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New insights into the mechanism of iron transport through the bacterial Ftr-system present in pathogens

Running title: Copper and iron-bound structure of FtrA-P19

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Abbreviations: *Rubrivivax gelatinosus*, Rg; EPR, electron paramagnetic resonance; WT, wild-type; RMSD, root mean square deviation; PDB, protein data bank; EDTA, ethylenediaminetetraacetic acid; ITC, isothermal titration microcalorimetry; DSC, Differential scanning calorimetry.

Abstract

Iron is an essential nutrient in bacteria. Its ferrous form, mostly present in low oxygen and acidic pH environments, can be imported using the specific Ftr-type transport system, which encompasses the conserved FtrABCD system found in pathogenic bacteria such as *Bordetella*, *Brucella* and *Burkholderia*. **The non-pathogenicity and versatile metabolism of *Rubrivivax gelatinosus* make it an ideal model to study the FtrABCD system.** Here, we report **a new aspect of its regulation** and the role of the periplasmic proteins FtrA and FtrB using *in vivo* and *in vitro* approaches. We investigated the metal binding mode and redox state of copper and iron to FtrA by crystallography and biophysical methods. An ‘as isolated’ FtrA protein from the bacterial periplasm contained a copper ion (Cu^+) identified by electron paramagnetic resonance (EPR). Copper is coordinated by four conserved side chains (His and Met) in the primary metal site. Structural analysis of *R. gelatinosus* FtrA and FtrA homologues revealed that copper binding induces a rearrangement of the His95 imidazole ring, releasing thereafter space, as well as both Asp45 and Asp92 side chains, for iron binding in the secondary metal site. EPR highlighted that FtrA can oxidize the bound ferrous ion into the ferric form by reducing the bound Cu^{2+} into Cu^+ , both metal sites being separated by 7 Å. Finally, we showed that FtrB binds iron and not copper. These results provide new insights into the mechanism of ferrous iron utilization by the conserved FtrABCD iron transporter for which we propose a new functional model.

Introduction

Bacteria have evolved various types of iron acquisition systems. In aerobic environments, iron is mostly found under its ferric form (Fe^{3+}) and imported by extracellular siderophores or iron-binding proteins produced by bacterial cells. The iron-loaded siderophores are internalized into the periplasm by a TonB-dependent transport system [1–3], and **the ferric iron is imported** into the cytoplasm by an ABC-type transporter such as the FbpABC system [4,5]. In low oxygen and acidic pH environments, iron is rather under its ferrous form (Fe^{2+}). Unlike the ferric iron, it is admitted that the ferrous iron is directly imported into the periplasm and uses different transport systems such as the ubiquitous FeoAB and the Ftr-type transporters [6].

The common element to all Ftr-type ferrous iron uptake systems is the conserved permease, named EfeU and FetM in *E. coli*, cFtr in *Campylobacter jejuni* and FtrC in *Bordetella*, *Brucella* and *Burkholderia* species [7–15]. This permease is also present in eukaryotes, namely Ftr1P in yeast, which is the only well characterized permease so far [16–18]. Besides this permease, the components of the Ftr systems vary depending on the studied organism. In *E. coli* O157:H7 strain and *Bacillus subtilis*, in addition to the EfeU permease, the EfeUOB system comprises two periplasmic proteins EfeO (cupredoxine-I type oxydase) and EfeB [19,20]. In *Bordetella*, *Brucella* and *Burkholderia spp*, the FtrABCD system consists of two periplasmic proteins FtrA and FtrB (putative cupredoxin-II type oxidase), a polyferredoxin FtrD and the permease FtrC [10,11,13]. In *Campylobacter jejuni* (Cj), the Ftr system includes only one periplasmic protein denoted P19 (FtrA homologue) equivalent to FetP in the uropathogenic *E. coli* strain F11 in addition to the permease [9,21]. In contrast to the well-known Ftr system in yeast where the multicopper ferroxidase Fet3p oxidizes the ferrous iron into ferric iron prior to its import through Ftr1P [22], no ferroxidase activity has been shown in the bacterial Ftr system yet. Instead, it has been proposed for the FrABCD system that the putative cupredoxin-II type oxidase FtrB [20] would oxidize Fe^{2+} bound to FtrA into Fe^{3+} [8,10].

The bacterium *Rubrivivax gelatinosus* from *Burkholderiales* family possesses the Ftr-type system encoding operon annotated *frAPBCD* (Fig. 1A and 1B) [23]. The periplasmic protein named here RgFtrA, shares around 60% sequence identity with both *Brucella melitensis* and *Bordetella pertussis*

FrtA and 50% sequence identity with CjP19 and EcFetP. Its non-pathogenicity and versatile energetic metabolism make this bacterium an interesting model for studying the FtrABCD transport system found in pathogenic bacteria. Moreover, *R. gelatinosus* has the ability to grow under different oxygen concentrations. In the presence of oxygen, this bacterium respire, whereas under anaerobic condition with light, it grows by anoxygenic photosynthesis [24]. In addition to the iron system FtrAPBCD, *R. gelatinosus* possesses an effective ferric transport system FbpABC [25] and in its genome, genes encoding the ferrous iron system FeoAB are also present. Expression of iron acquisition genes is usually regulated by the available iron in the medium: in the presence of iron, the transcriptional regulator Fur-Fe²⁺ complex represses genes encoding different iron uptake systems such as those coding for the Ftr1P-type system [7,26], whereas under iron depletion, Fur is inactive and iron uptake genes are expressed. Oxygen concentration is another factor involved in the regulation of some ferrous iron transporters through the [4Fe-4S] cluster of the transcriptional regulator FNR. It was shown that under low oxygen concentration, FNR activates the expression of *feo* genes in *E. coli* and *Shigella flexneri* [27,28].

In this work, we focused on the two periplasmic proteins FtrA and FtrB from the *R. gelatinosus* FtrAPBCD system. We provide evidence showing the regulation of *ftrAPBCD* operon by iron and by oxygen *via* FNR global regulator. We investigated the structural and biochemical properties of RgFtrA towards both copper and iron metal sites, their redox state and their binding mode. Structural comparison of high-resolution RgFtrA structures (apo, copper as well as copper and iron forms) revealed a key histidine residue, strictly conserved, which explains the sequential binding of Cu then Fe as first suggested for CjP19 [29]. We demonstrate that RgFtrA has a ferrous iron oxydase activity and that the putative cupredoxin-II type oxidase RgFtrB binds iron and not copper as a starting point for further investigations of its putative role in Fe³⁺ transport. Overall, our data revisit a previous proposed model by providing new insights into the mode of function of the conserved FtrABCD transporters.

Results

***ftrAPBCD* operon and its regulation.** The *ftr* operon (*ftrAPBCD*) of *R. gelatinosus* (Fig. 1B) is quite similar to that of *Brucella*, *Bordetella* and *Burkholderia* species except the presence of an additional gene encoding the outer membrane porin FtrP [10,11]. Analyses of the promoter sequence of *ftrA* gene with PRODORIC database (<https://www.prodoric.de/vfp/>) [30] revealed, in addition to the presence of a Fur box, an FNR binding site located upstream of the Fur box, suggesting that the expression of the *ftrAPBCD* operon could be under the dual transcriptional control by the iron-sensing Fur and the oxygen-sensing FNR regulators. To verify this hypothesis, we analyzed the transcription of *ftrAPBCD* genes of the wild-type (WT) and the Δfnr mutant, lacking FNR regulator, by semi-quantitative RT-PCR under three oxygen conditions (aerobic (AE), microaerobic (MA) and anaerobic photosynthesis (PS) conditions), in malate medium with iron or in iron-depleted malate medium. First, because the cDNA was synthesized using an *ftrD* reverse primer, this experiment confirmed that *ftrAPBCD* is an operon, since using this *ftrD*-amplified cDNA, we were able to amplify the distal gene *ftrA* (Fig. 1B-C). On the other hand, the RT-PCR data clearly showed that expression of the *ftrAPBCD* operon was induced under iron deficiency conditions. Moreover, it also showed that the expression of *ftrAPBCD* operon was oxygen dependent. Indeed, under iron depleted conditions, in the WT, expression of *ftrA* and *ftrD* genes was higher under microaerobic and anaerobic conditions than under aerobic condition (Fig. 1C) strongly suggesting a regulation by FNR. This hypothesis was validated by analyzing *ftrAPBCD* expression in the Δfnr mutant background. Δfnr mutant does not grow under strict photosynthesis conditions, we therefore compared the *ftrAPBCD* expression profile under aerobic *versus* microaerobic conditions. As shown in Figure 1C, expression of *ftrAPBCD* in the Δfnr mutant was no longer induced under microaerobic condition, in sharp contrast to the WT. In the WT, FtrA was more abundant under iron depleted condition as expected, while in Δfnr mutant, the abundance of FtrA was drastically reduced (data not shown). The expression level depends both on iron availability, very likely *via* Fur and on oxygen concentration *via* FNR regulator.

RgFtrA is involved in iron transport. For a previous study on iron import and resistance to heavy metals, we have constructed the $\Delta ftrATp$ and $fpbAKm$ single defective mutants as well as the $fpbAKm-\Delta ftrATp$ (ffa) double mutant. $fbpA$ codes for the periplasmic iron binding protein of the ferric iron ABC transporter FbpABC system [25]. In the malate medium containing iron, the same growth was observed for all strains under microaerobic condition regardless of the medium pH (Fig. S1A and S1B). In contrast, in iron depleted malate medium at pH 6 under microaerobic condition (Fig. 2), the $\Delta ftrATp$ mutant grew with two phases: a lag phase of 18 hours before regaining a growth rate similar to the WT whereas the $fpbAKm$ mutant grew quite similarly to the WT. At pH 7, no phenotype was observed for $\Delta ftrATp$ mutant and $fpbAKm$ growth was slowed down and did not grow as well as the WT.

The most important growth inhibition was observed with the double mutant ffa at both pHs revealing that Ftr and Fbp are active and complementary. Moreover, no other iron transporter seems to compensate the absence of both Ftr and Fbp systems, suggesting that in our tested conditions, the ubiquitous $feoAB$ system was not functional. To verify it, we constructed and observed the phenotype of the $feoBKm$ single mutant and the $feoKm-\Delta fbpATp$ (ffO) double mutant (Fig. S1). Because $feoBKm$ behaved as the WT and ffO showed a phenotype similar to $fpbAKm$, the Feo system seems non-functional in *R. gelatinosus*.

Structures of apo His-tag RgFtrA and RgFtrA, and comparison with apo structures of homologues. To obtain the soluble FtrA of *R. gelatinosus* in *E. coli* cytoplasm, the $ftrA$ gene was synthesized without the signal sequence encoding the first 23 amino acids, which serve for the translocation to the bacterial periplasm. An $ftrA$ gene coding for the His-tag RgFtrA protein was also synthesized with no signal peptide and with additional 11 amino acids including 6 histidine residues at the C-terminus. Both proteins (RgFtrA and His-tag RgFtrA) were purified from the cytoplasm of *E. coli* as a dimer in solution, according to gel filtration measurements. The numbering used for the description of residues corresponds to the mature protein of 162 amino acids (without the signal peptide) in order to simplify its comparison with the well studied P19 homologue from *C. jejuni* [9,29].

Using thermal denaturation experiments by differential scanning calorimetry (DSC), adding 0.5 mM or 1 mM ethylenediaminetetraacetic acid (EDTA, a strong chelating agent) to RgFtrA and His-tag RgFtrA had no effect on the T_m of 52.1 °C and 53.7 °C, respectively (Fig. S2), indicating that both proteins were in an apo form, confirmed by X-ray fluorescence scans on RgFtrA crystals at synchrotron beamline. They both crystallized under the same conditions leading to isomorphous crystals diffracting at 1.4 Å and 1.23 Å, respectively, and containing a very similar homodimer in their asymmetric unit with an average root mean square deviation (RMSD) of 0.12 Å for 312 C α atoms (Fig. 3A, Tables 1 and 2). A full RgFtrA monomer was built in the electron density maps, while for the three other monomers (RgFtrA and His-tag RgFtrA), between 154 and 157 amino acids were built because their C-terminus including the His-tag was not visible. FtrA forms a recognizable dimer like a prolate spheroid with an interface of 1863 Å² per subunit corresponding to an average of 20.5% of the total surface area of each monomer and 50 amino acids per monomer involved in 18 hydrogen bonds and 6 salt bridges. The dimeric interface, which encompasses two β sheets (β_6 and β_9) and 5 loops, is characteristic of biological interactions [31].

