

1 **bHLH11 inhibits bHLH IVc proteins by recruiting the**
2 **TOPLESS/TOPLESS-RELATED corepressors in Arabidopsis**

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19 **Running title:** bHLH11 acts an active repressor

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21 **One-sentence summary:** bHLH IVc proteins promote the bHLH11 protein
22 accumulation in the nucleus where bHLH11 inhibits the transcriptional
23 activation ability of bHLH IVc via its EAR motifs recruiting the
24 TOPLESS/TOPLESS-RELATED corepressors.

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26

27 **ABSTRACT**

28

29 Iron (Fe) homeostasis is essential for plant growth and development. Many
30 transcription factors play pivotal roles in the maintenance of Fe homeostasis.
31 bHLH11 was identified as a negative transcription factor regulating Fe
32 homeostasis, however, the underlying molecular mechanism remains elusive.
33 We generated two loss-of-function *bhlh11* mutants which display the enhanced
34 sensitivity to Fe excess, the increased Fe accumulation and the elevated
35 expression of Fe deficiency responsive genes. bHLH11 protein, localized in
36 both the cytoplasm and nucleus, decreases in response to Fe deficiency.
37 Coexpression assays indicate that bHLH IVc transcription factors (TFs)
38 (bHLH34, bHLH104, bHLH105, and bHLH115) facilitate the nuclear
39 accumulation of bHLH11 protein. Further analysis indicates that bHLH11
40 represses the transactivity of bHLH IVc TFs towards bHLH Ib genes (*bHLH38*,
41 *bHLH39*, *bHLH100*, and *bHLH101*). bHLH11 contains two EAR motifs which
42 are responsible for the repression function by recruiting the
43 TOPLESS/TOPLESS-RELATED (TPL/TPRs) corepressors. Correspondingly,
44 the expression of Fe uptake genes increases in the *tpr1 tpr4 tpl* mutant.
45 Moreover, genetic analysis reveals that bHLH11 has functions independent of
46 FIT. This study provides insights into the complicate Fe homeostasis signaling
47 network.

48

49 **Introduction**

50 Iron (Fe) is an indispensable microelement for plant growth and development.
51 Plants acquire Fe from the soil, which has low concentration of Fe available,
52 especially in alkaline environments (Jeong and Guerinot, 2009). As about
53 one-third of the world's cultivated land is calcareous (alkaline), iron deficiency
54 is common for plants (Grotz and Guerinot, 2006). Fe functions in many
55 physiological processes, such as photosynthesis, respiration, hormone
56 biosynthesis, and nitrogen fixation. Fe deficiency causes symptoms including
57 delayed growth and leaf chlorosis and can affect the yield and nutritional
58 quality of crops (Kobayashi and Nishizawa, 2014). Although Fe is required for
59 plant growth and development, Fe excess can be toxic to plants because Fe
60 can cause the production of reactive oxygen radicals that are harmful to plant
61 cells (Quinet et al., 2012). Therefore, maintaining Fe homeostasis in plant cells
62 is crucial for their normal growth and development.

63 Plants have evolved a set of molecular mechanisms for iron absorption,
64 transport, distribution, and storage that ensure appropriate Fe concentrations
65 in cells under low Fe availability. Dicotyledonous and non-gramineous
66 monocotyledonous plants take up Fe using a reduction strategy (strategy I). In
67 *Arabidopsis thaliana*, this strategy involves rhizosphere acidification, ferric iron
68 reduction, and ferrous iron transport. H⁺-ATPases such as the P-type ATPase
69 AHA2/AHA7 release protons into the soil, which improves the solubility of Fe in
70 the soil (Santi and Schmidt, 2009; Kobayashi and Nishizawa, 2012). Then, the
71 root surface Fe chelate reductase FERRIC REDUCTION OXIDASE2 (FRO2)
72 catalyzes the reduction of Fe³⁺ to Fe²⁺ (Robinson et al., 1999).
73 IRON-REGULATED TRANSPORTER1 (IRT1) transports Fe²⁺ into roots
74 (Henriques et al., 2002; Varotto et al., 2002; Vert et al., 2002). By contrast,
75 gramineous plants employ a chelation strategy (strategy II) in which
76 high-affinity Fe chelators of the mugineic acid family, also known as
77 phytosiderophores, are secreted into the rhizosphere and facilitate the uptake
78 of the Fe³⁺-phytosiderophore complex. Recent studies suggest that secretion

79 of Fe-chelating compounds is also important for the survival of
80 non-gramineous plants such as *Arabidopsis* in alkaline soil (Rodríguez-Celm
81 et al., 2013; Schmidt et al., 2014; Fourcroy et al., 2014, 2016; Siwinska et al.,
82 2018; Tsai et al., 2018).

83 To maintain Fe homeostasis, plants must sense the environmental Fe
84 concentration and fine-tune the expression of Fe uptake-associated genes
85 accordingly. BRUTUS (BTS) interacts with the basic helix-loop-helix
86 transcription factors bHLH105 and bHLH115 and promotes their degradation
87 (Selote et al., 2015). IRON MAN (IMA), a class of peptides, interact with and
88 inhibit BTS, facilitating the accumulation of bHLH105 and bHLH115 (Grillet et
89 al., 2018; Li et al., 2021). bHLH105 and bHLH115 belong to the bHLH IVc
90 group, which contains four members. The other two members are bHLH34 and
91 bHLH104. These four members regulate the expression of *FER-LIKE*
92 *IRON-DEFICIENCY-INDUCED TRANSCRIPTION FACTOR (FIT)*, bHLH Ib
93 genes (*bHLH38*, *bHLH39*, *bHLH100*, and *bHLH101*) and *POPEYE (PYE)*
94 (Zhang et al., 2015; Li et al., 2016; Liang et al., 2017). bHLH121 interacts with
95 bHLH IVc proteins and is required for the maintenance of Fe homeostasis (Kim
96 et al., 2019; Gao et al., 2020; Lei et al., 2020). Downstream of bHLH IVc and
97 bHLH121, FIT interacts with bHLH Ib TFs to promote the expression of
98 Fe-uptake associated genes *IRT1* and *FRO2* (Yuan et al., 2008; Wang et al.,
99 2013). In contrast, PYE and bHLH11 are negative regulators of Fe
100 homeostasis (Long et al., 2010; Tanabe et al., 2019). Additionally, bHLH105
101 also functions as a negative regulator when it interacts with PYE (Tissot et al.,
102 2019). There is also a similar Fe deficiency response signaling network in rice
103 (Ogo et al., 2007; Kobayashi 2013, 2019; Zhang et al., 2017, 2020; Wang et al.,
104 2020; Li et al., 2020).

105 The overexpression of *bHLH11* causes the dramatic decline of Fe uptake
106 genes including *IRT1* and *FRO2*, and the severe Fe deficiency symptoms
107 (Tanabe et al., 2019). However, the molecular mechanism by which bHLH11
108 regulates Fe homeostasis remains elusive. In the present study, we

109 characterized the roles of bHLH11 in the maintenance of Fe homeostasis in
110 Arabidopsis. bHLH11 is localized in both the cytoplasm and nucleus and is
111 exclusively in the nucleus when bHLH IVc TFs are abundant. bHLH11 also
112 interacts with and inhibits bHLH IVc TFs. bHLH11 exerts its transcriptional
113 repression function by its two EAR motifs recruiting the transcriptional
114 TOPLESS/TOPLESS-RELATED corepressors.

115

116 **Results**

117 **Loss-of-function of *bHLH11* impairs Fe homeostasis**

118 It was reported that the overexpression of *bHLH11* leads to the enhanced
119 sensitivity to Fe deficiency (Tanabe et al., 2019). To further explore the
120 molecular mechanism of bHLH11 regulating Fe homeostasis, we employed the
121 CRISPR-Cas9 system to edit *bHLH11*. Two single guide RNAs were designed
122 to specifically target exons 4 and 3 of *bHLH11* and respectively integrated into
123 the binary vector with a Cas9 (Liang et al., 2016) which were then used for
124 transformation of wild type plants. We identified two homozygous mutants
125 (*bhlh11-1* and *bhlh11-2*), the former containing a 1-bp insertion in exon 4 and
126 the latter containing a 2-bp deletion in exon 3 (Figure S1A), both of which
127 caused a frameshift mutation (Figure S1B). Expression analysis indicated that
128 *bHLH11* mRNA levels did not change in these two mutants (Figure S1C).
129 When grown on Fe0 (Fe free) or Fe100 (100 μ M Fe²⁺) medium, no visible
130 difference was observed between the *bhlh11* mutants and wild type (Figure
131 1A). By contrast, when grown on Fe300 (300 μ M Fe²⁺) medium, the *bhlh11*
132 mutants produced low shoot biomass (Figure 1A, B). The *bHLH11* driven by its
133 native promoter rescued the sensitivity of *bhlh11-1* to Fe excess (Figure S1D).
134 Fe content analysis suggested that the Fe concentration of *bhlh11* mutants
135 was higher than that of the wild type (Figure 1C). These data suggest that the
136 loss-of-function of *bHLH11* leads to the enhanced sensitivity to Fe excess.

137 To further investigate the effect of bHLH11 on the Fe signaling network, we
138 examined the expression of several Fe homeostasis associated genes (Figure

139 1D). The expression of *IRT1*, *FRO2*, *bHLH38*, *bHLH39*, and *bHLH100* was
140 higher in the *bhlh11* mutants than in the wild type regardless of Fe status.
141 These data further support the negative regulation function of *bHLH11* in the
142 Fe homeostasis.

