Does a low-pH microenvironment around phototrophic FeII-oxidizing bacteria prevent cell encrustation by FeIII minerals?

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Abstract
Neutrophilic FeII-oxidizing bacteria precipitate positively charged FeIII minerals that are expected to sorb to the negatively charged cell surface, leading to encrustation and thus limiting the cells’ accessibility to substrates and nutrients. However, electron-microscopy analysis of phototrophic iron-oxidizing Thiodictyon sp. strain F4 cells showed no encrustation, but mineral precipitation at a distance from the cell surface. In situ fluorescence microscopy analysis of F4 cells using a pH-sensitive fluorescent dye revealed a low cell surface pH (6.0 ± 0.1) in contrast to the bulk pH (6.6 ± 0.1). Biogeochemical modeling showed that the pH difference reduces FeIII sorption and FeIII precipitation rates at the cell surface, therefore directing mineral formation away from the cells. The results from this study therefore suggest that the establishment of a low cell surface pH could provide a mechanism for photoferrotrophs to successfully prevent FeIII mineral precipitation on the cell surface.

Introduction
While FeII is relatively soluble at a neutral pH, FeIII preferentially occurs in the solid phase as FeIII [FeIIIaq] of approximately \(1 \times 10^{-10}\) mol L\(^{-1}\) (Stumm & Morgan, 1995). Iron(III) (oxy)hydroxides [FeIIIppt] are produced chemically and biologically in the environment (Emerson, 2000; Ehrlich & Newman, 2009) by microbial and chemical iron(II) [FeII] oxidation at a neutral pH (Widdel et al., 1993; Hafenbradl et al., 1996; Straub et al., 1996; Emerson, 2000; Canfield et al., 2005). The phenomenon of microbial FeIII oxidation was discovered already in the early days of microbiology in the 19th century (Ehrenberg, 1836; Winogradsky, 1888). At that time, though, the interaction of iron(III) oxides [FeIIIoxides] with bacteria could only be observed visually, but neither the corresponding metabolism nor the underlying mechanism could be explained.

While aerobic neutrophilic FeII oxidizers Gallionella and Leptothrix thrive at low concentrations of oxygen, phototrophic FeII oxidizing bacteria, as well as nitrate-reducing FeII oxidizers, convert FeII under anoxic conditions. FeII-oxidizing bacteria can access the dissolved substrate (FeIIaq) easily at circumneutral pH. However, under such pH conditions, the cells may be harmed by the poorly soluble product FeIIIppt of their metabolism because FeIIIaq as well as the positively charged FeIIIppt strongly sorb to the negatively charged bacterial cell surfaces. Consequently, cells are expected to encrust with FeIIIppt, leading to a limited exchange of metabolites with their environment. In particular, the FeIII-oxidizing nitrate-reducing Acidovorax sp. strain BoFeN1 was shown to encrust heavily with FeIIIppt at circumneutral pH (Miot et al., 2009a; Schaedler et al., 2009).

In this study, we focus on a photoferrotrophic strain that oxidizes FeII according to the following equation (Widdel et al., 1993):

\[
4\text{Fe}^{2+} + h\nu + \text{HCO}_3^- + 10\text{H}_2\text{O} \rightarrow 4\text{Fe(OH)}_3^{\text{III}} + \text{CH}_2\text{O} + 7\text{H}^+ \tag{1}
\]

It has been shown that all electrons stemming from FeII oxidation are used by phototrophic FeII-oxidizing bacteria for CO2 fixation and thus biomass generation (Widdel et al., 1993). Recently, it was suggested for two different strains of
phototrophic FeII-oxidizing bacteria that the oxidation of FeII occurs in the periplasm of the cells (Croal et al., 2007; Jiao & Newman, 2007). In contrast to chemotrophic FeIII-oxidizing strains, none of the known photoferrotrophs encrusts with FeIII minerals (Miot et al., 2009b; Schaedler et al., 2009). It is particularly essential for phototrophic FeIII-oxidizing strains to avoid cell encrustation as the mineral coatings may not only limit the exchange of nutrients and metabolites of the cell with its environment. Additionally, the amount of light available for photosynthesis decreases due to the mineral crust and therefore reduces the total energy available to the cell.