RgFtrA resembles the *C. jejuni* P19 (CjP19; [9,29]) and the uropathogenic *E. coli* strain F11 FetP [32]. Indeed, SSM-EBI (<http://www.ebi.ac.uk/msd-srv/ssm>) reports RMSD of 0.85-1.13 Å between RgFtrA and the numerous structures of CjP19 over 146-148 C α atoms with 55% sequence identity, and RMSD of 0.91-1.13 Å with the structures of FetP over 146-147 residues with 52% sequence identity. The structures of RgFtrA and His-tag RgFtrA confirmed the absence of any metal. Instead, a water molecule is located in the primary metal binding site about 1.3 Å from the expected copper position represented by a Zn ion modelled at 10% occupancy in CjP19 having led the authors to describe this Zn-CjP19 structure as an apo form (PDB 3LZN). The water molecule makes hydrogen bonds with the side chains of three conserved residues: His41 (His42/His44 in CjP19/FetP), Glu43 (Glu44/Glu46 in CjP19/FetP) and His131 (His132/His127 in CjP19/FetP) (Fig. 3A). Remarkably, comparison of the apo structures of RgFtrA, CjP19 (PDB 3LZN) and EcFetP (PDB 3NRQ) revealed a major local difference concerning the conserved copper ligand, His95. Indeed, in RgFtrA only, the imidazole ring of His95 is rotated by about

60° placing the NE2 atom 4.4 Å away from the copper binding site (Fig. 3B). In this position, His95 interacts with the side chains of Asp45 and Asp92 (Fig. 3B).

The primary site of RgFtrA displays a strong preference for copper binding *in vitro*. Using thermal denaturation experiments by DSC, the primary metal site was investigated on the RgFtrA (30 μM) treated with EDTA to remove any metal traces (Table 3 and Fig. S3). DCS analyses revealed that Fe²⁺, Fe³⁺, and Mg²⁺ did not alter the *T_m* whereas Mn²⁺ has a modest effect. Significant increase of the *T_m* by ~6-7°C was observed with Co²⁺ and Zn²⁺, by ~10°C with Cd²⁺ and by ~20°C with Ni²⁺. Drastic shifts > 23°C, were observed for Cu⁺ and Cu²⁺. The same approach by nano-differential scanning fluorimetry (nanoDSF) on Tycho NT.6 instrument led to similar results as DSC (Fig. S4). Performing these experiments with the reductant agent Tris(2-carboxyethyl)phosphine (TCEP) added to ferrous iron gave the same result as ferrous iron alone without TCEP (data not shown).

Binding of Cu²⁺, Ni²⁺ and Zn²⁺ to the RgFtrA treated with EDTA was further explored using ITC (Fig. 4). As expected, the best affinity was measured with Cu²⁺ with a *K_D* value of 0.6 μM. Microcalorimetry data confirmed the 1:1 binding stoichiometry revealing a negative enthalpy change.

The secondary metal binding site of RgFtrA binds iron *in vitro*. We explored the second metal binding site by DSC using the RgFtrA treated with EDTA and loaded with equimolar CuCl₂ at acidic (6.5) and basic (8) pH values (Table 3 and Fig. S5). DCS analyses clearly revealed that RgFtrA-Cu has a strong preference for iron with large *T_m* shifts of ~15°C at pH 6.5 (Table 3). Mn²⁺ has a modest effect and no shift was observed with Mg²⁺ or Co²⁺. Similar results were obtained at pH 8 with *T_m* shifts less pronounced (Fig. S5). NanoDSF on Tycho NT.6 instrument confirmed DSC analyses (Fig. S6).

Binding of Fe²⁺ to the RgFtrA treated with EDTA and loaded with equimolar CuCl₂ was further explored using ITC at pH 6.5 and 8 (Fig. 5). The similar measured *K_D* values of 3.3 and 5.8 μM indicated that pH has not real effect on iron binding. The microcalorimetry data confirmed the 1:1 binding stoichiometry and revealed a negative enthalpy change.

Primary metal binding site

Cu⁺ bound. A Strep-tag RgFtrA protein was synthesized with a strep tag of 8 amino acids at the C-terminus (the signal peptide was conserved). The strep-tag *ftrA* synthesized gene under its own regulated promoter was cloned in a replicative plasmid pBBR1-MCS2 and expressed in the $\Delta ftrATp$ mutant. The 'as-isolated' Strep-tag RgFtrA was expressed and purified from the periplasm of *R. gelatinosus* to determine the presence of a physiologic metal if any and its redox state. The first batch of purified protein crystallized in conditions different from those used for the His-tag RgFtrA and RgFtrA expressed in the cytoplasm of *E. coli* (Tables 1 and 2). DSC experiments showed that the first batch of 'as-isolated' Strep-tag RgFtrA was already loaded with metal because addition of EDTA resulted in a significant decrease of the T_m by $\sim 9^\circ\text{C}$ to reach a value closer to that observed for the apo His-tag RgFtrA (Fig. 6A). In addition, an X-ray fluorescence spectrum on crystals measured at synchrotron beamline detected only one type of metal identified as being copper. Therefore, a SAD dataset was collected at the copper anomalous signal peak at 1.3 Å resolution (Table 1). The electron density maps revealed the presence of two very well defined copper ions located at the oligomeric interface (one copper per subunit, Fig. 3C). Their low temperature factors (B -factor) of 11.8/11.2 Å² in chain A/chain B imply full occupancy. Their coordination is tetrahedral with distances of 2.02-2.37 Å with the conserved protein side chains of His41, Met88, His95 and His131' from the neighbouring subunit (Table 4). The EPR spectrum recorded on the same batch of protein showed no detectable metal signal (Fig. 6B). In contrast, upon addition of an HCl drop ($\sim 18\%$), a Cu²⁺ signal was detected (Fig. 6B), demonstrating that the 'as-isolated' protein used for this study binds copper in its Cu⁺ redox state. However, more batches of 'as-isolated' Strep-tag RgFtrA were prepared and EPR on these different batches showed that the Strep-tag RgFtrA could be isolated with trace of Cu²⁺ (not shown). Superposition of the structures of Strep-tag RgFtrA-Cu⁺ and apo RgFtrA showed that in the apo RgFtrA, the water molecule in the primary metal binding site is at 1.8 Å from the bound Cu⁺ ion in the Strep-tag RgFtrA-Cu⁺, near the His95 side chain, which interacts with the Cu⁺ ion (Fig. 3D). The position of this water molecule was possible because His95 side chain was out of the copper site as described above in the paragraph of the apo structures.

Cu²⁺ bound. RgFtrA (expressed and purified from the cytoplasm of *E. coli*) loaded with equimolar CuCl₂ and 1 mM FeSO₄ crystallized in conditions containing at least 50 mM MgCl₂ (Table 2). Removing MgCl₂ prevented crystals growth. Two crystals denoted X1 and X2 grown with 100 and 50 mM MgCl₂,

respectively, were collected. X-ray fluorescence spectrum measured from the crystals did not indicate the presence of iron and data collected at the iron anomalous signal peak lacked anomalous signal. A full occupancy Cu^{2+} was modelled in each subunit in both structures taking into account that the protein was co-crystallized with CuCl_2 . The Cu^{2+} occupies the primary metal site as expected making the same protein contacts compared with the reducing copper form (Table 4).

Ni^{2+} bound. Because nickel ions have a stability effect on RgFtrA as strong as copper ions as shown by DSC, we investigated their binding mode. In the presence of 20 mM NiSO_4 , the His-tag RgFtrA crystallized in the same conditions as the apo His-tag RgFtrA and RgFtrA leading to isomorphous crystals (Table 2). The presence of nickel in the crystals was confirmed by X-ray fluorescence and data were collected at the nickel anomalous edge. The electron density maps revealed the presence of nickel in the primary metal site of only one subunit despite a protein:nickel ratio of 1:200. The side chain of His95 takes two alternative positions with similar occupancies, one with the imidazole ring bound to the metal, the other pointing away from it as observed in the apo form (Fig. 7A). The Ni^{2+} was thus refined with the assumed 50% metal occupancy resulting in a B-factor of 33.8 \AA^2 . It interacts with His41, His95 and His131' from the neighbouring subunit and makes a very short contact with His95 (Fig. 7A and Table 4). The Ni^{2+} , which is 0.6 \AA from the Cu^+ ion is close to Glu43 carboxylate and far away from Met88 (Fig. 7B and Table 4). The other subunit of the His-tag RgFtrA-Ni complex contains a water molecule as observed in the apo form (Fig. 7C).

Secondary metal binding site

Fe^{3+} bound. Crystals of 'as-isolated' Strep-tag RgFtrA (Cu^+ bound) purified from the periplasm of *R. gelatinosus* were soaked with 0.1 M FeCl_3 . Crystals turned orange suggesting ferric ion incorporation. An X-ray fluorescence spectrum measured from the crystal detected the presence of iron. A SAD dataset was collected at the iron anomalous signal peak at 2.75 \AA resolution (Table 1). The asymmetric unit contains one dimer and each subunit binds an iron ion in the secondary metal site adjacent to the copper site located at the protein surface (Fig. 8A and 8B). Fe^{3+} was modelled taking into account that the crystal was coloured before X-ray exposure. Each Fe^{3+} binds to the Glu2, Glu43 and Asp92 carboxylates

(Fe-O distances ranging from 1.7 to 2.3 Å) showing the shortest interaction with Asp92 (Table 5 and Fig. 8A). The two metal ions (Cu^+ and Fe^{3+}) are separated by ~ 7 Å (Fig. 8C).

Mg²⁺ bound. Co-crystallization with 4 mM FeSO_4 of the 'as-isolated' Strep-tag RgFtrA (Cu^+ bound) purified from the periplasm of *R. gelatinosus* led to crystals different from those obtained with the protein without FeSO_4 likely due to the presence of a very high concentration of 200 mM MgCl_2 in the crystallization conditions (Table 1). The asymmetric unit contains three very similar dimers, each subunit showing full occupancy of the primary metal site with essentially the same protein- Cu^+ interactions (Table 4). A magnesium ion was modelled in all secondary metal site subunits being ~ 7 Å apart from the Cu^+ ion (Table 5 and Fig. 8D). Chains A, B and F show an octahedral coordination with Glu43, Asp92 and four water molecules at distances of 1.92-2.4 Å compatible with a full ion occupancy (Table 5 and Fig. 8D). The Glu2 carboxylate occupies a different position in the magnesium-ligated complex *versus* the iron-ligated complex, where it makes water-mediated magnesium interactions instead of direct metal interaction (Fig. 8E). In the structures of RgFtrA denoted X1 and X2 (Table 2) obtained from crystallization conditions with much lower concentration of MgCl_2 (100 and 50 mM, respectively), the electron density maps suggest the presence of a Mg^{2+} in each secondary metal site due to a short contact with Asp92 side chain (Table 5). Each Mg^{2+} ion seems well coordinated in both X1 and X2 structures compared with those in the structure obtained with 200 mM MgCl_2 .

The bound-copper RgFtrA can oxidize the ferrous iron into ferric iron. The “as purified” RgFtrA treated with EDTA or not exhibited the EPR spectrum shown in Fig. 9 (black spectrum) indicating that the protein was devoid of EPR detectable metals. Upon reconstitution with Cu^{2+} (equimolar), the protein exhibited a Cu^{2+} EPR spectrum (red spectrum). Then, Fe^{2+} was added in a stoichiometric amount directly in the EPR tube. Similar results were obtained with either FeCl_2 or FeSO_4 (not shown). After this Fe^{2+} addition, the Cu^{2+} EPR signal decreased significantly and a spin $S = 5/2$ Fe^{3+} EPR signal between ~ 600 gauss and ~ 2000 gauss appeared (blue spectrum). This experiment showed that the bound Cu^{2+} has a redox potential high enough for oxidizing the Fe^{2+} into Fe^{3+} . After further addition of Fe^{2+} to reach a concentration twice that of the protein, the amplitude of the Fe^{3+} and Cu^{2+} signals remained unchanged

(green spectrum) showing that there was no detectable auto oxidation of the Fe^{2+} in these conditions. Moreover, this suggests that this was not the Fe^{2+} present in the bulk, which was responsible for the Cu^{2+} reduction bound to the protein. The copper reduction more likely occurs after the binding of Fe^{2+} in its site. The distance of 7 Å between the copper and iron binding sites given by the crystal structures is perfectly compatible with such an electron transfer, see for example [33]. With a redox potential similar for the $\text{Cu}^{2+}/\text{Cu}^+$ and the $\text{Fe}^{3+}/\text{Fe}^{2+}$ couples, and assuming that both the copper binding site and iron binding site are fully occupied in the EPR conditions, we would expect a two-fold decrease of the amplitude of the Cu^{2+} signal. The remaining Cu^{2+} signal amplitude was in fact closer to 30 % of the signal prior to Fe^{2+} addition. The binding to the protein increases the E_m of the $\text{Cu}^{2+}/\text{Cu}^+$ couple and decreases the E_m of the $\text{Fe}^{3+}/\text{Fe}^{2+}$ couple when compared to the E_m of these metals in solution so that the E_m of the $\text{Cu}^{2+}/\text{Cu}^+$ couple becomes slightly above that of the $\text{Fe}^{3+}/\text{Fe}^{2+}$ couple.

