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Mapping and Characterization of the \textit{fefe} Gene That Controls Iron Uptake in Melon (\textit{Cucumis melo} L.)

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Iron (Fe) deficiency in plants limits crop growth and productivity. Molecular mechanisms that plants use to sense and respond to Fe deficiency by coordinated expression of Fe-uptake genes are not fully understood. The C940-fe chlorotic melon (\textit{Cucumis melo}) mutant known as \textit{fefe} is unable to upregulate Fe-uptake genes, however, the \textit{FeFe} gene had not been identified. In this study, we used two F$_2$ mapping populations to map and identify the \textit{FeFe} gene as \textit{bHLH38}, a homolog of subgroup Ib \textit{bHLH} genes from \textit{Arabidopsis thaliana} that are involved in transcriptional regulation of Fe-uptake genes in partnership with the \textit{FIT} gene. A Ty1-copia type retrotransposon insertion of 5.056 kb within \textit{bHLH38} is responsible for the defect in \textit{bHLH38} in \textit{fefe}, based on sequencing and expression analysis. This retrotransposon insertion results in multiple non-functional transcripts expected to result in an altered and truncated protein sequence. Hairy root transformation of \textit{fefe} plants using wild-type \textit{bHLH38} resulted in functional complementation of the chlorotic \textit{fefe} phenotype. Using a yeast-2-hybrid assay, the transcription factor \textit{Fit} interacted with the wild-type \textit{bHLH38} protein, but did not interact with the \textit{fefe} \textit{bHLH38} protein, suggesting that heterodimer formation of \textit{Fit/bHLH38} to regulate Fe-uptake genes does not occur in \textit{fefe} roots. The second subgroup Ib \textit{bHLH} gene in the melon genome is not functionally redundant to \textit{bHLH38}, in contrast to \textit{Arabidopsis} where four subgroup Ib \textit{bHLH} genes are functionally redundant. Whereas the \textit{Arabidopsis} \textit{bHLH} transcript levels are upregulated by Fe deficiency, melon \textit{bHLH38} was not regulated at the transcript level. Thus, the \textit{fefe} mutant may provide a platform for studying \textit{bHLH38} genes and proteins from other plant species.

Keywords: iron uptake and metabolism, \textit{bHLH} transcription factor, mutant proteins, \textit{Cucumis melo}, gene expression regulation

INTRODUCTION

Iron (Fe) is crucial for plant growth, development, and productivity (Briat et al., 2015). Iron is involved in chlorophyll synthesis and is a constituent of certain enzymes involved in metabolism (Kobayashi and Nishizawa, 2012; Bashir and Nishizawa, 2013). Iron deficiency is a major limiting factor for crop production, especially in alkaline soils, which occur on approximately 30% of the earth (Chen and Barak, 1982). Plant species can be classified into two categories based on their Fe uptake mechanisms (Marschner et al., 1986). Iron uptake in graminaceous species, known as Strategy II, is characterized by production of high-affinity Fe(III) binding compounds called...
phytosiderophores, which are secreted into the rhizosphere to form phytosiderophore-Fe(III) complexes. These complexes are taken up by the root cells through a specific plasma membrane transport system (Römheld and Marschner, 1986; Curie et al., 2004). Iron uptake in dicotyledonous and non-graminaceous monocots, known as Strategy I, is characterized by soil transport system (Römheld and Marschner, 1986; Curie et al., 2004). In Arabidopsis thaliana, root cells through a specific plasma membrane form phytosiderophore-Fe(III) complexes. These complexes are phytosiderophores, which are secreted into the rhizosphere to uptake of Fe(II) by Fe transporter proteins (Kobayashi and Fe(II) by ferric chelate reductase (FCR) proteins and subsequent Reduction Oxidase 2 (supplemented with Fe, similar to and cannot survive under normal culture conditions unless it is gene expression (Waters et al., 2014; Hsieh and Waters, 2016), responses (Von Jolley et al., 1991; Welkie, 1996), and Fe-uptake resulting in the C940-fe germplasm (Nugent, 1994). The genetic basis for spontaneously in the melon cultivar Edisto (Nugent and Bhella, 1994). The AtFRO2, AtIRT1, AtNRAMP1 and various other genes are transcriptionally activated by the basic helix-loop-helix (bHLH) transcription factor AtFit1 (Colangelo and Guerinot, 2004; Yuan et al., 2005). The subgroup Ib genes of the bHLH superfamily, bHLH38, bHLH39, bHLH100 and bHLH101, are upregulated by Fe deficiency in Arabidopsis (Wang et al., 2007; Dinneny et al., 2008; Buckhout et al., 2009; Yang et al., 2010; Bauer and Blondet, 2011; Schuler et al., 2011; Stein and Waters, 2012; Waters et al., 2012; Andriankaja et al., 2014; Maurer et al., 2014). The AtFRO2 protein interacts with these subgroup Ib bHLH proteins to regulate Fe-uptake genes as a heterodimer complex (Yuan et al., 2008; Wang et al., 2013). The bHLH38, bHLH39, bHLH100 and bHLH101 genes are functionally redundant in Arabidopsis, as single, double, or triple loss-of-function bHLH mutations do not have a chlorotic Fe deficiency phenotype under Fe sufficient conditions (Wang et al., 2007; Sivitz et al., 2012; Andriankaja et al., 2014; Maurer et al., 2014), except in one report (Wang et al., 2013).