Various mechanisms have been suggested to explain why some FeII-oxidizing strains do not encrust: (1) organic exopolymers, similar to those observed for Gallionella spp., could help to localize precipitation away from the cell (Chan et al., 2009; Schaedler et al., 2009 and references therein). Still, this mechanism does not explain why the FeII released does not immediately bind to the negatively charged cell surface after FeII oxidation. (2) Shedding of parts of the cell surface covered with Fe precipitates could remove the precipitates from the surface (Emerson & Revsbech, 1994). (3) FeIII complexation by organic ligands allows FeIII to remain in solution and prevents precipitation (Croal et al., 2004a). However, in the case of autotrophic organisms, CO2 reduction for the synthesis of such ligands is quite costly in terms of the Fe requirement. Four atoms of FeII have to be oxidized to reduce a single CO2 to biomass (e.g. usable in a complexing molecule keeping FeIII in solution) Eqn. (1).

Several carbon atoms are usually required per ligand. For example, 36 FeII would have to be oxidized to produce a single ligand molecule with six carbon atoms (with a C oxidation state of 0) and this carbon would not be available for cell biomass while it could complex only one FeIII atom. Organic ligands (such as oxalate) requiring very few electrons for their synthesis from CO2 could potentially be suitable as complexing agents. However, although oxalic acid would fulfill this requirement by needing only two electrons to be synthesized from two CO2 molecules, an effective recycling of any of these ligands is unlikely, mainly because it does not penetrate the cell itself (Invitrogen, 2003; Marcotte & Brouwer, 2005) by Molecular Probes®. (2) Shedding of parts of the cell surface covered with Fe precipitates could remove the precipitates from the surface (Emerson & Revsbech, 1994). (3) FeIII complexation by organic ligands allows FeIII to remain in solution and prevents precipitation (Croal et al., 2004a). However, in the case of autotrophic organisms, CO2 reduction for the synthesis of such ligands is quite costly in terms of the Fe requirement. Four atoms of FeII have to be oxidized to reduce a single CO2 to biomass (e.g. usable in a complexing molecule keeping FeIII in solution) Eqn. (1). (4) The amount of negatively charged (and therefore FeIII binding) functional groups on the cell surface could be reduced or the cell surface charge could even be reversed from negative to positive to avoid primary precipitation (Beveridge & Murray, 1980; Urrutia Mera et al., 1992). This has been shown only for cyanobacteria (McConnaghey & Whelan, 1997; Martinez et al., 2010), but it is unknown whether photoferrotrophs are also able to change their surface charge. (5) FeIII precipitation at the cell surface could be delayed, lowered or even prevented by acidifying the cell microenvironment (Sobolev & Roden, 2001; Kappler & Newman, 2004; Schaedler et al., 2009).

An environment of enhanced proton activity around the cell (termed 'low-pH microenvironment' in this study) has the advantage for a cell that it does not require additional biomass to be produced for the complexation of FeIII by organic ligands, shedding of parts of the cell or localization of precipitates on exopolymers (e.g. stalks). In the case of phototrophic FeIII-oxidizing bacteria, acidification around colonies of cells has been observed previously (Kappler & Newman, 2004). However, it is unclear whether this acidification stems from the formation of Fe(OH)3 from Fe3+ that would release protons [according to Eqn. (1), where FeII oxidation is coupled to CO2 fixation] or whether the lower pH was actively caused by the cells via proton translocation.

Therefore, the objectives of this study were (1) to analyze the pH at the single-cell level compared with the bulk medium using the phototrophic FeIII-oxidizing bacterium Thiodictyon sp. F4, a gram-negative, γ-proteobacterium (Croal et al., 2004b), and (2) to evaluate whether and how the establishment of a single-cell pH-microenvironment influences FeIII sorption and FeIII mineral precipitation. Micrometer pH measurements were achieved using a combination of confocal laser scanning microscopy (CLSM) and a pH-dependent fluorescent dye. Geochemical modeling was used to determine the pH dependence of FeIII sorption and precipitation and to evaluate whether an actively sustained pH microenvironment might prevent encrustation.

