Modulation of iron responsive gene expression and enzymatic activities in response to changes of the iron nutritional status in *Cucumis sativus* L.

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**Key words:** *Cucumis sativus*, Fe(III)-chelate reductase, H⁺-ATPase, iron deficiency response, phosphoenolpyruvate carboxylase.

**Abstract**

Regulation exerted by the iron status of the plant on the iron deficiency responses was investigated in cucumber roots (*Cucumis sativus* L.) both at the biochemical and molecular level. Absence of iron induced the expression of the *CsFRO1*, *CsIRT1*, *CsHA1* and the *Cspepc1* transcripts that was followed by an increase in the corresponding enzymatic activities. Supply of iron repressed gene expression, in particular those of the Fe(III)-chelate reductase and for the high affinity iron transporter and reduce the enzymatic activities. Our results confirm and extend the hypothesis of a coordinate regulation of these responses. Besides these two activities strictly correlated with iron deficiency adaptation, we considered also the H⁺-ATPase and the phosphoenolpyruvate carboxylase, that have been shown to be involved in this response.
**Abbreviations:** BPDS, bathophenanthrolinedisulphonate; BTP, 1,3-bis[tris(hydroxymethyl)-methylamino]-propane; MES, 2-[N-morpholino]ethanesulphonic acid; MOPS, 3-(N-morpholino)propanesulfonic acid; PMSF, phenylmethylsulphonyl fluoride; PPFD, photosynthetic photon density flux

**Introduction**

Plants require iron to complete their life cycle. The importance of iron is due to the existence of two stable, but convertible forms, Fe(III), ferric, and Fe(II), ferrous, which take part in fundamental processes involving electron transfer reactions, mainly requested in both the oxidative (respiration) and biosynthetic (photosynthesis) pathways (Curie & Briat, 2003; Hell & Stephan, 2003). On the contrary, excess of iron can produce toxic oxygen compounds, for instance O$_2^-$, H$_2$O$_2$ and above all free hydroxyl radicals, produced by the Fenton reaction (Halliwell & Gutteridge, 1999; Briat, 2002). Consequently, balanced iron acquisition by the roots and control of the homeostatic mechanisms are necessary to prevent suffering or excess of this transition metal.

There is generally a high quantity of iron in the soil, but, in aerobic and sub-alkaline pH environment it is present mainly as Fe(III)-oxide and -hydroxide and its solubility is strongly restricted (Guerinot & Yi, 1994). To cope with this problem and to enhance the metal bioavailability, plants have evolved developmental and biochemical adaptation (Strategy I and Strategy II) to low iron concentration in the environment (Römheld & Marschner, 1986). Concerning Strategy I plants, evident responses reside at the root-soil interface involving morphological changes in the root architecture by increasing the number of secondary roots, root hair density and the formation of transfer cells at the root apex in order to enhance the absorbing root surface. In the meantime, primary
biochemical and molecular responses serve one main function: increase the rhizosphere iron availability and its uptake (Schmidt, 1999; Curie & Briat, 2003; Hell & Stephan, 2003).

Strategy I plants (dicotyledonous and non-graminaceous plants) are able to respond to lack of iron mainly by increasing the reduction, the acidification and the uptake activities by inducing trans-plasma membrane proteins present in the rhizodermal root cells {i.e. Fe(III)-chelate reductase [FC-R], H^+-ATPase and iron regulated transporters (IRT), respectively} directly involved in the iron uptake system (Curie & Briat, 2003). Extrusion of electrons and protons leads to an enhancement of soluble form of iron in the rhizosphere. It has been observed in Strategy I plants, that in response to iron starvation, there was an induction of the genes encoding for the Fe(III)-chelate reductase (AtFRO2, PsFRO1 and LeFRO1) which were already characterized (Robinson et al., 1999; Waters et al., 2002; Li et al., 2004). Enhancement of H^+ efflux, due to an increase in a P-type H^+-ATPase activity in response to iron deprivation, was demonstrated in many Strategy I plants (Schmidt, 1999; Zocchi, 2006). A multigene family encoding different isoforms for H^+-ATPase and tissue specific expression patterns have been demonstrated (Palmgren, 2001; Dell’Orto et al., 2002; Santi et al., 2005). After mobilization and reduction, the ferrous form, the unique form to be absorbed by these plants, needs to be taken up across the plasma membrane by a specific iron transporter (IRT1) that has been characterized in A. thaliana (Eide et al., 1996) and successively in pea and tomato (Cohen et al., 1998; Eckhardt et al., 2001).