The putative cupredoxin-II type oxidase FtrB binds iron and not copper. To investigate the role of RgFtrB, the single defective mutant ΔftrBTp and the double defective mutant $\text{fbpAKm}\Delta\text{ftrBTp}$ (*ffb*) were constructed. Growth assays with ΔftrBTp showed exactly the same phenotype as ΔftrATp demonstrating that FtrB and FtrA share a common role in iron transport (Fig. S1). Using the mature RgFtrB (30 μM) expressed and purified from *E. coli* cytoplasm and treated with EDTA to remove any metal traces, DSC analyses revealed that the presence of Cu^{2+} did not alter the T_m in contrast to Fe^{2+} (and Fe^{3+}) which drastically impacted the T_m by ~ 24.3°C (Fig. 10). The nano-differential scanning fluorimetry (nanoDSF) could not be performed due to the lack of tryptophan in RgFtrB. We showed that FtrB could bind iron and no copper.

Discussion

Regulation of iron transporter genes

Bacteria respond to iron-limiting conditions by inducing the expression of proteins involved in iron acquisition to capture and import the sparsely iron present in their environment. The activation of iron transporters relies on the availability of iron and its redox state and thus also on the oxygen concentration. In this work, we show that transcriptional regulator FNR (sensitive to oxygen) regulates the *ftrAPBCD* genes of *R. gelatinosus* and very likely in other bacteria. Indeed, we have identified an FNR binding site upstream of the Fur box in the promoter region of genes encoding a Ftr1P system in all analyzed bacteria including *E. coli*, *C. jejuni*, *B. pertussis*, *P. syringae*, *P. palustris* or *B. subtilis* (Fig. 11). It should be noted that in *B. abortus* (Fig. 11), which lacks Fur, the FNR binding site overlaps the binding site denoted ICE of the iron response transcriptional regulator Irr, involved in the regulation of the FtrABCD operon [11]. Overall, this strongly suggests that *ftr1P* operon expression could be under the dual transcriptional control by the iron-sensing Fur or Irr and the oxygen-sensing FNR regulators.

R. gelatinosus possesses three iron transporter systems in its genome, the Fbp, Feo and Ftr systems. The FbpABC systems are known to transport ferric iron while the FeoAB and Ftr1P systems bind ferrous iron. Our growth analyses in iron-depleted media, at acidic and neutral pH suggested that the Ftr iron transporter is required at acidic pH and microaerobic condition. This result is in agreement with the function of each transporter (Fe^{2+} for Ftr and Fe^{3+} for Fpb) because acidic pH and low oxygen favor the ferrous iron form.

FNR is known to activate the transcription of the ubiquitous iron transporter FeoABC [27]. A *feoAB* operon is present in *R. gelatinosus*, nonetheless, the growth inhibition phenotype of the double mutant *ffa* suggested that the Feo system is not active. It should be noted that in other bacteria possessing the FtrABCD system, no functional Feo system has been identified so far. This is the case for *Bordetella*, *Brucella* and for many *Burkholderia* [10,11,14].

A conserved histidine orchestrates the sequential metal binding order in FtrA

This work reveals the structural basis of FtrA displaying two conserved metal binding sites referred as primary and secondary metal sites for all orthologues. FtrA/P19 proteins are all dimeric and the two

adjacent binding metal sites are formed at the interface between the two monomers. When present, the physiological metal ion bound to the primary metal site is copper as demonstrated by EPR experiments on the ‘as isolated’ Strep-tag RgFtrA purified from the periplasm of *R. gelatinosus*. Therefore, the purified ‘as isolated’ Strep-tag RgFtrA could remain loaded with endogenous reduced Cu^+ ion although the overall purification process was done under aerobic conditions. This indicates that the four copper-coordinating amino acids (His41, Met88, His95, His131) forming a typical tetrahedral geometry have a remarkable ability to stabilize Cu^+ and protect it from natural oxidation. Of particular note is that while the copper ion is not required for RgFtrA stability and folding (as demonstrated by the purified apo form and crystal structures), DSC, nanoDSF and affinity measurement all showed copper as the best ligand of the primary metal site, in agreement with the physiological metal.

A remarkable outcome of this work is demonstrating that the primary metal site is not preformed. Our apo form versus copper-ligated structures clearly revealed the important role of the side chain of His95. Indeed, its imidazole ring plays a regulatory role in the sequential order of metal binding. In the absence of any metal, His95 interacts with both carboxylates of Asp45 and Asp92, which both form the secondary metal site. Copper binding induces a rotation of His95 side chain, which seals the primary metal binding site, by interacting tightly with the copper ion thus leaving the space for iron to bind to the secondary metal site. The ferrous iron interacts directly with the side chains of Glu2, Glu43 and Asp92 and through water molecules with Asp45. Glu43 is $\sim 4 \text{ \AA}$ distant from the copper and thus makes no direct interaction with this metal (Table 4). No obvious conformational change was observed for Glu43 in our eight-crystal structures (apo form *versus* metal complexes), however only a slight displacement could bring its side chain closer to copper site as observed in CjP19 structures [29]. The structure-based sequence alignment of RgFtrA with homologues such as CjP19 (PDB 3LZN) and EcFetP (PDB 3NRP) indicates that all residues involved in metal binding are conserved (Fig. 12A). Copper coordination occurs through His41/His42/H44, His95/His95/His97, His131/His132/H127 and Met88/Met88/Met90 in RgFtrA/CjP19/EcFetP and iron coordination via 3 side chains (Glu2/Glu3/Glu4, Glu43/Glu44/Glu46 and Asp92/Asp92/Asp94). The structures of RgFtrA and CjP19 (PDB 1I0V) in complex with Cu and Fe in oxidizing condition superimpose with a RMSD of 0.8 \AA for all $\text{C}\alpha$ atoms.

Their copper and iron bind at the same position with the protein side chains around each metal overlapping and making similar metal contacts (Fig. 12B). Overall, our structures point out that the apo structure of CjP19 (PDB 3LZN) should maybe be reconsidered as not an apo structure.

The structure of RgFtrA-Cu⁺Ni²⁺ revealed that the nickel binds only the three His of the four side chains of the primary binding site, suggesting that Met88 is specific for copper binding and positioning, in line with the results of the corresponding variant M88I on CjP19 (Chan et al 2020). Although DSC and nanoDSF showed that Mg²⁺ is not a ligand of RgFtrA, excess of MgCl₂ in the crystallization conditions lead to Mg²⁺ bound in the iron binding site. Thus, Mg²⁺ such as Mn²⁺ act as iron mimics. The metal coordination by Glu2/Glu3 observed in iron-bound structures of RgFtrA and CjP19, respectively, was not observed in the Mn²⁺-CjP19 (PDBs 3LZR and ELZQ) and our Mg²⁺-RgFtrA structures. This emphasizes the previous assumption that Glu2/Glu3 participates to iron selectivity and has an important role. DSC and nanoDSF confirmed iron as the preferred metal in agreement with the physiological ligand of the secondary metal site of FtrA-P19.

FtrABCD iron transport system

The four-component Ftr system FtrABCD is highly conserved among *Brucella*, *Bordetella* and *Burkholderia* gathering different species including *R. gelatinosus* [10,11,14,15] and suggesting similar functional mode for these iron transporters. Until now, no or few biochemical data is available on the four proteins of this Ftr system. As mentioned in the introduction, the best-characterized Ftr system is from yeast, composed of the permease Ftr1p and the multicopper ferroxidase Fet3p. Fet3p captures Fe²⁺ by conserved acidic residues and oxidizes it by a Cu²⁺ in Type-1 site defined by conserved HisHisCys residues, resulting in Fe³⁺ production. Fet3p interacts with Ftr1p and the iron oxidation occurs concomitantly to the transport of ferric iron [22]. The yeast Ftr model for the bacterial Ftr systems used as a reference and has not yet been validated in bacteria due to the lack of data on iron oxidation. Here for the first time, we have experimentally identified FtrA as a protein with ferrous iron oxydase activity able to oxidize the ferrous ion into ferric iron in a Cu dependent manner. Therefore, although FtrA contains only one atypical copper site (not a Type-1 site) and has no resemblance with Fet3p, FtrA seems to play a similar function as Fet3p. Glu43/Glu44 in RgFtrA/CjP19 located between the copper and iron

sites as described above has been proposed to facilitate electron transfer between the two metals [29]. For the bacterial two-component Ftr systems with P19/FetP and cFtr/fetM in *C. jejuni/E. coli*, based on structural data, we predict that CjP19 and EcFetP would also display a ferrous iron oxydase activity similar to FtrA.

FtrB until now was considered as a new cupredoxin based on sequence alignment, conservation of several amino acids and structural model analyses. Nonetheless, as highlighted recently [15], no biochemical analyses have so far confirmed copper binding in FtrB. Instead, FtrB has been suggested as a protein with a key function in copper-dependent iron oxidation and a model was proposed in which FtrA delivers Fe^{2+} to FtrB, which will then oxidize the iron and transfer it to the FtrC permease [8,10]. We have shown that RgFtrB does not bind copper but iron likely through conserved acidic residues at positions 42, 44, 51 and 76. Carboxylate at position 77 in *R. gelatinosus* is also conserved being a Glu in *R. gelatinosus* and an Asp in *Brucella* and *Bordetella* FtrB homologues based on multiple-sequence alignment (Fig. 13A). AlphaFold2 prediction model (ColaFold on google) showed that the conserved acidic residues are surface-exposed and could form an iron binding site (Fig. 13B). The FtrB protein, which plays an important role in the Ftr system, since ΔfirBTp and ΔfirATp mutants share the same phenotype, is very likely not a cupredoxin. We suggest an iron (Fe^{3+}) chaperon activity for FtrB.

In the two-component Ftr systems with P19/FetP (FtrA homologues) and cFtr/fetM in *C. jejuni/E. coli*, no cupredoxin has been identified yet. However, a large extra-periplasmic domain is present in FtrC from *C. jejuni* and *E. coli* and seems to interact with P19/FetP [29]. Based on these observations, we propose an alternative model to the one proposed in *Bordetella* where, first, FtrA is a Cu-dependent ferrous iron oxydase which oxidizes the Fe^{2+} to Fe^{3+} (Fig. 14) and then the oxidized iron is transferred to the permease FtrC through the iron chaperon FtrB. The reduced Cu^+ in FtrA could be regenerated by transferring the harvested electrons to the polyferredoxin FtrD predicted to act as an electron sink, allowing the recycling of electrons from Cu^+ to Cu^{2+} of FtrA. [8,10,11,14].

Conclusion

Our work brings new insights into the functional mode of the highly conserved bacterial four-component FtrABCD ferrous iron transport system through the study of FtrA and FtrB proteins. We highlighted a role of FtrA as a ferrous iron oxidase and FtrB as an iron chaperone, likely linking FtrA and FtrC. Our model functionally resembles the yeast Ftr system composed of Ftr1p-Fet3p. Our structural data showed that the side chain of His95 in FtrA guides the sequential order of metal binding where the primary copper site must be filled first before the iron site. The copper metal ligand of FtrA is thus essential for iron binding and oxidation. More biochemical data are required to validate our new Ftr-system model such as protein interactions.

Materials and methods

Bacterial strains and growth. *R. gelatinosus* was grown at 30°C, in the dark either in aerobically (high oxygenation: 20 ml medium in 250 ml flasks), or microaerobically (low oxygenation: 50 ml medium in 50 ml flasks) conditions or with light under anaerobic conditions (filled tubes) in malate or in malate iron depleted growth medium. *E. coli* was grown overnight at 37°C in LB medium. Antibiotics (50 µg/ml) such as ampicillin, kanamycin (Km), trimethoprim (Tp), were added when appropriate. One liter of malate medium consists of: 30 ml 100g.l⁻¹ malate NH₄, 20 ml of concentrate base (34 mM EDTA, 130 mM KOH, 58 mM MgSO₄, 22 mM CaCl₂, 7.5 µM (NH₄)₆Mo₇O₂₄, 0.4 mM nicotinic acid, 74 µM thiamine, 2 µM Biotin, 356 µM FeSO₄), 20 ml 1M phosphate buffer pH 7, 1g N-Z amine (Merck ; Ref :C7290), 1 ml of metals (0.85 mM EDTA, 0.28 mM ZnSO₄, 9.1 mM MnSO₄, 1.5 mM CuSO₄, 0.85 mM Co(NO₃), 0.46 mM Na₂B₄O₇, 18 mM FeSO₄) and 1g of yeast extract.

Bacterial strains and plasmids are listed in Table S1. Growth under microaerobic conditions of *R. gelatinosus* strains was measured at OD_{680nm} during 40 h using the Tecan Infinite M200 luminometer (Mannerdorf, Switzerland). For the photosynthetic growth, strains were grown with light in filled well plates cover by a sealed membrane.

