



CO₂/HCO₃⁻ Accelerates Iron Reduction through Phenolic Compounds

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ABSTRACT Iron is a vital mineral for almost all living organisms and has a pivotal role in central metabolism. Despite its great abundance on earth, the accessibility for microorganisms is often limited, because poorly soluble ferric iron (Fe³⁺) is the predominant oxidation state in an aerobic environment. Hence, the reduction of Fe³⁺ is of essential importance to meet the cellular demand of ferrous iron (Fe²⁺) but might become detrimental as excessive amounts of intracellular Fe²⁺ tend to undergo the cytotoxic Fenton reaction in the presence of hydrogen peroxide. We demonstrate that the complex formation rate of Fe³⁺ and phenolic compounds like protocatechuic acid was increased by 46% in the presence of HCO_3^- and thus accelerated the subsequent redox reaction, yielding reduced Fe²⁺. Consequently, elevated CO₂/HCO₃⁻ levels increased the intracellular Fe²⁺ availability, which resulted in at least 50% higher biomass-specific fluorescence of a DtxR-based Corynebacterium glutamicum reporter strain, and stimulated growth. Since the increased Fe²⁺ availability was attributed to the interaction of HCO_3^- and chemical iron reduction, the abiotic effect postulated in this study is of general relevance in geochemical and biological environments.

IMPORTANCE In an oxygenic environment, poorly soluble Fe^{3+} must be reduced to meet the cellular Fe^{2+} demand. This study demonstrates that elevated CO_2/HCO_3^{-} levels accelerate chemical Fe^{3+} reduction through phenolic compounds, thus increasing intracellular Fe^{2+} availability. A number of biological environments are characterized by the presence of phenolic compounds and elevated HCO_3^{-} levels and include soil habitats and the human body. Fe^{2+} availability is of particular interest in the latter, as it controls the infectiousness of pathogens. Since the effect postulated here is abiotic, it generally affects the Fe^{2+} distribution in nature.

KEYWORDS iron homeostasis, iron reduction, carbon dioxide, bicarbonate, DtxR, pathogens, *Corynebacterium glutamicum*, iron homeostasis

ron is a vital mineral for almost all living organisms and participates in inevitable electron transfer reactions since it serves a redox potential from -500 to +300 mV by switching between two oxidation states (Fe²⁺/Fe³⁺) (1). Despite its great abundance in the Earth's crust (representing the fourth most abundant element), its availability for organisms is limited, because it is mainly present as poorly soluble Fe³⁺ (10⁻¹⁸ M at pH 7.0) in an oxidative environment (2). Sophisticated strategies to increase its accessibility have evolved, including the secretion of siderophores and smaller iron chelators that enhance the solubility of Fe³⁺ (3). Iron complexes are then taken up via designated transport systems in an energy-dependent fashion (2). Since reduced Fe²⁺ is incorporated as a prosthetic group in a number of enzymes that belong to respiratory complexes or that participate in tricarboxylic acid (TCA) cycle reactions and stress response inside the cell, the reduction of Fe³⁺ is of central relevance. Several studies on the characterization of ferric reductase activity are reviewed by Schröder et **Citation** Müller F, Rapp J, Hacker A-L, Feith A, Takors R, Blombach B. 2020. CO₂/HCO₃⁻ accelerates iron reduction through phenolic compounds. mBio 11:e00085-20. https://doi .org/10.1128/mBio.00085-20.

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Received 13 January 2020 Accepted 31 January 2020 Published 10 March 2020 al. (4). It turned out that most designated ferric reductases are in fact flavin reductases that regenerate the substrate for chemical iron reduction. This is not limited to intracellular reactions, since it was shown recently that the human pathogen *Listeria monocytogenes* possesses an extracellular electron transfer apparatus, in which iron reduction is mediated by flavin (5).