Micro-array analysis using A. thaliana grown under iron deficiency (Thimm et al., 2001) has revealed that there were significant changes in the transcription of different genes, therefore reflecting the complexity of the molecular and metabolic response.
Many genes involved in Fe-starvation responses have been cloned and Fe responsiveness has demonstrated the importance of transcriptional control in the regulation of Strategy I mechanisms (Curie & Briat, 2003). To promote and sustain the increase in the release of electrons and protons in the rhizosphere it has been shown that significant metabolic changes occurred in roots: enhancement of glycolitic pathway rate, cytosolic dehydrogenase activities, as well as respiration rate (Rabotti et al., 1995; Espen et al., 2000). Moreover, organic acid synthesis and CO₂ dark fixation increase under Fe-starvation during which phosphoenolpyruvate carboxylase (PEPC) activity was shown to increase by four times or more (Rabotti et al., 1995; De Nisi & Zocchi, 2000; Lopez-Millan et al., 2000). The anaplerotic role of root PEPC has been characterized in root of cucumber grown under Fe-deficiency (De Nisi & Zocchi, 2000) and in other Strategy I species, both herbaceous and arboreous (Lopez-Millan et al., 2000; Ollat et al., 2003), which showed also an enhanced H⁺-ATPase activity. So it is possible to consider the inducible enhancement of PEPC activity as one of the more important markers regarding metabolic responses to Fe deficiency.

In this work we were interested in identifying the coordination of biochemical and molecular responses in Strategy I plants, which are, more likely, the result of tightly controlled homeostatic mechanisms.

**Materials and Methods**

Plant material and growth conditions

Seeds of cucumber (*Cucumis sativus* L. cv Marketer) were surface sterilized and sown in Agriperlite, watered with 0.1 mM CaSO₄, allowed to germinate in the dark at 26°C
for 3 d, and then transferred to a nutrient solution of the following composition: 2 mM Ca(NO)₃, 0.75 mM K₂SO₄, 0.65 mM MgSO₄, 0.5 mM KH₂PO₄, 10 µM H₃BO₃, 1 µM MnSO₄, 0.5 µM CuSO₄, 0.5 µM ZnSO₄, 0.05 µM (NH₄)Mo₇O₂₄ and 100 µM Fe(III)-EDTA (when added). The pH was adjusted to 6.0-6.2 with NaOH. Aerated hydroponic cultures were maintained in a growth chamber with a day/night regime of 16/8 h and a PPFD of 200 µmol m⁻² s⁻¹ at the plant level. The temperature was 18°C in the dark and 24°C in the light. Plants showed chlorotic symptoms after approximately seven days of culture in the absence of Fe.

In vivo measurement and localisation of the acidification and reduction capacities

Medium acidification capacity was measured directly in the nutrient solution by measuring the pH every day with a pHM64 (Radiometer, Copenhagen) pHmeter. Fe(III)-reductase activity was measured by using the bathophenantrolinedisulfonate (BPDS) reagent (Chaney et al., 1972). Ten apical root segments about 2 cm long were incubated in 10 ml of a solution containing 0.5 mM CaSO₄ and 0.5 mM K₂SO₄ pH 6.0, in the dark at 26°C under shaking. After 1 h incubation the solution was replaced with 5 ml of a solution with the following composition: 0.5 mM CaSO₄, 0.5 mM K₂SO₄, 0.1 mM Fe(III)-EDTA and 0.25 mM BPDS pH 6.0. After 3 h, 2 mL of the solution were withdrawn and the absorbance at 535 nm determined with a spectrophotometer. BPDS forms a stable, water soluble, red complex with Fe²⁺ and only a weak complex with Fe³⁺. The amount of reduced Fe was calculated by the concentration of the formed Fe²⁺ (BPDS)₃ complex (ε of BPDS is 22.1 mM⁻¹ cm⁻¹).
Visualization and localization of proton release and Fe(III) reduction was performed by embedding the roots in a agar medium as described in Marschner et al. (1982) in the presence of the pH indicator Bromocresole Purple and BPDS, respectively.

Isolation of plasma membrane vesicles

Enriched plasma membrane (PM) vesicles were purified using the two-phase partitioning procedure as previously described (Rabotti & Zocchi, 1994). Final pellets were resuspended in a medium containing 2 mM MES (2[N-morpholino]ethanesulphonic acid)-BTP (1,3-bis[tris(hydroxymethyl)-methylamino]-propane), pH 7.0, 1 mM PMSF and 330mM sucrose.

Assay of H+-ATPase and FC-R activities in plasmalemma-enriched vesicles

H+-ATPase activity was assayed with a spectrophotometric method (as described by Palmgren et al., 1990), coupling ATP hydrolysis to NADH oxidation, at 25°C as already reported (Rabotti & Zocchi, 1994). The reaction was started by the addition of 20-50μl of PM preparation and NADH oxidation was followed spectrophotometrically at 340 nm in a V550 spectrophotometer (Jasco, Tokyo, Japan) as already described (Rabotti & Zocchi, 1994).