Gene cloning, plasmid constructions and mutant strains construction. Standard methods were performed according to Sambrook et al. [34]. Primers used are listed in Table S2. To delete *ftrB* gene, two PCR fragments obtained with RG479-RG480KpnI and RG481KpnI-RG482 primers were digested by KpnI and cloned in pGemT-easy leading to pGe- Δ *ftrB*. Then the Tp cartridge was inserted at the KpnI site within Δ *ftrB* of pGe- Δ *ftrB*. The integrative plasmid pGe- Δ *ftrB*Tp was used to inactivate *ftrB* by electroporation in the WT and in the mutant *fbpAKm* in order to construct the single mutant Δ *ftrB*Tp and the double mutant *ffb* (*fbpAKm*- Δ *ftrB*Tp) of *R. gelatinosus*. KS-*feoBKm* was obtained from DNA isolated from *copA:: Ω -feoB::Tn5* (unpublished data) digested by PstI and cloned into Bluescript KS+. Insertion of the transposon in *feoB* in the *copA:: Ω* mutant [35] makes the strain more resistant to copper than the single mutant *copA:: Ω* (unpublished data). This plasmid was used to inactivate *feoB* by electroporation in the WT. To construct KS-*fbpATp*, we replaced the Km cartridge of KS-*fbpAKm* by the Tp cartridge by digesting the plasmid and p34S-TP with SmaI.

We ligated them and obtained KS-*fbpATp*. This plasmid was used to transform the *feoBKm* mutant to get the *feoBKm-fbpATp* (*ffO*) double mutant. Transformants were selected on malate plates supplemented with the appropriate antibiotics under aerobic conditions. Genomic DNA was prepared from the ampicillin sensitive transformants and confirmation of the presence of the antibiotic resistance marker at the desired locus was performed by PCR.

mRNA preparation and qPCR. Total RNA was purified from WT and FNR⁻ mutant cells grown under different oxygenic conditions (aerobic (AE), microaerobic (MA) and photosynthesis (PS) conditions) in malate medium (M) composed of malate NH₄, phosphate potassium buffer at pH 7, iron, vitamin free casamino acids, metal 44 and yeast extract or in an iron depleted malate medium (-Fe) as described in [36]. For semi-quantitative qPCR, cDNA were generated from 1 µg of total RNA with a specific reverse primer RG498 using the Superscript IV (Invitrogen) and by following the manufacturer's protocol. PCR was done with 2 µl of cDNA with specific primers (table SII) to amplify fragments of *ftrA* (RG489-RG490) and *ftrD* (RG497-RG498) genes. Amplified products were analysed on a 1.4% agarose gel.

Cloning, expression and purification of mature recombinant His-tag RgFtrA. The synthetic gene (Genscript) coding for RgFtrA without its signal peptide and fused to 11 amino-acids including a C-terminal His-tag was inserted into pET-26b. *E. coli* BL21 pLysS were transformed by pET26b-*ftrAH₆*. Cells were grown at 37°C in LB broth supplemented with 1 mM isopropyl β-D-1-thiogalactopyranoside to induce protein production. Cells were centrifuged, resuspended in a buffer of 50 mM Tris-HCl, pH 8, 20 mM imidazole and 300 mM NaCl, and disrupted by sonication. After centrifugation at 25,000 g for 30 min at 4°C, the supernatant was loaded onto a 5 ml His-Trap HP column (GE Healthcare). Protein elution was performed with 50 mM Tris-HCl pH 8, 300 mM imidazole and 300 mM NaCl. Protein fractions were loaded onto a gel filtration column (HiLoad 26/60 Superdex 200 prep grade, GE Healthcare) equilibrated with 50 mM Tris-HCl pH 8 and 150 mM NaCl. The protein fractions were pooled, concentrated and stored at -80°C. This protein led to the first structure of apo His-tag RgFtrA at 1.4 Å resolution.

Cloning, expression and purification of mature recombinant RgFtrA and RgFtrB. The synthetic gene (Genscript) coding for the mature RgFtrA (without the signal peptide) was inserted into pET-26b.

The synthetic gene (Genscript) coding for the mature RgFtrB was inserted into pET-29b. *E. coli* BL21 pLysS were transformed by pET26b-*ftrA* or pET29b-*ftrB*. Cells were grown at 37°C in LB broth supplemented with 1 mM isopropyl β -D-1-thiogalactopyranoside to induce protein production. Cells were centrifuged, resuspended in a buffer of 50 mM sodium acetate, pH 4.5 and disrupted by sonication. After centrifugation at 25,000 g for 30 min at 4°C, the supernatant was loaded onto a cation-exchange chromatography (HiTrap SP 5ml, GE Healthcare). Protein elution was performed with a linear gradient of 0-1 M NaCl. Protein impurities were further removed by gel filtration (HiLoad 26/60 Superdex 200 prep grade, GE Healthcare) equilibrated with 50 mM Tris-HCl pH 8 and 150 mM NaCl. The protein fractions were pooled, concentrated and stored at -80°C for crystallization assays. For RPE, DCS and ITC experiments, which are highly sensitive to metal trace, above purified RgFtrA and RgFtrB were treated with 50 mM ethylenediaminetetraacetic acid (EDTA) for 30 min. A desalting column allowed EDTA removing using 50 mM Tris-HCl pH 8 or MES pH 6.5 and 150 mM NaCl. The RgFtrA after EDTA treatment was loaded with equimolar CuCl₂ for DCS, ITC, RPE and crystallization assays.

Cloning, expression and purification of ‘as isolated’ mature Strep-tag RgFtrA. The synthetic gene (Genscript) coding for the mature RgFtrA protein fused to a C-terminal Strep-tag with its own promoter sequence was inserted in the puc57 plasmid at KpnI-XbaI sites. From this plasmid called puc57-*ftrAstreptag*, the tagged gene was subcloned into the replicative plasmid pBBR1MCS-2 at KpnI-XbaI sites. pBBR2-*ftrAstreptag* plasmid thus obtained was introduced by electroporation in the *Δ ftrATp* mutant. Strep-tag RgFtrA protein was purified from *R. gelatinous* *Δ ftrATp* mutant cells grown photosynthetically (2 L) for 48 h at 30°C in iron depleted medium. Cells were pelleted by centrifugation at 3,000 g for 15 min at 4°C. The bacterial pellet was washed with 0.1 M Tris-HCl pH 8 and 150 mM NaCl buffer and centrifuged. The bacterial pellet was fast frozen in liquid nitrogen before storage at -20°C. For cell lysis by sonication, the bacterial pellet was resuspended in 0.1 M Tris-HCl pH 8 and 150 mM NaCl, supplemented with 0.1 M of PMSF (phenylmethylsulfonyl fluoride) and DNase 50 μ g/mL. Lysed cells were centrifuged at 208,000 g for 90 min at 4°C. The soluble fraction was filtered before loaded onto a StrepTrapHP column (5ml, Merck). The proteins were eluted with 5 ml of 2.5 mM desthiobiotin. Protein fractions were loaded onto a gel filtration (HiLoad 26/60 Superdex 200 prep grade,

GE Healthcare) equilibrated with 50 mM Tris-HCl pH 8 and 150 mM NaCl. The protein fractions were concentrated on Amicon YM-10 (Millipore) concentrator pooled, and stored at -80°C .

Crystallization and structure determination. Crystallization conditions for His-tag RgFtrA, RgFtrA and Strep-tag RgFtrA were screened using QIAGEN kits (Valencia, CA) with a Mosquito nanodrop robot (SPT Labtech). His-tag RgFtrA and RgFtrA crystals were manually optimized in the same crystallization condition (Tables 1 and 2). Strep-tag RgFtrA crystals were obtained in three different conditions of polyethylene glycol with one condition containing 0.2 M MgCl_2 and were manually optimized. Few crystals of Strep-tag RgFtrA were soaked with 0.1 M FeCl_3 and turned orange. Co-crystallization of His-tag RgFtrA with 20 mM NiSO_4 was manually realized (Table S1). RgFtrA treated with EDTA and loaded with equimolar CuCl_2 and 1 mM FeSO_4 crystallized in the same condition containing MgCl_2 than the Strep-tag RgFtrA co-crystallized with 4 mM FeSO_4 . Unfortunately, no iron was present in these co-crystal structures, instead magnesium ions were observed. Crystals were transferred to a cryo-protectant solution (mother liquor supplemented with 20% PEG 400 or 30% glycerol) and flash-frozen in liquid nitrogen. Diffraction data were collected at 100 K on PROXIMA 1 and 2 beamlines at SOLEIL synchrotron (Saint-Aubin, France). Data processing was performed using the XDS package [37] (Table 1 and 2). We determined first the structure of His-tag RgFtrA at 1.4 Å resolution by molecular replacement using the structure of *Campylobacter jejuni* P19 (CjP19 PDB 3LZL) as a search model. Because, the crystals of RgFtrA were isomorphous to those of His-tag RgFtrA, phases determination at 1.23 Å resolution was straightforward. Both His-tag and RgFtrA structures were empty. The presence of nickel in co-crystals of His-tag RgFtrA with 20 mM NiSO_4 was confirmed by an X-ray fluorescence emission scan on the beamline and a nickel anomalous data set at 1.379288 Å wavelength was collected at 1.8 Å resolution. A nickel ion is 50% present in only one subunit of the dimer. The three structures of the endogenous Strep-tag RgFtrA were determined by molecular replacement with PHASER [38] using the high resolution refined structure of RgFtrA as a search model. They contain a fully occupied endogenous copper ion in each subunit of the asymmetric unit. No iron was detected using X-ray fluorescence emission scan in the Strep-tag RgFtrA-iron co-crystals. The structure composed of three dimers in the asymmetric unit revealed the presence of a magnesium ion

originated from the crystallization conditions bound to the second metal site. Removing MgCl_2 prevented crystals growth. Two crystals of RgFtrA treated with EDTA and loaded with equimolar CuCl_2 and 1 mM FeSO_4 denoted X1 and X2 grown with 100 and 50 mM MgCl_2 , respectively, were collected. Because of the anisotropy of the diffraction of X1 and X2 crystals as well as the Strep-tag RgFtrA crystal grown in MgCl_2 , the DEBYE and STARANISO programs developed by Global phasing Ltd were applied to the data scaled with AIMLESS using the STARANISO server (<http://staraniso.globalphasing.or/>). These programs perform an anisotropic cut-off of merge intensity data on the basis of an analysis of local $I/\sigma(I)$, compute Bayesian estimates of structures amplitudes, taking into account their anisotropic fall-off, and apply an anisotropic correction to the data. The corrected anisotropic amplitudes were used for further refinement.

The presence of iron in soaked crystals of Strep-tag RgFtrA with 0.1 M FeCl_3 was confirmed by an X-ray fluorescence emission scan on the beamline and an iron anomalous data set at 1.73925 Å wavelength was collected at 2.75 Å resolution. An iron ion is present in each subunit of the dimer of the asymmetric unit.

Refinement of each structure was performed with BUSTER-2.10 [39], employing TLS groups and NCS restraints when more than one molecule was present in the asymmetric unit. Inspection of the density maps and manual rebuilding were performed using COOT [40]. Refinement details of each structure are shown in Tables 1 and 2. Molecular graphics images were generated using PyMOL (<http://www.pymol.org>).

Electron paramagnetic resonance (EPR) measurement. X-band cw-EPR spectra were recorded with a Bruker Elexsys 500 X-band spectrometer equipped with a standard ER 4102 (Bruker) X-band resonator, a Bruker teslameter, an Oxford Instruments cryostat (ESR 900) and an Oxford ITC504 temperature controller.

Differential scanning calorimetry (DSC). Thermal stability of the His-tag RgFtrA, the RgFtrA, the RgFtrA treated with EDTA and loaded with equimolar CuCl_2 , the Strep-tag RgFtrA expressed in *R. gelatinosus* periplasm, and the RgFtrB treated with EDTA was performed by DSC on a MicroCal model

VP-DSC in a standard buffer with 30 μM and 300 μM for protein and metal, respectively. Divalent metals were used such as MgSO_4 , MnCl_2 , NiSO_4 , CuCl , CuCl_2 , CoCl_2 , CdCl_2 , ZnSO_4 , FeSO_4 , and FeCl_3 . Each measurement was preceded by a baseline scan with the standard buffer. All solutions were degassed just before loading into the calorimeter. Scans were performed at 1 $\text{K}\cdot\text{min}^{-1}$ between 20 and 90°C. The heat capacity of the buffer was subtracted from that of the protein sample before analysis. Thermodynamic parameters were determined by fitting the data to the following equation:

$$\Delta C_p(T) = \frac{K_d(T) \Delta H_{cal} \Delta H_{vH}}{[1 + K_d(T)]^2 RT^2}$$

where K_d is the equilibrium constant for a two-state process, ΔH_{vH} is the enthalpy calculated on the basis of a two-state process and ΔH_{cal} is the measured enthalpy.