### Materials and methods

#### Medium, chemicals and growth conditions

The phototrophic FeIII oxidizer Thiodictyon sp. strain F4 (Croal et al., 2004b) was grown in a mineral medium buffered with 22 mM bicarbonate at pH 6.8–6.9. The addition of 10 mM FeCl2 as electron donor decreased the pH slightly to pH 6.6. For a detailed description of the growth conditions, see Hegler et al. (2008). The fluorescent dye SNARF®-4F 5-(and-6)-carboxylic acid (SNARF4F) (Invitrogen, 2003; Marcotte & Brouwer, 2005) by Molecular Probes® was dissolved in water (2 mM) and aliquots were frozen and used not longer than 4 months to avoid the potential hydrolytic degradation of the dye. Samples of Thiodictyon sp. strain F4 cultures were taken at the end of the exponential growth phase and 500 µl of the sample was mixed with 1 µl of the dye solution in an anoxic glovebox (100% N2) in order to avoid oxygen exposure (final concentration of the dye: 4 µM). Although some association of the dye and the cell wall may occur, we chose this dye because it does not penetrate the cell itself (Invitrogen, 2003). The potential association of the dye with the cell...
surface was not expected to affect its pH-dependent fluorescence (Invitrogen, 2003).

**Microscopic analysis of a single-cell pH microenvironment**

An oxygen-tight microscopy chamber covered with a coverslip was used to avoid physical pressure to the minerals and to maintain the sample anoxic during microscopy. The chambers were prepared by gluing a coverslip onto the microscopy slide with epoxy-based glue (Uhu plus schnell-fest; UHU GmbH & Co. KG, Germany) including two syringe needles as the inlet and the outlet. They were washed with double-distilled water before use.

The chambers were filled with the culture/dye mixture and incubated for 45 min before microscopic analysis at light saturation. This duration of time allows CCCP to deenergize the membranes. As it was recently suggested that Fe$^{II}$ oxidation occurs in the periplasm of phototrophic Fe$^{II}$-oxidizing bacteria (Croal *et al.*, 2007; Jiao & Newman, 2007), no active (energy-requiring) uptake of Fe$^{II}$ into the cytoplasm is needed and therefore Fe$^{II}$ oxidation did not stop immediately after the addition of CCCP. However, reducing equivalents accumulate due to the oxidation of Fe$^{II}$, but cannot be reoxidized; hence, finally, Fe$^{II}$ oxidation will stop due to the lack of electron acceptors.

Microscopy was performed on a Leica TCS SP2 confocal scanning laser microscope. The dye was excited at 488 nm, and the fluorescence was measured at emission wavelengths of 580–590 and 650–660 nm (Fig. 1a–c). The ratio of the two wavelengths was used to calculate the pH according to a calibration curve. Continuous spectra for dye calibration were measured in 20 mM PBS buffer with a pH between 5 and 7.5 using a Jena Analytics FlashScan 550 fluorescence plate reader at an excitation of 488 nm. The ratio between the individual wavelengths (580–590 and 650–660 nm) was calculated depending on the pH. We fitted the curve to account for the nonlinear parts away from the pKₐ of the

