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4 5	Does a voltage-sensitive outer envelope transport mechanism contribute to the chloroplast iron uptake?	
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1 Main Conclusion

Based on the effects of inorganic salts on chloroplast Fe uptake, the presence of a voltagedependent step is proposed to play a role in Fe uptake through the outer envelope.

- 4
- 5 Abstract

Although iron (Fe) plays a crucial role in chloroplast physiology, only few pieces of
information are available on the mechanisms of chloroplast Fe acquisition.

8 Here, the effect of inorganic salts on the Fe uptake of intact chloroplasts was tested, assessing

9 Fe and transition metal uptake using bathophenantroline-based spectrophotometric detection

10 and plasma emission coupled mass spectrometry, respectively. The microenvironment of Fe

11 was studied by Mössbauer spectroscopy.

12 Transition metal cations $(Cd^{2+}, Zn^{2+}, Mn^{2+})$ enhanced, whereas oxoanions (NO_3^-, SO_4^{-2-}) and 13 BO_3^{-3-} reduced the chloroplast Fe uptake. The effect was insensitive to diuron (DCMU), an 14 inhibitor of chloroplast inner envelope associated Fe uptake. The inorganic salts affected 15 neither Fe forms in the uptake assay buffer nor those incorporated into the chloroplasts. The 16 significantly lower Zn and Mn uptake compared to that of Fe indicates that different 17 mechanisms/transporters are involved in their acquisition.

The enhancing effect of transition metals on chloroplast Fe uptake is likely related to outer envelope-associated processes since divalent metal cations are known to inhibit Fe^{2+} transport across the inner envelope. Thus, a voltage-dependent step is proposed to play a role in Fe uptake through the chloroplast outer envelope on the basis of the contrasting effects of transition metal cations and oxoaninons.

23

24 Abbreviations

apoLhcII, Light Harvesting Complex II apoprotein; BPDS, bathophenantroline disulphonate;
Chl, chlorophyll; CCCP, Carbonyl cyanide m-chlorophenyl-hydrazone; DCMU, 3-(3,4dichlorophenyl)-1,1-dimethylurea; ΔΨ, transmembrane electrochemical potential; EDTA,
Ethylenediaminetetraacetic acid; FRO, Ferric chelate Reductase Oxidase protein; HEPES, 4-

(2-hydroxyethyl)-1-piperazineethanesulfonic acid; ICP-MS, Inductively Coupled Plasma
 Mass Spectrometry; IE, inner envelope; OE, outer envelope; OEP, Outer Envelope Protein;
 PM, plasma membrane; PPFD, photosynthetic photon flux density; RbcL, Rubisco large
 subunit; VDAC1, Voltage-Dependent Anion Chanel 1; VHA, V-type H⁺ ATPase.

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6 Key words: chloroplast; envelope membrane; iron metabolism; Mössbauer spectrosopy;
7 voltage-dependent transport

1 Introduction

2 Metal ions are essential micronutrients for all living organisms including plants. Among the transition metals, iron (Fe) is the most abundant in plant tissues. In shoot tissues of plants with 3 4 normal Fe supply, up to 80–90% of the total Fe is found in the chloroplasts (Terry and Abadía 5 1986; Morrissey and Guerinot 2009), and thylakoid membranes themselves contain 6 approximately 60% of total leaf Fe (Castagna et al. 2009). The major Fe-sinks in the 7 chloroplasts are proteins binding non-heme Fe, Fe-S clusters and heme cofactors. The absence 8 of Fe induces strong deficiency symptoms, the so-called Fe chlorosis. The biosynthesis of 9 chlorophyll (Chl) and Fe-S clusters as well as the assembly of pigment-protein complexes is 10 strongly hampered by Fe deficiency, leading to a decrease in photosynthetic capacity and productivity of plants (Andaluz et al. 2006; Timperio et al. 2007; Abadía et al. 2011; Basa et 11 12 al. 2014). Fe is also required for the activity of Fe-containing enzymes involved in protection 13 against oxidative stress, and therefore anti-oxidative defence mechanisms can be affected 14 when Fe is in short supply (Latifi et al. 2005; Tewari et al. 2005).

15 In contrast with the wealth of data on Fe root acquisition (Abadía et al. 2011) and long 16 distance transport (Rellán-Álvarez et al. 2010), our knowledge on transition metal uptake by 17 leaf cells and organelles are still scarce (for review see: Krämer et al. 2007; Palmer and 18 Guerinot 2009; Abadía et al. 2011). In the mesophyll apoplast and symplast, Fe can occur in 19 the form of citrate or nicotianamine (NA) complexes (Weber et al. 2007; Álvarez-Fernández 20 et al. 2014), but their participation and importance in the leaf cell Fe uptake and transport processes is still not known. Leaf mesophyll cells are known to take up both Fe^{2+} and Fe^{3+} 21 (Nikolić and Römheld 2007), and although the whole process is not well understood yet, at 22 least part of the apoplasmic Fe³⁺ can undergo reduction mediated by FRO reductases for an 23 24 effective Fe uptake (Jeong et al. 2008) similarly to root Fe aquisition.