The NADH-dependent Fe(III)-reductase (FC-R) activity was assayed in the dark at 25°C in 1 ml volume containing 250 mM sucrose, 15 mM MOPS-BTP (pH 6.0), 0.25 mM FeEDTA, 0.25 mM NADH, 0.01% Lubrol. The reaction was started by the addition of 20-50μl of PM preparation and NADH oxidation was followed
spectrophotometrically at 340 nm in a V550 spectrophotometer (Jasco, Tokyo, Japan) as already described (Rabotti & Zocchi, 1994).

PEPCase assay

The PEPCase, soluble cytosolic enzyme, was extracted from roots of plants grown in the presence or in the absence of Fe as reported by De Nisi & Zocchi (2000). Reaction was started by adding aliquots of protein extracts and the enzymatic assays were performed at 25°C in 1.5 ml final volume. Oxidation of NADH was followed spectrophotometrically at 340 nm in a V550 spectrophotometer (Jasco, Tokyo, Japan) as already described (De Nisi & Zocchi, 2000).

Semiquantitative RT-PCR

Root and leaf tissues were pulverised in liquid nitrogen using mortar and pestle and total RNA was extracted using Trizol® reagent (Invitrogen), and first strand cDNA synthesis was carried out using iScript™ cDNA Synthesis Kit (Bio-Rad) according to the manufacturer’s instructions. The gene-specific primers used to amplify: CsFRO1 (accession No. AY590765) were 5’-GTATCACATATGCTTGGC-3’ (forward) and 5’-CTACGAATGCGAGGAATAG-3’ (reverse); CsIRT1 (accession No. AY590764) primers used were 5’-CGCAGCAGGTATCATTCTCG-3’ (forward) and 5’-TCTGCCTGAAGAATACAGCC-3’ (reverse); Cspep1 (accession No. AJ417435) the primers used were 5’-GGACACAGACGAGATTCCATC-3’ (forward) and 5’-CCAGTGTTTCTGCATTCCCGC-3’ (reverse); the primers used to amplify actin (accession No. AB010922) were 5’-GCCTGCTATGTATGTGTGGCCATC-3’ (forward)
and 5’-CAAGAGCAACATATGCCAGCT-3’ (reverse). To amplify CsHA1 (accession No. AJ703810) the primers used were 113F2 (5’-CTCCAACCAGCACCAGAAA-3’) and 113R1 (5’-TCCTTCATCTCTTCTGCAACA-3’) (Santi et al., 2005). The thermal cycle program was: one initial cycle of 94°C 5 min, followed by cycles of 94°C 30 sec; 56°-60°C 1 min (as annealing temperature we used 56°C for CsFRO1 and 60°C for Cspepc, CsIRT1, CsHA1, and Csactin), 72°C 1 min, with 28 cycles for CsFRO1, CsIRT1, CsHA1, Cspepc and 26 cycles for actin, all followed by a final 72°C elongation cycle for 5 min. RT-PCR was carried out on the first-strand cDNA using Taq DNA polymerase (Promega) and the identity of the amplified fragments were verified by sequencing both the strands.

Protein determination

Protein was determined by the using dye-binding method of Bradford (1976), using γ-globulin as a standard.

Results

Effect of iron availability on acidification and reduction activities

To investigate the control exerted by the iron status on the Strategy I plant responses, we started with the induction of Fe deficiency at the whole plant level. Plants were grown for 8 days in the absence of iron to induce all the deficiency responses (Rabotti & Zocchi, 1994) and after this period iron was supplied at a concentration of 100 µM as FeEDTA. The time course of acidification before and after iron supply is reported in
Figure 1A. After 7d in the absence of iron, roots sharply decrease the pH of the medium by almost two pH units, while the control roots do not show any acidification capacity. After iron supply, roots still retain the capacity to decrease the pH for the first 24 h and then the pH raises to higher values. We have assayed the H⁺-ATPase activity during this time course on a PM-enriched fraction isolated from the roots. The results are reported in Table 1. The H⁺-ATPase activity measured confirms previous results (Rabotti & Zocchi, 1994; Dell’Orto et al., 2000) showing a 50% increase under iron deficiency. After iron supply the this activity still remains higher for 48 h followed by a decrease, more or less, to the control level. This difference can also be visually appreciated in Figure 2A where roots are embedded in an agar containing the pH-sensitive dye Bromocresol Purple; the yellow colour around the – Fe roots denotes a decrease in the pH value (lower than 5.0).