![Fig. 1](image-url). Confocal laser scanning microscopy (CLSM) imaging of *Thiodictyon* sp. F4 cells stained with a pH-dependent fluorescent dye, demonstrating the presence of a cell pH microenvironment. (a–c) CLSM pictures of cells stained with SNARF4. (a) 580–590 nm, (b) 650–660 nm, (c) gray-scale image. The ratio of (a)/(b) allows calculating the pH. (d) pH microenvironment around cells of *Thiodictyon* sp. F4, the pH was 6.6 ± 0.1 in the background (yellow box) and mineral (red box), bulk measurements with a pH electrode confirmed the values determined with the pH-dependent dye. The pH at the cell level was 5.9 ± 0.1 (green cell) and 6.0 ± 0.1 (blue cell). (e) Red, background; green, cell; blue, mineral, CCCP used to decouple the proton gradient. The pH was 6.6 ± 0.1 at the cell level, the background also had pH 6.6 ± 0.1 for both areas, the pH at the mineral was 6.5 ± 0.1 while the entire frame had a pH of 6.6 ± 0.1. Bulk measurements with a pH electrode confirmed these values. Scale bar = 2 μm in all images (a)–(e). (f) Scanning electron micrograph of *Thiodictyon* sp. F4, scale bar = 1 μm. The cell is associated with Fe$^{III}$ minerals, but not encrusted.
dye. The ratio between the individual wavelengths and its correlation to the pH was used to calculate the pH values in the microscopy images.

In order to determine whether the establishment of the ΔpH in the cell microenvironment is an active or a passive process, carbonyl cyanide 3-chlorophenylhydrazone (CCCP; Sigma Aldrich, Germany) was used – a protonophore that decouples the proton-motive force (PMF) across the membrane (Harold, 1972).

Electron microscopy

For scanning electron microscopy, samples were prepared as described previously (Schaedler et al., 2008).

Calculation of the FeII oxidation rate per cell

In order to calculate the amount of FeII oxidized per cell per unit time, we quantified the decrease of total FeIII over time as described previously (Hegler et al., 2008). Additionally, cells were counted microscopically: 1 mL culture was fixed with glutaraldehyde (final concentration of 2.5%), followed by the dissolution of the minerals at pH 3 with oxalic acid (15 g L−1) and oxalate (28 g L−1). After the dissolution of the FeIII and staining of the cells with 4’,6-diamidino-2-phenylindol (DAPI), a subsample was passed through a Nucleopore-filter (0.22 μm pore size, Millipore) and the amount of cells per filter area was counted in an epi-fluorescence microscope. The data allowed calculating the maximum FeIII production rates per cell, which were used for calculating the sorption of FeIII onto the cell surface. The bulk FeIII formation rate was determined by following FeII oxidation over time (which is equal to FeIII formation) and calculating the turnover in the entire culture over the complete oxidation phase of 218 h. The release of FeIII per second was calculated to be 4.3 × 10−7 mM s−1.

Modeling

The resolution of the microscope did not allow resolving the expected pH gradient between the cell surface and the bulk medium. Thus, a gradient of proton activity was interpolated between the cell surface and the bulk medium for modeling. Additionally, geochemical modeling was applied to evaluate the sorption of ferric iron to the cell surface. Unfortunately, cell surface titrations could not be used to quantify the different functional groups involved in mineral sorption processes at the cell surface of the FeII-oxidizing strain because the FeIII minerals (produced during FeII oxidation) interfered with the titrations. Microscopic analysis revealed (data not shown) that dissolving the minerals killed the cells, partially destroyed the cell integrity and increased the number of (titratable) functional groups originating from the cell interior, likely falsifying the titration measurements. Additionally, sorption of FeIII ions to cell surfaces cannot be quantified directly at circumneutral pH due to the low solubility of the FeIII at this pH.

However, it has been shown that as an alternative approach, a general model for site densities and pK values can be used to model the adsorption of metals to cell surfaces (Fein et al., 2001). Yee & Fein (2003) showed that the adsorption behavior of different physiological and phylogenetic groups of bacteria, gram-positive as well as gram-negative, and cell mixtures is very similar for various metals. The most abundant cell surface moieties are carboxyls (Borrok et al., 2004). Other functional groups such as protonated amines deprotonate at pH values well above 7 and are thus positively charged at circumneutral pH values. Following the approach of Yee & Fein (2003), we quantified the cell surface interaction of FeIII with the most abundant functional group (carboxyl group).