Tycho NT.6 Nanotemper Differential Scanning Fluorimeter. This equipment measures the change in intrinsic fluorescence of tryptophan residues in protein samples. Protein unfolding was followed by tryptophan fluorescence intensity at 330 and 350 nm in the presence of different metals (same as for DSC). The inflection temperature (T_i) was determined by detecting the maximum of the first derivative of the fluorescence ratios (F_{350}/F_{330}) after fitting experimental data with a polynomial function. Data were measured in triplicate. Tycho NT.6 was used with a heating rate of 30°C min^{-1} .

Isothermal titration microcalorimetry measurements (ITC). Isothermal titration microcalorimetry experiments were performed with an ITC200 isothermal titration calorimeter from MicroCal (Malvern). The experiments were carried out at 25°C. Protein concentration in the microcalorimeter cell (0.2 ml) was 50 μM . Nineteen injections of 2 μl metal at concentrations ranging from 700 to 800 μM were performed at intervals of 180 s while stirring at 500 rpm. The experimental data were fitted to theoretical titration curves with software supplied by MicroCal (ORIGIN®). This software uses the relationship between the heat generated by each injection and ΔH (enthalpy change in Kcal/Mol), K_a (the association binding constant in M^{-1}), n (the number of binding sites), total protein concentration and free and total ligand concentrations [41].

Accession numbers. The atomic coordinates and structure factors have been deposited at the Protein Data Bank under PDB ID 7R3P and 7R4U for His-tag RgFtrA and RgFtrA respectively, PDB ID 7R3S for His-tag RgFtrA-Ni²⁺, PDB ID 7R4V for Strep-tag RgFtrA-Cu⁺, PDB ID 7R5P for Step-tag RgFtrA-Cu⁺Mg²⁺, PDB ID 7R4Z for Step-tag RgFtrA-Cu⁺Fe³⁺ and PDB ID 7R5E and 7R5G for RgFtrA-Cu²⁺Mg²⁺ denoted X1 and X2 in Table 1, respectively

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Author contributions. ASS and SO performed the experiments on *R. gelatinosus*. SP and AV performed the crystallization assays. AV and SM performed the crystallography work. MAN and AB performed the DSC/microcalorimetry and EPR experiments, respectively. ASS and SM wrote the manuscript. All the authors discussed the results and contributed to the writing of the manuscript.

Conflict of interest. The authors declare no conflict of interest.

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Table 1. Crystallographic data and refinement parameters.

	apo RgFtrA	Strep-tag RgFtrA-Cu ⁺	Strep-tag RgFtrA- Cu ⁺ Fe ³⁺
PDB code	7R4U	7R4V	7R4Z
Wavelength (Å)	0.979336	1.379288	1.73925
Crystallization conditions	2 M AS, 0.1 M NaCitrate pH 5.6	20% PEG 1500 0.1 M HEPES pH 7.5	20% PEG 2K MME 0.1 M tris-HCl pH 8.5 0.2 soaked with 0.1 M FeCl ₃ .
Za	2	2	2
Number of amino-acids Chains A/B	156/163	154/156	156/156
Space group Cell parameters (Å,°)	P2 ₁ 2 ₁ 2 ₁ a= 59.3 b= 79.9 c= 82.8	P2 ₁ 2 ₁ 2 ₁ a= 61.4 b= 65.9 c= 70.1	P2 ₁ 2 ₁ 2 ₁ a= 60.8 b= 71.2 c= 79.6
Resolution (Å)	48.25-1.23 (1.3-1.23)	49-1.3 (1.38-1.3)	2.75 (2.92-2.75)
No. of observed reflections	1460931 (179016)	676517 (14178)	52608 (3155)
No. of unique reflections	114401 (17537)	114500 (6218)	8992 (1137)
Completeness (%)	98.9 (94.9)	84.6 (28.5)	95.3 (76.7)
R _{sym} (%)	6.7 (200.9)	7.7 (54.9)	18.3 (93)
R _{pim} (%)	2.7 (112.4)	3.3 (51.8)	10.2 (87.8)
σ(I)	18.9 (1.05)	15.1 (1.15)	9.6 (1.21)
CC _{1/2}	1 (0.424)	0.998 (0.633)	0.993 (0.515)
R _{cryst} (%)	17	17.2	17.1
R _{free} (%)	18.6	19.2	22.8
rms bond deviation (Å)	0.01	0.009	0.01
rms angle deviation (°)	1.07	1.13	1.11
Average B (Å ²)			
Protein A/B	19.3/21.8	15.7/16.6	56/59
Cu ⁺ A/B		11.1/11.8	72.4/60.7
Fe ³⁺ A/B			60/58.8
solvent	36.7	27.3	47.5
^a Clashscore	1.14	2	0.83
MolProbity score	0.82	0.97	1.17
^a Ramachandran plot (%)			
Favoured	99.68	99.02	98.38
Outliers	0	0	0

Values for the highest resolution shell are in parentheses

CC_{1/2} = percentage of correlation between intensities from random half-dataset

^aCalculated with MolProbity

Table 2. Crystallographic data and refinement parameters.

	His-tag apo RgFtrA	His-tag RgFtrA-Ni ²⁺	RgFtrA-Cu ²⁺ Mg ²⁺ (X1)	RgFtrA-Cu ²⁺ Mg ²⁺ (X2)	Strep-tag RgFtrA-Cu ²⁺ Mg ²⁺
PDB code	7R3P	7R3S	7R5E	7R5G	7R5P
Wavelength (Å)	0.978566	1.48484	1.73114	1.73114	0.980115
Crystallization conditions	2 M AS, 0.1 M Citrate Sodium pH5.6	2 M AS, 0.1 M Citrate Sodium pH5.6 0.02 M NiSO ₄	25% PEG 4K, 0.1 M Tris-HCl pH 8.5 0.1 M MgCl ₂	25% PEG 4K, 0.1 M Tris-HCl pH 8.5 0.05 M MgCl ₂	25% PEG 4K, 0.1 M Tris-HCl pH 8.5 0.2 M MgCl ₂
Za	2	2	2	2	6
Number of aminoacids Chains A/B/C/D/E/F	157/157	157/157	156/156	156/155	156/156/154/156/156/155
Space group Cell parameters (Å, °)	<i>P</i> 2 ₁ 2 ₁ 2 ₁ <i>a</i> = 59.2 <i>b</i> = 80.3 <i>c</i> = 82.9	<i>P</i> 2 ₁ 2 ₁ 2 <i>a</i> = 59.3 <i>b</i> = 79.8 <i>c</i> = 82.4	<i>C</i> 2 <i>a</i> = 92.6 <i>b</i> = 48.9 <i>c</i> = 78.2 β = 123.1	<i>C</i> 2 <i>a</i> = 92.6 <i>b</i> = 48.4 <i>c</i> = 78.6 β = 123.8	<i>P</i> 2 ₁ <i>a</i> = 47.5 <i>b</i> = 77.9 <i>c</i> = 115.2 β = 95.8
Resolution (Å)	48.1-1.4 (1.48-1.4)	49.13-1.8 (1.9-1.8)	27.4-2.2 (2.27-2.2)	41.08-2.2 (2.23-2.2)	49-1.68 (1.86-1.68)
No. of observed reflections	1049112 (160830)	426698 (31633)	59199 (1111)	71689 (2428)	598262 (18238)
No. of unique reflections	78363 (12299)	68702 (10524)	14345 (523)	15041 (711)	92927 (4342)
Completeness (%)	99.7(97.9)	99.8 (56.1)	98.5 (69.4)	99.7 (75.3)	99.5 (94)
Completeness Staraniso (%)			98.5 (64.9)	99.7 (50.4)	94 (63.1)
<i>R</i> _{sym} (%)	8.6 (126.2)	7.8 (70.7)	12.8 (55.4)	6.7 (50.6)	7.7 (73.8)
<i>R</i> _{pim} (%)	2.4 (36.9)	7.5 (37.3)	6.7 (37.6)	3.3 (27.3°)	3.2 (36.9)
<i>I</i> / σ (<i>I</i>)	15.9 (1.58)	12.2 (1.17)	8.9 (2.5)	11 (2.8)	11.4 (1.7)
CC _{1/2}	0.999(0.718)	0.998 (0.561)	0.985(0.649)	0.997(0.873)	0.998 (0.725)
<i>R</i> _{cryst} (%)	17.8	17.6	17.6	17.2	17.9
<i>R</i> _{free} (%)	19.1	20.1	21.7	19.6	21.1
rms bond deviation (Å)	0.01	0.01	0.01	0.01	0.01
rms angle deviation (°)	1.1	1.07	1.11	1.08	1.04
Average B (Å ²)					
Protein A/B/C/D/EF	22.4/26	36/40.7	23.9/22.2 14.4/15.3	31.4/40.4 23.3/23.9	29/29/38.7/34/30.7/36 18.5/18.6/25.8/23.6/20.3/23.7
Cu A/B/C/D/E/F					
Ni ²⁺ A		33.8			
Mg ²⁺ A/B/C/D/E/F			34/40.1 25.4	61/65.9	27.4/47.5/55.6/55.6/37.5/25.3/46.9
Mg ²⁺ in solvent solvent	39	48.4	24.3	35	33.7
^a Clashscore	0.98	1.57	1.44	1.04	1.38
MolProbity score	0.79	0.93	1.16	0.8	0.98
^a Ramachandran plot (%)					
Favoured	99.03	98.4	99.35	99.35	98.69
Outliers	0	0	0	0	0

Values for the highest resolution shell are in parentheses

CC_{1/2} = percentage of correlation between intensities from random half-dataset

^aCalculated with MolProbity

Table 3. T_m of RgFtrA devoid of any metal without and with different cations in the primary metal binding site using DSC (see supplementary Figure S3). T_m of RgFtrA devoid of any metal and loaded with equimolar CuCl_2 with different cations in the secondary metal binding site using DSC at pH 6.5 (see supplementary Figure S6).

	T_m (°C)	ΔT_m (°C)	ΔH_1 (kcal/mol)	ΔHVH_1 (kcal/mol)
RgFtrA	52.06		167	199
+ MnCl_2	55.20	3.14	190	214
+ MgCl_2	52.45	0.39	175	204
+ FeSO_4	53.53	1.47	155	134
+ FeCl_3	53.37	1.31	152	166
+ CuCl	76.78	24.72	155	137
+ CuCl_2	75.94	23.88	110	111
+ NiSO_4	72.34	20.28	85.4	139
+ CdCl_2	61.96	9.9	101	132
+ ZnSO_4	59.02	6.96	78.6	136
+ CoCl_2	58.59	6.53	81.2	120
RgFtrA-CuCl_2	72.98		145	93.9
+ MnCl_2	75.44	2.46	152	109
+ MgCl_2	72.44	-0.54	118	86.5
+ CoCl_2	74.21	1.23	137	111
+ FeSO_4	87.78	14.8	66	228
+ FeCl_3	87.38	14.4	66	207

Table 4. Distances in Å between copper (or nickel) ion and RgFtrA amino acids of the primary metal site. H131' belongs to the neighbouring subunit. Distances (below 3Å) between metal ion (Fe^{3+} or Mg^{2+}) in the secondary metal site and RgFtrA amino acids or water molecules (W).

	His41	Met88	His95	His131'	Glu43
Cu⁺ in strep-tag RgFtrA	2.0	2.3	2.1	2.1	4
(chain A)	2.0	2.3	2.0	2.1	4
(chain B)					
Cu⁺ in strep-tag RgFtrA-CuFe					
(chain A)	2.4	2.3	2.2	2.2	4.3
(chain B)	2.3	2.3	2.0	2.2	4.4
Cu⁺ in strep-tag RgFtrA-CuMg					
(chain A)	2.1	2.4	2.0	2.1	3.8
(chain B)	2.1	2.3	2.1	2.1	4.0
(chain C)	2.2	2.3	2.2	2.1	4.0
(chain D)	2.2	2.3	2.1	2.1	3.8
(chain E)	2.1	2.3	2.0	2.2	3.8
(chain F)	2.1	2.3	2.1	2.1	3.8
Cu²⁺ in RgFtrA-CuMg (X1)					
(chain A)	2.1	2.4	2.2	2.2	3.9
(chain B)	2.2	2.2	2.2	2.2	4.8
Cu²⁺ in RgFtrA-CuMg (X2)					
(chain A)	2.1	2.3	2.1	2.2	3.7
(chain B)	2.1	2.3	2.1	2.2	5.4
Ni²⁺ in his-tag RgFtrA	2.4	3.2	1.7	2.4	2.7

Table 5. Distances (below 2.5 Å) between metal ion (Fe^{3+} or Mg^{2+}) in the secondary metal site and RgFtrA amino acids or water molecules (W).