The chlorotic melon C940-Fe (fefe) mutant originated spontaneously in the melon cultivar Edisto (Nugent and Bhella, 1988). The genetic basis for fefe was retained by outcrossing the original mutant plant to the cultivar Mainstream and self-pollinating chlorotic mutants until the F3 generation, resulting in the C940-fe germplasm (Nugent, 1994). The fefe mutant plants are incapable of inducing Strategy I Fe-uptake responses (Von Jolley et al., 1991; Welkie, 1996), and Fe-uptake gene expression (Waters et al., 2014; Hsieh and Waters, 2016), and cannot survive under normal culture conditions unless it is supplemented with Fe, similar to FIT mutants in Arabidopsis. 82 genes, including Fe-uptake genes and riboflavin synthesis genes, were not regulated by Fe-deficiency in fefe plants compared to their WT counterpart (Waters et al., 2014; Hsieh and Waters, 2016), suggesting that the fefe gene could be a transcription factor. Since the fefe lesion is not in the melon FIT gene, FeFe was predicted to act upstream of FIT or as a partner of Fit (Waters et al., 2014). The main objective of this work is to map and characterize the fefe gene. We used genetic, genomic, transcriptomic and molecular approaches to map and functionally characterize the fefe gene. The results of this research will provide increased understanding of Fe-homeostasis in Strategy I plant species.

MATERIALS AND METHODS

Genetic Mapping

An F2 mapping population consisted of 269 individuals from a cross between “snake melon” (PI 435288) and C940-fe. The population was genotyped and scored, and the chlorosis trait associated with the fefe mutation was mapped to an 8 cM region of linkage group 8 (LG8) (Ramamurthy and Waters, 2015). To fine map the fefe mutation, a second, 288 individual F2 mapping population was developed from a cross between “pocket melon” (PI 536481) and C940-fe. This F2 mapping population was grown in hydroponics as described below prior to scoring leaf chlorosis in F2 plants as “0” (chlorotic) or “1” (normal). The normal and chlorotic fefe mutant F2 plants were distinguishable 1–2 weeks after planting in nutrient solution. DNA was isolated from a single young leaf from each plant (Kang et al., 1998), and quantity and purity of DNA was assessed using spectrophotometry at 260 and 280 nm. DNA was diluted to 25 ng/µl and stored at −20°C until genotyping.

A total of 112 SSR markers for LG8 from the consensus genetic map (Diaz et al., 2011) and markers for LG8 provided by Syngenta on the ICUGI website1 were tested, and in total, 27 markers were polymorphic and were used for genotyping the F2 mapping population (Table 1). PCR reactions were performed in a final volume of 10 µl with 1 Taq buffer [(16 mM (NH4)2SO4, 67 mM Tris-HCl (pH 8.8 at 25°C), 0.1% stabilizer), 2 mM MgCl2, 0.15 mM dNTP, 1 µM each primer, 0.2 U Taq DNA polymerase (Bioline USA Inc., Taunton, MA, United States), and 20 ng DNA. The cycling conditions were: an initial cycle at 94°C for 3 min, followed by 40 cycles at 94°C, 30 s, 55–58°C, 30 s and 72°C, 30 s, and a final extension step at 72°C, 5 min. PCR products were visualized with UV light after electrophoresis in 3% superfine resolution agarose (Amresco LLC, Solon, OH, United States) gels with 1 TBE (0.9 M Tris-borate, 0.002 M EDTA, pH 8.0), stained with ethidium bromide.