Calculation of FeIII solubility

Geochemists Workbench 6.04 by Rockware® was used to calculate the speciation and thus the solubility of FeIII in the medium. A matrix including all the chemical compounds present in the medium was set up, see Table 1. For the thermodynamic equilibrium calculations, the dissolved ferric iron concentration (FeIII) was defined as follows (including the most dominant FeIII species):

$$\text{Fe}^{III}_{\text{aq}} = \text{Fe(OH)}_3^0 + \text{Fe(OH)}_2^+ + \text{FeHPO}_4^+ + \text{Fe(OH)}_4^-$$
$$+ \text{FeOH}^{2+} + \text{FeCO}_3^+ + \text{Fe}_3(\text{OH})_4^{3+}$$
$$+ \text{FeH}_2\text{PO}_4^{2+} + \text{Fe}_2(\text{OH})_3^{2+} + \text{Fe}\text{SO}_4^{2-} + \text{Fe}^{3+} + \cdots$$

(2)

Table 1. Settings and concentrations of all the chemical compounds included in the matrix used to calculate the solubility of FeIII in mineral medium over the pH range from 6 to 7. The resulting graph depicts all dissolved FeIII species (Fig. 2)

| Component added | 1 kg H2O | 4.4 mmol L−1 H3PO4 | 22 mmol L−1 HCO3 | 0.68 mmol L−1 Ca2+ | 4.4 mmol L−1 K+ | 2 mmol L−1 Mg2+ | 22 mmol L−1 Na+ | 30.2 mmol L−1 Cl− | 2 mmol L−1 SO4 | 6.6 mmol L−1 NH4 | 10 mmol L−1 Fe3+ | 1e-50 mmol L−1 O2(aq) | 6.0 pH slide pH to 7.0 | Decouple FeII and HSO4 | Suppress H3S (aq) and FeHS | Balance off |
|-----------------|----------|----------------------|------------------|------------------|-----------------|-----------------|-----------------|-----------------|-----------------|----------------|----------------|----------------|------------------|-----------------|----------------|----------------|----------------|

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Calculation of iron–cell-surface interaction

The following equation was applied to calculate the sorption of Fe$^{III}$ to the carboxylic groups of the cell surface:

$$[ML] = \frac{[\text{site}_{\text{max}}]K_aK_{\text{ads}}(M/L)}{1 + K_aK_{\text{ads}}(M/L)}$$  \hspace{1cm} (3)

where ML is the ligand–metal complex, $K_a$ is the carboxyl-equilibrium constant, $K_{\text{ads}}$ represents the adsorption constant while $M$ is the metal concentration and $H^+$ the proton concentration; site$_{\text{max}}$ gives the maximal surface site concentration. The maximal site concentration (site$_{\text{max}}$) represents approximately the total amount of −COOH surface sites at pH > 5.

Here, we briefly derive Eqn. (3) with the already introduced abbreviations: deprotonation of carboxylic groups is described by

$$R – COOH \rightleftharpoons R – COO^- + H^+$$  \hspace{1cm} (4)

Replacing $R – COO^-$ by $L^-$ and $R – COOH$ by $L$ and calculating the deprotonated ligand concentration, $L^-$ yields

$$L^- = [K_a][L]/[H^+]$$  \hspace{1cm} (5)

This formula determines the concentration of the deprotonated ligands at the cell surface with respect to pH. The adsorption equilibrium can be expressed by

$$M + L^- \rightleftharpoons ML$$  \hspace{1cm} (6)

which can be used to calculate the metal–ligand concentration

$$[ML] = K_{ads}[M][L^-]$$  \hspace{1cm} (7)

The substitution of $L^-$ of Eqn. (5) into Eqn. (7) leads to

$$[ML] = \frac{K_{ads}K_a[M][L]}{[H^+]}$$  \hspace{1cm} (8)

which – after constraining the maximal sorption with a Langmuir isotherm – yields Eqn. (3). A list of all the parameters used in the calculations is compiled in Table 2.

In the calculations, we did not consider abiotic processes that may initiate or stimulate precipitation such as for example the initial presence of nucleation sites (e.g. by the presence of siderite and vivianite crystallites in the growth medium).