	Glu2	Glu43	Asp92	W	W	W	W
Fe^{3+} (chain A)	2.3	2.2	1.7	2.6	3		
(chain B)	2.3	2.2	1.9				
Mg^{2+} (PDB 7R5E) (chain A)	3	2.7	2.0	2.7			
(chain B)	2.5	2.2	2.3				
Mg^{2+}(PDB 7R5G) (chain A)	3	2.6	2.2	2.6			
(chain B)		2.8	2.1	2.7			
Mg^{2+} (PDB 7R5P) (chain A)		2.3	2.1	2.2	2.0	2.0	2.2
(chain B)		2.4	2.2	2.2	1.9	1.9	2.3
(chain E)		2.2	2.2	2.1	2.1	1.9	2.3
(chain D)		2.3	2.3	2.2	2.2		2.4
(chain C)		2.2	2.1				
(chain F)		2.3	2.2	2.3		2.0	2.3

Figure legends

Figure 1. (A) A schematic representation of the ferrous iron uptake Ftr1-like system from *Rubrivivax gelatinosus*. **(B)** Organization of the *ftrAPBCD* operon presented as orange arrows and position of Fur and FNR binding sites in the promoter region. The operon is composed of 5 genes coding for the periplasmic FtrA, the porine FtrP, the CupII-type ferroxidase FtrB, the transporter FtrC (Ftr1p homolog) and the putative membrane polyferredoxin FtrD involved in the electron transfer. **(C)** Transcript level of *ftr* genes in the WT and Δfnr mutant of *R. gelatinosus* grown with (M) and without iron (-Fe) under aerobic (AE), microaerobic (MA) and photosynthesis anaerobic (PS) conditions. RT-PCR was performed to examine (i) the co-transcription of the *ftr* genes (ftrD reverse primer was used for the CDNA synthesis) (ii) genes regulation by oxygen concentration through FNR and (iii) by iron through Fur.

Figure 2. Growth comparison between the WT (red), *fbpAKm* (blue), $\Delta ftrATp$ (green), *ffa* (*fbpAKm*- $\Delta ftrATp$) (purple) at pH 6 and pH 7 grown in iron-depleted malate medium microaerobic conditions. Results are the average of 3 independent experiments.

Figure 3. (A) Ribbon representation of the RgFtrA dimer and a close-up view of the water molecule (red sphere) bound to the primary metal site. Amino-acids are labelled and water-side chains interactions are shown as dashed red lines. **(B)** Superposition of the primary metal site of apo structures: RgFtrA (cyan), CjP19 (grey) with a 10% bound Zn^{2+} (grey sphere) described as an apo form and apo EcFetP (yellow). The conserved amino acids forming the primary metal site are labelled and metal-CjP19 interactions are shown as plain grey lines. His95 in RgFtrA adopts a different position with the side chain out of the copper site. **(C)** Ribbon representation of the ‘as isolated’ Strep-tag RgFtrA dimer with an endogenous Cu^+ (purple sphere) bound the primary metal site and a close-up view of the RgFtrA- Cu^+ interactions. Amino-acids are labelled and metal-side chains interactions are shown as dashed lines. **(D)** Superposition of the copper binding site of ‘as isolated’ Strep-tag RgFtrA (purple) and RgFtrA (cyan). The conserved amino acids forming the primary metal site are labelled. The water molecule in

the RgFtrA is shown in red sphere and is at 1.8 Å of the Cu⁺ ion. Molecular graphics were generated using PyMOL (<http://www.pymol.org>).

Figure 4. Isothermal titration microcalorimetry experiments for K_D measurement of the RgFtrA-Cu²⁺ complex towards Fe²⁺ ions bound to the secondary metal site in MES at pH 6.5 (A) and in Tris-HCl at pH 8 (B). The top panel shows heat differences upon injection of metal ion and low panel shows integrated heats of injection with the best fit (solid line) to a single binding model using Microcal ORIGIN. ITC experiments were performed twice.

Figure 5. Isothermal titration microcalorimetry experiments for K_D measurement of the RgFtrA-Cu²⁺ complex towards Fe²⁺ ions bound to the secondary metal site in MES at pH 6.5 (A) and in Tris-HCl at pH 8 (B). A control experiment of 1mM FeSO₄ without protein was measured in MES at pH 6.5 (A). The top panel shows heat differences upon injection of metal ion and low panel shows integrated heats of injection with the best fit (solid line) to a single binding model using Microcal ORIGIN. ITC experiments were performed twice.

Figure 6. Differential scanning calorimetry thermograms of ‘as isolated’ Strep-tag RgFtrA expressed in *R. gelatinosus* with and without ethylenediaminetetraacetic acid (EDTA). The table below indicates the T_m . DSC experiments were performed twice. (B) Electron paramagnetic resonance spectrum recorded on ‘as isolated’ Strep-tag RgFtrA (80 μM) expressed in *R. gelatinosus* periplasm. The inset spectrum was recorded after the addition of a concentrated HCl (~ 18%) directly into the EPR tube. Temperature, 15 K; modulation amplitude, 25 G; microwave power, 20 mW; microwave frequency, 9.49 GHz; modulation frequency, 100 kHz. Inset: Temperature, 30 K; modulation amplitude, 25 G; microwave power, 5 mW; microwave frequency, 9.49 GHz; modulation frequency, 100 kHz.

Figure 7. (A) A close-up view of the His-tag RgFtrA-Ni²⁺ interactions. Amino-acids are labelled and metal-side chains interactions are shown as plain green lines. Ni²⁺ (green sphere) occupancy is 50% in line with the His95 side chain, observed in two positions as shown in the 2Fo-Fc electron density map contoured at 1σ: one allowing metal interaction and the other observed in apo RgFtrA. (B) Superposition of the copper binding site of ‘as isolated’ Strep-tag RgFtrA (purple) and His-tag RgFtrA-Ni²⁺ (green).

The conserved amino acids forming the primary metal site are labelled. **(C)** Structural comparison of the two subunits of His-tag RgFtrA-Ni²⁺ dimer. A nickel ion (green sphere) occupies the primary metal site at 50% occupancy in subunit A (lime green) while a water molecule (red sphere) is present in subunit B (pink). Amino acids forming the primary metal site are labelled. Interactions between His-tag RgFtrA and nickel ion are shown as plain green lines whereas water-side chains interactions are shown as dashed red lines. **Molecular graphics were generated using PyMOL (<http://www.pymol.org>).**

Figure 8. **(A)** Ribbon representation of ‘as-isolated’ Strep-tag RgFtrA dimer in complex with ferric iron and a close-up view of the Fe³⁺ (orange sphere) bound to the secondary metal site. Amino-acids are labelled and metal-side chains interactions are shown as plain orange lines **(B)** A different view of the same dimer as in **(A)** showing the Fe³⁺ exposed at the protein surface. **(C)** Close up view of the two metal binding sites of the Strep-tag RgFtrA-Cu⁺Fe³⁺ complex (purple and orange spheres for Cu and Fe respectively). The two metal sites are separated by 7 Å. **(D)** Close up view of the two metal binding sites of the ‘Strep-tag RgFtrA-Cu⁺Mg²⁺ complex. The two metal sites are separated by 6.7 Å. **(E)** Close-up view of the superimposed iron binding sites of the Strep-tag RgFtrA-Cu⁺Fe³⁺ and Strep-tag RgFtrA-Cu⁺Mg²⁺ complexes. **Molecular graphics were generated using PyMOL (<http://www.pymol.org>).**

Figure 9. EPR spectra recorded in the RgFtrA protein in a medium containing 50 mM MES and 150 mM NaCl at pH 6.5. In the four spectra, the RgFtrA concentration was 118 µM. The black spectrum corresponds to the “as purified” protein and was similar to that from the RgFtrA treated with EDTA. The red spectrum was recorded with the copper reconstituted protein. The blue spectrum was recorded with the copper reconstituted protein after the addition of 118 µM FeSO₄ directly in the EPR tube (from a stock at 1 mM FeSO₄ dissolved in water). The green spectrum was recorded with the copper reconstituted protein after the addition of 236 µM FeSO₄ directly in the EPR tube. The amplitude of the Fe³⁺ EPR signal did not increase upon the addition of 236 µM FeSO₄ when compared to the addition of 118 µM FeSO₄ showing that there was no detectable auto oxidation of the Fe²⁺ in these conditions. Instrument settings: Temperature, 15 K; modulation amplitude, 25 G; microwave power, 5 mW; microwave frequency, 9.49 GHz; modulation frequency, 100 kHz.

Figure 10. Differential scanning calorimetry thermograms of the RgFtrB with CuCl₂ and FeSO₄. The table below indicates the *T_m*. DSC experiments were performed twice.

Figure 11. Fnr (grey boxes) were found with PRODORIC database (<https://www.prodoric.de/vfp/>) in the promoter region of genes encoding the Ftr1P system from analyzed bacteria including *R. gelatinosus* (*R. gel*), *Magnetospirillum magneticum* (*M. mag*), *Escherichia coli* (*E. coli*), *Campylobacter jejuni* (*C. jeju*), *Brucella abortus* (*B. abo*), *Bordetella pertussis* (*B. per*), *Pseudomonas syringae* (*P. syr*), *Rhodopseudomonas. palustris* (*R. pal*) or *Bacillus subtilis* (*B. sub*).

Figure 12. (A) Structure-based sequence alignment generated using Clustal Omega (<https://www.ebi.ac.uk/Tools/msa/clustalo>), manually corrected using COOT [40] and finalized using ESPript3 (<https://esript.ibcp.fr/ESPript/ESPript>) of RgFtrA (PDB 7R4Z) with CjP19 (PDB 3LZN) and EcFetP (PDB 3NRP). The secondary structure elements of RgFtrA (conserved in CjP19 and EcFetP) are labelled and indicated by coils for α -helices, arrow for β -strands and η for short 3_{10} helices. Identical residues are shown as white letters on a red background. The positions of the RgFtrA amino acids involved in copper (His41, Met88, His95 and His131) and in iron (Glu2, Glu43 and Asp92) binding are indicated by green and red triangles, respectively. These residues are all conserved. (B) Structural comparison between ‘as isolated’ Strep-tag RgFtrA and CjP19 (PDB 5I0V) in complex with copper and ferric iron. Copper and iron are represented by sphere and indicated. Molecular graphic was generated using PyMOL (<http://www.pymol.org>).

Figure 13. (A) Sequence alignment generated using Clustal Omega (<https://www.ebi.ac.uk/Tools/msa/clustalo>) and ESPript3 (<https://esript.ibcp.fr/ESPript/ESPript>) of RgFtrB with Bortedella and Brucella homologues shown with their accession number from Protein Blast-NCBI (<https://blast.ncbi.nlm.nih.gov>). Identical residues are shown as white letters on a red background. The conserved acidic residues at positions 42, 44, 51, 77 and 76 are indicated by red stars. (B) Ribbon representation of the predicted structure of RgFtrB from AlphaFold2 (ColaFold on google). The conserved acidic residues at positions 42, 44, 51, 77 and 76 are labelled and their side chains are shown in red. Molecular graphic was generated using PyMOL (<http://www.pymol.org>).

Figure 14. A proposed model for the ferrous iron transporter, FtrAPBCD from *R. gelatinosus*. Ferrous ion enters the periplasm through the porin FtrP localized in the outer membrane of the cells. It is sequestered by the periplasmic copper dependent ferrous iron oxydase FtrA. Then, the oxidized iron is taken in charge by the chaperon FtrB, which delivers the ferric iron to the permease FtrC for transport into the cytoplasm. The electrons from the iron oxidation by copper in FtrA are recycled by the polyferredoxin FtrD acting as the electron sink and resetting the ferrous uptake system.

Supporting Information:

Table S1. Bacterial strains and plasmids.

Table S2. Primers used in this work.

Figure S1. Growth of different single and double mutants.

Figure S2. Differential scanning calorimetry thermograms of RgFtrA and His-tag RgFtrA

Figure S3. Differential scanning calorimetry thermograms of RgFtrA devoid of any metal without and with different cations.

Figure S4. Effect of metal presence on the thermal unfolding of RgFtrA devoid of any metal without and with different cations.

Figure S5. Differential scanning calorimetry thermograms of the apo RgFtrA loaded with equimolar CuCl₂ with different cations.

Figure S6. Effect of metal presence on the thermal unfolding RgFtrA-Cu²⁺ with different cations measured by nanoDSF.