Phenotype scores for 288 pocket melon × C940-fe F2 individuals and their corresponding genotypes across 27 loci in population were entered into a spreadsheet as an input file for QTL analysis in R/qtl software (Broman et al., 2003). Genetic positions were deduced for markers that were not present in the consensus map (from Syngenta and Li et al., 2011). Data checking steps for genotyping were performed using standard R/qtl functions (Broman et al., 2003). To obtain a better resolution of the fefe locus, a joint linkage map consisting of 577 individuals in both snake melon X fefe and pocket melon X fefe populations was constructed using polymorphic markers (N = 35) on LG8. For joint map construction, genotypic (LG8) and phenotypic information from the two mapping populations was input for QTL analysis in R/qtl software (Broman et al., 2003). The genetic maps for individual and joint analysis were constructed using est.rf and est.map functions of R/qtl Interval

1http://www.icugi.org/cgi-bin/cmmap/map_set_info?species=acc=CM
Seeds were imbibed in germination paper soaked with 0.2 mM RNA-seq growing condition of plants used for the binary phenotype (i.e., possible allowance for covariates, with a binary model to analyze detects a single QTL by performing genome wide scan with mapping was performed using the “scanone” function which joint map (I).

**TABLE 1** | Linkage mapping of the fefe gene using two different F_2 mapping populations and a joint linkage map.

| F2-population | N | LG | Number of markers | Map length (cM) | Position of peak (cM) | Linked marker | LOD | PVE |
|---|---|---|---|---|---|---|---|---|
| Snake melon X fefe population (I) | 269 | 8 | 14 | 173 | 81 | CMN038 | 53.5 | 59 |
| Pocket melon X fefe population (II) | 288 | 8 | 27 | 96 | 79 | CMACC146 | 62.4 | 63 |
| Joint map (I+II) | 557 | 8 | 35 | 121 | 81 | CMN038 | 115.5 | 62 |

*N- Number of individuals, *PVE- Percentage of phenotypic variation explained by each locus.

**Growing Condition of Plants Used for RNA-seq**

Seeds were imbibed in germination paper soaked with 0.2 mM CaSO_4 for 4 days, then were transferred to hydroponic containers. Seedlings were placed in sponge holders in lids of black plastic containers, four plants per 750 ml solution, with continuous aeration. The nutrient solution had the following composition: 0.8 mM KNO_3, 0.4 mM Ca(NO_3)_2, 0.3 mM NH_4H_2PO_4, 0.2 mM MgSO_4, 25 uM CaCl_2, 25 uM H_3BO_3, 2 uM MnCl_2, 2 uM ZnSO_4, 0.5 uM CuSO_4, 0.5 uM Na_2MoO_4, 100 mM MES buffer (pH 5.5) and 10 uM Sprint 138 (Becker-Underwood, Ames, IA, United States). Plants were grown in a growth chamber with lighting provided by a mixture of incandescent and fluorescent sources at 250 µmol m^-2 s^-1 for a photoperiod of 16 h (on at 06:00 and off at 22:00). For the +/-Fe RNA-seq experiment, Edisto and fefe mutants were pretreated for 9 days on -Cu solution, and fefe mutants that had green leaves were used for treatments of 3 days duration in -Fe nutrient solution or 20 uM Sprint 138. The purpose for the -Cu pretreatment was to use only healthy fefe plants [since the fefe chlorotic phenotype can be rescued using -Cu treatment (Waters et al., 2014)].

**RNA-seq and Differential Expression Analysis**

Total RNA was extracted from roots using the Plant RNeasy kit (Qiagen, Hilden, Germany). RNA quality and concentration was determined by UV spectrophotometry. Sources of RNA samples were as described in the previous section. RNA-seq was performed at the University of Nebraska Medical Center Next Generation Sequencing Core Facility using an Illumina HiSeq 2000 instrument. Barcoded libraries were constructed from 3 µg of root total RNA, with three biological replicate libraries per treatment. Replicates were run in separate lanes, with a total of six samples from different treatments in each lane. The short reads are available as NCBI BioProject: PRJNA371826. Because melon and cucumber genomes are orthologous and the cucumber genome sequence and annotation is complete (Huang et al., 2009; González et al., 2010), the cucumber transcriptome was used as a reference for read mapping. Trimming of primers and adapters was performed using Trimmomatic (V0.32), read mapping was performed using BOWTIE2 (Langmead and Salzberg, 2012) with -local-N1 option, conversion of mapped reads into sam format was performed using SAMtools (Li et al., 2009) and extraction of read counts from sam files was performed using perl scripts, as previously reported (Waters et al., 2014). For gene expression analysis, the data matrix was imported into R and analyzed using the Bioconductor package DESeq (Anders and Huber, 2010). The count data was normalized for library size and then transformed using variance stabilization. Poisson distributions of normalized counts for each transcript were compared for different conditions using a negative binomial test. Differential expression was called for genes with a false discovery rate moderated q-value < 0.05 (Benjamini and Hochberg, 1995), and also showed a 1.0 log fold-change in expression and >10 reads in at least one treatment. De novo reconstruction of the bHLH38 transcript was performed using Trinity software (V.r20131110) (Haas et al., 2013). IGVviewer 2.3 was used to view the reads that are mapped onto the reference sequence as applicable.