Calculation of the precipitation kinetics of Fe$^{III}_{\text{ppt}}$

For the kinetic precipitation calculation, the same speciation setup (Table 1) was used while the reaction rate constant for precipitation ($k_{\text{precipitation}} = 2.0 \times 10^7$ M$^{-1}$ s$^{-1}$, for comparison reaction rate constant for diffusion $k_{\text{diffusion}} = 7.9 \times 10^6$ M$^{-1}$ s$^{-1}$) was taken from Pham et al. (2006). In order to determine the precipitation kinetics, we measured the rate of Fe$^{III}$ production for Thiodictyon sp. strain F4 (4.3 $\times$ 10$^{-7}$ mM s$^{-1}$) in a growth experiment (Table 2). The rate of precipitation at the cell surface (at pH 6.0) and in the bulk (at pH 6.6) was calculated accounting for a constant Fe$^{III}_{\text{aq}}$ release of 4.3 $\times$ 10$^{-7}$ mM (per second).

Results and discussion

Absence of encrustation and establishment of a pH microenvironment around phototrophic Fe$^{II}$-oxidizing Thiodictyon sp. strain F4 cells

Scanning electron microscopy showed that cells of the phototrophic Fe$^{II}$-oxidizing Thiodictyon strain sp. F4 do not become encrusted when oxidizing Fe$^{II}$ at neutral pH (Fig. 1). This suggests that this strain might be able to minimize or even avoid the sorption of Fe$^{III}_{\text{aq}}$ to the cell surface in order to avoid the precipitation of Fe$^{III}_{\text{ppt}}$ on its cell surface. To determine whether a low cell surface pH exists for F4 cells and is possibly involved in preventing cell encrustation, we incubated the cells with a dye that shows pH-dependent variations in fluorescence emission.

We determined the pH in the bulk medium with the fluorescent dye in comparison with the values obtained using a standard pH electrode and obtained a value of 6.6 ± 0.1 using both independent methods. As can be seen in Fig. 1, for most active cells, we detected that the cell surface pH was at least 0.6 ± 0.1 units lower compared to the bulk medium of the phototrophic Fe$^{II}$-oxidizing strain Thiodictyon sp. F4. The same effect was observed for approximately 16 out of 20 investigated cells. However, we even found some cells with a larger shift of 0.8 ± 0.1 pH units between surface and bulk pH, although cells with such a larger pH shift were less frequent (about 10–15% of all cells). A few cells (~10%) showed only a very small pH difference from the bulk pH, possibly representing inactive microbial cells. The cells with the highest and the lowest pH variation were not considered in the subsequent geochemical calculations.

In order to test whether an active cell metabolism is necessary to establish the low cell surface pH, we decoupled the membranes with CCCP, a protonophore (Harold, 1972). As a consequence of the exposure to CCCP, the cells were no
ultimately used to fix CO$_2$ and thus produce biomass Eqn. carboxyl- and phosphoryl-functional groups. Calculations iron, compete directly with protons for binding sites at groups. In the initial adsorption step, metal ions, such as released initially by the cell) to cell surface functional groups. If enough light is present, a single electron can theoretically be used several times in the cyclic electron flow to generate the pH gradient across the membrane by pumping protons and creating a lower pH outside of the cell. For photoferro-trophs, the electrons stem from the oxidation of Fe$^{II}$ and are ultimately used to fix CO$_2$ and thus produce biomass Eqn. (1). Because continuous pumping of protons out of the cell interior (cytoplasm) to the periplasm would ultimately lead to an increasing internal pH being lethal in the end, the protons necessary for the low pH at the cell surface likely stem – at least to a significant extent – from the hydrolysis of Fe$_{aq}^{III}$ by H$_2$O molecules after the release of Fe$_{aq}^{III}$ from the Fe$^{II}$/Fe$^{III}$ oxidase. The hydrolysis leads first to the formation of [Fe$^{III}$(OH)(H$_2$O)$_5$]$^{2+}$ and finally to Fe(OH)$_3$ monomers, which then finally precipitate as Fe$^{III}$ hydroxide minerals.