143

144 **bHLH11 expression and subcellular localization**

145 To investigate the response of *bHLH11* to Fe status, RT-qPCR was used to
146 determine the expression of *bHLH11* in response to Fe status, showing that
147 *bHLH11* mRNA increased in the roots with an increase of Fe concentration in
148 the growth medium (Figure 2A), which is in consistence with the previous
149 study (Tanabe et al., 2019). To examine the response of *bHLH11* protein to Fe
150 status, the *bHLH11* overexpression construct was introduced into wild type
151 plants. In agreement with the previous report (Tanabe et al., 2019), *bHLH11*
152 overexpression plants were more sensitive to Fe deficiency compared with
153 wild type (Figure S2). One-week-old *bHLH11* overexpression plants grown on
154 Fe100 medium were transferred to Fe0 or Fe300 medium, and root samples
155 were harvested after 1, 2, and 3 days. Immunoblot analysis showed that
156 *bHLH11* increased with an increase in Fe concentration and decreased with a
157 decrease in Fe concentration (Figure 2B).

158 Several Fe-homeostasis associated bHLH TFs were found outside the
159 nucleus (Gratz et al., 2019; Trofimov et al., 2019; Lei et al., 2020; Wang et al.,
160 2020; Liang et al., 2020). To determine the subcellular localization of *bHLH11*,
161 we generated the 35S:*bHLH11-mCherry* construct, in which the mCherry tag
162 was fused in frame with the C end of *bHLH11*. When this construct was
163 transiently expressed in tobacco leaves, mCherry was mainly observed in the
164 cytoplasm and nucleus, which is very similar to that of free mCherry (Figure
165 2C). The cytoplasmic localization of *bHLH11* was unexpected because
166 transcription factors are known to function in the nucleus. Thus, we examined
167 whether *bHLH11* can be retained in the cytoplasm due to a lack of a nuclear
168 localization signal (NLS). NLS prediction was conducted by cNLS Mapper with

169 a cutoff score = 4 (Kosugi et al., 2009;
170 http://nls-mapper.iab.keio.ac.jp/cgi-bin/NLS_Mapper_y.cgi). No NLS was
171 found in bHLH11. When bHLH11 was fused with NLS-mCherry, which contains
172 an NLS from the SV40 virus, bHLH11-NLS-mCherry was exclusively localized
173 in the nucleus (Figure 2C). These data suggest that the lack of an NLS causes
174 the cytoplasmic localization of bHLH11.

175

176 **bHLH11 interacts with bHLH IVc TFs in the nucleus**

177 Considering that TFs usually function in the nucleus and an NLS allows
178 bHLH11 to remain in the nucleus, we hypothesized that bHLH11 might be
179 recruited to the nucleus by its nuclear-localized interaction partners. Recent
180 studies revealed that bHLH121, the closest homolog of bHLH11, interacts with
181 bHLH IVc TFs (Kim et al., 2019; Gao et al., 2020; Lei et al., 2020). Therefore,
182 we employed the yeast two-hybrid system to test whether bHLH11 interacts
183 with bHLH IVc TFs. The bHLH11 protein was fused with the GAL4 DNA
184 binding domain in the pGBKT7 vector as the bait (BD-bHLH11). bHLH IVc TFs
185 were cloned to the pGADT7 vector as the preys. As expected, bHLH11
186 interacts with all four bHLH IVc TFs in yeast (Figure 3A).

187 To confirm that bHLH IVc TFs interact with bHLH11 in plant cells, we
188 employed the tripartite split-GFP system (Liu et al., 2018). The GFP10
189 fragment was fused with bHLH IVc proteins in their N-end (GFP10-bHLH IVc)
190 and the GFP11 was fused with bHLH11 in the C-end (bHLH11-GFP11). As a
191 control, bHLH121 was fused with the GFP10. When GFP10-bHLH IVc and
192 bHLH11-GFP11 were transiently co-expressed with GFP1-9 in tobacco leaves,
193 the GFP signal was visible in the nucleus of transformed cells (Figure 3B). By
194 contrast, the combination GFP10-bHLH121/bHLH11-GFP11/GFP1-9 did not
195 result in visible GFP signal. The negative controls did not result in a GFP signal
196 in the cells (Figure S3).

197 We next performed a coimmunoprecipitation (Co-IP) assay to confirm the
198 interactions between bHLH IVc TFs and bHLH11 (Figure 3C). MYC tag-fused

199 bHLH IVc TFs and HA tag-fused bHLH11 were transiently co-expressed in
200 tobacco leaves. The total proteins were incubated with MYC antibody and
201 A/G-agarose beads and then separated on SDS-PAGE for immunoblotting
202 with HA antibody. Consistent with the results from the yeast two-hybrid and
203 tripartite split-GFP assays, bHLH IVc and bHLH11 were present in the same
204 protein complex. These data suggest that bHLH IVc TFs physically interact
205 with bHLH11.

206

207 **bHLH IVc TFs affect the subcellular localization of bHLH11**

208 Having confirmed that bHLH11 interacts with bHLH IVc TFs in the nucleus, we
209 wondered whether the bHLH IVc TFs have an impact on the subcellular
210 localization of bHLH11. When any one of these four GFP tagged proteins was
211 respectively co-expressed with bHLH11-mCherry, bHLH11-mCherry
212 accumulated exclusively in the nucleus (Figure 4A). By contrast, co-expression
213 of the free GFP did not affect the subcellular localization of bHLH11-mCherry
214 (Figure 4A).

215 To further confirm the distribution of bHLH11 in the cytoplasm and nucleus,
216 we used immunoblot to measure the expression of the bHLH11 protein. As
217 shown in Figure 4B, bHLH11 protein was detected both in the nucleus and
218 cytoplasm, and both its nuclear and cytoplasmic counterparts were responsive
219 to Fe status.

220

221 **bHLH11 antagonizes the transactivity of bHLH IVc TFs**

222 The bHLH Ib genes are activated directly by the bHLH IVc TFs (Zhang et al.,
223 2015; Li et al., 2016; Liang et al., 2017). Our expression analyses also
224 suggested that the bHLH Ib genes were upregulated in the *bhlh11* mutants,
225 implying that bHLH11 is a negative regulator of bHLH Ib genes. Because
226 bHLH11 interacts with the bHLH IVc TFs, we proposed that bHLH11 could
227 antagonize the functions of the bHLH IVc TFs. To confirm this hypothesis,
228 transient expression assays were conducted in *Arabidopsis* protoplasts (Figure

229 5A). The reporter construct *ProbHLH38:LUC*, in which the LUC reporter was
230 fused with the promoter of *bHLH38*, and different effectors in which the 35S
231 promoter was used to drive GFP, bHLH11 or bHLH IVc, were used in the
232 transient assays. Compared with GFP, bHLH IVc TFs activated the expression
233 of *ProbHLH38:LUC*, whereas bHLH11 had no significant effect. When bHLH11
234 and bHLH IVc were co-expressed, the LUC/REN ratio declined significantly.
235 These data suggest that bHLH11 inhibits the transactivity of bHLH IVc TFs
236 towards *bHLH38*.

237 To further investigate whether bHLH11 inhibits the functions of bHLH IVc
238 TFs by direct protein–protein interaction, we employed the *pGAL4* promoter. In
239 the reporter construct, GFP fused with an NLS sequence was driven by *pGAL4*
240 containing the minimal CaMV 35S promoter with five repeats of the GAL4
241 binding motif (Figure 5B). In the effectors, the DNA binding domain (BD) of
242 GAL4 was fused in frame with either bHLH104 or bHLH105 and driven by the
243 35S promoter. Consistent with the fact that bHLH IVc TFs are transcriptional
244 activators, the chimeric BD-bHLH104 or bHLH105 activated the expression of
245 *GFP*. When bHLH11 was co-expressed with BD-bHLH104 or BD-bHLH105,
246 the expression of *GFP* was significantly repressed. These data suggest that
247 bHLH11 antagonizes the transcriptional activation ability of bHLH IVc TFs
248 through direct protein interaction.

249 To further confirm the antagonistic role of bHLH11 to bHLH IVc TFs, we
250 generated *bHLH11-OX/bHLH104-OX* plants by crossing *bHLH11-OX-20* with
251 *bHLH104-OX*. Compared with *bHLH104-OX*, the tolerance of
252 *bHLH11-OX/bHLH104-OX* to Fe deficiency was reduced (Figure S4). Taken
253 together, our data suggest that bHLH11 can antagonizes the functions of bHLH
254 IVc TFs.

255 It is reported that *bHLH11* overexpression causes the increased expression
256 of bHLH Ib genes (Tanabe et al., 2019), which seems to be contrary to the
257 results above that bHLH11 represses the expression of *bHLH38* (Figure 5A).
258 We reasoned that the upregulation of bHLH Ib genes was not a direct result

259 from the *bHLH11* overexpression, but caused by a secondary effect of the
260 disrupted Fe homeostasis in *bHLH11*-OX plants. To avoid the secondary effect,
261 we generated transgenic plants containing a *pER8-bHLH11* construct, in which
262 the *HA-bHLH11* fusion gene was under the control of an inducible promoter,
263 activated by estradiol. Under Fe deficient conditions, the *pER8-bHLH11* plants
264 grew as well as the wild type, and the *bHLH11* transcript level was similar
265 between the wild type and *pER8-bHLH11*. After treatment with estradiol, the
266 *bHLH11* gene was overexpressed in the *pER8-bHLH11* plants (Figure S5A).
267 As expected, the *pER8-bHLH11* plants displayed the enhanced sensitivity to
268 Fe deficiency when grown on Fe0 + estradiol medium (Figure S5B). To
269 examine the expression of Fe deficiency-responsive genes, seven-day-old
270 plants grown on Fe0 medium were transferred to Fe0 medium with or without
271 estradiol for 6 h and roots were isolated for analysis. We found that the
272 expression of bHLH Ib genes was downregulated in the *pER8-bHLH11* plants
273 after treatment with estradiol (Figure 5C). Taken together, our data suggest
274 that *bHLH11* antagonizes the transcriptional activation ability of bHLH IVc TFs
275 to bHLH Ib genes.