**Reverse Transcription-PCR**

One microgram of DNase treated RNA (RNase-free DNase I, New England Biolabs, Ipswich, MA, United States) from roots of -Fe and -Fe/-Cu treated Edisto and fefe and from eight normal and eight mutant snake melon X fefe F2 roots was used for cDNA synthesis, using the High Capacity cDNA Reverse Transcription kit (ABI, Foster City, CA, United States) with random hexamers at 2.5 µM final concentration. The cDNA templates were PCR amplified using primers spanning the insertion: fefe_mrkr_F-5’-AAGCTCTAGATATATGGTTAATAA-3’ and fefe_mrkr_R-5’-TCCAGTTGCAGAATTATCGA-3’.

**Edisto bHLH38 Cloning and Hairy Root Transformation**

Edisto bHLH38 (MELO3C019065) full length genomic sequence (2.334 kb promoter +2.32 kb gene) was PCR amplified using the primers bHLH38_Promoter_F 5’-TCCCTTTGAAACCATATGG-3’ and bHLH38_XbaI_R 5’-GCATGATCTAGAACCACATTTGATATATATGTTAATAA-3’. Phusion High-fidelity DNA polymerase (Thermo Scientific) was used for PCR amplification of bHLH38 following manufacturer’s instructions. An XbaI restriction site was present in the promoter region at position 379 bp of 2.334 kb, and an XbaI site was in the reverse primer, underlined above. After XbaI restriction (NEB Biolabs), the resulting bHLH38 XbaI fragment was cloned into the pHairyRed (Lin et al., 2011) destination.

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2https://trace.ncbi.nlm.nih.gov/Traces/study/?acc=PRJNA371826&go=go

3http://www.broadinstitute.org/igv
vector. The Edisto-\textit{bHLH38} genomic construct was transformed into the K599 strain of \textit{Agrobacterium rhizogenes} (generously provided by Dr. Christopher Taylor, The Ohio State University) using freeze-thaw transformation (Wise et al., 2006). K599 containing pHairyRed-Edisto \textit{bHLH38} genomic fragment or K599 containing pHairyRed (empty vector) was grown in YEP plates with streptomycin and kanamycin selection. Agrobacteria suspension was prepared as previously described (Kereszt et al., 2007), and plug preparation and inoculation of Agrobacteria into rock wool plugs was performed as described (Chabaud et al., 2006), except we used hydroponic liquid media (described above) instead of half-strength MS. Stem sections with one or two axillary nodes from 1-month-old \textit{fefe} plants were cut and inserted into the hole in the rock wool plug, and the plants were covered in a humid chamber for 4–5 days under ambient light. The humid chamber was opened for dehydration treatment for several hours until the leaves were not turgid, and the humid chamber was closed. Hairy roots developed 2–3 weeks after transformation. Transgenic roots were distinguished from non-transgenic roots, based on the presence of DsRed fluorescence. Z-series images were acquired on a Nikon A1+ CLSM mounted on a Nikon 90i compound microscope. Excitation of DsRed was at 561 nm and emission was detected at 575–625 nm. Image series were projected to form a single image. The transmitted light images were simultaneously acquired, but only a single image plane is presented. At least two biological experiments were performed to obtain the transgenic plants.

**Ferric-Chelate Reductase Activity**

Root ferric reductase assays were performed for 50 min on transgenic roots (positive for DsRed fluorescence) and non-transgenic roots of \textit{fefe} plants that were grown on -Fe solution for 2 days, using 20 ml of an assay solution. The assay solution was composed of 0.1 mM ferrozine (3-(2-pyridyl)-5,6-diphenyl-1,2,4-triazine-4,4'-disulfonic acid sodium salt; Sigma–Aldrich), 0.1 mM Fe(III)-EDTA and 1 mM MES buffer (pH 5.5) (Fisher Scientific, Fair Lawn, NJ, United States). Change of assay solution from colorless to purple indicates ferric-chelate reductase activity.