In the Gram-positive soil bacterium *Corynebacterium glutamicum*, which is used in amino acid production on an industrial scale, little is known about iron uptake mechanisms because most annotated genes lack experimental validation (6, 7). During the development of a suitable minimal medium for *C. glutamicum*, the addition of citrate (8) or small amounts of the diphenolic compounds catechol or protocatechuic acid (PCA) (3,4-dihydroxybenzoic acid) revealed a growth-promoting effect, which was attributed to the improved iron uptake of Fe³⁺ chelates (9). To ensure sufficient initial iron uptake, PCA became a component of the widely used CgXII medium (10).

Excessive amounts of intracellular iron are deposited in specialized storage proteins (ferritin [Ftn] and Dps) and can be remobilized from there in times of iron starvation (2). A novel mode of iron remobilization was identified recently in *C. glutamicum*. Pupylation of a surface lysine residue of the Ftn and Dps proteins initiates the unfolding and subsequent Fe^{3+} release in the cytosol. However, the mechanism responsible for reduction of Fe^{3+} remains unclear (11).

Diphenolic substances (e.g., 2,3-dihydroxybenzoic acid [2,3-DHB] and derivatives) are secreted under iron-restricted growth conditions by a variety of organisms such as *Bacillus subtilis* and *Paracoccus (Micrococcus) denitrificans* (3, 12, 13). It was suggested that 2,3-DHB and derivatives are involved in iron uptake, as mutants deficient in the biosynthesis of 2,3-DHB and derivatives imported less iron, and adding these compounds enhanced iron transport (14). PCA is a structural relative of the common catecholate-type siderophore precursor 2,3-DHB and represents the basic compound of the rather unusual siderophore petrobactin (15). *Bacillus anthracis* and *Bacillus cereus* secrete great amounts of PCA in response to iron limitation (16–18). As a diphenolic Fe³⁺ chelator (19), it was expected to serve a function analogous to 2,3-DHB enhancing iron uptake, although evidence for this had not been provided (16).

Besides chelating iron, diphenols have the potential to reduce Fe^{3+} (20). The redox reaction in Fe^{3+} -PCA chelates (19, 21)—and generally speaking in Fe^{3+} -monocatecholate complexes—proceeds via the sequential oxidation of two adjacent hydroxyl groups to the respective semiquinone and quinone, providing two electrons per molecule for the reduction of Fe^{3+} (20). The reaction depends on an acidic pH, as iron and PCA bind in an 1:1 stoichiometry at a pH of <4.5 (19), and the redox reaction is inhibited by the presence of (at least) the biscatecholate complex (20, 21).

The phenotypical response of bacteria to different CO₂/HCO₃⁻ levels has been extensively reviewed with a biotechnological, pathogenic, or environmental focus (22–25). On the one hand, the effect on cell viability might be detrimental, considering that CO₂-based sterilization operating at high pressure is used in food processing. However, on the other hand, a class of bacteria called capnophiles demonstrates that HCO_3^- can be a vital substrate as they are unable to grow unless sufficiently high CO_2 levels are established (26). Carboxylation reactions might generally benefit from the increased availability of CO₂/HCO₃⁻⁻. C. glutamicum is especially well equipped with enzymes catalyzing these reactions, e.g., at the phosphoenolpyruvate (PEP)-pyruvateoxaloacetate node (27). As a consequence, higher product (e.g., succinate) and biomass yields per substrate were reported in C. glutamicum at elevated CO_2 levels and could be attributed particularly to an increased flux via the anaplerotic reactions (28-30). Anaerobic growth on glucose and tryptone could be enhanced by increasing concentrations of CO_2 . The authors suggest that this might enhance acetyl coenzyme A (acetyl-CoA) carboxylation reactions, yielding greater levels of fatty and mycolic acids (31). Although the exponential growth rate of C. glutamicum could not be promoted under aerobic conditions, elevated CO₂ in the inlet air of a bioreactor cultivation provoked the transcriptional response of almost the entire DtxR regulon (30). The master regulator of iron homeostasis in C. glutamicum was originally named after its