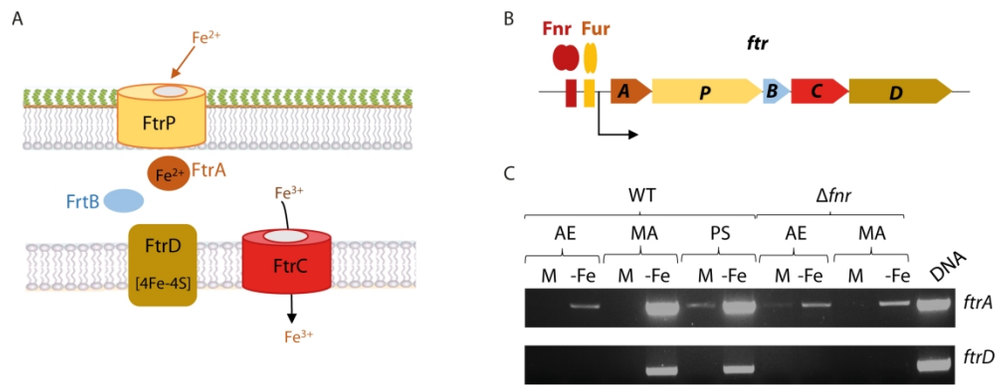


Figure 1

210x80mm (300 x 300 DPI)

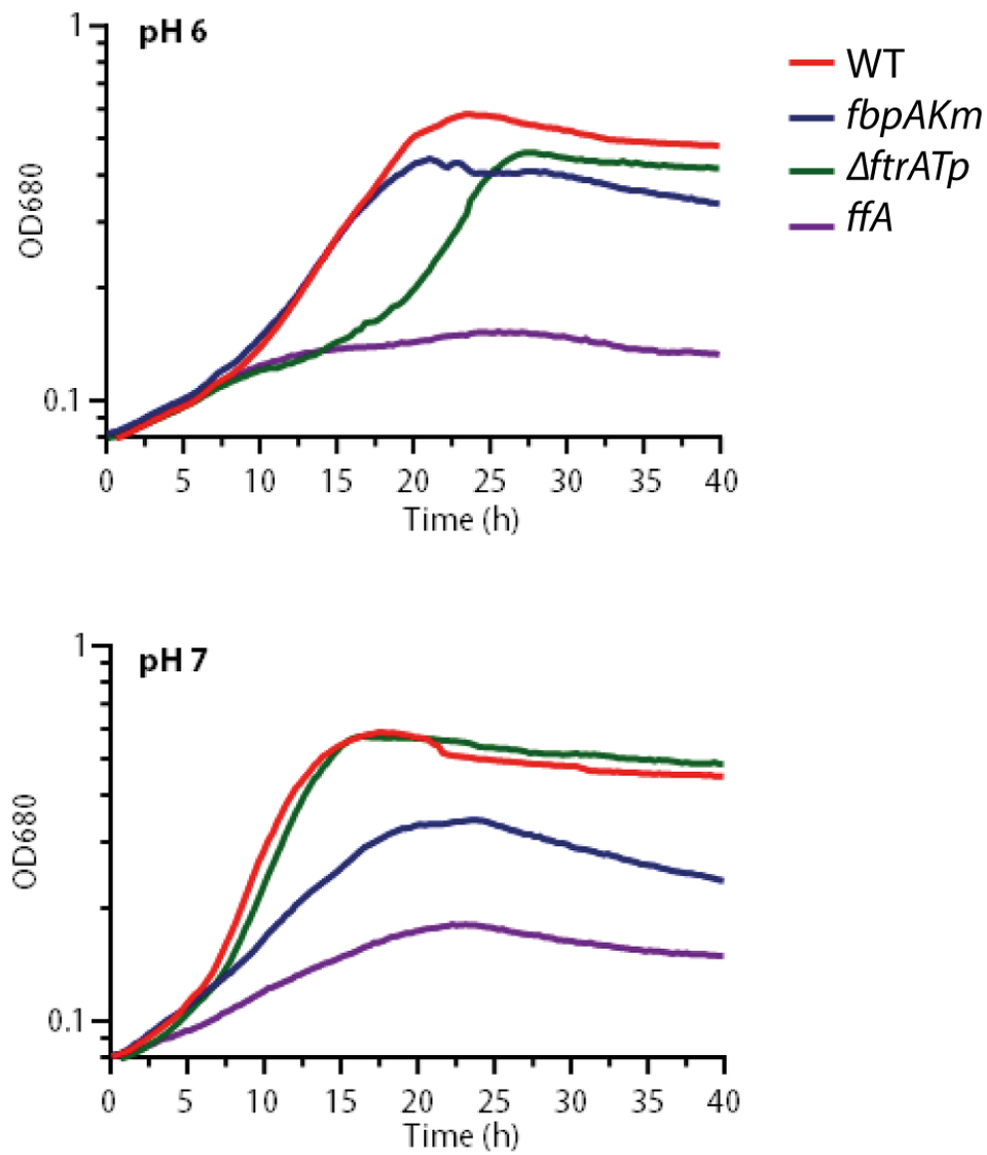


Figure 2

81x98mm (300 x 300 DPI)

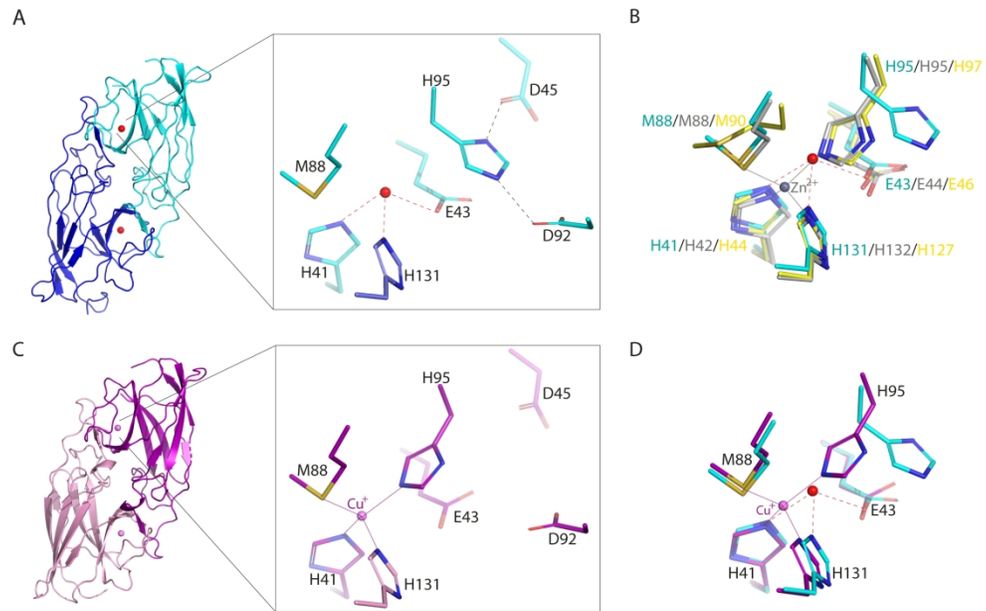
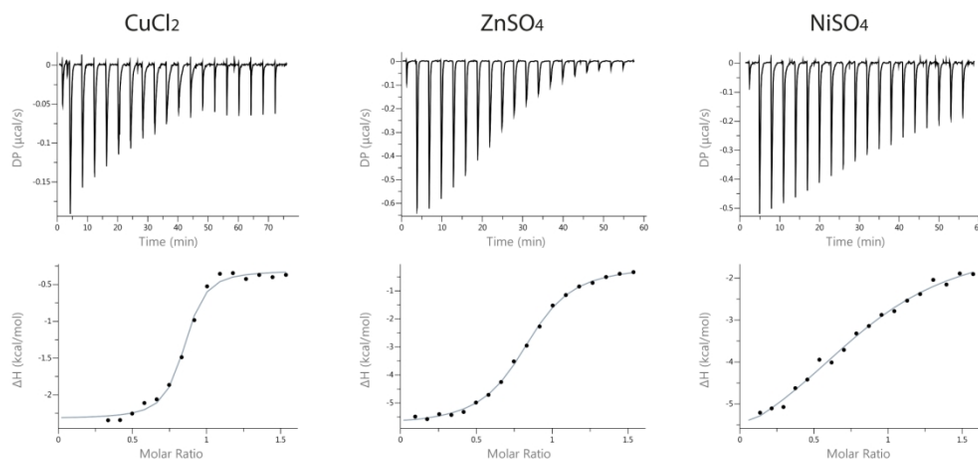


Figure 3

186x117mm (300 x 300 DPI)



25°C	n	K _D (µM)	ΔH (kcal/mol)	-TΔS (kcal/mol)	ΔG (kcal/mol)
RgFtrA / CuCl₂	0.7	0.51	-2.3	-6.29	-8.59
RgFtrA / ZnSO₄	0.7	2.55	-6.54	-1.09	-7.53
RgFtrA / NiSO₄	0.7	21.9	-6.42	-1.27	-6.36

Figure 4

161x119mm (300 x 300 DPI)

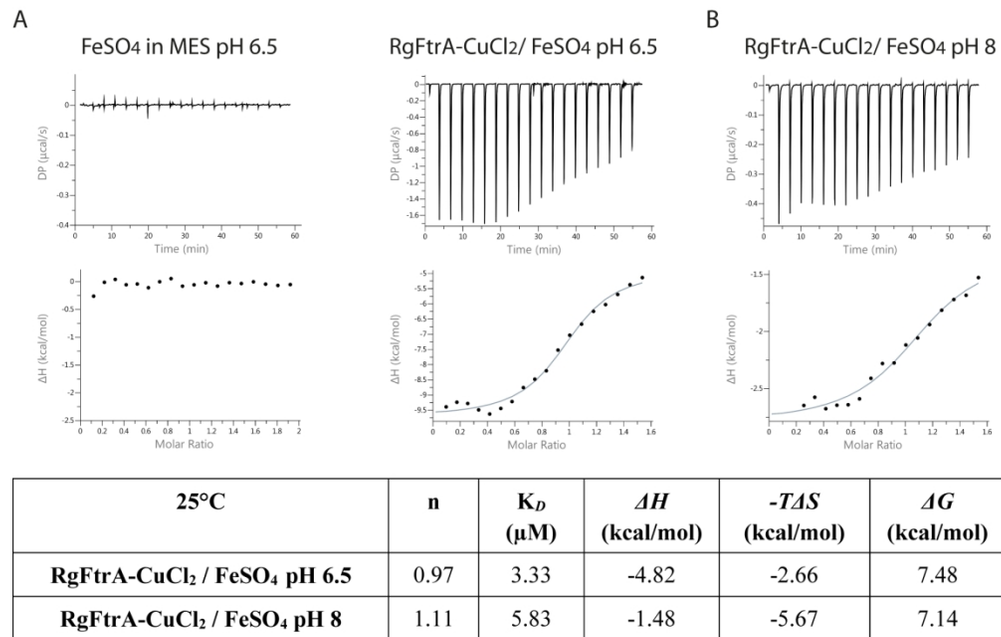


Figure 5

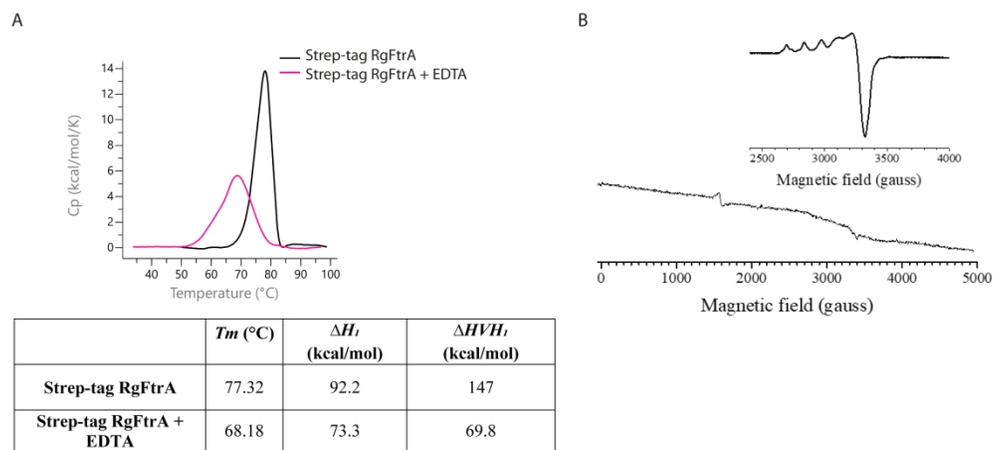


Figure 6

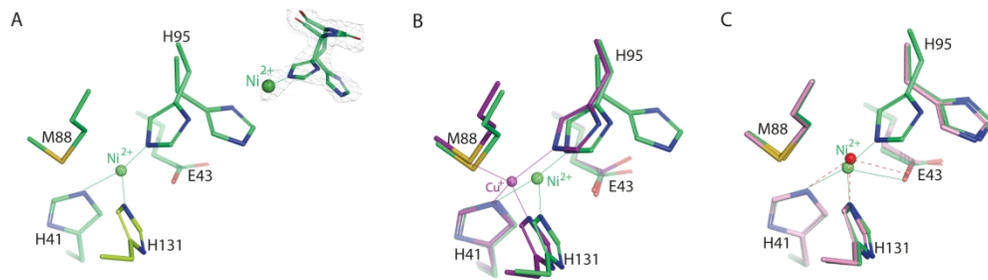


Figure 7

189x54mm (300 x 300 DPI)