**Effect of the low cell surface pH on the sorption, precipitation and dissolution of Fe$^{III}$**

After oxidation of Fe$^{II}$ by the cell and the release of Fe$_{aq}^{III}$, the cation either adsorbs to the negatively charged cell surface functional groups or diffuses away from the cell. The Fe$^{III}$-oxidizing enzymes likely release Fe$_{aq}^{III}$ and not a hydrolyzed Fe$^{III}$ species. The interaction of metals and metalloids with cell surfaces can be described as sorption, followed by precipitation of the metal ions as minerals (Beveridge & Murray, 1976). Thus, we calculated the sorption of Fe$_{aq}^{III}$ (as released initially by the cell) to cell surface functional groups. In the initial adsorption step, metal ions, such as iron, compete directly with protons for binding sites at carboxyl- and phosphoryl-functional groups. Calculations for carboxyl functional groups showed that the sorption of Fe$^{III}$ to carboxylic surface functional groups decreases by > 80% with a decline in pH of 0.6 units (Fig. 2). These results agree well with other studies that showed that Co$^{III}$, Nd$^{III}$, Ni$^{III}$, Sr$^{III}$, Zn$^{III}$ (Fein et al., 2001) and Fe$^{III}$ (Warren & Ferris, 1998) adsorb less to cell surfaces at a lower pH. Additionally, (Urrutia Mera et al., 1992) showed that cells of *Bacillus subtilis* bind heavy metals such as U and Sc substantially less to their cell surface when the cells were alive (with a functioning pmf). Less adsorption of metals due to competition with H$^{+}$ leads to fewer potential nucleation sites for mineral precipitation (Urrutia Mera et al., 1992; Fein et al., 2001).

The lower pH not only influences the competition of the Fe$_{aq}^{III}$ ions with protons for the cell surface-binding sites, but it also changes the solubility of Fe$^{III}$ itself. Because the concentration of Fe$_{aq}^{III}$ ions and therefore the solubility of the Fe$^{III}$ determine the kinetics and the extent of precipitation (Grundl & Delwiche, 1993; Pham et al., 2006; Rose & Waite, 2007), we determined the effect of a pH change of 0.6
from pH 6.6 to 6.0 on the solubility of Fe$^{III}$ (Fig. 2). Geochemical modeling showed that a lower pH of 6.0, which is present at the cell surface, will increase the fraction of Fe$^{III}$ that is in solution (Fe$_{aq}^{III}$) approximately fivefold compared with the bulk pH of 6.6, i.e. the concentration of Fe$^{III}$$_{aq}$ is only 20% at pH 6.6 compared with pH 6.0. Similar to that for chemical Fe$^{III}$ oxidation, probably also during microbial Fe$^{II}$ oxidation, the rate-limiting step in the transition from Fe$^{III}$$_{aq}$ to Fe$^{III}$$_{ppt}$ via Fe$^{III}$$_{eq}$ is the nuclease and precipitation of the Fe$^{III}$ mineral (Grundl & Delwiche, 1993). The precipitation rate of Fe$^{III}$ therefore decreases with decreasing pH. At pH 6.0 (cell), the precipitation rate was determined to be $8.3 \times 10^{-19}$ mM s$^{-1}$ whereas at pH 6.6 (bulk) the rate increased to $1.3 \times 10^{-9}$ mM s$^{-1}$. All in all, equilibrium calculations showed that the Fe$^{III}$$_{aq}$ concentration is higher at a lower pH. Additionally, precipitation of Fe$^{III}$ minerals in the bulk medium is faster because the pH in the bulk is higher than that at the cell surface. Therefore, a concentration gradient of Fe$^{III}$$_{aq}$ establishes between the cell surface and the bulk, with a higher Fe$^{III}$$_{aq}$ concentration at the lower pH at the cell surface and a lower Fe$^{III}$$_{aq}$ concentration at the higher pH in the bulk medium. The concentration gradient of Fe$^{III}$$_{aq}$ drives diffusion of Fe$^{III}$$_{aq}$ from low-pH zones at the cell surface (low precipitation, higher Fe$^{III}$$_{aq}$) to high-pH regions further away from the cell (faster precipitation, low Fe$^{III}$$_{aq}$, high mineral content) (Fig. 3).