Concerning the chloroplasts, photosynthetic organelles of endosymbiotic origin, Fe uptake may differ from that of eukaryotic cells, since Fe should cross two different membranes, the chloroplast outer (OE) and inner envelopes (IE). The first protein found to be involved in chloroplast Fe acquisition was PIC1/TIC21 (Duy et al. 2007a). PIC1 is localised in the IE of chloroplasts in *Arabidopsis*, was shown to be a component of the IE protein translocon machinery (Teng et al. 2006), and is a member of a larger 'Fe-import' complex together with the NiCo protein (Duy et al. 2011). Based on results obtained with PIC1 overexpressing lines,

1 it also seems to regulate chloroplast Fe metabolism (Duy et al. 2011). Another important 2 member of the Fe uptake machinary is the chloroplast ferric chelate oxidoreductase (FRO7), 3 in the absence of which chloroplasts were not able to take up Fe from the cytoplasm (Jeong et 4 al. 2008). The chloroplast FRO has been recently localised in the IE (Solti et al. 2014). In fact, the chloroplast FRO and IE uptake machinery must work in a close cooperation, because no 5 6 free Fe^{2+} accumulates during the uptake process (Solti et al. 2012). Bughio et al. (1997) and Solti et al. (2012) showed that Fe uptake of barley (Hordeum vulgare) and sugar beet (Beta 7 8 vulgaris) chloroplasts, respectively, were light/photosynthesis dependent, since both was blocked by a PSII inhibitor. Shingles et al. (2002) found the importance of inwardly directed 9 proton gradient in Fe^{2+} movement across the chloroplast IE membrane. 10

Despite some Fe metabolism related transporters have been discovered in the past few years, 11 no transport protein participating in the movement of Fe^{3+} across the chloroplast OE has been 12 described by now (Inoue 2011; Breuers et al. 2011, López-Millán et al. 2016). OE membrane, 13 14 the first barrier that regulates solute transport in and out of the chloroplasts, includes various 15 transport proteins and protein complexes (Gutierrez-Carbonell et al. 2014). Most of them are 16 descendant of prokaryotic ancestors (Reumann and Keegstra 1999), and similar to those occuring in Gram-negative bacteria. Gram-negative bacteria take up Fe in Fe³⁺-siderophore 17 chelated forms across the plasma membrane (PM). This uptake of Fe³⁺-siderophores is 18 19 voltage sensitive (Braun 2003). In Escherichia coli, the main component in the Fe uptake across the outer membrane is the FecA, a TonB-dependent Fe³⁺-citrate receptor/gated 20 channel, which is energised via a proton gradient across the cytoplasmatic membrane through 21 22 cytoplasmatic membrane integrated proteins (Duy et al. 2007b; Braun and Herrmann 2007; 23 Marshall et al. 2009). β -barrel, pore-forming transport proteins, such as TOC75, are abundant 24 in the OE (Inoue 2007; Duy et al. 2007b; Breuers et al. 2011; Gutierrez-Carbonell et al. 2014). Among OE proteins, the presence of a specific channel for amino acids (OEP16), an 25 26 ATP- and substrate-regulated channel (OEP21), a cation-selective channel (OEP37) and an unspecific channel (OEP24) have been already approved (Duy et al. 2007b; Breuers et al. 27 28 2011; Gutierrez-Carbonell et al. 2014). The outer envelope proteins and the function of OEPs 29 have been discussed several times (Soll et al. 2000; Bölter and Soll 2001; Duy et al. 2007b; 30 Breuers et al. 2011), but the significance of voltage-regulation in the Fe uptake of intact 31 chloroplasts is not clear yet.

The aim of this study was to test whether voltage-sensitive transport through OE may have a
 function in the Fe acquisition of intact chloroplasts. In particular, we studied how transition
 metal cations and oxoanions influence the Fe uptake of intact chloroplasts.

4

5 Materials and Methods

6

7 Plant material

8 Sugar beet (*Beta vulgaris* L. cv. Orbis) plants were grown in hydroponics in a climate 9 chamber with 14/10 h light (160-200 μ mol m⁻² s⁻¹ photosynthetic photon flux density, 10 PPFD)/dark periods, 24/22 °C and 70/75% relative humidity in modified ¹/₄ strength Hoagland 11 solution with 10 μ M Fe³⁺-citrate (Fe:citrate = 1:1, Reanal Kft., Hungary) as Fe source (Solti et 12 al. 2012). Mature leaves of plants having 7-8 leaves were used for isolation of chloroplasts.

13

14 Chloroplast isolation, determination of purity and intactness

15 Sugar beet chloroplasts were isolated and purified on a stepwise sucrose gradient as described in Solti et al. (2012), and see also the Supplementary Material 1. Chloroplast density was 16 determined by counting in a Nikon Optiphot-2 microscope. Chloroplasts were solubilised in 17 18 62.5 mM Tris-HCl, pH 6.8, 2% SDS, 2% DTT, 10% glycerol, and 0.001% bromophenol blue 19 at room temperature for 30 min. Proteins were separated in 10–18% gradient polyacrylamide 20 gels in a MiniProtean apparatus (BioRad) using a constant current of 20 mA per gel at 6 °C. 21 Protein concentration of samples was determined by comparing the area density with that of a 22 standard mixture using Phoretix 4.01 software (Phoretix International, Newcastle upon Tyne, 23 UK).

To detect mitochondrial contamination and estimate the integrity of the chloroplasts, protein
 blots were carried out. Membrane proteins separated by SDS-PAGE were transferred to
 Amersham[™] Protran[™] Premium 0.2 um NC blotting membranes (Amesham-Pharmacia,

1 Germany) in a 25 mM Tris, pH 8.3, 192 mM glicine, 20% (v/v) methanol and 0.02% (m/v)

2 SDS at 4 °C using 90 V constant voltage (<0.4 A) for 3 h.

3 The purity of chloroplast preparations was checked with rabbit polyclonal antibody against 4 mitochondrial alternative oxidase (AOX 1/2, a mitochondrial inner envelope marker, Lang et al. 2011) (Agrisera AG, Vännäs, Sweden). Mitochondrial sign at ~ 34 kDa was identificated 5 6 according the manufacturer's informations (for more information, please visit: 7 http://www.agrisera.com/en/artiklar/aox1_2-plant-alternative-oxidase-1-and-2.html). In order to estimate the integrity of the chloroplasts, membranes were decorated with rabbit polyclonal 8 9 antibodies against apoLHCII (a gift from Dr. Udo Johanningmeier, Bohum Universität, 10 Germany) and RbcL (Rubisco large subunit, form I and form II; Agrisera AG, Vännäs, Sweden). Antibodies were dissolved in 20 mM Tris-HCl (pH 7.5), 0.15 M NaCl, 1% gelatine 11 12 following the manufacturer's instructions. Horseradish peroxidase- (HRP-) conjugated goatanti-rabbit IgG (BioRad, Inc.) was used to detect bands following the manufacturer's 13 14 instructions. Chloroplast integrity was estimated by comparing the RbcL/apoLhcII ratio in 15 solubilized leaf tissues and chloroplast samples as in Solti et al. (2012).