276

277 **The repression function of *bHLH11* requires its EAR motifs**

278 Considering that *bHLH11* negatively regulates the expression of Fe
279 deficiency-responsive genes, we wanted to know how *bHLH11* acts as a
280 transcriptional repressor. There are two typical ethylene-responsive element
281 binding factor-associated amphiphilic repression (EAR) motifs (LxLxL) in the
282 C-terminal region of *bHLH11* (Figure 6A). It is known that EAR motifs account
283 for the repression function of many transcription factors (Kagale et al., 2010).
284 To investigate whether the EAR motifs are required for the repression function
285 of *bHLH11*, we conducted reporter–effector transient expression assays in
286 which *bHLH105* was used as an effector to activate *ProbHLH38-LUC* (Figure
287 6B). We compared the effects of GFP, *bHLH11*, *bHLH11dm* (a version
288 containing two mutated EAR motifs), and *bHLH11dm-VP16* (VP16, an

289 established activation domain) on bHLH105. Compared with the significant
290 repression effect of bHLH11 on bHLH105, bHLH11dm had no significant effect,
291 whereas bHLH11dm-VP16 enhanced the transactivation function of bHLH105
292 (Figure 6C). These data suggest that bHLH11 functions as a transcriptional
293 repressor and that this function is dependent on its EAR motifs.

294 To assess the consequences of disrupting the repression functions of
295 bHLH11 *in vivo*, we generated *bHLH11dm-VP16* transgenic plants. The
296 *bHLH11dm-VP16* plants showed the enhanced tolerance to Fe deficiency
297 compared to *bHLH11-OX* plants (Figure 6D). Correspondingly, the expression
298 of *IRT1* and *FRO2* was activated in the *bHLH11dm-VP16* plants (Figure S6).
299 Taken together, the EAR motifs are needed for the repression function of
300 bHLH11.

301

302 **bHLH11 interacts with the transcription corepressors TPL/TPRs**

303 The EAR motif is a characteristic of proteins interacting with the TPL/TPRs
304 which function as transcription corepressors (Szemenyei et al., 2008; Pauwels
305 et al., 2010; Causier et al., 2012). Thus, we determined whether bHLH11
306 interacts with TPL/TPRs. Yeast two-hybrid assays indicated that bHLH11
307 interacts with TPL/TPRs (Figure 7A). To further investigate whether the EAR
308 motifs are required for the interaction, the various EAR-mutated versions,
309 bHLH11m1, bHLH11m2, and bHLH11dm, were tested. The results indicated
310 that the interaction between bHLH11 and TPL/TPRs was dependent on the
311 EAR motifs, as the mutation of both EAR motifs abolished the interaction
312 between bHLH11 and TPL/TPRs.

313 Next, we wanted to know whether TPL/TPRs participate in the regulation of
314 Fe homeostasis. To this aim, the *tpr1 tpr4 tpl* triple mutant plants were used for
315 phenotypic analysis. When grown on Fe0 or Fe100 medium, no visible
316 difference was observed. In contrast, when grown on Fe300 medium, the
317 shoot biomass of the *tpr1 tpr4 tpl* triple mutant plants was higher than that of
318 wild type (Figure 7B, C). The measurement of Fe concentration showed that

319 the *tpr1 tpr4 tpl* triple mutant plants had higher Fe concentration than wild type
320 (Figure 7D). We also examined the expression of Fe deficiency responsive
321 genes, finding that the expression of *IRT1* and *FRO2* was higher in the *tpr1*
322 *tpr4 tpl* than that in the wild type (Figure 7E). These data suggest that
323 TPL/TPRs negatively regulate the expression of Fe uptake genes.

324

325 **Genetic relationship between *bHLH11* and *FIT***

326 Tanabe et al. (2019) reported that *bHLH11* negatively regulates Fe uptake by
327 repressing *FIT* transcription. To explore whether the function of *bHLH11*
328 depends on *FIT*, we conducted genetic analysis. The *bHLH11*-OX-20 line was
329 crossed into the *fit-2* mutant, and homozygous *bHLH11*-OX-20/*fit-2* plants
330 were identified. When grown on Fe100 or Fe300 medium, no visible difference
331 was observed among the wild type, *fit-2*, *bHLH20*-OX-20 and
332 *bHLH11*-OX-20/*fit-2* plants (Figure 8A; Figure 7A). However, when grown on
333 Fe0 medium, the *bHLH11*-OX-20/*fit-2* plants were more sensitive to Fe
334 deficiency than *fit-2* and *bHLH11*-OX-20 plants, as shown by the shorter roots
335 and bleached leaves (Figure 8A; Figure S7B). These data suggest that
336 *bHLH11* has roles independent of *FIT*.

337 Subsequently, we examined the expression of *IRT1*, *FRO2* and *bHLH Ib*
338 (Figure 8B). Under Fe deficient conditions, the expression of *IRT1* and *FRO2*
339 was lower in the *fit-2* than that in the *bHLH11*-OX-20, but as low as that in the
340 *bHLH11*-OX-20/*fit-2*. In contrast, the expression of *bHLH Ib* genes was similar
341 among *fit-2*, *bHLH20*-OX-20 and *bHLH11*-OX-20/*fit-2* under Fe deficient
342 conditions. Under Fe sufficient conditions, although the expression of *bHLH Ib*
343 genes was higher in the *fit-2*, the introduction of *bHLH11*-OX-20 significantly
344 repressed their expression. These data suggest that *bHLH11* represses the
345 expression of *bHLH Ib* genes independently of *FIT* under Fe sufficient
346 conditions.

347

348 **Discussion**

349 Plants sense Fe-deficient environments and activate a signal transduction
350 cascade that ultimately results in the transcriptional regulation of downstream
351 effector genes of the Fe uptake system. The expression of Fe
352 homeostasis-associated genes is tightly regulated by Fe availability, including
353 environmental Fe availability and local Fe availability in developing tissues and
354 organs. However, this mechanism is not an on-off process but rather a
355 fine-tuned one, with multiple layers of transcription regulations. Considerable
356 progress has been made in deciphering the signal transduction pathways that
357 maintain Fe homeostasis, leading to the identification of many signaling
358 components. Here, we show that *bHLH11* acts an active repressor by
359 recruiting TPL/TPRs. *bHLH11* contributes to Fe homeostasis by repressing
360 *bHLH IVc* TFs.

361 The antagonistic regulation between positive and negative TFs is prevalent
362 in plants. For example, the positive TFs MYC2/MYC3/MYC4 and the negative
363 TFs *bHLH3/bHLH13/bHLH14/bHLH17* antagonistically modulate jasmonic
364 acid signaling (Fernandez-Calvo et al., 2011; Song et al., 2013). In Fe
365 homeostasis signaling, the *bHLH IVa* TFs (*bHLH18*, *bHLH19*, *bHLH20*, and
366 *bHLH25*) antagonize the *bHLH Ib* TFs to regulate FIT protein stability under Fe
367 deficiency (Cui et al., 2018). We show the antagonistic function of *bHLH11* to
368 *bHLH IVc* TFs, which may explain why *bHLH11*-OX plants display the severe
369 Fe deficiency phenotypes similar to those of *bHLH IVc* mutants (Liang et al.,
370 2017). In addition to transcriptional regulation, the protein degradation is
371 another type of regulation in Fe homeostasis signaling. As reported previously,
372 *bHLH105* and *bHLH115* are degraded by BTS (Selote et al., 2015), and FIT by
373 BTSL1 and BTSL2 (Rodríguez-Celma et al., 2019). We found that *bHLH11*
374 protein decreased under Fe deficient conditions (Figure 2B), which may
375 benefit plants by alleviating the repression of *bHLH11* to Fe uptake associated
376 genes. Further investigation is required to understand the post-transcription
377 regulation of *bHLH11* in response to Fe deficiency. These coordinated
378 regulations of transcription and post-transcription may help plants adapt to

379 their various Fe-nutrition habitats.

380 Although bHLH Ib genes were upregulated in both the *bHLH11-OX* and
381 *bhlh11* mutant plants, we provide evidence supporting that bHLH11 negatively
382 regulates bHLH Ib genes: (1) bHLH11 represses the promoter of *bHLH38* in
383 the transient expression assays (Figure 5A); (2) the transient induction of
384 *bHLH11* reduced the abundance of bHLH Ib genes in the *pER8-bHLH11* plants
385 (Figure 5C); (3) the overexpression of *bHLH11* reduced the levels of bHLH Ib
386 genes in the *fit-2* under Fe sufficient conditions (Figure 8B). We proposed that
387 the upregulation of bHLH Ib genes may result from a feedback regulation that
388 the severe Fe deficient status of *bHLH11-OX* plants activates bHLH Ib genes.
389 In fact, feedback regulations are universal in Fe homeostasis. For example,
390 bHLH Ib genes are also activated in the *irt1*, *frd3*, *opt3* and *fit* which are
391 defective in Fe uptake or transportation (Wang et al., 2007). Therefore, our
392 data suggest that the abundance of bHLH Ib genes are balanced by bHLH IVc
393 TFs and bHLH11. We demonstrated that bHLH11 negatively regulates the
394 expression of *IRT1* and *FRO2*, which is in consistence with the conclusion that
395 bHLH11 represses the FIT-dependent Fe uptake (Tanabe et al., 2019).
396 Meanwhile, bHLH11 also plays roles independent of FIT since its
397 overexpression made *fit-2* more sensitive to Fe deficiency (Figure 8A), which is
398 reasonable because bHLH11 can inhibit bHLH IVc TFs.