**Yeast 2-Hybrid Interactions**

The Matchmaker\textsuperscript{TM} GAL4 Two-Hybrid System 3 (Clontech) was used for the yeast 2-hybrid experiment. Coding sequences for Edisto \textit{FIT}, Edisto \textit{bHLH38} and the \textit{fefe} \textit{bHLH38} with its 14 bp insertion were amplified from cDNA and were cloned into pGEM-T vector and were later cloned as EcoRI fragment into pGADT7 and pGBK77 yeast two-hybrid vectors. Drop assays were performed by growing AH109 transformants on synthetic
dropout (SD) liquid media without leucine (L) and tryptophan (T) at 29°C until an OD$_{600}$ of >0.5 is reached. Cultures were then diluted to an OD$_{600}$ of 0.1 and diluted in a 10× dilution series. For each dilution, 5 µl of cell suspension was spotted on SD media without leucine, tryptophan, histidine (H) and adenine (A).

RESULTS

**FeFe Gene Mapping**

The chlorotic phenotype (Figure 1) mapped to a single locus on LG8 using the snake melon X fefe F$_2$ mapping population (Ramamurthy and Waters, 2015) (Figure 2A). Using Syngenta marker positions, the fefe locus was mapped to a 6 cM (2-LOD support) (Figure 2A) region in the snake melon X fefe population. The percentage of variation explained by LG8 locus was approximately 59% (Figure 2A and Table 1). Due to the low polymorphism rate of 12% in the snake melon X fefe population, we could not further narrow down the fefe genetic interval with the markers available. Therefore, we also mapped the chlorosis phenotype in a pocket melon X fefe F$_2$ mapping population. Of 288 F$_2$ plants, 196 had normal green leaves and 92 had chlorotic leaves. 27 markers on LG8 were polymorphic in the parents. The distribution of the parental genotypes in the F$_2$ population was almost equal based on 27 polymorphic loci on LG8, with 20.7% pocket melon genotype, 30.1% C940-fe genotype and 42.2% heterozygous. The fefe locus was mapped to a 4 cM interval, with a LOD score of 62 (Figure 2B and Table 1), with 63% of the variation explained by the peak (Table 1). Since both populations in this study had a common parent, C940-fe (fefe), we used joint linkage analysis to refine the fefe genetic interval to a 1 cM peak on LG8 with a LOD score of 115 (Figure 2C and Supplementary Data S1A). The closest marker at the peak explained ~61% of the variation (Figure 2C and Table 1).

**FeFe Candidate Genes**

Since the fefe gene was predicted to be a transcription factor (Waters et al., 2014), we explored possible candidates within the 1 cM confidence interval. The genetic interval spanning the fefe gene, between the markers DM0766 and DM0640, corresponded to two scaffolds (scaffold0068 and scaffold0036) of the melon genome draft (Garcia-Mas et al., 2012). There were 186 predicted genes (Supplementary Data S1B) in this interval. There were six genes annotated as transcription factors in the mapped interval, based on homology to Arabidopsis thaliana: AT3G56970.1 BHLH038; AT5G04150.1 BHLH101; AT3G57390.2 AGL18; AT5G62470.1 MYB96; AT3G57040.1 ARR9 (RESPONSE REGULATOR 9); and AT3G10760.1 myb family transcription factor. We ruled out AGL18, MYB96 and ARR9 as unlikely candidates for fefe based on their function (To et al., 2004; Verelst et al., 2007; Seo et al., 2011). The AT3G10760.1 myb family transcription factor is putatively involved in the fruit ripening process (Pillet et al., 2015). Two transcription factors in the interval, bHLH38 and bHLH101, are associated with regulation of Fe-uptake genes (Wang et al., 2007). To determine if these genes were polymorphic between fefe and WT Edisto, or were differentially regulated, we sequenced the cDNAs and observed...
FIGURE 3 | Gene expression (normalized read counts ± SD) of bHLH38 in wild-types Edisto and snake melon and the fefe mutant under Fe deficient and Fe sufficient control conditions in two independent RNA-seq experiments. Experiment 1 is from Waters et al. (2014); experiment 2 is from this study. ∗ Indicates significant difference between Fe-deficient and control treatment means at FDR < 0.05.

FIGURE 4 | RNA-seq reads of melon bHLH38 mapped onto cucumber reference transcript Csa4M434480.1. (A) Read mapping of bHLH38 full length transcript in Edisto and fefe under Fe sufficient or deficient conditions; (B) Detail of sequence around site of the abrupt decrease in read counts in fefe bHLH38.