Consequences of the low cell surface pH for cell encrustation and metal binding in photoferrotrophs and implications for chemotrophic Fe$^{II}$ oxidizers

In this study, we demonstrated the establishment of a single-cell low-pH microenvironment at the cell surface of the phototrophic Fe$^{II}$-oxidizer *Thiobacillus* sp. strain F4 using an in-situ approach with a pH-dependent fluorescent dye in combination with confocal laser scanning microscopy. The observed pH decrease at the cell surface has three consequences for the cell: (1) a successful competition of protons with the Fe$^{III}$$_{aq}$ for sorption sites at the cell surface leading to less binding of Fe$^{III}$ to the cell surface, (2) an increase in Fe$^{III}$ solubility and (3) a lower precipitation rate of Fe$^{III}$$_{ppt}$ at the cell surface (Fig. 3). This suggests that the establishment of a cell pH microenvironment could provide a mechanism for photoferrotrophs to successfully prevent Fe$^{III}$ mineral precipitation on the cell surface. The results of this study may therefore explain why none of the known phototrophic Fe$^{II}$-oxidizing strains encrusts (Fig. 1). An exception is the purple nonsulfur bacteria, *Rhodopseudomonas palustris*, which differs from the other photoferrotrophs in that it is a mixotrophic Fe$^{II}$ oxidizer, i.e. it needs an organic substrate in addition to the Fe$^{II}$, and it is not clear whether this strain benefits from Fe$^{II}$ oxidation at all (Heising & Schink, 1998; Schaedler et al., 2009). The benefits for a cell to avoid encrustation are obvious because (1) more light can reach the cell surface, leading to a higher metabolic rate, and (2) the uptake of nutrients from the environment and the release of toxic metabolites to the surrounding is not hampered.

For chemotrophic Fe$^{II}$-oxidizing strains, other mechanisms to prevent cell encrustation are expected, because these microorganisms depend solely on the energy provided by the chemotrophic oxidation of Fe$^{II}$ with either O$_2$ or nitrate as electron acceptor. Protons therefore cannot be pumped using light energy as by the phototrophs in cyclic photophosphorylation where a single electron can be used multiple times to pump protons as long as light energy is available. In chemotrophic Fe$^{II}$ oxidizers, proton translocation is directly coupled to chemotrophic Fe$^{II}$ oxidation and electron transport phosphorylation, where one electron stemming from the oxidation of Fe$^{II}$ runs through the electron transport chain once, finally being used to reduce O$_2$ or nitrate. It has been shown that some nitrate-reducing mixotrophic Fe$^{II}$-oxidizing bacteria encrust with Fe$^{III}$$_{ppt}$ on the cell surface (Miot et al., 2009a; Schaedler et al., 2009). Although these bacteria also utilize the PMF to generate ATP and to produce reducing equivalents, the energy available to these bacteria is much less compared with the phototrophic strains because phototrophic organisms can tap the energy source of the solar radiation. In contrast, chemotrophic organisms (such as nitrate-reducing and microaerophilic Fe$^{II}$-oxidizing bacteria) are limited by the energy that can be obtained from Fe$^{II}$ oxidation, which makes the generation of a large proton gradient less likely or even impossible.

The results presented here shed light on a long lasting enigma: already in 19th century Ehrenberg (1836) and...
Winogradsky (1888) had observed biogenic FeIII minerals and speculated on their formation and the influence of bacteria. Although many open questions remain, today, almost 200 years later, we understand the complexity of the interaction between bacteria and the FeIII minerals better. The presence of pH microenvironments around phototrophic FeIII oxidizers provides a possible explanation as to why and how phototrophic FeIII-oxidizing bacteria avoid encrustation by FeIII minerals. Additionally, the light dependence of this metabolism potentially explains why phototrophic FeIII oxidizers use a different strategy to avoid cell encrustation than the microaerophilic FeII-oxidizing bacteria, such as Gallionella and Leptothrix.

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