399 bHLH11 has no canonical NLS sequence. bHLH11 protein exists in the
400 nucleus and cytoplasm, and it accumulates in the nucleus with the assistance
401 of bHLH IVc TFs (Figure 4A). This interaction dependent nuclear localization of
402 bHLH11 might contribute to the maintenance of Fe homeostasis. bHLH11
403 inhibits the activation ability of bHLH IVc TFs and restricts the expression of Fe
404 uptake-associated genes. The repression function of bHLH11 may help plants
405 avoid Fe toxicity and adapt to environments with excessive Fe by reducing the
406 rate of Fe uptake. This putative protein shuttling strategy enables plants to
407 respond quickly to Fe fluctuation.

408 Two types of transcriptional repressors exist: active and passive (Krogan

409 and Long, 2009). Generally, active repressors repress transcription by
410 recruiting transcriptional repression components, whereas passive repressors
411 indirectly influence transcription by competitively interfering with activators.
412 TPL/TPRs are a class of corepressors and usually recruited by the EAR motif
413 containing transcriptional repressors (Kagale et al., 2010; Causier et al., 2012).
414 bHLH11 functions negatively and has two EAR motifs, both of which can
415 interact with TPL/TPRs, suggesting that bHLH11 is an active repressor. The
416 observation that the EAR motif is conserved in bHLH11 homologs across
417 different plant species, such as *Zea mays*, *Oryza sativa*, and *Brassica rapa*
418 (Figure S8), implies that different plant species employ a conserved repression
419 strategy to fine-tune Fe homeostasis. In Fe homeostasis signaling pathway,
420 PYE is a negative regulator of Fe homeostasis associated genes *ZIF1*, *FRO3*
421 and *NAS4* (Long et al., 2010), and an EAR motif was found in its C-terminal
422 region (Kagale et al., 2010). ZAT12 (ZINC FINGER OF ARABIDOPSIS
423 THALIANA 12) functions as a negative regulator of Fe uptake, and contains an
424 EAR motif responsible for the interaction with FIT (Le et al., 2016). By contrast,
425 EIN3 (ETHYLENE-INSENSITIVE 3), also containing an EAR motif (Kagale et
426 al., 2010), is a positive regulator of Fe homeostasis by interacting with FIT
427 (Lingam et al., 2011). It is likely that these TFs also recruit TPL/TPRs. In
428 support of this hypothesis, the *tpr1 tpr4 tpl* plants are also sensitive to Fe
429 excess to a lesser extent compared with the *bh1h11* plants, as shown by the
430 shoot biomass (Figure 1B; Figure 7C).

431 This study expands our knowledge of the Fe homeostasis transcription
432 network mediated by bHLH Ib and IVc proteins. Based on our findings, we
433 propose a putative working model for bHLH11 (Figure 9). When Fe is sufficient,
434 bHLH11 mRNAs increase and its proteins are stable. To limit Fe uptake, bHLH
435 IVc proteins facilitate the accumulation of bHLH11 in the nucleus, where
436 bHLH11 recruits TPL/TPR corepressors to repress the activation of bHLH IVc
437 proteins to bHLH Ib genes, and then the reduction of bHLH Ib genes results in
438 the down-regulation of Fe uptake genes *IRT1* and *FRO2*. When Fe is limited,

439 bHLH11 proteins decrease rapidly and few bHLH11 proteins enter the nucleus
440 to inhibit bHLH IVc. Finally, the bHLH Ib proteins accumulate and promote the
441 expression of *IRT1* and *FRO2*. This enables plants to control Fe uptake and
442 maintain Fe homeostasis. Our study provides experimental support for the
443 existence of an elaborate system that allows plants to respond dynamically to
444 Fe status. This mechanism is based on an equilibrium between the activation
445 of Fe uptake-associated genes by bHLH IVc and their repression by bHLH11.

446

447 **Materials and methods**

448 **Plant materials and growth conditions**

449 *Arabidopsis thaliana* ecotype Col-0 was used as the wild type in this study. *fit-2*
450 was described previously (Lei et al., 2020). *tpr1 tpr4 tpi* (N72353) was obtained
451 from NASC. Plants were grown in long photoperiods (16-hour light/8-hour dark)
452 or short photoperiods (10-hour light/14-hour dark) at 22°C. Surface sterilized
453 seeds were stratified at 4°C for 2 d before being planted on medium. Half
454 Murashige and Skoog (MS) medium with 1% sucrose, 0.8% agar A and the
455 indicated Fe²⁺EDTA concentration were used. Fe0 (0 μM Fe²⁺), Fe50 (50 μM
456 Fe²⁺), Fe100 (100 μM Fe²⁺) and Fe300 (300 μM Fe²⁺).

457

458 **Generation of CRISPR/Cas9-edited *bHLH11***

459 For CRISPR/Cas9-mediated editing of *bHLH11*, two target sites were
460 designed by CRISPR-GE (Xie et al., 2017) to target the third and fourth exon
461 of *bHLH11*, which were driven by the AtU3b promoter and respectively cloned
462 into the pMH-SA binary vector carrying the Cas9 (Liang et al., 2016). The wild
463 type plants were transformed and positive transgenic plants were selected on
464 half-strength MS medium containing 20 μg/mL hygromycin. The positive
465 transformants were sequenced and the homozygous mutants without the Cas9
466 were selected for further analysis.

467

468 **Generation of transgenic plants**

469 HA-tag or VP16 domain were fused in frame with the full-length coding
470 sequence of *bHLH11* to generate 35S:HA-*bHLH11* and 35S:*bHLH11dm-VP16*
471 in the pOCA30 binary vector. HA-*bHLH11* was cloned into pER8 vector (Zuo et
472 al., 2001). These constructs were introduced into *Agrobacterium tumefaciens*
473 (EHA105) respectivley and then used for transformation in the wild type
474 Arabidopsis. For complementation of *bhlh11-1*, the 3kb DNA fragment
475 upstream of *bHLH11* translation start site was used to drive HA-*bHLH11-GFP*
476 and then introduced into the *bhlh11-1* by *A. tumefaciens* mediated
477 transformation.

478

479 **Yeast-two-hybrid assays**

480 Full-length bHLH11 was cloned into pGBT7 as a bait. The full-length of bHLH
481 IVc in the pGADT7 was described previously (Li et al., 2016). Growth was
482 determined as described in the Yeast Two-Hybrid System User Manual
483 (Clontech).

484

485 **Subcellular localization**

486 For the construction of 35S:*bHLH11-mCherry*, mCherry-tag was fused with
487 bHLH11. 35S:*bHLH34-GFP*, 35S:*bHLH104-GFP*, 35S:*bHLH105-GFP*,
488 35S:*bHLH115-GFP*, and 35S:*GFP* were described previously (Lei et al., 2020).
489 35S:*bHLH11-mCherry* was co-expressed with various GFP-containing vectors
490 in tobacco cells. Epidermal cells were recorded on an OLYMPUS confocal
491 microscope. Excitation laser wave lengths of 488 nm and 563 nm were used
492 for imaging GFP and mCherry signals, respectively.

493

494 **Fluorescence complementation assays**

495 The tripartite split-GFP fluorescence complementation assay was described as
496 previously (Lei et al., 2020). The C-end of bHLH11 was fused with the GFP11
497 tag. The N-end of bHLH IVc and bHLH121 was fused with the GFP10 tag. All
498 vectors were introduced into *A. tumefaciens* (strain EHA105) and the various

499 combinations of Agrobacterial cells were infiltrated into leaves of *Nicotiana*
500 *benthamiana* by an infiltration buffer (0.2 mM acetosyringone, 10 mM MgCl₂,
501 and 10 mM MES, PH 5.6). Gene expression was induced 1 day after
502 agroinfiltration by injecting 20 µM β-estradiol in the abaxial side of the leaves.
503 Fluorescence of epidermal cells was recorded on a Carl Zeiss Microscopy.

504

505 **Co-immunoprecipitation assay**

506 HA-bHLH11 and MYC-bHLH IVc or MYC-GFP were transiently expressed in
507 the *N. benthamiana* leaves and the leaves were infiltrated with MG132 12
508 hours before harvesting. 2 g leaf samples were used for protein extraction in 2
509 ml IP buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM MgCl₂, 20%
510 glycerol, 0.2% NP-40, 1 X protease inhibitor cocktail and 1 X phosphatase
511 inhibitor cocktail from Roche). Lysates were clarified by centrifugation at 20,
512 000 g for 15 min at 4 °C and were immunoprecipitated using MYC antibody. IP
513 proteins were analyzed by immunoblot using anti-HA and anti-MYC antibody
514 respectively (Affinity Biosciences).

515

516 **Gene expression analysis**

517 Total root RNA was extracted by the use of the Trizol reagent (Invitrogen). For
518 the reverse transcription reaction, 1 µg total RNA was used for cDNA synthesis
519 by oligo(dT)18 primer according to the manufacturer's protocol (Takara). The
520 resulting cDNA was subjected to relative quantitative PCR using the ChamQ™
521 SYBR qPCR Master Mix (Vazyme Biotech Co.,Ltd) on a Roche LightCycler
522 480 real-time PCR machine, according to the manufacturer's instructions. For
523 gene expression analysis in Arabidopsis plants, the relative level of genes was
524 normalized to *ACT2* and *TUB2*. For the quantification of each gene, three
525 biological replicates were used. The primers used for quantitative reverse
526 transcription-PCR are listed in Table S1.

527

528 **Fe concentration measurement**

529 To determine Fe concentration, rosette leaves from three-week-old seedlings
530 grown in soil were harvested and dried at 65 °C for 3 days. About 100 mg dry
531 weight was wet-ashed with 5 ml of 11 M HNO₃ and 1 ml of 12 M HClO₄ for 20
532 min at 220°C. Each sample was diluted to 16 ml with 18 MΩ water and Fe
533 concentration was analyzed on a Thermo SCIENTIFIC ICP-MS(iCAP6300).