the normalized RNA-seq read counts from two independent experiments. Sequencing the bHLH101 gene showed that it was not polymorphic between the mutant and WT plants. The expression of bHLH101 was extremely low in both WT and mutant plants, ranging from 0 to 3 total raw read counts (compared to average read counts of approximately 450) under Fe replete or Fe-deficient conditions, suggesting that bHLH101 can be considered not expressed. The bHLH38 transcript levels were much higher than bHLH101 both in WT and fefe, ranging from 965 to 3979 read counts under Fe replete or deficient conditions, respectively. Transcript abundance did not change significantly under Fe deficiency in Edisto and snake melon.
WT roots. Although the trend of Fe regulation appeared similar in WT and fefe, the fefe $bHLH38$ expression differed in the two RNA-seq experiments (Figure 3). In the first experiment, $bHLH38$ was not significantly up-regulated by Fe deficiency, but in the second experiment, $bHLH38$ abundance was about fourfold higher under Fe deficiency due mainly to an unusually low read count in the +Fe fefe sample. We also checked abundance of $bHLH38$ transcript by RT-PCR (data not shown) and confirmed that the transcript is not regulated by Fe deficiency.

To better understand the expression of the $bHLH38$ gene in fefe and WT plants, we visualized read mapping of fefe and Edisto $bHLH38$ against the reference cucumber CsbHLH38 transcript (Csa4M434480.1). The reads mapped uniformly to the reference CsbHLH38 transcript in Edisto (Figure 4A), but the fefe read counts were much higher at the beginning of the transcript and decreased abruptly at position 370 bp of the reference transcript (Figure 4B), for both Fe deficient and Fe sufficient roots, suggesting the presence of a transcript variant. We performed de novo assembly of the Edisto and
**FIGURE 6 |** PCR amplification of genomic *bHLH38* from wild-type varieties and the *fefe* mutant. **(A)** From left to right, two individual plants of the wild-type Edisto, *fefe* mutant, wild-type Mainstream, wild-type snake melon, and two individual plants of the wild-type pocket melon. **(B)** Diagram depicting the melon *bHLH38* gene structure and insertion site of the 5.057 kb retrotransposon in *fefe* *bHLH38* (triangle). Arrows indicate the location of primers used for amplification of genomic *bHLH38* in **(A)**.

*fefe* transcriptomes to reconstruct the *bHLH38* transcripts. The assembled WT *bHLH038* transcript was 943 bp, which includes predicted start and stop codons to include 309 deduced amino acids. The *de novo* assembly of the *fefe* mutant resulted in eight unique transcripts, which were longer than the WT transcript by 277–4789 bp. The extra length occurred beginning at 549 bp in the WT *bHLH038* transcript, where we saw read mapping anomalies relative to the cucumber transcript. Both before 549 bp and after the insertion, from 550 bp onward, the *fefe* transcript sequence matched the WT transcript sequence. Using RT-PCR with primers spanning the insertion site to visualize cDNA, a single band was present in Edisto, whereas the *fefe* parent contained multiple bands, in agreement with the *de novo* assembly results, in -Fe, and -Fe/-Cu treatments (**Figure 5A**).

Similar to Fe deficiency, Cu deficiency did not change the *bHLH38* banding pattern. These results suggest that the *fefe* mutant was producing multiple insertion-containing *bHLH38* transcripts, or one large transcript that had been differently or partially spliced. A cDNA laddering pattern was also observed in the snake melon X *fefe* F$_2$ mapping population (**Figure 5B**), but only in individual plants with the chlorotic *fefe* phenotype.

The *fefe* cDNA band that was closest in size to the Edisto *bHLH38* cDNA was sequenced, to reveal the presence of a 14 bp insertion. Relative to the start codon, the *fefe* transcript had a reading frame shift followed by a premature in-frame stop codon, and would produce a different deduced amino acid sequence (Supplementary Data S2) that could negatively affect protein structure and function.
FIGURE 7 | Complementation of the chlorotic fefe phenotype by hairy root transformation. (A) Plant mock-transformed with Agrobacterium rhizogenes K599, transformed with empty pHairyRed vector, and transformed with pHairyRed-Edisto-bHLH038. (B–D) DsRed expression in hairy roots of fefe plants viewed by confocal microscopy, (B) mock transformation, (C) pHairyRed and (D) pHairyRed-Edisto-bHLH038. Bright field view of same roots in (B–D). (E) mock transformation, (F) pHairyRed and (G) pHairyRed-Edisto-bHLH038 roots; (H) Ferric chelate reductase activity of rescued fefe plant roots transformed with pHairyRed-Edisto-bHLH038 compared to non-transgenic roots.

