent, 1994). Our characterization of the fefe mutant showed that 82 genes that were up-regulated by Fe deficiency in wild-type roots were not regulated by Fe deficiency in the fefe mutant, including key Fe uptake genes (Waters et al., 2014). Thus, both fit and fefe mutants have blocked Fe uptake responses upstream of Fe uptake genes.

Less is known about molecular Fe deficiency responses in leaves than in roots, as only a few studies have profiled genome-wide gene expression in leaves (Waters et al., 2012; Ivanov et al., 2012; Rodríguez-Celma et al., 2013a; Moran Lauter et al., 2014). Several Fe regulated genes respond to Fe deficiency in both leaf and root tissues, whereas some are specific to roots or leaves. While AtFIT is only expressed and regulated by Fe in roots, the transcripts of AtbHLH38, AtbHLH39, AtbHLH100, and AtbHLH101 are up-regulated in both roots and leaves of Fe-deficient Arabidopsis (Wang et al., 2007; Rodriguez-Celma et al., 2013a). In our previous Arabidopsis leaf microarray study (Waters et al., 2012), At1G47400 (named iron responsive protein 1 (AtIRPI) in Rodriguez-Celma et al., 2013a) and AtKCS17 (3-ketoacyl-CoA synthase, At4G34510) were among the most strongly up-regulated genes in Fe-deficient Arabidopsis leaves (10.2-fold for AtIRPI and 36.0-fold for AtKCS, respectively). A more detailed knowledge of leaf Fe deficiency responses is needed to understand whole-plant adaptations to low Fe conditions, since a leaf-originated signal is thought to be necessary for normal regulation of Fe deficiency responses in roots (De Nisi et al., 2012; García et al., 2013).

Although Fe is abundant in soils, it has low solubility, especially in calcareous, alkaline soils, which occur on 30% of the earth (Chen and Barak, 1982). Plants can show iron deficiency chlorosis (IDC) on alkaline soils (Mengel and Geurtzen, 1986), resulting in reduced growth and yield (Hansen et al., 2004; Rogovska et al., 2007; Briat et al., 2015). Uptake of other metal micronutrients, such as Mn and Zn, can also be inhibited in alkaline soils (George et al., 2012). Soil alkalinity is largely due to bicarbonates (HCO$_3^-$) and carbonates (CO$_3^{2-}$) (Coulombe et al., 1984; Mengel et al., 1984), and therefore, bicarbonate has been commonly used to induce IDC symptoms in hydroponic Fe nutrition studies (Coulombe et al., 1984; Chaney et al., 1992; Romera et al., 1992a; Lin et al., 1997; Waters and Troupe, 2012). However, whether IDC results from low Fe supply, alkaline stress, or a combination of these factors is still unclear. Most IDC studies to date have not treated Fe supply and alkalinity as separate variables. Studies that applied bicarbonate to both Fe-deficient and Fe-sufficient plants (Fleming et al., 1984; Romera et al., 1992a; Alcántara et al., 2000) were carried out prior to modern molecular methods, and thus it is not clear whether low Fe supply and alkaline stress cause equivalent molecular responses. In Strategy I species, bicarbonate-treated Fe-deficient plants had low root expression of FIT, FRO2, and IRT1, and had inhibited root FCR activity compared with Fe-deficient plants grown without bicarbonate (Romera et al., 1997; Lucena et al., 2007; García et al., 2014). However, we found that cucumber FCR activity was stimulated by bicarbonate treatment in plants supplied with Fe (Waters and Troupe, 2012). Except for Romera et al. (1992a), pH-matched control treatments have not been included in these studies. In pilot studies in our lab with sodium bicarbonate, potassium bicarbonate, and HEPES buffer, the FCR activity response of plants in pH buffered treatments was not distinguishable from the FCR response of bicarbonate-treated plants, regardless of the counter-ion. As such, we use the term alkalinity to refer to bicarbonate treatments. One of our objectives was to determine how pH influences physiological and molecular responses to Fe deficiency in roots and leaves.

Iron deficient Strategy I plant species have long been known to increase efflux of root exudates (Cesco et al., 2010). Some species, such as Arabidopsis thaliana, produce phenolic compounds (Fourcroy et al., 2014; Schmid et al., 2014; Schmidt et al., 2014) while other species, including cucumber and melon, produce flavin compounds (Susin et al., 1993; Welkie, 2000; Rodriguez-Celma et al., 2011b). Although the function of flavin compounds in plant Fe deficiency is not well defined, they may function in reduction or complexation of extracellular Fe to facilitate Fe acquisition (Cesco et al., 2010; Sisó-Terraza et al., 2016). Proteins involved in riboflavin synthesis increased in abundance in response to Fe deficiency or Fe deficiency in alkaline conditions (Rellán-Alvarez et al., 2010; Rodriguez-Celma et al., 2011b) and genes involved in riboflavin biosynthesis were up-regulated in iron-deficient roots in normal or alkaline conditions (Rellán-Alvarez et al., 2010; Rodriguez-Celma et al., 2013b). However, the expression of riboflavin synthesis genes by alkaline stress separately from Fe deficiency has not been studied. Thus, another objective was to determine how riboflavin synthesis genes respond to Fe deficiency and alkaline stress in roots and leaves.