16

17 Measurements of Fe uptake

18 Iron uptake was assessed from the total chloroplast Fe content before and after a 30 min Fe 19 uptake assay. Total chloroplast Fe was measured with BPDS according to Solti et al. (2012). The assay was carried out with 0.5 ml chloroplast suspension (100 μ g chlorophyll (Chl) ml⁻¹; 20 approximately 76000 \pm 9500 chloroplasts μ l⁻¹) in uptake buffer (50 mM HEPES-KOH, pH 21 7.0, 330 mM sorbitol, 2 mM MgCl₂), and 100 µM Fe³⁺-citrate (Fe:citrate 1:1; Reanal Kft., 22 23 Hungary) was used as Fe source. In order to test the effects of inorganic salts on the Fe uptake 24 process, one of the following inorganic salts: KCl, K₃BO₃, KNO₃, K₂SO₄, CdSO₄, CdCl₂, 25 ZnSO₄, ZnCl₂ and MnCl₂ was also added to the Fe uptake medium at a concentration of 500 26 μM. Transition metal cations were also tested as chloride salts at a concentration of 200 μM. 27 K^+ and Cl^- were present both in the isolation buffer and the uptake assay medium in mM 28 concentrations. To uncouple chloroplast envelope membrane $\Delta \Psi$, 5 µM of the ionophore 29 CCCP (carbonyl cyanide m-chlorophenyl-hydrazone) was added to the Fe uptake assay 30 medium. The reduction-based Fe uptake across the chloroplast IE membrane was disrupted by 1 using 10 μ M of the photosynthetic electron transport inhibitor DCMU (3-(3,4-2 dichlorophenyl)-1,1-dimethylurea), which blocks NADPH production by the photosynthetic 3 electron transport chain. Values obtained with the Fe uptake medium free of any added 4 inorganic salts and inhibitors are referred to as 'control' values throughout the paper. Iron 5 uptake was initiated by illuminating the samples with 160 μ mol m⁻² s⁻¹ PPFD white light, and 6 terminated by placing the samples in ice to the dark. Fe uptake values are expressed in 7 attomol (amol; 10⁻¹⁸ mol) Fe taken up chloroplast⁻¹.

8

9 Determination of element concentrations in chloroplasts

10 Chloroplast samples, taken before and after the Fe uptake assay, were washed in washing buffer containing 50 mM HEPES-KOH (pH 7.0), 330 mM sorbitol, 2 mM EDTA, 2 mM 11 MgCl₂. The total Fe content of chloroplasts was determined after reduction by 100 µM 12 ascorbic acid and Fe^{2+} complex formation with 300 µM BPDS as in Solti et al. (2012), using 13 an absorption coefficient of 22.14 mM⁻¹ cm⁻¹ for the Fe(II)-BPDS complex (Smith et al. 14 15 1952). For determination of the concentrations of other elements, chloroplast samples taken 16 before and after the uptake assays were washed in washing buffer, resuspended in uptake buffer and dried for one week at 60 °C. Samples were digested by HNO₃ for 30 min at 60 °C 17 18 and then in H₂O₂ for 90 min at 120 °C. After filtration by MN 640W paper, ion contents were 19 measured using ICP-MS (Inductively Coupled Plasma Mass Spectrometer, Thermo-Fisher, 20 USA).

22 Mössbauer spectroscopy

Changes in the chemical microenvironment of ⁵⁷Fe were assessed using Mössbauer 23 spectroscopy (Solti et al. 2012). After 30 min incubation in uptake buffer supplemented with 24 ⁵⁷Fe³⁺-citrate, chloroplasts were washed in washing buffer to remove any excess Fe adsorbed 25 26 on the organelle surface. Concentrated chloroplast suspensions were placed in a conventional 27 constant acceleration type Mössbauer spectrometer (Wissel) in a liquid nitrogene bath cryostat at 80 K. A 57 Co(Rh) source of ~10⁹ Bq activity was used, and the spectrometer was calibrated 28 29 with α -Fe at room temperature. Evaluation of spectra was carried out using the MOSSWIN 30 code (Klencsár et al. 1996). The Mössbauer parameters calculated for the spectral components

²¹

1 were: isomer shift (δ , mm s⁻¹), quadrupole splitting (Δ , mm s⁻¹) and partial resonant 2 absorption areas (S_r , %). These parameters provide information on the electron densities at the 3 Mössbauer nuclei (including also the valence state) and on the magnitude of any electric field 4 gradients (indicating the coordination number of the resonant atom). Quantitative analytical 5 information for the different species found can be obtained from the relative spectral areas 6 (Greenwood and Gibb 1971).

7

8 Statistical analysis

9 Fe uptake measurements were carried out with three technical repetitions in each of three to 10 four biological repetitions. To analyse statistical differences between means a Student's t-test 11 was applied. To compare multiple treatments, one-way ANOVA was performed with a 12 Tukey-Kramer multiple comparison *post hoc* test, using InStat v. 3.00 (GraphPad Software, 13 Inc.).

14

15 Results

16 Intactness of chloroplasts

17 To determine the intactness of chloroplasts, the ratio of RbcL to apoLhcII was followed and compared during the whole isolation process, i.e. in leaves, leaf homogenates, first chloroplast 18 19 pellets and in class I and class II chloroplast fractions (Fig. 1). Purified class I chloroplast 20 fractions were free of mitochondrial contamination (no sign of AOX 1/2 was detected 21 according to westen blots). Although the first chloroplast pellet contained larger amount of 22 damaged chloroplast, intact class I chloroplasts could be purified by sucrose gradient 23 centrifugation. Comparing the RbcL/apoLhcII ratio of chloroplast samples to that of leaves, 24 the intactness of class I chloroplast was 96.8±8.3% (for calculations, see Supplementary Material 1), while the soluble Rubisco escaped from the chloroplast stroma of damaged class 25 26 II chloroplast. When chloroplasts were subjected to Fe uptake assay, inorganic salts did not 27 influenced significantly the intactness of the chloroplasts (see Supplementary Material 2).