To further investigate the source of the transcript insertions in fefe bHLH38, we amplified genomic bHLH38 in normal and fefe plants by PCR and sequenced the products. All WT lines produced an amplicon of 1.019 kb, while fefe produced a 6.076 kb fragment (Figure 6). The fefe bHLH38 contained a 5.057 kb insertion relative to WT Edisto (Supplementary Data S3). The sequence of the insertion had identical 278 bp sequences at both extremes of the insertion (Supplementary Data S3 and Figure S1). A BLAST search of the fefe genomic bHLH38 sequence against the melon reference genome had hits to six melon genomic scaffolds, unassembled sequences and scaffold 36. Length of the hits for fefe bHLH38 using BLAST search against the melon genome ranged between 272 and 1368 bp, and the total scores from BLAST search ranged between 608 and 2510. An NCBI conserved domain search identified the insertion as a long terminal repeat (LTR) Ty1-copia type retrotransposon. The fefe bHLH38 contained helix-loop-helix, polypurine tract, RNAseH1-RT-Ty1, Reverse transcriptase, Integrase, gag-polypeptide and primer binding site of LTR-copia type domains (Supplementary Figure S1). There were also large genomic bands in WTs Mainstream and pocket melon, of a similar but not identical size as the band in fefe (Figure 6). From the banding pattern and BLAST search, it appears that the retrotransposon in fefe-bHLH38 could be present in other loci in the melon genome, potentially as an intact sequence. A global BLAST search for sequences similar to fefe bHLH38 in the NCBI nucleotide database indicated that the fefe bHLH38 retrotransposon was specific to melon.

Complementation of the fefe Phenotype Using Hairy Root Transformation

Since the FeFe gene is necessary for normal plant growth and Fe uptake only in roots (Waters et al., 2014), we tested whether the chlorotic fefe phenotype could be complemented with a normal copy of bHLH38. We transformed fefe plants with Agrobacterium rhizogenes to generate hairy roots containing the WT bHLH38...
gene (Figure 7). The chlorotic phenotype of the fefe plants was rescued in the pHairyRed-Edisto-bHLH38 treated plants, but not in mock or empty vector treated plants (Figure 7A). The rescued fefe plant roots were positive for the dsRed reporter gene (Figures 7B–G), and were able to initiate ferric reductase activity under Fe deficiency (Figure 7H), suggesting that normal root Fe uptake responses were recovered in the transgenic fefe plant roots expressing Edisto-bHLH38.

**Yeast Two-hybrid Assays for Protein–Protein Interactions**

To determine whether the bHLH38 protein interacts with the Fit protein, as it does in Arabidopsis (Yuan et al., 2008), we performed a yeast two-hybrid experiment. Yeast cells transformed with different combinations of bait and prey plasmids were tested for auxotrophic growth. The melon Fit protein tested positive for interacting with Edisto bHLH38, but did not interact with fefe bHLH38 (Figure 8). The Edisto bHLH38 protein was capable of forming a homodimer, although yeast growth was less robust than in other combinations (Supplementary Figure S2A). The fefe bHLH38 did not form homodimers (Supplementary Figure S2B). Yeast growth suggested that there was some degree of interaction between Edisto bHLH38 and fefe bHLH38. This test could not be used to determine whether melon Fit forms a homodimer, since Fit was capable of auto-activation, as indicated by growth of yeast transformed with pAD+pBD-FIT (Figure 8).

**DISCUSSION**

In this paper, we seek new understanding of how Strategy I plants respond to Fe-deficiency stress by mapping the fefe gene that controls Fe uptake in melon. We mapped the fefe gene to bHLH38, which contains a 5.056 kb Ty1-copia type retrotransposon insertion. Multiple length transcripts were observed in fefe-bHLH38 (Figures 5A,B), apparently due to the full retrotransposon being incorporated into the transcript and partially spliced out to varying degrees. The altered RNA-seq read mapping we observed, with about three times higher reads before the retrotransposon insertion site in fefe-bHLH38 (Figures 3, 4), may arise from promoters within LTR regions ofLTR retrotransposons (Kumar and Bennetzen, 1999). Coincidentally, the loss of Fe homeostasis in the tomato fer mutant (Ling et al., 2002) is due to an insertion of a copia-type retrotransposon, called Rider, in the first exon of the FER gene (Cheng et al., 2009). Rider replicates by reverse transcribing an aberrant and novel cDNA that can include nearby genes, and this novel cDNA is then integrated into a new location in the genome (Lisch, 2012). We saw evidence for a pseudogene in
some varieties of melon (Mainstream and pocket melon) that was amplified by PCR primers located in the exons of the bHLH38 gene (Figure 6), suggesting that the retrotransposon in fefe exists in other loci in the melon genome. To our knowledge this retrotransposon has not been described in detail, since BLAST searches of the reference melon genome only hit unassembled scaffolds, however, it could be an important feature in melon evolution and diversity (Kumar and Bennetzen, 1999).