We addressed the objectives above using molecular and physiological approaches, by measuring leaf chlorosis, root FCR activity, Fe and other metal micronutrient accumulation, and expression of Fe uptake genes and riboflavin synthesis genes in cucumber roots and leaves. Our third objective was to use the melon fefe mutant to determine if alkaline-stimulated root gene expression depends on the fefe Fe deficiency regulatory pathway. The results of this research will
lead to increased understanding of how alkaline stress inhibits Fe uptake and causes IDC, and will allow improved design of future studies to develop alkaline stress-tolerant crop varieties.

Materials and methods

Plant materials and growth conditions

Cucumber seeds of cv Ashley were purchased from Eden Brothers (Asheville, NC, USA) and seeds of Miniature White were purchased from Jonny’s Selected Seeds (Winslow, ME, USA). Melon seeds of cv Edisto were purchased from Victory Seed Company (Molalla, OR, USA), and seeds of C940-fe (fefe; Nugent, 1994) were a gift from Michael A. Grusak (USDA-ARS Children’s Nutrition Research Center, Houston, TX, USA). Seeds were germinated in germination paper soaked with 0.1 mM CaSO₄ and incubated in the dark at 30 °C for 3 days. Seedlings were transferred to black tubs (four seedlings per tub) with 750 mL of nutrient solution made with 1.5 mM KNO₃, 0.8 mM Ca(NO₃)₂, 0.3 mM (NH₄)₂HPO₄, 0.2 mM MgSO₄, 25 μM CaCl₂, 25 μM H₃BO₃, 2 μM MnCl₂, 2 μM ZnSO₄, 0.5 μM Na₂MoO₄, 0.1 μM CuSO₄, and 1 mM MES buffer (pH 5.5). Iron was supplied as Fe(III)–ethylenediamine-N,N′-bis(2-hydroxyphenylacetate) (EDDHA) (Sprint 138, Becker-Underwood) at concentrations indicated below. The Fe(III)–EDDHA chelate is stable at the mildly acidic and alkaline pH used in this study (Chaney et al., 1972; Halvorsen and Lindsay, 1972). Plants were grown in a growth chamber at 22 °C with a 16 h photoperiod and photosynthetic photon flux density of 300 μmol m⁻² s⁻¹ photosynthetically active radiation. Bicarbonate was supplied as potassium bicarbonate (Fisher Scientific) or sodium bicarbonate (Arm & Hammer) with no difference in results. Plants were grown for 4 days in complete solution with 0.5, 1.0, 2.5, or 10 μM Fe (pretreatment), followed by 3 days of treatment with bicarbonate (10 mM) at the same Fe concentration as during the pretreatment period. At the end of the treatment, the first true leaf was still growing and the second leaf was emerging (see Supplementary Fig. S1 at JXB online for plant DW). Final nutrient solution pH was determined using a pH meter at the end of the treatment period.

Ferric chelate reductase activity and chlorophyll content

Ferric chelate reductase activity was measured using whole roots of individual cucumber plants after 3 days of treatment. Roots were excised, rinsed in deionized water, and submerged in 20 mL assay solution (1 mM MES buffer, pH 5.5, 150 μM Fe(III)-EDTA, and 200 μM ferrozone) for 30–60 min. Ferrozone–Fe(II) was measured by absorbance at 562 nm (subtracting blanks of assay solution with no plants) and reduced Fe was calculated using the extinction coefficient of 28.16 mM⁻¹ cm⁻¹. Chlorophyll of the first true leaf was determined using a SPAD-502 chlorophyll meter (Minolta). Significance of differences between Fe treatments at each bicarbonate level, and between bicarbonate treatments at each Fe level were determined by Student’s t-test.

Quantification of dry weight and mineral content

Note that with Fe(III)EDDHA as an Fe source, Fe does not accumulate in the apoplast, in contrast to Fe sources such as Fe(III)EDTA (Longnecker and Welch, 1990; Strasser et al., 1999) or FeSO₄ (Waters and Bleivins, 2000). Plants were dissected into roots, first true leaf, and the remainder of the plant (stem+cotyledons) after 3 days treatments and dried at 70 °C in a drying oven. After measuring dry weight (DW), tissue samples were digested in concentrated nitric acid–hydrogen peroxide with stepwise heating at 100, 125, 150, and 165 °C to dryness, and then resuspended in 5 mL 1% HNO₃ (Guttieri et al., 2015). Iron, Cu, Zn, and Mn contents for plant parts were quantified by inductively coupled plasma mass spectrometry in the University of Nebraska Redox Biology Center Spectroscopy Facility. Total plant DW or mineral content for each replicate plant was determined by summing plant parts. Significant differences between treatments for DW and each mineral were determined by one-way ANOVA.