FIG 1 Biomass formation of *C. glutamicum* FEM3 (A) and biomass-specific fluorescence after 24 h of cultivation in a parallel bioreactor setup (B). *C. glutamicum* FEM3 was cultivated aerobically in minimal medium with 20 g glucose liter⁻¹ with synthetic air containing 20% CO_2 (21% O_2 , 59% N_2) or ambient air (0.04% CO_2). Data points (A) and bars (B) represent mean values, and error bars indicate standard deviations for three biological independent replicates.

functional homologue in the pathogenic *C. diphtheriae*, where it controls the toxin production in response to the iron availability (32). In fact, to date, many pathogens that induce the expression of toxin genes under iron starvation conditions and typically integrate this signal via the transcriptional regulators Fur, DtxR and homologues thereof are known (33–35). In analogy with the iron limitation, elevated CO_2/HCO_3^- levels represent another suitable indicator for the host environment (22). The expression of toxin-encoding genes in *Vibrio cholerae* or *Pseudomonas aeruginosa* is mediated by the transcriptional regulators ToxT and RegA in response to elevated HCO_3^- concentrations (23). Interestingly, *regA* expression in *P. aeruginosa* is in turn repressed in an iron-dependent manner by a Fur homologue (33). Hence, understanding the interaction between CO_2/HCO_3^- and iron availability is particularly interesting with regard to toxin production by pathogenic bacteria (23).

RESULTS

Elevated levels of CO₂/**HCO**₃⁻ **increase intracellular Fe**²⁺ **availability.** To monitor intracellular Fe²⁺ availability, we constructed the fluorescence-based reporter strain *C. glutamicum* FEM3 which responds to the activation state of DtxR (see Fig. S1 in the supplemental material). Therefore, we chromosomally integrated the *lacl* gene under the control of the *ripA* promoter (P_{*ripA*}), which is controlled by DtxR. High Fe²⁺ concentrations induce DtxR binding to P_{*ripA*}, thus provoking repression of *lacl* and ultimately resulting in *egfp* expression under the control of the strong *tac* promoter (P_{*tac*}). Thus, the genetic circuit of *C. glutamicum* FEM3 constitutes a signal amplifier and a converter of the native repression mechanism.

Under all conditions tested, *C. glutamicum* FEM3 and the wild type (WT) showed identical growth. Cultivating *C. glutamicum* FEM3 under iron starvation (1 μ M) resulted in a marginal increase of the fluorescence per biomass [78 ± 4 arbitrary units (a.u.) · (g_{CDW} liter⁻¹)⁻¹ where g_{CDW} is the cell weight (dry weight) in grams] over the autofluorescence of the WT [64 ± 3 a.u. · (g_{CDW} liter⁻¹)⁻¹] after 24 h. At iron excess (100 μ M), *C. glutamicum* FEM3 yielded a fluorescence per biomass which was about fourfold higher [313 ± 20 a.u. · (g_{CDW} liter⁻¹)⁻¹] at the end of the cultivation compared to conditions under iron starvation (Fig. S2).

Having proven the functionality at different iron concentrations, we cultivated *C*. *glutamicum* FEM3 aerobically in a parallel bioreactor system to evaluate the intracellular Fe²⁺ availability in response to altering CO₂/HCO₃⁻ levels. With 20% CO₂ in the inlet air, *C. glutamicum* FEM3 grew exponentially (growth rate $[\mu] = 0.36 \pm 0.02 \text{ h}^{-1}$) until the carbon source glucose was exhausted (Fig. 1A). In contrast, aeration with an ambient CO₂ concentration (0.04%) provoked biphasic growth of *C. glutamicum* FEM3 with a growth rate of 0.26 \pm 0.05 h⁻¹ and 0.32 \pm 0.02 h⁻¹ in growth phase one and two, respectively (Fig. 1A). Biomass-specific fluorescence started at similar levels in both