1 Influence of inorganic salts on the chloroplast iron uptake

Chloroplasts were able to take up Fe from a medium containing Fe³⁺-citrate independently of 2 3 the inorganic ions present. Though a 30-min incubation in the Fe uptake medium in the presence of Cd^{2+} , Mn^{2+} , Zn^{3+} , NO_3^{-} , SO_4^{2-} or BO_3^{2-} did not cause significant changes in the 4 chloroplast intactness (Fig. S2, Table S2), the Fe uptake varied in the presence of different 5 ions. When salts were applied at 500 μ M concentrations, transition metal cations (Cd²⁺, Zn²⁺, 6 Mn^{2+}) generally enhanced the Fe uptake whereas anions added as K⁺ salts (NO₃⁻, SO₄²⁻ and 7 BO_3^{3-}) decreased it (Fig. 2). Among the metal cation chloride salts, ZnCl₂ was the most 8 effective in stimulating Fe uptake. It was followed by CdCl₂, whereas MnCl₂ had a weakly 9 significant effect (P<0.05). Using K^+ salts, effects of anions were independent of their 10 valence: Cl⁻ did not influence the chloroplast Fe uptake and the inhibition by NO_3^{-} and BO_3^{3-} 11 was similar, whereas the presence of SO_4^{2-} decreased strongly the uptake of Fe. When the 12 transition metal cations and SO_4^{2-} were present together in the uptake medium (transition 13 14 metals were added in forms of sulphate salts) they apparently had antagonistic effects, leading to an intermediate Fe uptake, i.e. to values between those measured with K₂SO₄ and the metal 15 16 chloride salts.

17 The chloroplast Fe uptake was also affected by the concentration of transition metal cations in 18 the uptake medium, Fe uptake being more enhanced at a concentration of 200 μ M than at 500 19 μ M (Fig. 3). The differences in Fe uptake using 200 and 500 μ M metal concentrations was 20 22±4% in average.

21 To uncouple chloroplast envelope membrane $\Delta \Psi$, CCCP was added to the Fe uptake assay 22 medium. The ionophore CCCP eliminated the Fe uptake capacity of chloroplasts both in 23 darkness (not shown) and in light (Fig. 4), since hardly any measurable changes were found in 24 the Fe content of chloroplasts irrespectively of the inorganic metal salt used. The reduction-25 based Fe uptake was disrupted by the photosynthetic electron transport inhibitor DCMU, 26 which blocks NADPH production. In the absence of any additional inorganic salts, DCMU 27 significantly decreased the Fe uptake of control chloroplasts under light conditions during a 30-min incubation, which dropped from 570 ± 78 to 120 ± 15 amol Fe chloroplast⁻¹ (a 79%) 28 29 decrease, Fig. 4). Using inorganic metal salts, a DCMU-induced decrease in Fe uptake was also found (Fig. 4) but the tendency of changes was the same as without DCMU (Fig. 2): KCl 30 and KNO₃ induced no singificant change, transition metal cations (except Mn^{2+}) and 31

oxoanions significantly increased and decreased the Fe uptake, respectively. The percentage
 of changes were also similar to the case when inorganic salts were applied in the absence of
 DCMU.

4

5 ICP studies on the element content of chloroplasts and uptake of transition metals

6 Numerous elements were detected in isolated chloroplasts, with Ca, Mg and S being present 7 in high amounts (Table 1). Potassium content was somewhat higher but comparable to that of 8 Na. Among the essential transition metals, two groups could be distinguished, with Fe and Zn 9 being approximately five- to ten-fold more abundant when compared to Cu, Mn and Mo. 10 Among non-essential metals, the chloroplast contents of Al and Ba were high and comparable 11 to those of Fe and Zn. Other non-essential transition metals were present in lower amounts. 12 Incubation of chloroplasts in the control Fe uptake medium for 30 min in the light did not 13 change the element contents considerably (the Cu and Cr content showed a small but 14 significant increase only), except that a 3.4 fold-increase in their Fe content.

15 The uptake of transition metals into the chloroplasts was also monitored by ICP-MS. The 16 incubation of chloroplasts in Fe uptake medium supplemented with transition metals in the 17 form of chloride at 500 µM concentrations for 30 min in the light led to a significant uptake of 18 each metal (Fig. 5). The uptake of Zn and Mn were significantly lower than that of Fe, 19 whereas the Cd uptake was comparable to that of Fe uptake in the control assay. After the 30 20 min incubation period in the presence of 500 µM ZnCl₂, MnCl₂ and CdCl₂, the final metal contents taken up were 112±4, 146±5 and 515±17 amol chloroplast⁻¹ for Zn, Mn and Cd 21 22 respectively, whereas that of Fe ranged from 883 ± 3 to 1168 ± 4 amol Fe chloroplast⁻¹.

23 Mössbauer analysis of Fe forms

The Fe chemical forms in the assay medium were not altered in the presence of the applied inorganic salts. In the uptake medium, the 57 Fe³⁺-citrate solution showed only one component, with hyperfine parameters δ =0.47(1) mm s⁻¹ and Δ =0.64(1) mm s⁻¹, typical of high spin Fe³⁺carboxylate complexes (Solti et al. 2012) (Fig. 6A). The presence of anions did not have any effect on the Mössbauer spectrum, with no additional quadrupole doublets appearing on the spectra; as an example, when adding 500 μ M SO₄²⁻ the only component had hyperfine 1 parameters $\delta = 0.48(1)$ mm s⁻¹, $\Delta = 0.65(1)$ mm s⁻¹ (Fig. 6B). Similar results were found with 2 other anions (not shown).