RNA preparation, RT-PCR and real-time PCR analyses

Total RNA was extracted from approximately 80 mg of frozen tissues of cucumber or melon plants using the RNAeasy Plant Mini Kit (Qiagen). RNA samples were treated with DNaseI (Promega, USA) and RNA quality and concentration were determined by A₂₆₀/A₂₈₀ ratio. For reverse transcription reactions, 3 μg of total RNA was used with the High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, CA, USA). Single-stranded cDNA corresponding to 15 ng of total RNA was used as a template in 15 μL total volume for real-time PCR assay. Real-time PCRs were carried out with 667 nM of gene-specific primers and GoTag qPCR Master Mix (Promega, USA) using an IQ5 MyiQ detection system (Bio-Rad, Hercules, CA, USA). Coding sequences for cucumber and melon bHLH38, bHLH101, NRAMP1, RIBAI, PYRD, PHS1, DMRLs and Fe6 H1 (Table 1) were identified from Cucurbet Genomics Database (http://www.icugi.org/cgi-bin/CuGI/index.cgi) or from the Melonomics database (https://melenomics.net/) by BLAST searches and phylogenetic analysis. Primer sequences (Supplementary Table S1) were designed using the NCBI primer design tool (http://www.ncbi.nlm.nih.gov/tools/primer-blast/index.cgi). The thermal cycler program was one initial cycle of 95 °C, 8 min 30 s; followed by cycles of 95 °C, 10 s; 57 °C, 30 s; 72 °C, 15 s, with 40 cycles; followed by melt curve analysis for all genes. The relative gene expression considered the 10 μM Fe–0 bicarbonate-treated roots as control, and was calculated using the equation Y = 2⁻ΔΔCt, where

| Gene name | Cucumber ID | Melon ID | Arabidopsis ID | Description | WT +Fe | WT -Fe | WT FC | fefe +Fe | fefe -Fe | fefe FC |
|-----------|-------------|----------|----------------|-------------|-------|-------|-------|---------|---------|--------|
| RIBA1     | Csa4M111580 | Melo3C0024826 | AT5G64300 | GTP cyclohydrolase II | 656 | 11 118 | 17.0 | 118 | 110 | 0.9 |
| PYRD      | Csa6M003430 | MUS1870 | AT4G20960 | Diaminohydroxypyrophosphorylaminopyrimidine deaminase | 369 | 505 | 1.4 | 415 | 394 | 0.9 |
| PHS1      | Csa1M655920 | Melo3C010048 | AT3G47390 | Pyrimidine reductase | 177 | 1670 | 9.4 | 27 | 25 | 1.0 |
| DMRLs     | Csa6M266300 | MUS9012 | AT2G44050 | Dimethyl-ribityllumazine synthase | 1838 | 6234 | 3.4 | 1690 | 1142 | 0.7 |
| RIBC      | Csa6M128550 | MUS45607 | AT2G20690 | Riboflavin synthase α chain | 220 | 385 | 1.8 | 302 | 233 | 0.8 |
\[ \Delta C_i = \text{test gene} (C_{i, \text{treatment}} - C_{i, \text{control}}) - \text{ubiquitin} (C_{i, \text{treatment}} - C_{i, \text{control}}). \]

At least three independent RNA extractions (from different plants) and RT-qPCR reactions with two technical replicates per sample were performed. Significance of differences between Fe treatments at each bicarbonate level, and between bicarbonate treatments at each Fe level were determined by Student's t-test.

Results

Physiological Fe deficiency and alkaline stress responses in cucumber

To understand the plant response to Fe supply at normal hydroponic pH and at alkaline pH, we grew plants with or without bicarbonate over a range of Fe supply, using pH-stable Fe(III)-EDTA to ensure that Fe was soluble at all pH treatments (Chaney et al., 1972; Halvorson and Lindsay, 1972). Without bicarbonate, final solution pH was lower (pH 4.9) at the lowest Fe supply (0.5 μM) compared with final solution pH of the 10 μM Fe control (pH 6.0; Fig. 1A), likely reflecting increased H⁺-ATPase activity at low Fe supply. With bicarbonate, the final solution pH did not vary by Fe supply, and was 8.3–8.4 in all treatments. The first leaf of cucumber plants was slightly more chlorotic at the lowest Fe supply without bicarbonate, indicating that none of the Fe treatments caused severe Fe deficiency at normal pH. With alkaline stress (10 mM bicarbonate), the chlorophyll content of the first leaf was significantly lower at each Fe concentration than in plants without bicarbonate (Fig. 1B). The root FCR activity was highest at 0.5 μM Fe supply without bicarbonate, and was also significantly up-regulated at 1.0 and 2.5 μM Fe (Fig. 1C). However, plants supplied with bicarbonate had similar FCR activity at all Fe concentrations. The FCR activity of plants treated with bicarbonate was significantly higher than that of plants without bicarbonate at 2.5 and 10 μM Fe, but was lower than that of plants without bicarbonate at 0.5 μM Fe. Thus, while low Fe supply and alkaline stress both resulted in elevated FCR activity in cucumber roots, the patterns were quite different. Comparing bicarbonate treatments at specific Fe supplies, bicarbonate caused either inhibition (0.5 μM Fe) or stimulation (10 μM Fe) of FCR activity. We obtained similar chlorophyll and FCR activity results with another variety of cucumber (Miniature White; Supplementary Fig. S2). These results indicated that alkaline stress elevated the root FCR activity relative to Fe-replete plants, but abolished the normal response to Fe supply.