534

535 **Transient expression assays in *Arabidopsis* protoplasts**

536 *Arabidopsis* mesophyll protoplasts preparation and subsequent transfection
537 were performed as described previously (Wu et al., 2009). The promoter
538 sequence of *bHLH38* was amplified from genomic DNA and cloned into
539 pGreenII 0800-LUC vector which contains a renillia luciferase encoding gene
540 *REN* driven by the 35S promoter. The coding sequences of GFP and various
541 kinds of bHLHs (bHLH34, bHLH104, bHLH105, bHLH115, bHLH11,
542 bHLH11dm and bHLH11dm-VP16) were respectively cloned into the pGreenII
543 62-SK vector under control of 35S promoter. For the reporter and effectors, 10
544 µg plasmid for each construct was used. After protoplast preparation and
545 subsequent transfection, firefly luciferase (LUC) and REN activities were
546 measured using the Dual-Luciferase Reporter Assay System (Promega)
547 following the manufacturer's instructions. Relative (LUC) activity was
548 calculated by normalizing against the REN activity.

549

550 **Transient expression assays in tobacco**

551 *Agrobacterium tumefaciens* strains EHA105 was used in the transient
552 expression experiments in tobacco. *pGAL4* promoter and BD domain were
553 described previously (Li et al., 2016). *pGAL4* promoter was fused with
554 NLS-GFP and cloned into the pOCA28 binary vector. 35S:BD,
555 35S:BD-bHLH104, 35S:BD-bHLH105 and 35S:HA-bHLH11 were constructed
556 in the pOCA30 binary vector. For co-infiltration, different constructs were mixed
557 prior to infiltration. Leaf infiltration was conducted in 3-week-old *N.*
558 *benthamiana*. *NPTII* gene in the pOCA28 vector was used as the internal

559 control. *GFP* transcript abundance was normalized to that of *NPTII*.

560

561 **Immunoblotting**

562 For total protein extraction, roots were ground to a fine powder in liquid
563 nitrogen and then resuspended and extracted in RIPA buffer (50 mM Tris, 150
564 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS, 1 mM PMSF, 1 x
565 protease inhibitor cocktail [pH 8.0]). Isolation of cytoplasmic and nuclear
566 proteins was performed as described previously (Li et al., 2018). Sample was
567 loaded onto 12% SDS-PAGE gels and transferred to nitrocellulose membranes.
568 The membrane was blocked with TBST (10 mM Tris-Cl, 150 mM NaCl, and
569 0.05% Tween 20, pH8.0) containing 5% nonfat milk (TBSTM) at room
570 temperature for 60 min and incubated with primary antibody in TBSTM
571 (overnight at 4°C). Membranes were washed with TBST (three times for 5 min
572 each) and then incubated with the appropriate horseradish
573 peroxidase-conjugated secondary antibodies in TBSTM at room temperature
574 for 1.5 h. After washing three times, bound antibodies were visualized with
575 ECL substrate.

576

577 **Supplemental data**

578 **Supplemental Figure 1.** Identification of *bhlh11* mutants.

579 **Supplemental Figure 2.** Identification of *bHLH11* overexpression plants.

580 **Supplemental Figure 3.** Negative controls of tripartite split-sfGFP
581 complementation assays.

582 **Supplemental Figure 4.** Antagonism between *bHLH104* and *bHLH11*.

583 **Supplemental Figure 5.** Phenotypes of *pER8-bHLH11* transgenic plants.

584 **Supplemental Figure 6.** Expression of *IRT1* and *FRO2* in the
585 *bHLH11dm-VP16* plants.

586 **Supplemental Figure 7.** Analysis of *bHLH11-OX-20/fit-2* plants.

587 **Supplemental Figure 8.** Conserved EAR motifs in the *bHLH11* homologs from
588 various plants species.

589 **Supplemental Table 1.** Primers used in this paper.

590

591 **Acknowledgements**

592 We thank the Biogeochemical Laboratory and Central Laboratory
593 (Xishuangbanna Tropical Botanical Garden) for assistance in the
594 determination of metal contents. We also thank Germplasm Bank of Wild
595 Species in Southwest China for confocal laser scanning microscopy.

596

597 **Finding**

598 This work was supported by the National Natural Science Foundation of China
599 (31770270) and the Applied Basic Research Project of Yunnan Province
600 (2019FB028 and 202001AT070131).

601

602 **Author contributions**

603 G.L. conceived the project. Y.L., R.L., M.P., C.L., Z. L., and G.L. constructed
604 plasmids, M.P. and Y.C generated transgenic plants, and Y.L. and R.L.
605 characterized plants, determined gene and protein expression, and conducted
606 cellular assays. Y.L. and G.L. wrote the manuscript and all authors discussed
607 and approved the manuscript.

608

609 *Conflict of interest statement. None declared.*

610

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802

Figure Legends

Figure 1. *bhlh11* mutants are sensitive to Fe excess.

(A) Phenotypes of *bhlh11* mutants. Two-week-old seedlings grown on Fe0, Fe100 or Fe300 medium.

(B) Shoot biomass of *bhlh11* mutants. Fresh weight of two-week-old shoots grown on Fe0, Fe100 or Fe300 medium. Three biological duplicates, each of which contains 15 plants, were analyzed.

(C) Fe concentration of rosette leaves of three-week-old wild type and *bhlh11* plants grown in soil.

(D) Expression of *IRT1*, *FRO2* and bHLH Ib genes. Four-day-old plants grown on Fe100 medium were transferred to Fe0, Fe100 or Fe300 medium for three days, and root samples were harvested and used for RNA extraction and RT-qPCR.

(B-D) The different letters above each bar indicate statistically significant differences as determined by one-way ANOVA followed by Tukey's multiple comparison test ($P < 0.05$).

Figure 2. Response of bHLH11 to Fe status

(A) RT-qPCR analysis of *bHLH11* expression. Four-day-old plants grown on Fe100 medium were shifted to Fe0, Fe50, Fe100 or F300 medium for 3 days. Roots were used for RNA extraction and RT-qPCR. The different letters above each bar indicate statistically significant differences as determined by one-way ANOVA followed by Tukey's multiple comparison test ($P < 0.05$).

(B) Degradation of bHLH11 in response to Fe deficiency. Seven-day-old wild type and *bHLH11*-OX-20 seedlings grown on Fe100 medium were transferred to Fe0 or Fe300 medium, and root samples were harvested after 1, 2, and 3 days. anti-HA was used to detect HA-bHLH11. β -tubulin was used as a loading control.

(C) Subcellular localization of bHLH11. The free mCherry, bHLH11-mCherry or bHLH11-NLS-mCherry were respectively expressed in *N. benthamiana* leaves.

Figure 3. bHLH11 physically interacts with bHLH IVc TFs.

(A) Yeast two-hybrid analysis of the interaction between bHLH11 and bHLH IVc TFs. Yeast cotransformed with different BD and AD plasmid combinations was spotted on synthetic dropout medium lacking Leu/Trp (SD-W/L) or Trp/Leu/His/Ade (SD -W/L/H/A).

(B) Interaction of bHLH11 and bHLH IVc TFs in plant cells. Tripartite split-sfGFP complementation assays were performed. bHLH34, bHLH104, bHLH105, bHLH115, and bHLH121 were fused with GFP10, and bHLH11 was fused with GFP11. The constructs were introduced into agrobacterium respectively, and the indicated combinations were co-expressed in *N. benthamiana* leaves.

(C) Co-IP analysis of the interaction between bHLH11 and bHLH IVc TFs. Total proteins from different combinations of HA-bHLH11 and MYC-GFP,

MYC-bHLH34, MYC-bHLH104, MYC-bHLH105, or MYC-bHLH115 were immunoprecipitated with anti-MYC followed by immunoblotting with the indicated antibodies. MYC-GFP was used as a negative control. Protein molecular weight (in kD) is indicated to the left of the immunoblot.

Figure 4 Change of bHLH11 subcellular localization.

(A) Location of bHLH11 in the absence or presence of bHLH IVc. bHLH11-mCherry was co-expressed with bHLH IVc TFs. The combination of bHLH11-GFP and free mCherry was used as a negative control. Transient expression assays were performed in tobacco leaves.

(B) Immunoblot analysis of bHLH11 protein distribution in the cytoplasm and nuclear fractions. Seven-day-old *bHLH11-OX-20* seedlings grown on Fe100 medium were transferred to Fe0, Fe100 or Fe300 medium. Root samples were harvested after 3 days, and cytoplasmic and nuclear proteins were extracted and subjected to immunoblot analysis with the indicated antibodies. Ratio indicates the relative protein abundance of HA-bHLH11.

Figure 5. bHLH11 antagonizes the transcriptional activation ability of bHLH IVc TFs.

(A) bHLH11 represses the functions of bHLH IVc TFs. Schematic diagram of the constructs transiently expressed in *Arabidopsis* protoplasts. The LUC/REN ratio represents the LUC activity relative to the internal control REN. The asterisk indicates a significant difference as determined by Student's t Test.

(B) bHLH11 inhibits the functions of bHLH IVc TFs by direct protein-protein interaction. The schematic diagram shows the constructs used in the transient expression assays in tobacco leaves. The abundance of *GFP* was normalized to that of *NPTII*. The asterisk indicates a significant difference as determined by Student's t Test.

(C) Expression of bHLH Ib genes in *pER8-bHLH11* plants. Seven-day-old plants grown on Fe0 medium were transferred to Fe0 medium with or without 4 μ M estradiol for 6 h, and root samples were harvested and used for RNA extraction and RT-qPCR. The different letters above each bar indicate statistically significant differences as determined by one-way ANOVA followed by Tukey's multiple comparison test ($P < 0.05$).

Figure 6. bHLH11 acts as a repressor dependently on its two EAR motifs.

(A) Schematic diagram of the various mutated versions of bHLH11. The mutated amino acid is indicated in red. bHLH11m1, the first EAR mutated. bHLH11m2, the second EAR mutated. bHLH11dm, both double EARs mutated. bHLH11dm-VP16, bHLH11dm fused with the VP16 domain.