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FIG 2 (A) Shaking flask cultivations of wild-type (WT) *C. glutamicum* in minimal medium with 20 g glucose liter⁻¹ without supplement (reference), with 195 μ M PCA, 50 mM NaHCO₃ or a combination of both supplements. (B) Biomass-specific fluorescence of *C. glutamicum* FEM3 after 24 h of cultivation with the indicated supplements. Data points and bars represent mean values with error bars indicating standard deviations for 3 to 10 independent biological replicates. Values that are significantly different by a two-sample *t* test are indicated by bars and asterisks as follows: *, *P* < 0.05; **, *P* < 0.01.

conditions but after 24 h on average reached almost twofold-higher values at 20% CO₂ content, whereas it remained at a low level at the ambient CO₂ concentration (Fig. 1B). Biphasic growth of *C. glutamicum* FEM3 and the WT was also observed on minimal medium containing glucose in shaking flasks at an ambient CO₂ concentration (Fig. 2A). In accordance with the bioreactor cultivations, growth was stimulated by adding 30 mM NaHCO₃, which shortened the initial (nonexponential) growth phase from about 7 h under reference conditions to 4 h and led to a growth rate of 0.40 \pm 0.03 h⁻¹ during the second growth phase. The addition of 195 μ M of the iron chelator PCA restored exponential growth throughout the cultivation (Fig. 2A) and resulted in a slightly higher growth rate compared to the rate for the second growth phase of the reference condition (reference, $\mu = 0.37 \pm 0.01$ h⁻¹; PCA, $\mu = 0.43 \pm 0.01$ h⁻¹).

The biomass-specific fluorescence of the shaking flask approach showed a 0.5- and 3.0-fold increase in response to the supplementation of NaHCO₃ or PCA compared to the reference cultivation after 24 h, respectively (Fig. 2B). Although growth of *C. glutamicum* FEM3 was not further improved by the combined addition of NaHCO₃ and PCA (Fig. 2A), there might be a synergistic effect on fluorescence level (4.2-fold compared to the reference; Fig. 2B), which was, however, not significantly higher than in the PCA-supplemented condition (by Student's *t* test). Supplementing with PCA or NaHCO₃ did not show a growth-stimulating effect in iron-depleted medium (Fig. S3), where biomass stagnated at a final 0.9 to 1.1 g liter⁻¹ regardless of supplementation. In summary, these results demonstrate that in the presence of the extracellular iron source (FeSO₄), elevated levels of CO_2/HCO_3^- and/or the addition of PCA to the minimal medium results in an increased intracellular Fe²⁺ availability in *C. glutamicum*.

CO₂/HCO₃⁻ does not increase thermal stability of the DtxR protein or interact with iron storage and mobilization. The DtxR protein of *C. glutamicum* shares 72% sequence identity with its extensively characterized homologue from *Corynebacterium diphtheriae*, including conserved ligand binding sites. Dimerization of the latter and subsequent DNA recognition are induced by sequential binding of two divalent metal ions per monomer. Besides the physiological effector of DtxR, Fe²⁺, *in vitro* activation was demonstrated with Ni²⁺, Mn²⁺, and Co²⁺ at various affinities (36). As they are stable in an oxidative environment, they are commonly found in crystallographic structures of DtxR. An anion binding site is located in close proximity to the low-affinity binding site and essential for coordination of the metal ion (37–39). To address the question whether binding of HCO₃⁻ to the anion binding site of DtxR could cause the higher activation, we performed differential scanning fluorimetry. The thermal stability of purified DtxR protein was enhanced by increasing concentrations of divalent metal ions (Ni²⁺ and Mn²⁺) and dissociation constants (K_D) were calculated to 5.3 ± 1.1 μ M



FIG 3 (A) Kinetic analysis of the Fe²⁺-BPS complex formation at 534 nm with 19.5 μ M PCA and/or 50 mM HCO₃⁻. (B) Relative PCA degradation over time in the presence and absence of 50 mM NaHCO₃. Data points and bars represent mean values with error bars indicating standard deviations for three to six independent replicates.

and 21.8 \pm 3.6 μ M, respectively. However, addition of up to 100 mM NaHCO₃ did not stabilize DtxR further at any concentration of metal ions (data not shown).