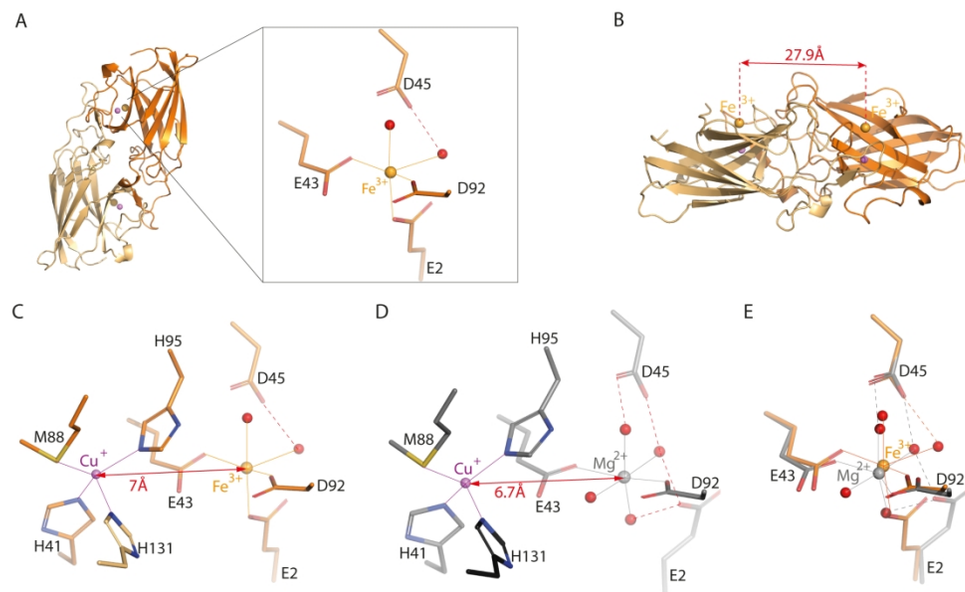


Figure 8

187x113mm (300 x 300 DPI)

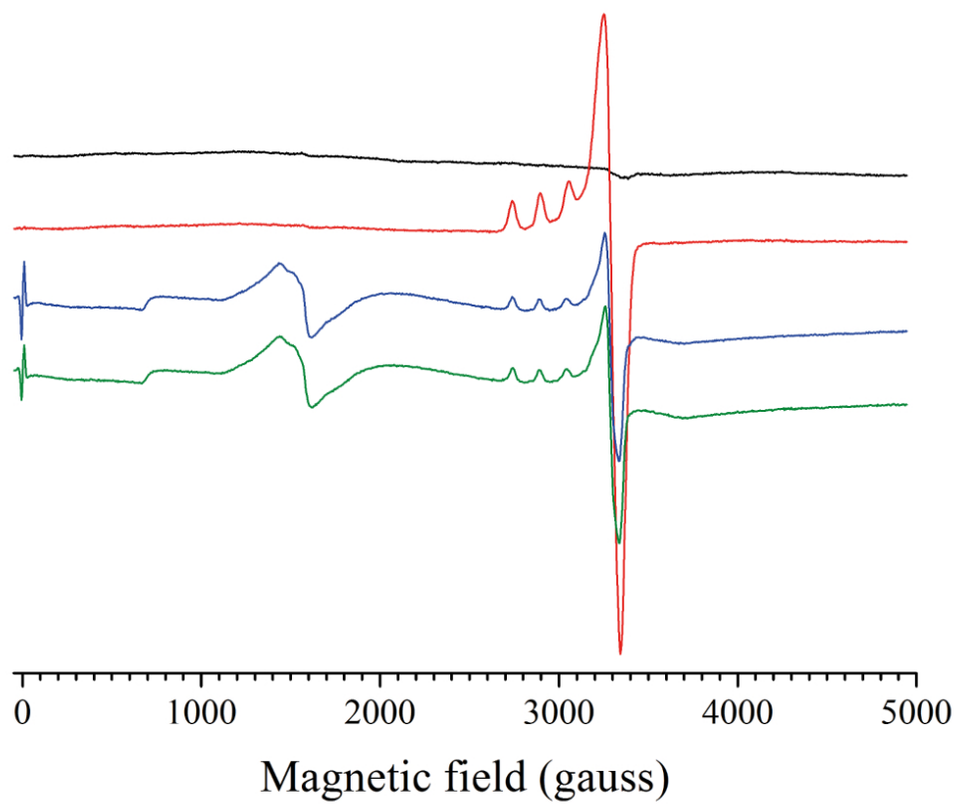
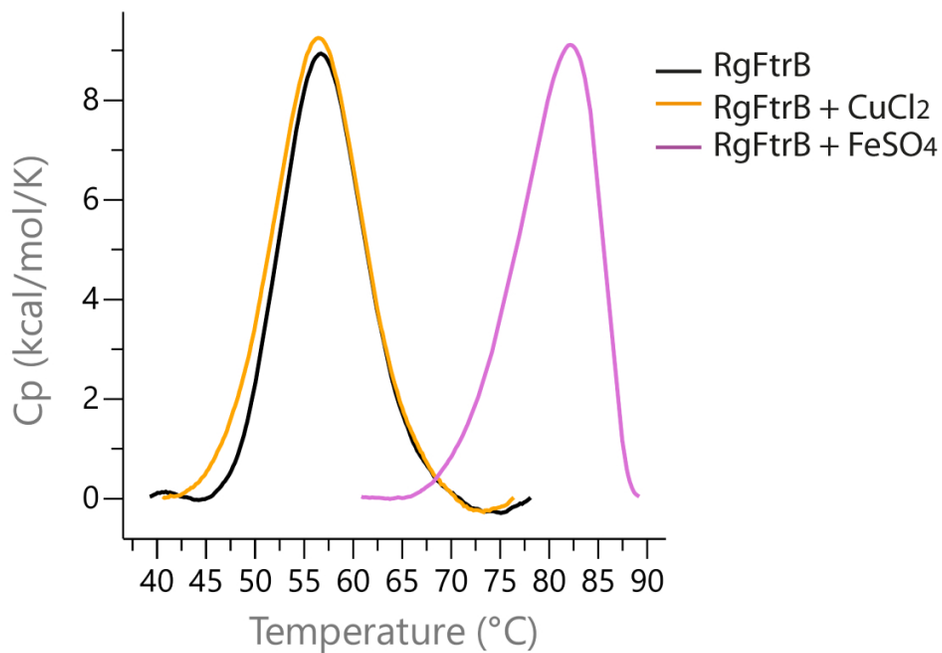


Figure 9

86x72mm (300 x 300 DPI)



	T_m (°C)	ΔT_m (°C)	ΔH_f (kcal/mol)	ΔHVH_f (kcal/mol)
RgFtrB	56.70		96.6	81.5
+ CuCl ₂	56.27	-0.43	108	74.3
+ FeSO ₄	81.01	24.32	96.1	94.9

Figure 10

99x102mm (300 x 300 DPI)

<i>R. gel</i>	ccgc TTGA tgcagg GCAA gtc-N ₉₅ -atg
<i>M. mag</i>	gttc TTGA tcagca GGAT cac-N ₃₆₂ -atg
<i>E. col</i>	cgag TTTG tcgcca CCAA cat-N ₂₁₅ -atg
<i>C. jeu</i>	tttt TTGA ttttta TTAA gct-N ₂₈₁ -atg
<i>B. abo</i>	agaa TTGA tctaaa CTAT tgt-N ₁₂₀ -atg
<i>B. per</i>	gtag TTGT tcatca TCTT cata-N ₃₀₂ -atg
<i>P. syr</i>	gtag TTGT ttttta TCGG gat-N ₉₃ -atg
<i>B. sub</i>	tttt TTGC tgtcca TCAA ggt-N ₉₆ -atg

Figure 11

100x41mm (300 x 300 DPI)

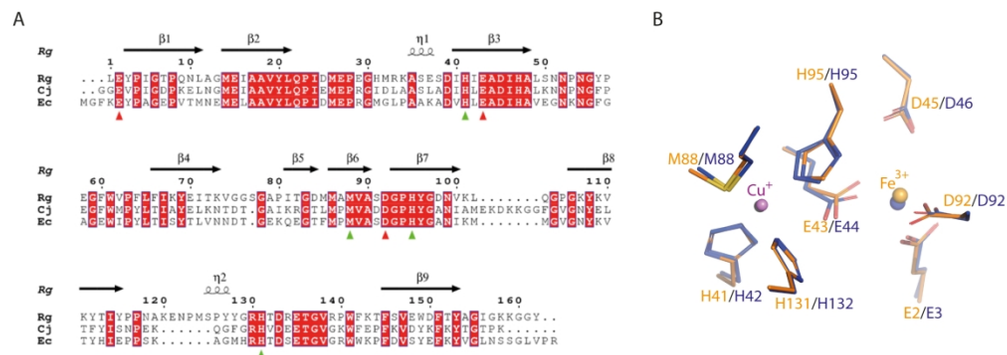


Figure 12

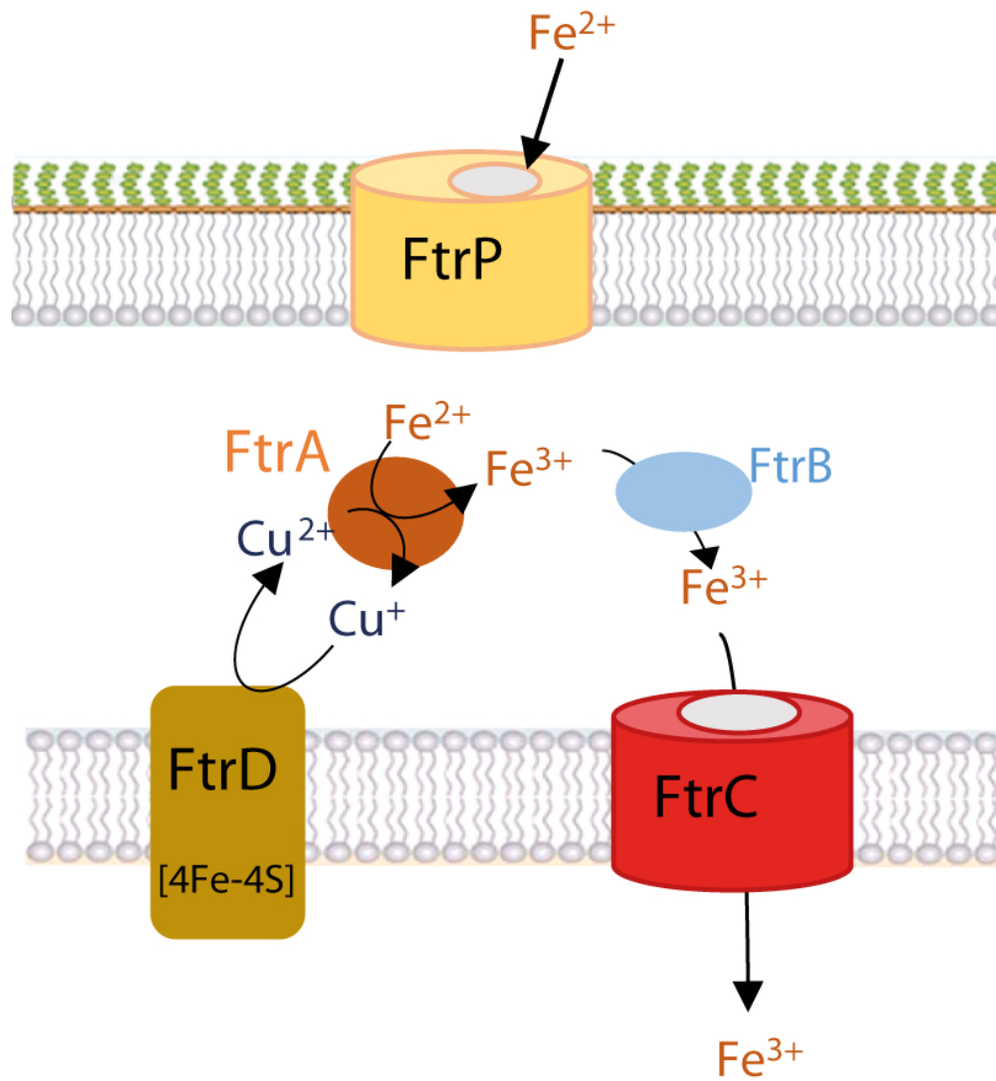


Figure 14

68x76mm (300 x 300 DPI)