(B) The schematic diagram shows the constructs used in the transient expression assays in (C).

(C) The EAR motifs are required for the repression of bHLH11. *Arabidopsis* protoplasts were used for transient expression assays. The abundance of *GFP*

was normalized to that of *NPTII*. The different letters above each bar indicate statistically significant differences as determined by one-way ANOVA followed by Tukey's multiple comparison test ($P < 0.05$).

(D) Phenotypes of *bHLH11dm-VP16* and *bHLH11-OX* plants. Seven-day-old seedlings grown on Fe0 or Fe100 medium are shown.

Figure 7. Enhanced Fe deficiency response in *tpr1 tpr4 tpl*.

(A) The EAR motifs are required for the interaction between bHLH11 and TPL/TPRs. Yeast cotransformed with different BD and AD plasmid combinations was spotted on synthetic dropout medium lacking Leu/Trp (SD –T/L) or Trp/Leu/His/Ade (SD –T/L/H/A).

(B) Phenotypes of *tpr1 tpr4 tpl*. Two-week-old seedlings grown on Fe0, Fe100 or Fe300 medium are shown.

(C) Shoot biomass of *tpr1 tpr4 tpl*. Fresh weight of two-week-old shoots grown on Fe0, Fe100 or Fe300 medium. Three biological duplicates, each of which contains 15 plants, were analyzed.

(D) Fe concentration of rosette leaves of 3-week-old wild type and *tpr1 tpr4 tpl* plants grown in soil.

(E) Expression of *IRT1*, *FRO2* and bHLH Ib genes. Four-day-old plants grown on Fe100 medium were transferred to Fe0, Fe100 or Fe300 medium for three days, and root samples were harvested and used for RNA extraction and RT-qPCR.

(C-D) The asterisks indicate that the values are significantly different from the corresponding wild type value by Student's t Test ($P < 0.05$).

Figure 8. Genetic interaction between *bHLH11* and *FIT*.

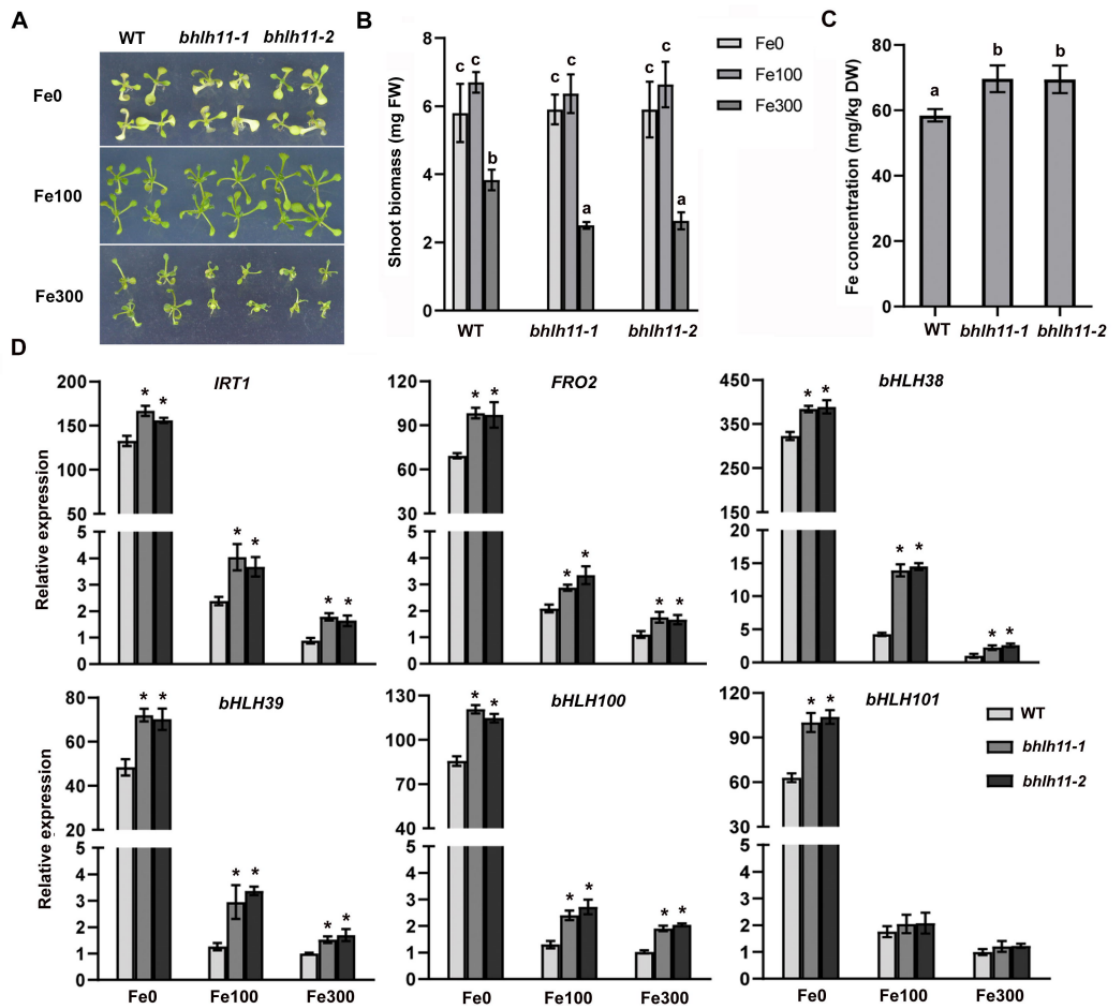
(A) Phenotypes of *bHLH11-OX-20/fit-2* plants. Seven-day-old seedlings grown on Fe0 or Fe100 medium are shown.

(B) Expression of *IRT1*, *FRO2* and bHLH Ib genes. Four-day-old plants grown on Fe100 medium were transferred to Fe0 or Fe100 medium for three days, and root samples were harvested and used for RT-qPCR. The different letters above each bar indicate statistically significant differences as determined by one-way ANOVA followed by Tukey's multiple comparison test ($P < 0.05$).

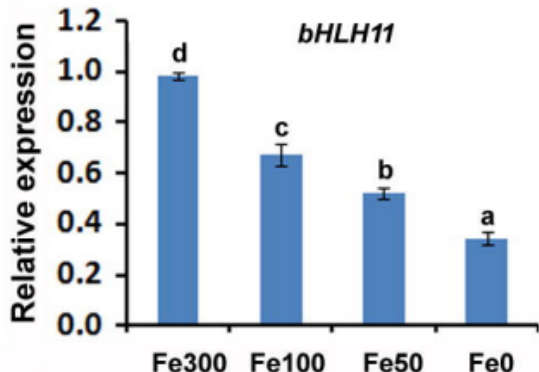
Figure 9. A working model of bHLH11 in Fe homeostasis.

bHLH11 functions as an active repressor by recruiting TPL/TPR corepressors, and bHLH IVc TFs promote the nuclear accumulation of bHLH11. Under Fe-sufficient conditions, *bHLH11* message is activated and its protein accumulates. bHLH11 inhibits the transactivity of bHLH IVc TFs to bHLH Ib genes. The repression function of bHLH11 allows plants to avoid Fe toxicity. Under Fe-deficient conditions, unknown proteins repress the transcription of *bHLH11* and its protein decreases, which alleviates the bHLH11-mediated repression to bHLH IVc TFs. bHLH IVc TFs promote the transcription of bHLH Ib genes, and then bHLH Ib proteins interact with FIT to activate the expression of the Fe uptake genes *IRT1* and *FRO2*. The question signs