To investigate whether elevated CO_2/HCO_3^{-} levels have an impact on iron storage or mobilization, we created deletion mutations (Δftn , Δdps , $\Delta ftn \Delta dps$, and Δpup) in *C. glutamicum* FEM3 and the WT. The deletion mutants lack either one (Δftn , Δdps) or both iron storage proteins ($\Delta ftn \Delta dps$) or are unable to initiate remobilization of the stored iron by pupylation (Δpup). With the exception of *C. glutamicum* Δpup , none of the deletion mutants showed a difference regarding growth and fluorescence in any of the conditions tested (data not shown). *C. glutamicum* Δpup revealed an even stronger growth defect than the WT under reference conditions. However, supplementation of PCA or NaHCO₃ restored the respective growth phenotype with the result that *C. glutamicum* Δpup and the WT grew identically again (Fig. S4).

CO₂/HCO₃⁻ accelerates Fe³⁺ reduction through phenolic compounds. Since we could not attribute the improved Fe³⁺ reduction capacity in the presence of elevated CO_2/HCO_3^{-} levels to an interaction with stored intracellular iron, we focused on the chemical reduction by PCA and HCO_3^- using the Fe^{2+} -specific iron chelator bathophenanthroline disulfonic acid (BPS) for detection. After 5 h of incubation, the amount of Fe²⁺-BPS complexes was about sevenfold higher in the presence of 19.5 μ M PCA compared to a sample without additives (blank control; Fig. 3A). Although the addition of 50 mM NaHCO₃ alone increased the final absorbance only marginally, it promoted the iron reduction in the presence of PCA significantly. Differences caused by the presence of NaHCO₃ were most remarkable during the initial reduction phase with a >50% greater amount of Fe²⁺-BPS complexes up to 100 min after the start of the assay (Fig. 3A). The sample pH was maintained at 7.4 throughout the incubation in 200 mM 3-(N-morpholino)propanesulfonic acid (MOPS) buffer, thus proving that a shift in pH was not causing the increase in absorbance. We also replaced NaHCO₃ with KHCO₃, which led to the identical improvement in iron reduction (Fig. 3A). An equimolar NaCl concentration did not trigger iron reduction (data not shown).

We monitored the degradation of PCA by liquid chromatography-quadrupole time of flight mass spectrometry (LC-MS-QTOF) analysis and found only about half of the residual amount of PCA when samples were incubated for 5 h in the presence of HCO_3^- compared to samples incubated without HCO_3^- (Fig. 3B). Furthermore, we identified oxidation products with the chemical formula $C_7H_6O_6$ and $C_7H_8O_6$.

Concluding that PCA serves as electron donor in the reduction of Fe³⁺, we tested other aromatic compounds at identical concentrations that could be associated with iron reduction.

Catechol and 2-amino-3-hydroxybenzoic acid (3-hydroxyanthranilic acid [3-HAA]) carrying two adjacent hydroxyl groups or a mix of amino and hydroxyl groups, respectively, exhibited reduction capacities that were at least as high as PCA. Iron



FIG 4 Kinetic analysis of the $Fe^{2+}-BPS$ complex formation at 534 nm with different functionalized aromatic compounds. Data points represent mean values with error bars indicating standard deviations of three to six independent replicates.

reduction by 4-amino-2-hydroxybenzoic acid (*para*-aminosalicylic acid), where the amino and hydroxyl groups are not positioned adjacent to each other, was only intermediate, whereas benzoic acid lacking hydroxyl and amino groups did not reduce iron at all (Fig. 4). The addition of 50 mM NaHCO₃ alongside with either of the above compounds (except for benzoic acid) accelerated the iron reduction in the same way as observed with PCA.