Figure 1B shows the time course of the Fe³⁺ reduction before and after iron supply. Also in this case, the reduction activity is sharply increased under iron deficiency reaching the maximum after 7d of iron starvation, then the activity slowly decreases. After iron supply there is an second peak of increase in the reducing capacity of Fe³⁺ even greater with respect to the previous one. This sharp increase last for 24 h and is followed by a rapid decrease in the reduction capacity down to the value of the control. As for the H⁺-ATPase activity, we have measured the Fe³⁺-chelate reductase (FC-R) activity on a PM-enriched fraction. The results are reported in Table 1. This in vitro result confirms what was demonstrated in the time course experiment with a second peak of activity greater than that obtained in the -Fe roots. The increase in the reduction activity after Fe supply to Fe-starved roots, is a well known response and is referred to as a substrate induction effect, i.e. iron acts as a local inducer (Vert et al, 2003).
difference in the activity of Fe(III) reduction is well visible in Figure 2B where the roots are embedded in an agar containing Fe$^{3+}$-EDTA and BPDS as a chelating Fe$^{2+}$ agent [(the red colour is due to the formation of the complex Fe$^{2+}$(BPDS)$_3$]. Fig. 2 also shows the morphological changes occurring at the root level under Fe deficiency; starved plants show an increase in the amount of lateral roots and swollen tips (left part of each plate).

In previous papers we have demonstrated the implication of the PEPC in the response to Fe deficiency and several hypothesis were proposed to explain the strict correlation existing between the induction of iron deficiency responses and the PEPC activation (Rabotti et al., 1995; De Nisi & Zocchi, 2000). Table 1 show the results of the time course of the PEPC activity. Under Fe deficiency the PEPC activity is increased by 4 time and also in this case there is an increase 24 h after Fe supply, consistent with those shown by the H$^+$-ATPase and FC-R activities.

Expression of iron deficiency response genes in cucumber plants

Gene expression analysis was performed using CsFRO1, CsIRT1, CsHA1 and Cspepc1 sequences. CsFRO1 and CsIRT1 were recently characterised by Waters et al. (2007) and shown to encode the ferric reductase and the iron transporter proteins, respectively. CsFRO1 is hortologous to AtFRO2 (Robinson et al., 1999), LeFRO1 (Li et al., 2004) and PsFRO1 (Waters et al. 2002), while CsIRT1 is hortologous to AtIRT1 (Eide et al., 1996) and LeIRT1 (Eckhardt et al., 2001). Concerning to the H$^+$-ATPase, we considered the differential expression level of CsHA1 gene (Santi et al., 2005). For PEPC we considered the Cspepc1 gene expression level. In the database (http://www.ncbi.nlm.nih.gov) two partial mRNAs for PEPC are present for Cucumis
sativus (submitted by Santi et al, unpublished): Cspepc1 (AJ417435) and Cspepc2 (AJ417436). Preliminary phylogenetic analysis carried out on amino acidic sequences of cucumber and Arabidopsis PEPC isoforms [Cspepc1 (CAD10147), Cspepc2 (CAD10148), Atppc1 (CAD58725), Atppc2 (CAD58726), Atpp3 (AAC24594), Atppc4 (CAC86034)] showed that only the Atppc4 was not related to all the other sequence considered (data not shown). For this reason, we performed the nucleotide sequence alignment of Cspepc1, Cspepc2 and of the genes encoding for the three PEPC isoforms of Arabidopsis, and namely: Atppc1 (AJ532901), Atppc2 (AJ532902), Atppc3 (AF071788) (Sanchez & Cejudo, 2003). Cspepc1 showed 79% identity with Atppc3, 74%, with Atppc2 and 78% with Atppc1, while Cspepc2 showed 76% identity with Atppc3, 73% with Atppc2 and 78% with Atppc1. We decided to use in this work the Cspepc1 gene because its sequence showed the highest identity value with Atppc3, which is almost exclusively expressed in roots (Sanchez & Cejudo, 2003).