For an additional comparison of plant response to Fe deficiency and alkaline stress, we measured plant biomass and mineral accumulation in plant tissues. Biomass of the total plant or plant parts was not significantly affected by the treatments, except for roots of 0.5 Fe–10 mM bicarbonate-treated plants, which were slightly larger than roots of the other treatments (Supplementary Fig. S1). The control plants grown with 10 μM Fe had the highest total Fe content, which decreased by about half under mild Fe deficiency (0.5 μM Fe; Fig. 2A). Bicarbonate treatment at 10 μM Fe supply resulted in an approximately 40% decrease in total Fe content, indicating that alkaline stress inhibited Fe uptake, despite using a pH-stable Fe source. The low Fe, bicarbonate-treated plants had the lowest Fe content in leaves, at 19% of the control value. Iron content decreased in roots and stem+cotyledon with low Fe and/or bicarbonate treatment (Fig. 2A). For other metal micronutrients, Cu, Zn, and Mn contents decreased in leaf tissue of bicarbonate-treated plants in both Fe supply regimes (Fig. 2B–D). In stem+cotyledon, Zn and Mn decreased under bicarbonate treatment, while Cu content was unchanged. In root, Mn and Zn increased in low Fe
Alkaline stress disrupts iron deficiency signaling and bicarbonate treatments, while Cu increased only in low Fe treatments.

**Molecular responses to Fe deficiency and alkaline stress in cucumber**

To gain a molecular understanding of similarities and differences between molecular responses to Fe deficiency and alkaline stress, we measured gene expression of known Fe deficiency responsive genes in root and first leaf tissue of cucumber plants. The Fe uptake transcription factor genes CsFIT and CsHLH38 were up-regulated by Fe deficiency in roots (Fig. 3A, B), but CsFIT was not further induced by bicarbonate treatment at low Fe and was not induced by bicarbonate at replete Fe. Transcripts of CsHLH38 and CsbHLH101 were greatly induced by bicarbonate in roots with low Fe supply, but there was no stimulation by bicarbonate in roots with normal Fe supply (Fig. 3B, C). The expression of CsIRT1 and CsbFRO1 was increased under Fe deficiency (Fig. 3D, E), and their expression was increased further by bicarbonate at low Fe concentration. Bicarbonate also stimulated expression of these genes at normal Fe supply. In contrast, while the expression of CsNRAMP1 was strongly induced by Fe deficiency (Fig. 3F), bicarbonate treatment at low Fe supply resulted in much lower induction compared with the treatment without bicarbonate. Two additional genes based on previous leaf microarray results from Fe-deficient Arabidopsis (Waters et al., 2012) were studied. The 5960 gene (Csa2M005960, homologous to AtIRP1) had an expression pattern similar to bHLH38. The CsKCS gene (Csa2M361630) was not consistently regulated by Fe or bicarbonate in roots, although some treatments were statistically significant.

In leaf tissue, two patterns of gene expression were apparent, both of which were substantially different from root gene expression patterns. CsFIT, CsbFRO2, or CsNRAMP1 transcripts were not detected in leaf tissue. None of the other genes' expression increased in leaf in response to bicarbonate at full Fe supply (Fig. 4). The transcripts of CsbHLH38 and Cs5960 were highly induced in leaf tissue under Fe deficiency, but this induction was completely abolished in the leaf of bicarbonate-treated plants (Fig. 4A, D). However, these transcripts were synergistically increased by Fe deficiency and alkaline stress in roots (Fig. 3B, G). CsbHLH101 and CsIRT1 were induced by Fe deficiency, to a lower extent than in roots, but the addition of bicarbonate diminished their expression in the Fe-deficient leaf to control levels (Fig. 4B, C), whereas their expression was stimulated in roots (Fig. 3C, D). The expression of CsKCS was induced by Fe deficiency in leaf, but not in root (Figs. 3H and 4E). Thus, alkaline stress interfered with the molecular leaf response to low Fe supply.

**Fig. 2.** Mineral content in cucumber (cv. Ashley) plants. Cucumber seedlings were transferred into hydroponics with 0.5 or 10 μM Fe for 4 days pretreatment, and then treated with or without 10 mM bicarbonate for 3 days. The roots, first leaf, and stem+cotyledons of each plant were harvested separately. (A) Fe content, (B) Cu content, (C) Zn content, and (D) Mn content in different parts of plants and whole plants. Bars represent mean±SD (n=8). Different letters indicate significant differences (P<0.05) based on ANOVA using the Holm–Sidak method; NS indicates no significant differences between treatments.
Expression of riboflavin synthesis pathway genes

We identified homologs of genes known to be involved in the riboflavin synthesis pathway (Table 1) and measured their gene expression in response to Fe deficiency and alkaline stress. The riboflavin synthesis genes were up-regulated by Fe deficiency in roots, except for CsRIBC (Fig. 5). The addition of bicarbonate to the low Fe treatment did not greatly change the transcript levels, but CsRIBA1 and CsPHSI riboflavin synthesis genes were significantly induced by bicarbonate under 10 μM Fe supply. Thus, plant Fe status seems to be a more important factor than pH for induction of riboflavin synthesis genes, in contrast to the Fe uptake genes. In the leaf, none of the riboflavin synthesis genes were up-regulated by bicarbonate treatment at full Fe supply. Three of the genes were up-regulated by Fe deficiency, but this up-regulation was abolished when bicarbonate treatment was also applied (Fig. 5F–J), similar to what we observed for other leaf genes (Fig. 4).