Regulation of Fe-uptake genes, such as FRO1, Nramp1, and IRT1, was abolished in the fefe mutant (Waters et al., 2014), similar to the Fit mutant of Arabidopsis (Colangelo and Guerinot, 2004) and the fer mutant of tomato (Ling et al., 2002), however, the lesion in the fefe mutant is not in the FIT gene (Waters et al., 2014). The formation of a heterodimer between the Fit protein and subgroup Ib bHLH proteins is a hallmark of transcriptional regulation of Fe-uptake genes (Yuan et al., 2008; Du et al., 2015). The fefe-bHLH38 transcript that was closest in size to the normal bHLH38 transcript would be translated to a protein that has an altered sequence after the first 120 aa, and terminates at 144 aa instead of 249 aa, due to the 14-bp insertion (Supplementary Data S2). Using the yeast 2-hybrid technique, the WT-bHLH38 interacted with the Fit protein, however, the fefe-bHLH38 protein did not form a heterodimer with the Fit protein (Figure 8). We propose that this lack of Fit-bHLH38 heterodimer formation is the cause of abolished upregulation of Fe-uptake gene expression in fefe. The severity of the fefe phenotype under Fe sufficient conditions from mutation of a single bHLH gene is surprising considering that in Arabidopsis, single, double, or even triple bHLH mutants (with bHLH38 or bHLH39 remaining and the other three subgroup Ib bHLH genes knocked out) had no Fe deficiency phenotype under Fe sufficient conditions (Wang et al., 2007; Sivitz et al., 2012; Andriankaja et al., 2014; Maurer et al., 2014), suggesting that one of the four bHLH proteins is adequate for Fe-uptake gene regulation. The tomato genome has three subgroup Ib bHLH genes (Sun et al., 2015), and the soybean genome has two subgroup Ib bHLH genes. A 12 bp deletion in one of the soybean bHLH genes was suggested to cause increased sensitivity to alkalinity-induced Fe deficiency chlorosis (Peiffer et al., 2012). But, like the Arabidopsis bHLH mutants, the soybean lines with this deletion are not chlorotic under Fe sufficient conditions (O’Rourke et al., 2007). A knockout line for the soybean bHLH genes has not been reported, and quadruple Arabidopsis bHLH lines have not been generated. So far, fefe is the only subgroup Ib bHLH mutant with an Fe uptake phenotype as severe as the fit or fer mutants. While the melon genome has a second subgroup Ib bHLH gene, bHLH101 was not polymorphic between WT and the fefe mutant, its transcript abundance was strikingly low, which together with genetic results suggests that melon bHLH101 is not functionally redundant with melon bHLH38.

Another key difference between melon bHLH38 and subgroup Ib bHLH genes in other plant species is their transcriptional regulation in roots by Fe status. In Arabidopsis and tomato, subgroup Ib bHLH genes are upregulated upon Fe deficiency in roots (Wang et al., 2007; Dinneny et al., 2008; Buckhout et al., 2009; Yang et al., 2016; Bauer and Blondet, 2011; Schuler et al., 2011; Stein and Waters, 2012; Waters et al., 2012; Andriankaja et al., 2014; Maurer et al., 2014; Sun et al., 2015). However, bHLH38 was not upregulated in Fe deficient melon roots (Figure 3; Waters et al., 2014; Hsieh and Waters, 2016). Copper deficiency also did not change melon bHLH38 expression, or its transcript pattern in fefe (Figure 5A), suggesting that the rescue of the chlorotic phenotype of fefe under simultaneous Fe and Cu deficiency (Waters et al., 2014) was not due to a change in bHLH38 expression, splicing, or protein function. However, since the bHLH38 protein is crucial to Fe homeostasis, its regulation may be entirely at the post-transcriptional or post-translational level. Arabidopsis FIT is regulated at both the transcriptional and post-transcriptional levels (Meiser et al., 2011; Sivitz et al., 2011).

We confirmed that the fefe defect in root Fe-uptake is due to loss of function of bHLH38 by complementation of the fefe chlorotic phenotype with WT-bHLH38 (Figure 7). The mapping and identification of the fefe mutation as bHLH38 has given new insight into regulation of Fe homeostasis in Strategy I plants. The fefe mutant may prove to be a valuable platform for studying bHLH genes and proteins from other plant species, since it can be complemented by hairy root transformation. Further characterization of bHLH38 protein regulation in melon is a needed future direction to help provide understanding of Fe-uptake control mechanisms.

**AUTHOR CONTRIBUTIONS**

RR and BW planned and designed experiments. RR conducted experiments and performed statistical and bioinformatics analysis. RR and BW wrote the manuscript. All authors read and approved the whole manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2017.01003/full#supplementary-material
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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