The presence of iron is believed to be an induction signal for the expression of the iron responsive genes, in particular for AtFRO2 and AtIRT1, but these transcripts are often hardly detectable in the presence of iron (Vert et al, 2001, Connolly et al, 2002; Vert et al, 2003, this paper). On the contrary, it is in the absence of iron that genes encoding for FRO2 and FROI are up regulated (Waters et al., 2002; Connolly et al., 2003; Li et al., 2004) and it is also in this condition that IRT1 mRNA and protein accumulates in A. thaliana (Eide et al., 1996; Connolly et al., 2002; Vert et al., 2002). In this work we confirm these data in cucumber as well and we extended them also to the expression of the CsHA1 and the Cspepc1 transcripts. In control and starved roots we observe a close coordination concerning the expression of these four activities. In fact, all of them are increased under Fe deficiency condition (Fig. 3A) and for the
CsFRO1, CsIRT1 and CsHA1, in particular, the highest level of the mRNAs expression coincided with the maximum of their enzymatic activity (compare the –Fe lanes 5d and 8d, Fig. 3A and Table 1). For AtFRO2 and AtIRT1 a coordinate control of the expression was formulated (Connolly et al., 2003; Vert et al., 2003). We may extend this hypothesis also to the CsFRO1, CsIRT1, CsHA1 and the Cspepc1 (Fig. 3). In fact, when starved roots are re-supplied with iron there is a decrease in the expression of all the transcripts and after 48 h they are almost undetectable (Fig. 3B). This is particularly true for what concern the CsFRO1 and CsIRT1, that are directly involved in iron acquisition, while for CsHA1 and Cspepc1 even after 48 h the transcripts are still present. On the other hand, these two last enzymes are not involved solely in the Fe deficiency responses but participate in many other cellular events. Western blot analysis of the protein extracted from plants grown in the same conditions had shown a similar pattern of accumulation for the H+−ATPase and the PEPC (Dell’Orto et al., 2000; De Nisi & Zocchi, 2000) and IRT1 (Connolly et al., 2002; Vert et al., 2003).

**Discussion**

Cucumber roots respond to iron deficiency by inducing acidification of the culture medium and reduction of Fe(III) within 7 d (Fig. 1A and 1B). Gene expression analysis carried out by semi-quantitative RT-PCR revealed that CsFRO1, CsIRT1, CsHA1 and Cspepc1 transcripts (Fig. 3A) are increased during this period accordingly with the iron deficiency induction of the iron uptake system. This induction is well correlated with the increase in the relative enzymatic activities (Table 1). This work for the first time put in relation the induction of specific genes for iron uptake as CsFRO1 and CsIRT1 and that of the CsHA1 and Cspepc1. From these experiments it is clear that the whole
iron deficiency response is under the same gross control and that supply of iron rapidly de-induces the expression of these transcripts altogether (Fig. 3B), greater for the CsFRO1 and CsIRT1 (they serve specifically for iron uptake), which reach the level seen in the control roots within 48 h, than for the CsHA1 and Cspepc1, that are less specific and also serve for other important cellular functions (Chollet et al., 1996; Palmgren, 2001). A further possible explanation of this different response could relay on the fact that for CsFRO1 and CsIRT1 we can assume a primary coordinate regulation, both local and systemic in response to a direct event (presence or absence of iron). For what concern the CsHA1 and the Cspepc1 a kind of secondary regulation can be supposed in view of a less direct involvement of these two activities in the iron-deficiency responses. It seems possible to hypothesise a sequential coupled regulation which involves these four genes: direct or primary for the response of CsFRO1 and CsIRT1, to promote iron uptake, and secondary or metabolic for CsHA1 and Cspepc1. What is intended for metabolic is the necessity to increase the production of NAD(P)H and ATP for the FC-R and the H⁺-ATPase activities, respectively, that brings to an increase in the rate of glycolysis and perhaps of the pentose phosphate pathway (Rabotti et al. 1995; Espen et al. 2000), along with the necessity to extrude protons which tend to accumulate as the glycolysis rate increases in a sort of pH-stat mechanism (Sakano, 1998). Thus, the activation of CsHA1 and Cspepc1 transcripts should seem to be stimulated as a metabolic consequence of iron starvation rather than by a direct system. This seems to be in agreement with the microarray analysis shown by Colangelo and Guerinot (2004) where there is no evidence that these two last genes are targets of FIT1 regulation. Of course, we can not ruled out the possibility that other regulatory mechanisms operate, such as posttranscriptional modifications, as suggested for FRO2
and IRT1 in Arabidopsis (Connolly et al., 2003), that could involve also the $H^+$-ATPase and the PEPC. Furthermore, even the possibility of posttranslational modifications, that might finely regulate the activities of the enzymes correlated with the iron deficiency responses, can not also be ruled out. This might explain why the $H^+$-ATPase and the PEPC activities can be reduced though their transcripts are still consistently presents (Table 1 and Fig. 3B).

In addition to the activity of $CsFRO1$ and $CsIRT1$, we show that also other activities considered to be linked to the iron deficiency response, the $CsHA1$ (in particular) and the $Cspepc1$, may be regulated throu

What kind of signals are involved in the response to iron deficiency is not yet known. We can assume that iron itself, through its movements in the xylematic and phloematic saps, may signal the iron status of the plant. It can act, according to the dual regulation model proposed by Vert et al. (2003), either as a local inducer signal and as a repressive systemic signal once its concentration inside the plant reaches a satisfactory level. Whether iron acts directly or in association with other molecules is not yet
known. On the other hand, iron deficiency itself cannot be considered a promotive systemic signal, but other molecules or mechanisms could act to induce such responses.