Alkaline stress responses in fefe melon plants

Since our results with cucumber indicated that plant Fe uptake genes were up-regulated by both alkaline stress and Fe deficiency, we tested whether up-regulation of their melon counterparts requires a functional FeFe gene. The WT line Edisto was not strongly chlorotic under low Fe (0.5 μM Fe) or alkaline pH (10 mM bicarbonate) stress. The fefe mutant was more sensitive to low Fe supply, and fefe roots had a yellow coloration under low Fe with bicarbonate (Fig. 6A, right).
Alkaline stress disrupts iron deficiency signaling

Edisto FCR activity was stimulated by bicarbonate at 10 μM Fe supply. FCR activity was stimulated by Fe deficiency (0.5 μM), but this activity was repressed by adding bicarbonate to the low Fe treatment (Fig. 6B, left). FCR activity was not up-regulated in fefe melon at low Fe supply, but FCR was significantly stimulated by bicarbonate at 10 μM Fe, but to a lower extent than in Edisto (Fig. 6B, right). This indicated that alkaline stress stimulation of FCR activity is not entirely dependent on the FeFe protein.

In the WT roots, the CmFIT gene was up-regulated by Fe deficiency, but its expression was greatly reduced by bicarbonate treatment (Fig. 7A). CmbHLH38 was induced by bicarbonate in WT roots at low Fe supply (Fig. 7C), and CmbHLH101 was induced by bicarbonate at both low Fe and high Fe supply (Fig. 7E). The expression pattern of CmFRO1 corresponded to FCR activity in WT melon (Figs. 6B and 7G). The transcript patterns of both CmIRT1 and CmNRAMP1 in Edisto were consistent with expression in cucumber (Figs 2D, F and 7I, K). In fefe roots, CmFIT, CmbHLH38, CmFRO1, and CmNRAMP1 did not respond to Fe deficiency or to bicarbonate treatment. However, both CmbHLH101 and CmIRT1 had higher expression in bicarbonate-treated, Fe-sufficient roots. These results suggested that the alkaline stress response was mostly, but not entirely, dependent on fefe.

Expression of riboflavin synthesis pathway genes in fefe melon

To test whether Fe deficiency regulation of riboflavin synthesis genes requires the FeFe gene, we gathered RNAseq expression data from our previous study (Waters et al., 2014) (Table 1). In the WT, three (RIBA1, PHS1, and DMRLs) of the five genes were significantly up-regulated under Fe deficiency, but none of these genes were up-regulated by Fe deficiency in the fefe mutant. To test whether up-regulation of these genes in response to alkaline stress depends on the FeFe gene, we measured their expression in WT and fefe mutant roots by RT-qPCR. Consistent with the RNAseq results, CmRIBA1, CmPHS1, and CmDMRLs were induced by Fe deficiency in WT. Transcript levels of CmRIBA1 and CmPHS1 were lower in bicarbonate-treated Fe-deficient Edisto roots (Fig. 8A, C), similar to CmFIT expression, but CmDMRLs expression was increased (Fig. 8G). In the fefe mutant, expression of CmPYRD was somewhat higher in low Fe, both with or without bicarbonate treatment, and CmDMRLs was up-regulated under low Fe without bicarbonate (Fig. 8). However, expression of CmRIBA1 and CmPHS1 was almost completely abolished in the fefe mutant, suggesting that expression of these genes strongly depends on the FeFe regulatory pathway.
Although it is well known that IDC occurs in alkaline soils with low Fe availability, the physiological and molecular basis for this phenomenon is not well understood. In this study, we treated Fe supply and pH as separate variables. Using a range of Fe supply with two nutrient solution pHs, we found that root FCR activity responded to Fe supply only at normal pH, not at alkaline pH (Fig. 1C). We then used low and normal Fe supply, with or without bicarbonate, to determine whether gene expression in roots and leaves responded to Fe supply at each pH, and whether gene expression responded to pH at each Fe supply. Whether root FCR activity is inhibited or induced by alkaline pH depends on the point of reference. At low Fe supply, bicarbonate treatment inhibited FCR activity, relative to the treatment without bicarbonate. However, at alkaline pH the FCR activity at 10 and 2.5 μM Fe was elevated, relative to activity at normal pH. We found similar results for melon FCR activity (Fig. 6B), in that bicarbonate treatment inhibited FCR activity in Fe-deficient plants, while in Fe-supplied plants, bicarbonate stimulated FCR activity. These results are consistent with a previous study in cucumber and sunflower (Romera et al., 1992a). Together, these results explain conflicting reports in the literature, where bicarbonate treatment decreased FCR activity (Romera et al., 1997; Lucena et al., 2007; García et al., 2014) or stimulated FCR activity (Waters and Troupe, 2012), depending on Fe supply.