To further elucidate the role of HCO_3^{-} in the reduction of iron, we analyzed the complex formation between Fe³⁺ and the iron chelator PCA that is required prior to reduction. In accordance with iron reduction, the initial rate of Fe³⁺-PCA complex formation (0 to 35 min) was increased by 46% through the addition of 50 mM NaHCO₃ (5.2 ± 1.0 Δ mA₅₆₀ min⁻¹ [change in milli absorbance units at 560 nm per minute] versus 7.6 ± 1.3 Δ mA₅₆₀ min⁻¹; Fig. 5A). In addition to the altered kinetics, the maximum absorbance of those complexes shifted to a shorter wavelength (λ_{max}). The assay pH was again not affected by the addition of NaHCO₃, and intentional pH perturbation with 50 mM KOH or HCl failed to reproduce the λ_{max} shift (Fig. 5B).

DISCUSSION

Recently, Blombach et al. (30) initially disclosed a link between a high CO_2 proportion in the inlet air and a transcriptional response of almost the complete DtxR regulon in *C. glutamicum*. In this study, we demonstrate that elevated CO_2/HCO_3^- levels increased the intracellular Fe²⁺ availability and thus stimulated growth of *C. glutamicum*. The higher Fe²⁺ concentration was not coupled to biological activity. Instead, we



FIG 5 Complex formation between Fe³⁺ and PCA in the presence and absence of 50 mM NaHCO₃. (A) The kinetics of complex formation was monitored by an increase of the absorbance at 560 nm. (B) Wavelength of the maximum absorbance (λ_{max}) of the Fe³⁺-PCA complexes formed in the presence and absence of NaHCO₃ correlated with pH. Data points represent mean values, and error bars indicate standard deviations for three to five independent replicates.

identified an abiotic interaction of CO_2/HCO_3^- with chemical iron reduction through phenolic compounds.

Growth retardation had not been reported previously, when C. glutamicum cultures were aerated with pressurized air and has been introduced in this study presumably by the insertion of an additional minimal medium preculture, since all other process parameters were kept the same (30). The growth defect could be transferred to a shaking flask approach and was apparently caused by a limitation of intracellular iron availability, as suggested (i) by the lower fluorescence of FEM3, (ii) by the stronger growth retardation of C. glutamicum Δpup in the reference culture, which was found to exhibit a growth retardation only in iron starvation (11) and (iii) by the fact that PCA supplementation restored exponential growth (9). Analogously, an extended lag phase of C. glutamicum in minimal medium had been observed when the main culture was inoculated from a preculture in minimal medium or when the inoculum was washed excessively and, consequently, the addition of iron chelators like catechol, PCA, or citrate to minimal medium was suggested (8, 9). Interestingly, this requirement for iron chelators became apparent only when carbonate was omitted from the medium (8, 9, 40). However, the role of CO_2 with regard to the iron availability was not addressed further until noting a transcriptional response of the DtxR regulon (30).

The cometabolization of glucose and PCA can improve the growth rate of *C*. *glutamicum* to about 0.61 h⁻¹. However, this requires essentially higher PCA concentrations and lower cell densities (40). At standard PCA concentrations (as applied in this study), growth was not found to be increased over the maximum of $\mu = 0.42$ h⁻¹ when glucose was utilized as the sole carbon and energy source (41). Thus, growth of PCA-supplemented and nonsupplemented shaking flask cultures recorded herein was in qualitative and quantitative accordance with the literature (40, 41).