In a study on iron deficiency responses in Arabidopsis by using microarray analysis (Thimm et al., 2001) it was found that in the shoot, several genes involved in the metabolism and export of carbohydrate, are strongly up-regulated in this condition, in particular the phosphate/triose phosphate translocator and the sucrose transporter, suggesting an increased energy requirement outside the shoot. In fact, the energy demand in the roots under iron stress deficiency is very high since they necessitate an increased amount of reducing equivalents, energy and tricarboxylic cycle intermediates to sustain all the processes induced by this condition (Zocchi, 2006 and reference therein). Indeed, an increase in the sugar concentration has been demonstrated in the phloem of iron deficient bean plants (De Vos et al., 1986). Whether sugars, or other molecules transported along with them in the phloem, are responsible for a systemic signal in iron deficiency response is still unknown. The promotive signal has been assigned to several molecules until now, such as IAA (Landsberg, 1984; Römheld & Marschner, 1986), ethylene (Lucena et al., 2006), sugar (Bienfait et al., 1987), iron complexed by a ligand (Kruger et al., 2002) and recently to nitric oxide (Graziano et al., 2002). Regulation of the transcription factors that control the expression of genes involved in iron uptake and metabolism are characterized in bacteria (Escolar et al., 1999), in yeast (Saccharomyces cerevisiae (Yamaguchi-Iwai et al., 2002) and in vertebrate (Papanikolaou & Pantopoulos, 2005). Recently, also in tomato and Arabidopsis transcription factors controlling the iron deficiency responses and the iron uptake has been described (Colangelo & Guerinot, 2004; Jacoby et al., 2004;
Brumbarova & Bauer, 2005; Yuan et al., 2005) suggesting that an analogous system might operate at the plant level.

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References

Bienfait HF, de Weger LA, Kramer D (1987) Control of the development of iron-efficiency reactions in potato as a response to iron deficiency is located in the roots. Plant Physiol 83:244-247

Bradford MM (1976) A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Anal Biochem 72:248-254

Briat JF (2002) Metal ion-activated oxidative stress and its control. In: Montagu M, Inzé D (eds) Oxidative stress in plants. Taylor & Francis Publishers, London, pp 171-190

Brumbarova T, Bauer P (2005) Iron-mediated control of the basic helix-loop-helix protein FER, a regulator of iron uptake in tomato. Plant Physiol 137:1018-1026

Chaney RL, Brown JC, Tiffin LO (1972) Obligatory reduction of ferric chelates in iron uptake by soybeans. Plant Physiol 50:208-213
Chollet R, Vidal J, O’Leary LH (1996) Phosphoenolpyruvate carboxilase: a ubiquitous, highly regulated enzyme in plants. Ann Rev Plant Physiol Plant Mol Biol 46: 273-298

Cohen K, Fox TC, Garvin DF, Kochian LV (1998) The role of iron-deficiency stress responses in stimulating heavy-metal transport in plants. Plant Physiol 116:1063–1072

Colangelo EP, Guerinot ML (2004) The essential basic helix-loop-helix protein FIT1 is required for the iron deficiency response. Plant Cell 16:3400-3412

Connolly EL, Fett JP, Guerinot ML (2002) Expression of the IRT1 metal transporter is controlled by metals at the levels of transcript end protein accumulation. Plant Cell 14:1347–1357

Connolly EL, Campbell NH, Grotz N, Prichard CL, Guerinot ML (2003) Overexpression of the FRO2 ferric chelate reductase confers tolerance to growth on low iron and uncovers posttranscriptional control. Plant Physiol 133:1102–1110

Curie C, Briat JF (2003) Iron transport and signaling in plants. Annu Rev Plant Biol 54:183-206

Dell’Orto M, Santi S, De Nisi P, Cesco S, Varanini Z, Zocchi G, Pinton R (2000) Development of Fe-deficiency responses in cucumber (Cucumis sativus L.) roots: involvement of plasma membrane H^+-ATPase activity. J Exp Bot 51:695-701

Dell’Orto M, Pirovano L, Villalba JM, Gonzalez-Reyes JA, Zocchi G (2002) Localization of the plasma membrane H^+-ATPase in Fe-deficient cucumber roots by immunodetection. Plant Soil 241:11–17
De Nisi P, Zocchi G (2000) Phosphoenolpyruvate carboxylase in cucumber (\textit{Cucumis sativus} L.) roots under iron deficiency: activity and kinetic characterisation. J Exp Bot 352:1903-1909

de Vos CR, Lubberding HJ, Bienfait HF (1986) Rhizosphere acidification as a response to iron deficiency in bean plants. Plant Physiol 81:842-846