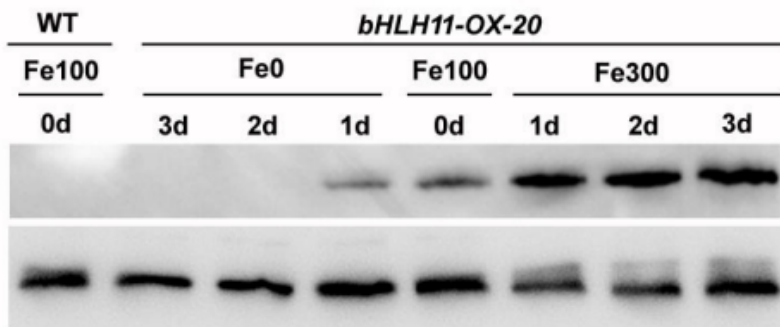
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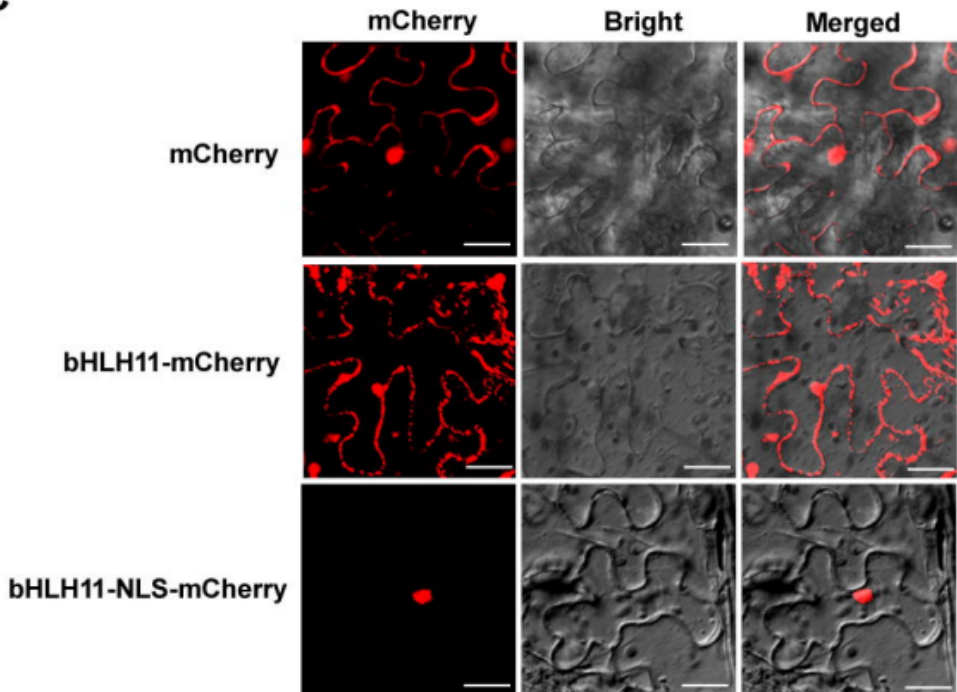
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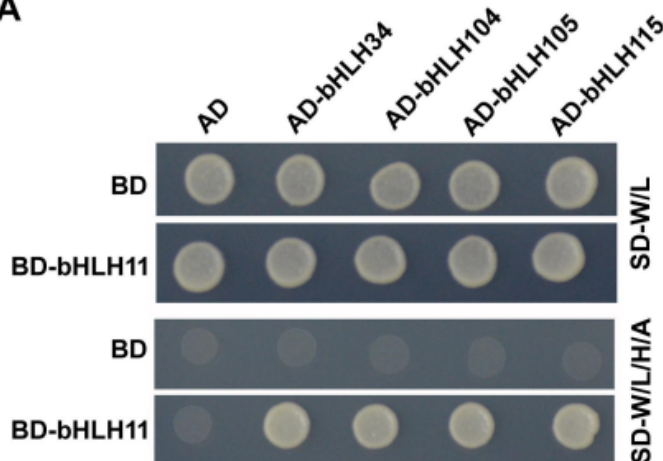
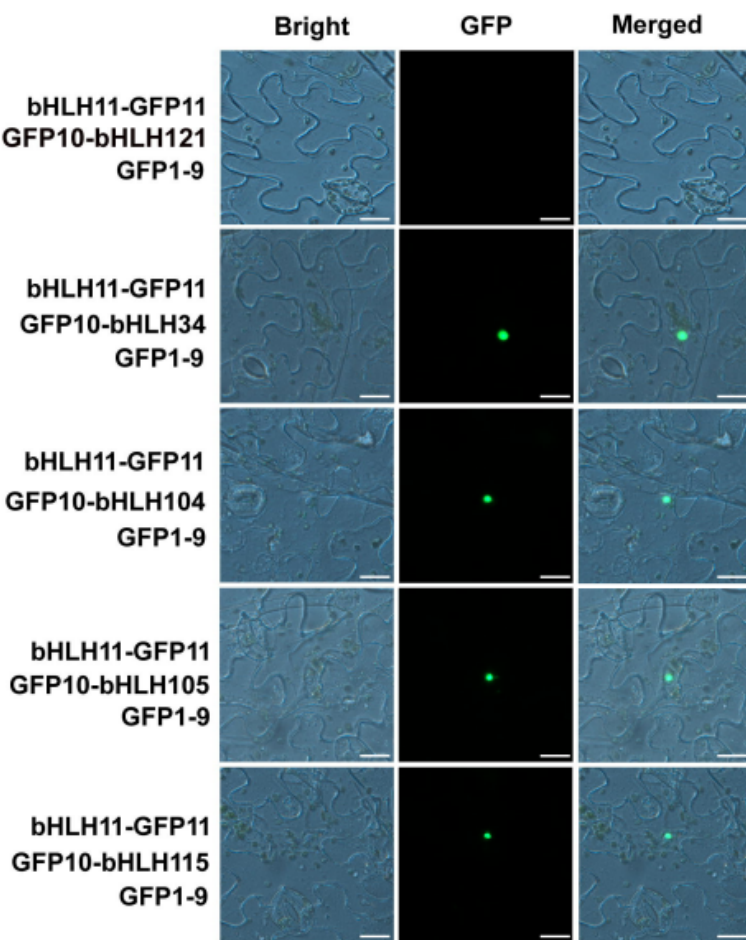
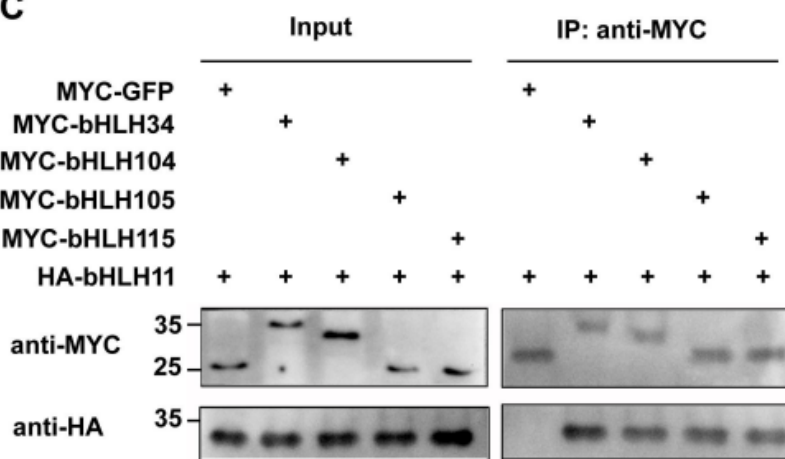


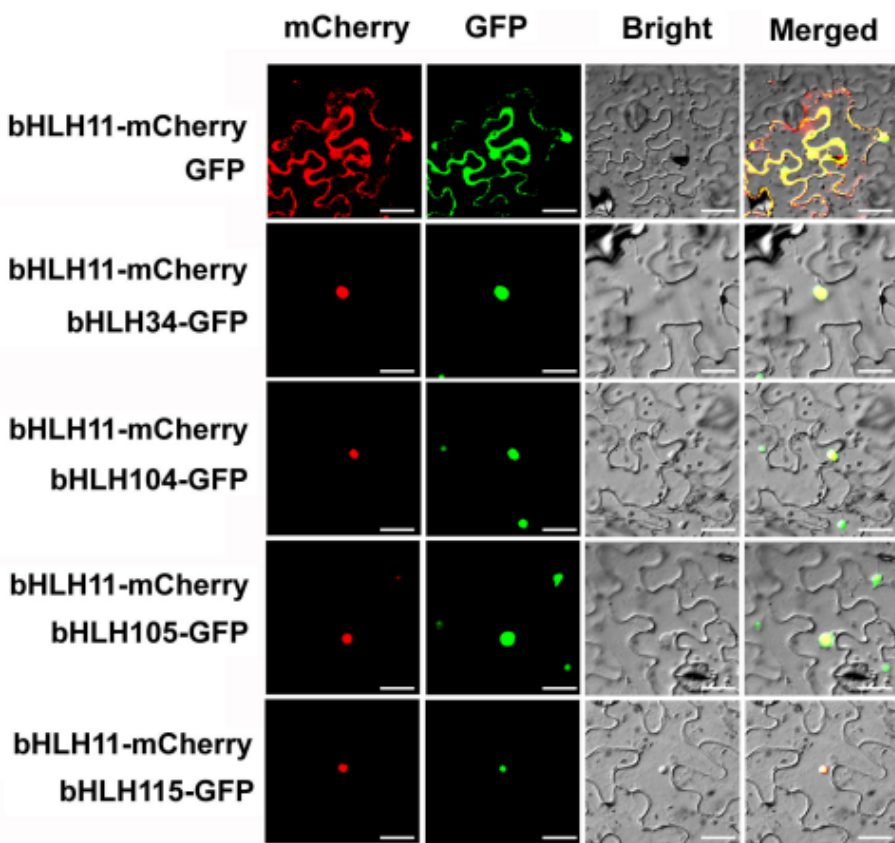
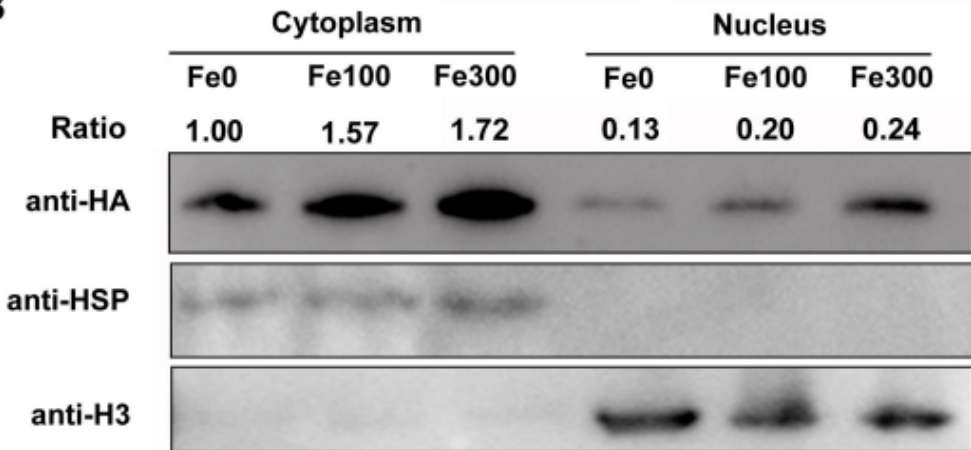
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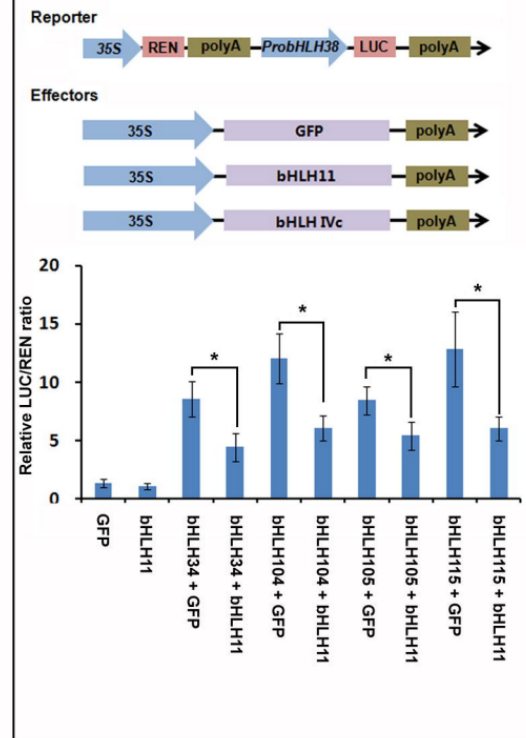
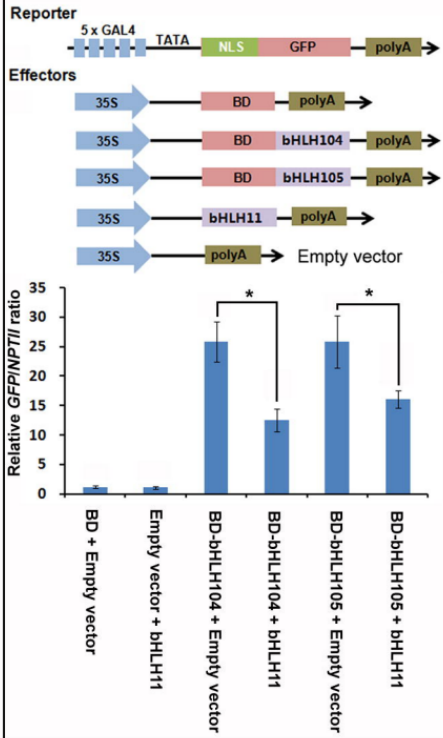
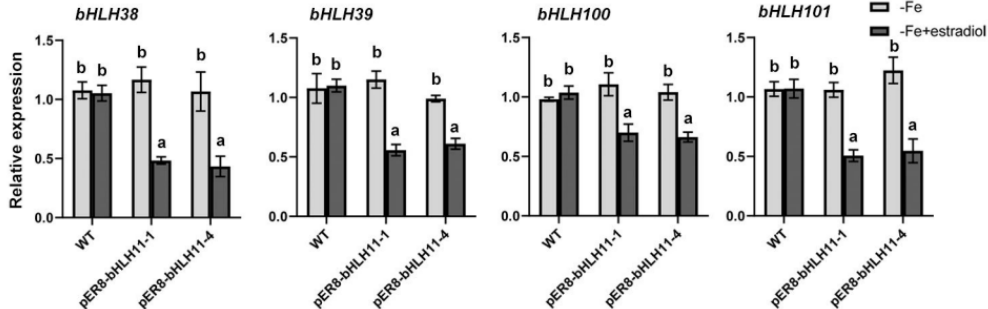