**Discussion**

Although it is well known that IDC occurs in alkaline soils with low Fe availability, the physiological and molecular basis for this phenomenon is not well understood. In this study, we treated Fe supply and pH as separate variables. Using a range of Fe supply with two nutrient solution pHs, we found that root FCR activity responded to Fe supply only at normal pH, not at alkaline pH (Fig. 1C). We then used low and normal Fe supply, with or without bicarbonate, to determine whether gene expression in roots and leaves responded to Fe supply at each pH, and whether gene expression responded to pH at each Fe supply. Whether root FCR activity is inhibited or induced by alkaline pH depends on the point of reference. At
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Root molecular response to alkalinity and Fe supply

Alkalinity up-regulated many of the Fe deficiency responses even when plants were grown with normal Fe supply. While the cucumber transcription factor genes did not respond to alkaline pH at 10 μM Fe, the Fe uptake genes were up-regulated (Fig. 3). In Fe-deficient Arabidopsis, AtbHLH38 and AtbHLH39 proteins interact with AtFIT, and the resulting heterodimer regulates transcription of downstream genes like AtFRO2 and AtIRT1 (Yuan et al., 2008). The difference in expression between transcription factors and their targets suggests that there could be regulators in addition to FIT/bHLH that respond to alkaline pH to up-regulate the expression Fe uptake genes in Fe replete, alkaline stressed plants. Alternatively, FIT or subgroup Ib bHLH protein activity (Meiser et al., 2011; Hindt and Guerinot, 2012), rather than transcriptional regulation, may respond to alkaline pH at normal Fe supply.

While melon and cucumber FCR activity responded to Fe supply only at normal pH (Figs 1 and 6), cucumber Fe uptake gene expression responded to Fe deficiency at both normal and alkaline pH (Fig. 3). At low Fe supply, most of the cucumber Fe uptake genes (except CsFIT and CsNRAMP1) responded to alkaline treatment by a further increase in transcript abundance (Fig. 3). This was not the case in melon, which had diminished Fe uptake gene expression in response to Fe deficiency at alkaline pH (Fig. 6), similar to previous results in Arabidopsis (García et al., 2014). In a field setting in which plants suffer from IDC, both low Fe supply and alkaline pH occur. In a prior study, low levels of bicarbonate increased expression of CsFRO1 and CsIRT1 in Fe-deficient roots (as seen here), while higher levels decreased expression of these Fe uptake genes (Lucena et al., 2007; García et al., 2014). The bicarbonate concentration we chose is in the stimulatory range for cucumber, but is in the inhibitory range for melon. In Fe-deficient plants higher concentrations of bicarbonate were required to inhibit Fe uptake gene expression in Arabidopsis than in cucumber (Lucena et al., 2007). Extending this difference in response between Arabidopsis and cucumber, and cucumber and melon to other dicot species, these results provide insight into why some species of plants are more sensitive to IDC than others.

Fig. 6. Melon plant growth and ferric chelate reductase activity with Fe deficiency and bicarbonate treatments. Wild-type (Edisto) and fefe mutant seedlings were grown in hydroponics with 0.5 or 10 μM Fe for pretreatment for 4 days, and then supplied without or with 10 mM bicarbonate for 3 days. (A) Photograph of the plants. The final nutrient solution pH of each treatment (means±SE, n=6) is indicated below each plant. (B) Ferric chelate reductase activities of roots after 4 days. * and ** indicate statistical significance at P<0.05 and P<0.01, respectively, comparing bicarbonate treatments within the same Fe supply; T and TT indicate statistical significance at P<0.05 and P<0.01, respectively, comparing Fe supply treatments within the same bicarbonate treatment.
Fe deficiency and alkaline stress regulation of riboflavin synthesis genes