Eckhardt U, Marques AM, Buckhout TJ (2001) Two iron-regulated cation transporters from tomato complement metal uptake–deficient yeast mutants. Plant Mol Biol 45:437-448

Eide DJ, Broderius M, Fett J, Guerinot ML (1996) A novel iron-regulated metal transporter from plants identified by functional expression in yeast. Proc Natl Acad Sci USA 93:5624-5628

Escolar R, Perez-Martin J, de Lorenzo V (1999) Opening the iron box: transcriptional metalloregulation by the FUR protein. J Bacteriol 181:6223-6229

Espen L, Dell’Orto M, De Nisi P, Zocchi G (2000) Metabolic responses in cucumber (\textit{Cucumis sativus} L.) roots under Fe-deficiency: a $^{31}${P}-nuclear magnetic resonance in-vivo study. Planta 210:985-992

Graziano M, Beligni MV, Lamattina L (2002) Nitric oxide improves internal iron availability in plants. Plant Physiol 130:1852–1859

Guerinot ML, Yi Y (1994) Iron: nutritious, noxious, and not readily available. Plant Physiol 104:815-820

Halliwell B, Gutteridge JMC (1999) Free Radicals in Biology and Medicine, 3rd edn. Oxford University Press, Oxford

Hell R, Stephan UW (2003) Iron uptake, trafficking and homeostasis in plants. Planta 216:541–551
Jacoby M, Wang H-Y, Reidt W, Weisshaar B, Bauer P (2004) FRU (BHLH029) is required for induction of iron mobilization genes in Arabidopsis thaliana. FEBS Lett 577: 528-534

Krüger C, Berkowitz O, Stephan UW, Hell R (2002) A metal-binding member of the late embryogenesis abundant protein family transports iron in the phloem of *Ricinus communis* L.. J Biol Chem 277:25062–25069

Landsberg EC (1984) Regulation of iron-stress-response by whole plant activity. J Plant Nutr 7:609-621

Li L, Cheng X, Ling HQ (2004) Isolation and characterization of Fe(III)-chelate reductase gene *LeFRO1* in tomato. Plant Mol Biol 54:125–136

López-Millán AF, Morales F, Andaluz S, Gogorcena Y, Abadia A, De Las Rivas J, Abadía J (2000) Responses of sugar beet roots to iron deficiency. Changes in carbon assimilation and oxygen use. Plant Physiol 124:885-897

Lucena C, Waters BM, Romera FJ, García MJ, Morales M, Alcantara E, Pérez-Vicente R (2006) Ethylene could influence ferric reductase, iron transporter, and H⁺-ATPase gene expression by affecting FER (or FER-like) gene activity. J Exp Bot 57: 4145-4154

Marschner H, Römheld V, Ossenberg-Neuhaus H (1982) Rapid method for measuring changes in pH and reducing process along roots of intact plants. Zeit Pflanz Boden 105S:407-416

Ollat N, Laborde B, Neveux M, Diakou-Verdin P, Renaud C, Moing A (2003) Organic acid metabolism in root of various grapevine (*Vitis*) rootstocks submitted to iron deficiency and bicarbonate. J Plant Nutr 26:2165-2176
Palmgren MG (2001) Plant plasma membrane H⁺-ATPases: powerhouses for nutrient uptake. Annu Rev Plant Physiol Plant Mol Biol 52:817-845

Palmgren MG, Askerlund P, Fredrikson K, Widell S, Sommarin M, Larsson C (1990) Sealed inside-out and right-side-out plasma membrane vesicles: optimal conditions for formation and separation. Plant Physiol 92:871-880

Papanikolaou G, Pantopoulos K (2005) Iron metabolism and toxicity. Toxicol Appl Pharmacol 202:199-211

Rabotti G, Zocchi G (1994) Plasma membrane-bound H⁺-ATPase and reductase activities in Fe-deficient cucumber roots. Physiol Plant 90:779-785

Rabotti G, De Nisi P, Zocchi G (1995) Metabolic implications in the biochemical responses to iron deficiency in cucumber (Cucumis sativus L.) roots. Plant Physiol 107:1195-1199

Robinson J, Procter CM, Connolly E, Guerinot ML (1999) A Ferric-Chelate Reductase for iron uptake from soils. Nature 397:694-697

Römheld V, Marschner H (1986) Mobilization of iron in the rhizosphere of different plant species. Ad Plant Nutr 2:155-204

Sakano K (1998) Revision of the biochemical pH-stat: involvement of alternative pathway metabolism. Plant Cell Physiol 39: 467-473