The chemical forms of the ⁵⁷Fe taken up by chloroplasts from the 100 μ M ⁵⁷Fe³⁺-citrate assay 3 medium during 30-min incubation in the light were also studied by Mössbauer spectroscopy 4 (Fig. 7). The spectra consisted of a broadened quadrupole doublet which was fitted to the 5 superposition of two doublets with the following hyperfine parameters: δ =0.46(1) mm s⁻¹, 6 $\Delta = 1.06(4) \text{ mm s}^{-1}$ (Fe_A component) and $\delta = 0.48(1) \text{ mm s}^{-1}$, $\Delta = 0.61(3) \text{ mm s}^{-1}$ (Fe_B component) 7 (Fig. 6A). These components have been assigned to heme groups or Fe_4S_4 (Fe_A) and high spin 8 Fe^{3+} -carboxylate complexes, respectively, with the latter being assigned to Fe^{3+} -citrate (Fe_B) 9 having passed the OE membrane but not yet metabolized (Solti et al. 2012). The relative 10 abundance of Fe_A component accounted for approximately one third of the total Fe present in 11 12 the sample $(35\pm10\%)$. The addition of anions or cations to the uptake medium did not result in any change in the Mössbauer spectra when compared to control samples containing only 13 ⁵⁷Fe³⁺-citrate: as an example, the Fe chemical forms in the chloroplasts were not altered by 14 the addition of 500 μ M Zn²⁺ (Fig. 7B). Similar results were found with other metals (not 15 shown). Signals that could be assigned to high spin Fe^{2+} compounds (e.g., $[Fe(H_2O)_6]^{2+}$ or 16 Fe²⁺-carboxylates) or ferritins were not found in any of the chloroplast samples analyzed. 17

18

19 Discussion

20 Though chloroplasts contain a large part of the total shoot Fe, and Fe plays a crucial role in chloroplast structure and function, only few pieces of information are available about the in 21 22 vivo mechanisms and regulation of Fe acquisition by chloroplasts (Nouet et al. 2011). In particular, no information is available on the role of the chloroplast OE in the chloroplast Fe 23 24 uptake process. Since chloroplasts are organelles of endosymbiotic origin, and Fe transport 25 across the outer membrane of the evolutionally related Gram-negative bacteria is a membrane 26 potential driven step, we tested whether voltage-sensitive transport through OE may also have a function in the Fe acquisition of intact chloroplasts. 27

28

29 Transition metals and oxoanions affect Fe uptake in chloroplasts

1 The presence of inorganic salts in the assay medium influenced the chloroplast Fe uptake but 2 did not affect chloroplast integrity. The Fe forms in the uptake buffer were unchanged as judged by Mössbauer spectroscopy. This latter is in line with the fact that Fe³⁺-citrate has a 3 much higher stability constant (K_i=11.5) than Zn^{2+} -citrate (K_i=5.0), Mn²⁺-citrate (K_i=4.2) or 4 Cd^{2+} -citrate (K_i=3.8) under the conditions used (Fodor 2002), which makes the occurrence of 5 6 metal-citrate complexes other than Fe^{3+} -citrate unlikely. Furthermore, the presence of divalent metal cations, as well as that of NO_3^{-} , SO_4^{2-} and BO_3^{3-} , had no effect on the Fe forms 7 incorporated into chloroplasts as judged by Mössbauer spectroscopy, since i) only two signals 8 9 were found, corresponding to Fe^{3+} -carboxylates (Fe_B) and Fe-S centers or heme (Fe_A), the same signals as reported previously in untreated chloroplasts, and ii) the ratio of Fe_A/Fe_B 10 component was similarly unchanged (Solti et al. 2012). These results, obtained in the 11 12 presence of relatively high concentrations of transition metal ions that are known to interact 13 with Fe homeostasis in vivo (Pilon et al. 2009; Kobayashi and Nishizawa 2012), suggest that 14 the effects of these metals on Fe uptake may reside mainly on transport processes through the chloroplast envelope, but not on a different allocation of Fe forms inside the chloroplast. 15

16 Iron uptake in intact chloroplasts was enhanced by transition metal cations, whereas it was 17 hampered by oxoanions. These results were the opposite to those of Shingles et al. (2002) who reported that divalent transition metal cations such as Zn^{2+} inhibited the movement of 18 Fe²⁺ across the chloroplast IE membrane. In addition, the effect of cations and anions were 19 20 similar in the absence and presence of DCMU. Iron uptake into the chloroplasts is 21 light/photosynthesis dependent, and photosynthesis inhibitors such as DCMU, are known to 22 eliminate the majority of chloroplast Fe uptake capacity (Bughio et al. 1997; Solti et al. 2012). The reason for the inhibitory effect of DCMU is that photosynthetically produced NADPH is 23 necessary to fuel the chloroplast IE Fe^{3+} -chelate reductase enzyme (Solti et al. 2014), which is 24 an essential component in the chloroplast Fe uptake process (Jeong et al. 2008). Based on 25 26 Mössbauer spectroscopy studies, the DCMU-insensitive Fe uptake has been postulated as a 27 Fe-pool that has moved across the OE membrane and accumulated between the OE and IE 28 membranes (Solti et al. 2012). Taken together the opposite effect of divalent cations on Fe 29 uptake of intact chloroplasts compared to IE (Fig. 2 versus Shingles et al. 2002), the 30 inhibitory effects of DCMU on the Fe movement across IE membrane (Solti et al. 2012) and 31 the similar influence of transition metal cations and oxoanions on the DCMU-sensitive and insensitive Fe uptake (Fig. 4), necessarily, an additional regulatory role of OE in Fe uptake is 32 33 strongly supported.

May a bacterial type, voltage-dependent Fe uptake mechanism exist in the chloroplast OE?