Cucumber and melon roots produce and release flavin compounds in response to Fe deficiency (Welkie, 2000; Rodríguez-Celma et al., 2011a). In cucumber and WT melon roots, most of the riboflavin synthesis genes were up-regulated by Fe deficiency in both normal pH and alkaline media (Figs 5 and 8, and Table 1). Bicarbonate treatment slightly increased expression of two or three (depending on species) of the riboflavin synthesis genes, but only at full Fe supply, and there was no synergistic up-regulation at the combined low Fe and alkaline treatment as was seen for Fe uptake genes. These results indicate that although alkaline stress induces some of the same gene expression responses as Fe deficiency, alkaline stress is not precisely equivalent to Fe deficiency stress. The overlapping but not equivalent effect of Fe deficiency and alkaline stress was seen in results of a recent metabolomics study that showed that abundance of some common metabolites changed in roots treated with low Fe or with alkaline solution, while abundance of other metabolites changed only in one of these conditions (Schmidt et al., 2014). In Medicago truncatula, several genes and proteins of the riboflavin synthesis pathway were up-regulated in Fe-deficient roots (Rodríguez-Celma et al., 2011a, b, 2013b). Iron-deficient Medicago combined with alkaline stress had increased abundance for one riboflavin synthesis protein and decreased abundance for another riboflavin synthesis protein, relative to Fe-deficient plants at normal pH (Rodríguez-Celma et al., 2011a), and accumulated more flavins than Fe-deficient plants at normal pH, although release of flavins from roots was decreased at alkaline pH (Rodríguez-Celma et al., 2011b). When the Arabidopsis bHLH38 and bHLH39 genes were overexpressed in tobacco, the plants produced more riboflavin (Vorwieger et al., 2007). Our results provide a mechanism for increased riboflavin synthesis in melon and cucumber under Fe deficiency, and suggest that plants that are under alkaline stress but with normal Fe supply may also produce more riboflavin.
Alkaline stress disrupts iron deficiency signaling

Alkaline stress stimulation of Fe uptake gene expression depends on the FeFe regulatory pathway

Most of the Fe uptake-related genes were not regulated by Fe deficiency in the fefe mutant, as expected, and most of these genes also were not responsive to alkaline stress (Fig. 7B, D, H, L), suggesting that the alkaline stress signal depends mainly on the FeFe regulatory pathway. However, CmHLH101 and CmIRT1 were up-regulated in the fefe mutant by alkaline stress under full Fe supply (Fig. 7F, J), suggesting that alkaline stress can at least partially regulate these genes independently of the FeFe regulatory pathway. These genes were not induced in fefe mutants by alkaline stress under low Fe supply, which suggests that, as for other genes, the combination of Fe deficiency and alkaline stress may result in inhibition of Fe uptake gene expression.

Many of the Fe uptake genes that were up-regulated by alkaline stress in melon and cucumber have homologs in Arabidopsis that are targets of the primary Fe homeostasis transcription factor FIT (Colangelo and Guerinot, 2004). The expression of the F6'H1 gene that is required for synthesis of phenolic root exudates also depends on FIT (Schmid et al., 2014). The melon FeFe gene has not been identified, but is predicted to be a transcription factor that is functionally upstream of CmFIT (Waters et al., 2014). Since melon is a flavin producer rather than a phenolics producer like Arabidopsis, we were able to test whether this primary Fe homeostasis regulator is required for expression of riboflavin synthesis genes, and whether alkaline stress responses require the FeFe gene. The three riboflavin synthesis genes that were up-regulated by Fe deficiency in wild-type melon (CmRIBA1, CmPHS1, and CmDMRLs) were dependent on FeFe for their up-regulation by Fe deficiency (Fig. 8). Combined with results for Fe uptake genes, these results show that FeFe is a master regulator of both Fe uptake genes and Fe deficiency-induced riboflavin synthesis. These results are consistent with an early characterization of the fefe mutant, where riboflavin efflux into the nutrient solution was increased by Fe deficiency in the wild-type, but was not increased in the fefe mutant (Welkie, 1996), and with our previous RNAseq results (Waters et al., 2014). Thus, both flavin (melon) and phenolic (Arabidopsis) producing Strategy I species depend on master Fe uptake regulators for increased synthesis and efflux of...
root exudates. Up-regulation of riboflavin synthesis genes in melon roots by alkaline stress in Fe-sufficient plants also was abolished in the fefe mutant (Table 1 and Fig. 8), demonstrating that, like the Fe deficiency signal, alkaline stress regulation of riboflavin synthesis genes depends on the FeFe regulatory pathway. There is evidence for a role for both phenolics and flavins in Fe uptake under alkaline stress (Rodriguez-Celma et al., 2013b; Schmidt et al., 2014; Schmid et al., 2014), which may explain why the phenolic and riboflavin synthesis genes are regulated by Fe uptake transcription factors.

Whole plant responses to Fe supply and alkaline stress: implications for intraplant signaling and IDC

Alkaline stress may have increased expression of Fe deficiency up-regulated genes because alkaline treatment resulted in lower plant Fe accumulation. Despite increased expression of CsFRO1 and CsIRT1 (Fig. 3D, E) and FCR activity in roots of plants treated with full Fe supply and bicarbonate, whole plant Fe content was lower (Fig. 2A) and was similar to that of Fe-deficient plants, indicating that Fe uptake was inhibited by alkaline stress, consistent with previous studies (Fleming et al., 1984; Alhendawi et al., 1997). Leaf chlorophyll was also lower at all Fe levels in alkaline stressed plants (Figs. 1B and 2A) and so was Fe in stem+cotyledon and roots, while root Zn and Mn increased. These results suggest that normal Strategy I root Fe uptake processes (e.g. FCR activity, Fe(II) transporter function) do not function properly when plants are growing at alkaline pH. FCR activity of sugar beet was inhibited by alkaline pH of the assay medium (Susin et al., 1996). While our FCR assay at pH 5.5 indicated that FCR activity was increased, the alkaline pH of the growth medium may disrupt the ability of the FCR protein to reduce Fe, and may inhibit other aspects of root Fe uptake, such as specificity of Fe over Zn and Mn (George et al., 2012). These results extend the previous evidence that alkaline stress blocks Fe uptake and translocation.