Sanchez R, Cejudo FJ (2003) Identification and expression analysis of a gene encoding a bacterial-type phosphoenolpyruvate carboxylase from arabidopsis and rice. Plant Physiol 132:949–957

Santi S, Cesco S, Varanini Z, Pinton R (2005) Two plasma membrane H⁺-ATPase genes are differentially expressed in iron-deficient cucumber plants. Plant Physiol Biochem 43:287–292
Schmidt W (1999) Mechanisms and regulation of reduction-based iron uptake in plants. New Phytol 141:1-26

Thimm O, Essigmann B, Kloska S, Altmann T, Buckhout TJ (2001) Response of arabidopsis to iron deficiency stress as revealed by microarray analysis. Plant Physiol 127:1030–1043

Vert G, Briat JF, Curie C (2001) Arabidopsis IRT2 gene encodes a root-periphery iron transporter. Plant J 26:181-189

Vert G, Grotz N, Dédaldéchamp F, Gaymard F, Guerinot ML, Briat JF, Curie C (2002) IRT1, an arabidopsis transporter essential for iron uptake from the soil and for plant growth. Plant Cell 14:1223–1233

Vert G, Briat JF, Curie C (2003) Dual regulation of the Arabidopsis high-affinity root iron uptake system by local and long-distance signals. Plant Physiol 132:796-804

Waters BM, Blevins DG, Eide DJ (2002) Characterization of FRO1, a pea ferric-chelate reductase involved in root iron acquisition. Plant Physiol 129:85–94

Waters BM, Lucena C, Romera FJ, Jester GG, Wynn AN, Rojas CL, Alcantara E, Perez-Vicente R (2007) Ethylene involvement in the regulation of the H+-ATPase CsHA1 gene and of the new isolated feric reductase CsFRO1 and iron transporter CsIRT1 genes in cucumber plants. Plant Physiol Biochem 45:293-301

Yamaguchi-Iwai, Ueta R, Fukunaka A, Sasaki R (2002) Subcellular localization of AFT1 transcription factor respond to iron status in Saccaromyces cerevisiae. J Biol Chem 277:18914-18918

Yuan YX, Zhang J, Wang DW, Ling H.-Q (2005) AtbHLH29 of Arabidopsis thaliana is a functional ortholog of tomato FER involved in controlling iron acquisition in strategy I plants. Cell Res 15: 613-621
Zocchi G (2006) Metabolic changes in iron-stressed dicotyledonous plants. In: Barton LL, Abadía J (eds) Iron Nutrition in Plants and Rhizospheric Microorganisms. Springer, Dordrecht pp 359-370
Table I. Effect of iron nutritional status on plasma membrane and PEPC root activities of plant grown in the presence or absence of iron and after iron resupply.

$H^+$-ATPase, Fe(III)-chelate reductase (FC-R) and PEPC activities were determined in the root apical segments of 8-day-old plants grown in iron sufficient or iron deficient nutrient solution and after Fe resupply to the -Fe roots. Data are the mean of three independent experiments. SE never exceeds 8%. Data are expressed as nmol NADH mg prot$^{-1}$ min$^{-1}$

| Time  | H$^+$-ATPase | FC-R | PEPC |
|-------|--------------|------|------|
|       | + Fe | - Fe | + Fe | - Fe | + Fe | - Fe |
| 7 d   | 103  | 157  | 40   | 142  | 82   | 235  |
| Iron resupply | 24 h | 160  | 196  | 310  |
|        | 48 h | 140  | 151  | 220  |
|        | 72 h | 116  | 96   | 103  |
Figure 1.

Time courses of acidification (A) and reduction (B) capacity of cucumber roots grown in the presence (100 µM Fe) (closed circle) or in the absence of Fe (closed square). Arrow indicates the Fe resupply (open square) (100 µM Fe-EDTA). A representative experiment is shown.
Figure 2.

Visualization of medium acidification (A) and iron reduction (B) capacity along cucumber primary roots. Excised primary roots were incubated in 0.1% agar medium and the acidification was detected as pH change of the indicator Bromocresole Purple (yellow); the reduction was determined as the Fe$^{2+}$-(BPDS)$_3$ complex formation (red).
Figure 3.
Iron deficiency-dependent expression of Strategy I responsive and PEPC genes. Total RNA was extracted from roots grown in the presence (+ Fe 8d) or in the absence of iron (-Fe 5d, 8d) and 12h, 24h and 48h after iron resupply. The transcript levels of CsFRO1, CsIRT1, CsHa1, Cspepc1 and actin of roots grown under different iron nutritional status were monitored by semi-quantitative RT-PCR. In A, expression pattern of transcripts after 5 and 8 days of iron deficiency respect to the control. In B, expression pattern after 12h, 24h and 48h of iron resupply to Fe-deficient 8-day-old plants.