3 To the best of our knowledge, no results have been published so far on the mechanism of Fe 4 movement across the chloroplast OE. Data obtained by Mössbauer spectroscopy indicated that Fe crossed the OE membrane in a chelated, Fe^{3+} -citrate form, which accumulated in the 5 6 inter-envelope space before reduction (Solti et al. 2012). The similarities that Gram-negative 7 bacteria, including cyanobacteria, share with chloroplasts may aid to understand the Fe uptake 8 mechanism of chloroplasts. In Gram-negative bacteria, voltage sensitivity of the uptake of 9 Fe³⁺-siderophores was found (Braun 2003), where changes in the PM $\Delta\Psi$ regulates the pore opening of the Fe³⁺-siderophore transporter in the outer membrane (Braun and Hantke 2011). 10 11 In Gram-negative bacteria, the Ton system contributes to the transfer of the energy, originates 12 from the polarization of the cytoplasmic membrane, to the OM transporters (Braun, 2014). A 13 voltage-sensitive mechanism may be also expected to facilitate the Fe uptake in chloroplasts, 14 since chloroplasts polarise their membrane systems similarly to free living Gram-negative 15 bacteria (Shingles et al. 2002). Chloroplasts are known to maintain $\Delta \Psi$ and ΔpH across the IE 16 membrane (Shingles and McCarty 1994; Pottosin and Dobrovinskaya 2015). $\Delta \Psi$ (positive intermembrane space) was shown to be an inwardly rectifying driving force for the Fe²⁺ 17 movement across the IE membrane (Shingles et al. 2002). Nevertheless, the polarization of 18 19 the IE membrane (accumulation of positive charges in the intermembrane space) necessarily 20 polarizes the OE membrane as well. Here, using the uncoupling ionophore CCCP, the Fe 21 uptake of chloroplasts was fully abolished, which supports the previous observations on the 22 voltage-dependency of Fe uptake mechanism. Nevertheless, CCCP, being a hydrophobic 23 compound, not only uncouples the $\Delta \Psi$ of the chloroplast IE membrane but can also 24 incorporate into chloroplast OE, thus eliminating any possible additional effects of the 25 inorganic salts. In the chloroplast OE, the OEP24 voltage-sensitive β -barrel protein, similar to 26 the mitochondrial voltage-dependent anion channel (VDAC) proteins (Röhl et al. 1999; 27 Clausen et al. 2004), have been reported This may have a number of distinct functions 28 (Homblé et al. 2012). The presence of these proteins in the OE is also supported by the recent 29 chloroplast envelope proteome analysis work of Gutierrez-Carbonell et al. (2014). Therefore, 30 a voltage-dependent Fe-complex transport mechanism can also be postulated for the 31 chloroplast OE. We hypothesize that the presence of transition metal cations and oxoanions 32 can change the polarisation of the OE membrane (depolarize and hyperpolarize it,

respectively), possibly due to the lower permeability of transition metal cations or oxoanions
compared to the ions normally present, such as K⁺ and Cl⁻, which have relatively low surface
charge and thus a smaller hydrate coat. A similar effect of inorganic ions was also observed
on the PM solute transport activity of root cells (Wang et al., 2011).

5

6 The effect of transition metals on chloroplast Fe uptake is quality- and concentration-7 dependent

8 Chloroplasts require transition metals such as Zn, Mn and Cu to be functional (Shcolnick and 9 Keren 2006). In fact, our results indicate that chloroplasts take up available transition metals from the assay medium, and their uptake increasing at higher concentrations. Whereas Cu and 10 Zn are known to be taken up by P-type ATPases (Abdel-Ghany et al. 2005; Finazzi et al. 11 2014) and by HMA1 (Kim et al. 2009), respectively, no data have been published yet on the 12 mechanism of chloroplast uptake of Mn, as well as for non-essential metals such as Cd (Nouet 13 et al. 2011). The significantly higher uptake of Cd^{2+} compared to Zn^{2+} and Mn^{2+} supports that 14 different chloroplast uptake mechanisms should be involved in both cases. Cadmium was 15 shown to be taken up by plant cells in competition to Ca^{2+} (Perfus-Barbeoch et al. 2002; 16 Rodríguez-Serrano et al. 2009), and thus chloroplasts may also take up Cd²⁺ mediated by Ca²⁺ 17 transporters. In spite of the fact that Zn^{2+} and Mn^{2+} inhibit Fe²⁺ uptake by the IE competitively 18 (Shingles et al. 2002), the relatively low uptake of Zn^{2+} and Mn^{2+} even when they are present 19 in high concentrations in the uptake medium makes unlikely that chloroplast Zn^{2+} and Mn^{2+} 20 21 uptake may occur through the Fe transport system. Among the transition metal cations tested, Zn^{2+} has the highest surface charge (i.e. the smallest ionic radius, the ionic radius for Cd^{2+} : 22 0.97 Å, for Mn^{2+} : 0.80 Å, and for Zn^{2+} : 0.74 Å), so that the size of its hydrate coat is the 23 largest. A larger hydrate coat in the case of free Zn^{2+} may cause a (higher) size-exclusion in 24 25 the movement across the OE membrane leading to a longer lasting depolarisation, which in 26 turn results in a higher enhancement of the chloroplast Fe uptake.

Higher transition metal concentrations in the uptake medium resulted in a relatively lower enhancement of Fe uptake (Fig. 3). While transition metal cations stimulated chloroplast Fe uptake, they are known to inhibit Fe²⁺ movement across the chloroplast IE membrane competitively (Shingles et al. 2002). A similar inhibitory effect was also found on the process of Fe uptake through bacterial PM (Moreau et al. 1998). The reason for the lower stimulating effect at higher metal concentrations may be the sum of the stimulation of Fe uptake through
 the OE and the competitive inhibition of the Fe uptake through the IE.