The results discussed so far show that cucumber roots up-regulated certain Fe uptake and riboflavin synthesis genes in response to Fe limitation and alkaline stress in roots. In contrast, in cucumber leaf none of the Fe uptake genes or riboflavin synthesis genes were up-regulated in response to alkaline stress. The expression of CsIRT1, CsbHLH101, and CsbHLH38 was up-regulated by Fe deficiency in leaf tissue, but this up-regulation was completely inhibited by bicarbonate (Fig. 4A–C), indicating that alkaline stress blocks Fe deficiency responses in leaves. It has long been suggested that alkaline stress inhibits Fe uptake into leaf cells (Mengel, 1994). Our results suggest that this lack of uptake could be caused by a lack of Fe uptake gene expression rather than simply by physical effects, such as alkalization of xylem sap pH. This abolition of gene up-regulation by Fe deficiency under alkaline stress was also true for homologs of Arabidopsis genes that are not known to be involved in Fe uptake but were up-regulated by Fe deficiency in leaves (5960 and KCS; Fig. 4D, E). Thus, this inhibition of Fe deficiency up-regulated genes by alkaline stress occurs more generally than only for Fe uptake genes. These results show that the leaf responded to alkaline stress quite differently from the root, and this may have important implications for whole-plant Fe sensing.

Intraplant signaling of Fe status is not well understood, but both local sensing of root Fe status and shoot-to-root signaling of leaf Fe status are important for up-regulation of root Fe uptake responses (Romera et al., 1992b; Vert et al., 2003; Lucena et al., 2006; Gayomba et al., 2015) through the FIT regulatory pathway. Our results give new clues into why alkaline stress may stimulate root Fe uptake responses in Fe-sufficient plants, but inhibit these responses in Fe-deficient plants. In bicarbonate-treated cucumber plants with full Fe supply, lower root Fe concentration may stimulate a local signal to increase root FCR activity and Fe uptake gene expression. The lack of response to Fe deficiency in bicarbonate-treated cucumber leaf (Fig. 4) suggests that alkaline stress interferes with leaf Fe sensing, which would be expected to inhibit shoot-to-root signaling. Since leaf signaling may be a major factor in up-regulation of root Fe uptake responses (Enomoto and Goto, 2008; Garcia et al., 2013), a lack of signal from alkaline-stressed, Fe-deficient leaves may explain why plants grown at alkaline pH did not increase root FCR activity at low Fe supply beyond the activity at full Fe supply.

![Fig. 9](image_url). Model of responses to Fe deficiency and Fe deficiency plus alkaline stress. A local root signal for Fe deficiency acts through the FeFe regulatory pathway upstream of FIT and a bHLH partner, which activate Fe uptake genes and subsequent Fe uptake activity. An Fe deficiency signal from the leaf up-regulates the same pathway in an additive manner (left). When alkaline stress is combined with Fe deficiency (right), it blocks the normal Fe deficiency response in leaf, resulting in a loss of shoot-to-root signal. Alkaline stress combined with Fe deficiency also weakens root Fe uptake responses at the transcript and protein activity levels.
That is, the local Fe status root signal was already fully activated by decreased root Fe concentration resulting from alkaline stress, but the leaf signal to further increase root FCR activity was absent. This model (Fig. 9) would also explain why both alkaline and Fe deficiency signals rely mainly on the FeFe regulatory pathway upstream of FIT (Figs. 7 and 8).

What is not clear from our current knowledge is why leaves exposed to alkaline stress fail to respond appropriately to low Fe supply, or why Fe-deficient roots cannot maintain up-regulated FCR activity and Fe uptake gene expression (Figs. 3, 4 and 7, and Lucena et al., 2007) under alkaline stress. Because a slight induction of FCR activity (Fig. 6) and CmbHLH101 and CmlIRTI up-regulation occurred in fefe mutant plants treated with full Fe supply and bicarbonate, there may be a direct pH signal that interacts with Fe status signaling. Further research into mechanisms of leaf Fe sensing and shoot-to-root signaling, and plant sensing of rhizosphere pH will be necessary to fully understand the IDC phenomenon that occurs in low Fe availability, alkaline soils. However, since plants respond to Fe deficiency differently when also exposed to alkaline stress, this study indicates that knowledge of plant responses to low Fe supply alone, or alkaline stress alone, will not be adequate to understand IDC. Future molecular physiological IDC studies will need to include Fe supply and pH treatments singly and in combination to fully understand the IDC phenomenon, and will need to incorporate leaf and root measurements simultaneously.

Supplementary data

Supplementary data are available at JXB online.

Figure S1. DW of cucumber plants.

Figure S2. Cucumber (cv Miniature White) leaf chlorophyll and ferric chelate reductase activity in response to Fe supply in normal pH or alkaline nutrient solution.

Table S1. Primers used in this study.

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