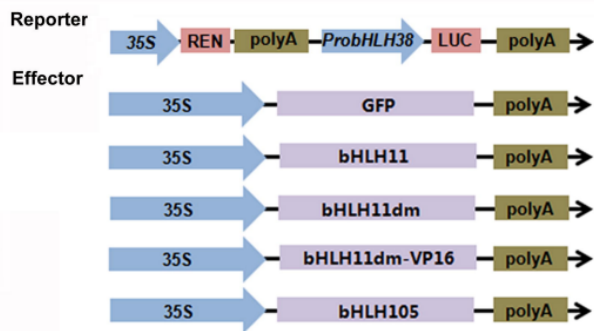
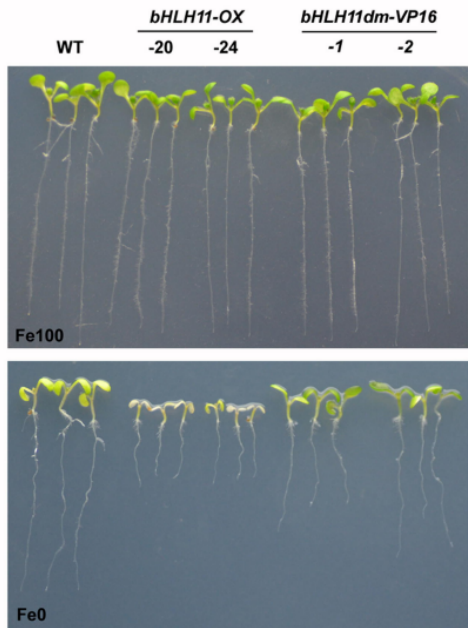
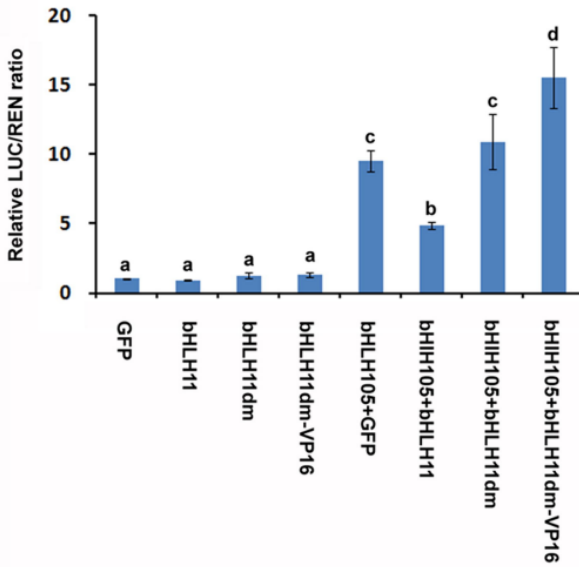
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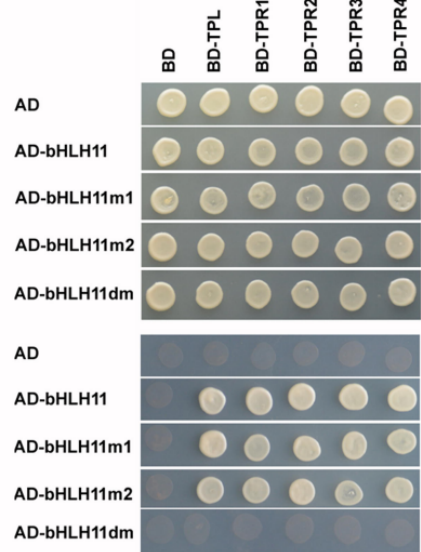
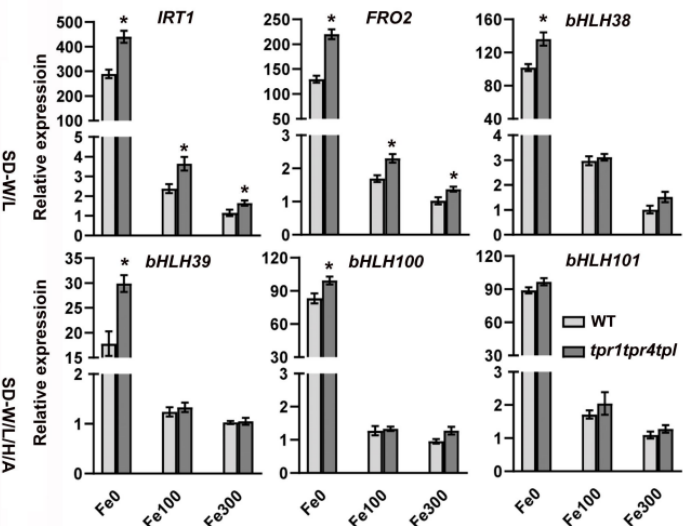
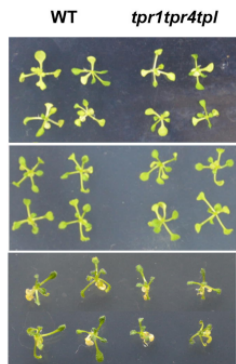
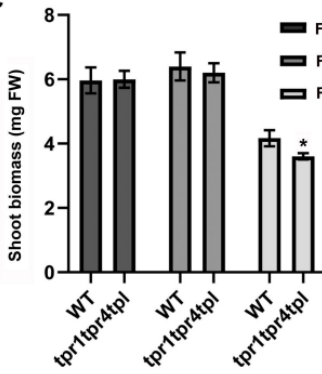
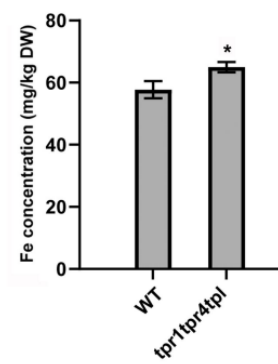


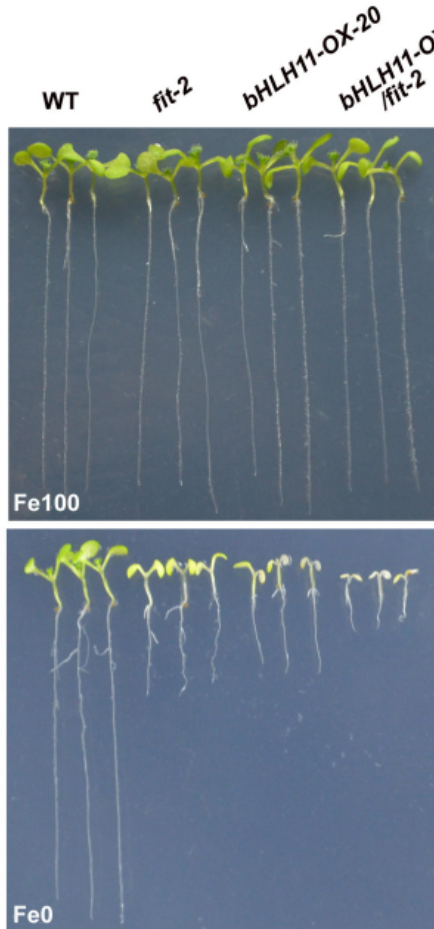
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