3

4 Conclusion

We propose that a voltage-dependent Fe³⁺-complex transport system is involved in the Fe³⁺-5 6 citrate transport across the chloroplast OE. Our proposal is based on the DCMU-insensitive 7 and uncoupling ionophore (CCCP)-sensitive stimulating effects of transition metal cations $(Cd^{2+}, Zn^{2+}, Mn^{2+})$ and inhibitory effects of oxoanions $(NO_3^-, SO_4^{-2-}, BO_3^{-2-})$ on the Fe uptake 8 of intact chloroplasts. The enhancement effect of transition metals on the Fe uptake is quality-9 10 and concentration dependent. The lower stimulation at higher concentrations may be 11 connected with the transition metal uptake into the inter-envelope space, and thus their inhibitory effect on the Fe^{2+} uptake system of the IE. 12

13

14 Author contribution statement

15 Á.S. conceptualized the study, designed the experiments, participated in most of the experimental work and wrote the manuscript. KK. performed the Mössbauer spectroscopy 16 17 studies. B.M. grew the experimental plants, contributed to the chloroplast number 18 calculations, protein polyacrylamide gel electrophoresis and western blots. S.V. contributed to 19 the experiments focusing on the effects of inorganic ions. É.H. and H.D.P. contributed to the 20 experiments focusing on the effect of inhibitors on the iron uptake of chloroplasts. B.T. 21 carried out the determination of element concentrations in chloroplasts by ICP-MS. J.A. and 22 F.F. contributed in the conceptualization of the study, design of the experiments and helped to 23 edit the manuscript. All authors have read and approved the article.

24

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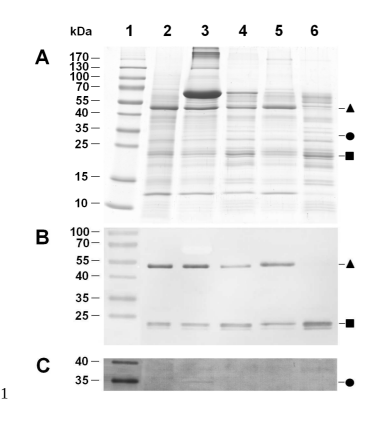
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14

Table 1 Element contents (in amol chloroplast⁻¹) in freshly isolated chloroplasts (A) and in
chloroplasts after incubation in the Fe uptake assay medium in the light for 30 min (B).
Statistical differences between means (Student's t-test, *P*<0.05) are indicated (*).

	Α	В
Al	317.3±72.3	317.3±43.3
B	359.4±20.9	323.2±2.7*
Ba	244.7±40.7	241.2±3.2
Ca	134873.3±19728.3	127210.6±27189.4
Cd	2.4 ± 0.6	2.1 ± 0.7
Cr	6.1 ± 0.8	7.8±0.5*
Cu	20.9 ± 8.8	26.9±1.0*
Fe	233.7±12.2	803.0±7.8*
K	7652.2±442.5	8584.8±3079.3
Li	17.2 ± 1.9	17.3±2.2
Mg	20027.2±2348.7	20859.1 ± 2674.8
Mn	59.5±7.7	61.8±10.7
Мо	53.0±5.2	48.3±3.8*
Na	4484.0±631.6	4026.6±721.0
Р	2239.8±148.5	2326.1±174.0
S	27101.1±1063.7	26928.0±2292.0
Sr	87.3±16.0	84.2±3.1
Zn	342.7±25.6	320.7±51.1



3 Figure 1 Coomassie stained solubilized proteins on polyacrylamide gel (A), combined 4 immunoblot against RbcL and apoLhcII (B), and immunoblot against AOX 1/2 (C). Samples 5 were 1 – molecular weight standard; 2 – leaf; 3 – leaf homogenate; 4 – first chloroplast pellet; 5 - class I chloroplasts; 6 - class II chloroplasts. As for molecular weight standards, 6 7 Fermentas Page Ruler Prestained Protein SM0671 was used. Marks are: triangle - RbcL; 8 circle - AOX 1/2; square - apoLhcII. Lanes on protein gels and immunoblots were loaded 9 with 20 µg solubilised protein except sample (3), where the lane was loaded with 20 µg solubilised proteins over the bovine serum albumin (at ~66 kDa) originating from the 10 11 isolating buffer.

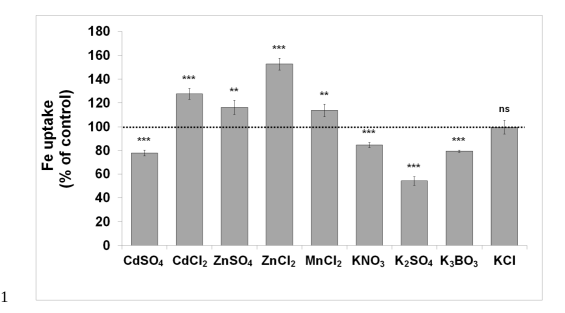
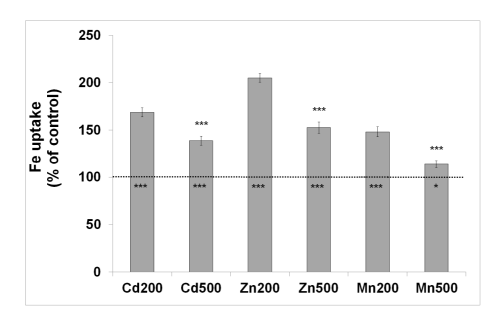
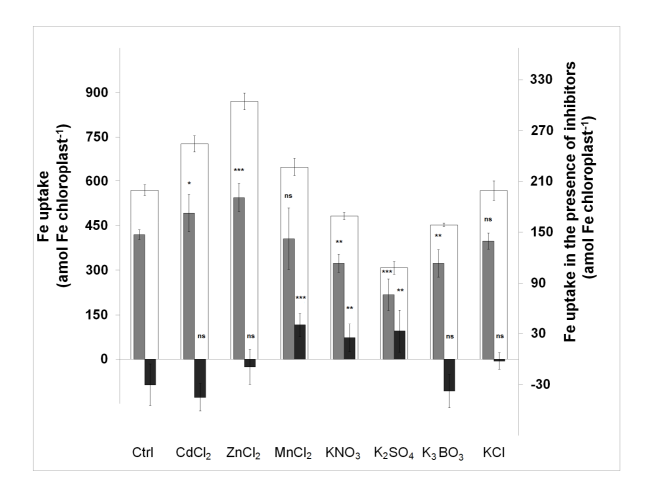


Figure 2 Chloroplast Fe uptake in the presence of inorganic salts at 500 μ M concentrations during a 30-min incubation period in the light. The chloroplast Fe uptake in the control, free of any added inorganic salts, was 570±78 amol Fe chloroplast⁻¹. Statistical differences between each treatment and the control, free of any added inorganic salts, are marked above columns (Student's t-test); ***: *P*<0.001, **: *P*<0.05, ns: not significant. To compare the effects of the different salts, one-way ANOVA was performed with Tukey-Kramer *post-hoc* test, and changes among treatments were found to differ significantly (*P*<0.05).



3 Figure 3 Effect of the transition metals Cd, Zn and Mn, used as chloride salts and at two 4 different concentrations (200 and 500 µM, indicated as Me200 and Me500) on the uptake of 5 Fe by chloroplasts during a 30-min incubation period in the light. The chloroplast Fe uptake in the control, free of any added inorganic salts, was 570±78 amol Fe chloroplast⁻¹. Statistical 6 7 differences between each treatment and the control, free of any added inorganic salts, are marked within each column (Student's t-test); *: P<0.10, ***: P<0.01. Also, significant 8 9 differences in the 500 µM treatments vs. the 200 µM ones (Student's t-test) are marked by 10 arterisks above the columns; ***: P<0.01. To compare the effects of the different salts, one-11 way ANOVA was performed with Tukey-Kramer post-hoc test, and changes among treatments were found to differ significantly (P < 0.05). 12



1

Figure 4 Effect of DCMU (grey columns, right y axis) and CCCP (black columns, right y 2 3 axis) on the Fe uptake of chloroplasts in the presence of inorganic salts at 500 µM 4 concentrations during 30-min incubation in the light compared to the samples containing no 5 inhibitors (white columns, left y axis). Statistical differences between each treatment and the 6 control, free of any added inorganic salts but containing the given inhibitor (grey and black 7 columns) are marked above the column (Student's t-test); ***: P<0.001, **: P<0.05, ns: non 8 significant. To compare the effects of the different salts in the presence of inhibitors, one-way 9 ANOVA was performed with Tukey-Kramer post-hoc test, and Fe uptake in the 10 corresponding DCMU and CCCP treatments were found to differ significantly (P<0.05).

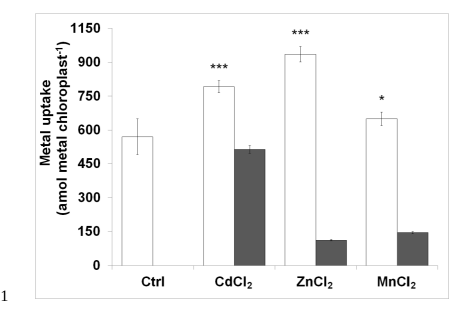


Figure 5 Chloroplast Fe uptake (white columns) and transition metal uptake (black columns)
from a Fe uptake medium without and with 500 μM transition metal cations during a 30-min
incubation period in the light. Significant differences from the control (free of any added
inorganic salts) values (Student's t-test) are marked by asterisks above columns; *: *P*<0.1,
***: *P*<0.01. To compare the uptake of different metals, one-way ANOVA was performed
using a Tukey-Kramer *post-hoc* test, and they were found to differ significantly (P<0.05).

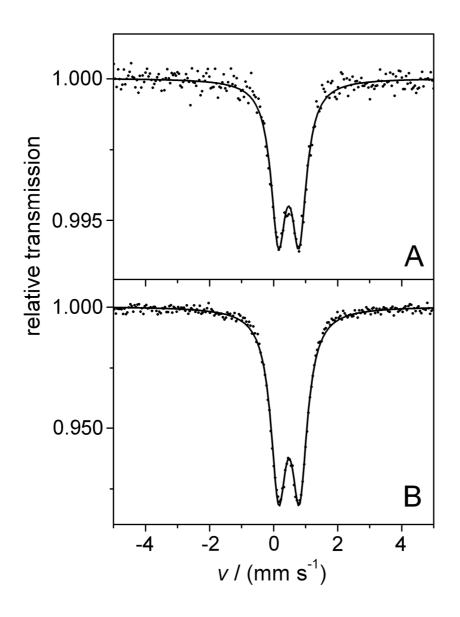




Figure 6 Mössbauer spectra of ${}^{57}\text{Fe}^{3+}$ -citrate 1:1.1 complexes in uptake medium without (A) and with a five times higher amount of K₂SO₄²⁻ over Fe (B). Evaluation and calculations of parameters for the spectral components, including isomer shift, quadrupole splitting and partial resonant absorption areas were calculated and fitted the MOSSWIN code.

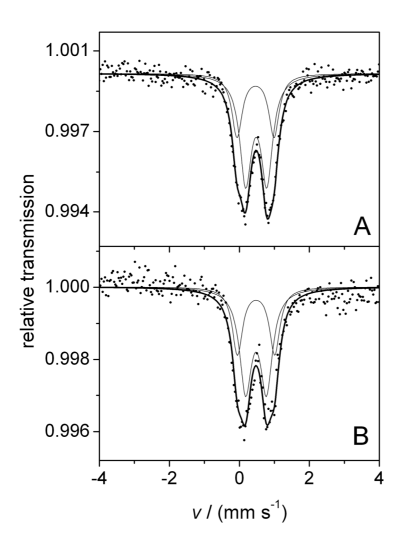


Figure 7 Mössbauer spectra of chloroplasts after 30-min incubation in the light in Fe uptake assay medium , in the absence (A) and in the presence (B) of 500 μ M ZnCl₂. Evaluation and calculations of parameters for the spectral components, including isomer shift, quadrupole splitting and partial resonant absorption areas were calculated and fitted the MOSSWIN code.