If NaHCO₃ supplementation were required for an increased flux via the anaplerotic reactions, and thus explaining the growth stimulation over a reference cultivation, the effect should persist in the simultaneous addition of PCA and NaHCO₃, because the small amount of PCA is readily consumed and cannot provide a long-lasting surplus of necessary precursors. However, supplementing with both PCA and NaHCO₃ did not result in a further growth improvement, and we found instead that the growth-stimulating effect of PCA and NaHCO₃ was related to the presence of the extracellular iron source. In iron-depleted medium, growth ceased after two to three cell divisions as reported previously (42).

Upon dissolution of the iron source FeSO₄ in CgXII medium, iron is present as reduced Fe²⁺. However, it will be quickly oxidized under aerobic conditions, yielding 99% of the initial iron concentration as Fe³⁺ after 15 min (43). Considering the standard preparation times of a shaking flask experiment and estimating the initially dissolved oxygen concentration at 7.5 to 8 mg liter⁻¹ (22), the entire amount of iron provided in the medium will be oxidized before inoculation. This highlights the need of enhanced iron reduction mechanisms to obtain increased intracellular Fe²⁺ levels. Since the increased Fe²⁺ availability in the presence of elevated CO₂/HCO₃⁻ levels could not be attributed to a biological function, we analyzed the chemical reduction capacity of PCA and HCO₃⁻⁻. Results of iron reduction assays and the LC-MS-QTOF approach show that PCA served as an electron donor for the redox reaction. Having performed the experiments at physiological pH (7.4), our results are in contrast with the common understanding that these redox reactions are restricted to acidic pH values (19-21). At pH 7.4, PCA coordination with Fe³⁺ represents a mix of 2:1 and 3:1 stoichiometry (20) and was expected to inhibit redox reactions of the iron monocatecholate complexes, which are favored at pH < 4.5 (21). Furthermore, our results indicate that the primary oxidation products of PCA (quinone and semiquinone) are further degraded. The chemical formula $C_7H_6O_6$ and $C_7H_8O_6$ obtained by the LC-MS-QTOF approach might represent β -carboxy-(*cis-cis*)-muconic acid or β -carboxy- γ -carboxymethyl- $\Delta^{\alpha,\beta}$ butenolide and butene-1,2,3-tricarboxy acid that were postulated to be degradation products, when PCA was oxidized with peroxyacetic acid (44). Interestingly, these

products also form part of the enzymatic degradation of PCA via the β -ketoadipate pathway (45).

Beyond PCA, other phenolic compounds can promote bacterial growth and have been associated with iron reduction and transport in the past. Examples are the reductive mobilization of ferritin iron through PCA and derivatives (46) as well as 2-amino-3-hydroxybenzoic acid secretion by *Cryptococcus neoformans* which mediated the iron reduction required for Fe^{2+} uptake (47). Supplementing with catechol stimulated growth of *C. glutamicum* comparably to PCA (9), and the chemical reduction of Fe^{3+} through catechol oxidation is well-known and has been reviewed (20). Our results provide evidence that the postulated iron reduction is not unique to PCA but characteristic of other functionalized phenolic compounds. Reduction assays with benzoic acid and 4-amino-2-hydroxybenzoic acid suggest that the carboxyl group was not required for iron reduction and that the stabilization of the semiquinone in adjacent hydroxyl (or mixed functional) groups was advantageous for the reduction of iron (drastically reduced in 4-amino-2-hydroxybenzoic acid).

The beneficial effect of HCO_3^- on the reduction of iron has not been postulated before. Remarkably, the iron reduction capacity of all aromatic compounds was enhanced in the presence of HCO_3^- , although iron reduction in the presence of HCO_3^- alone was negligible. The promoting effect of HCO_3^- on iron reduction was apparently related to the complex formation between Fe^{3+} and chelates, as exemplarily shown for PCA. Since we could not clarify the precise mechanism of interaction, one could assume that H^+ and OH^- concentrations were the reason for the λ_{max} shift in analogy with an overall alteration of the absorbance due to the formation of Fe(OH)-PCA complexes (19). However, we carefully performed pH buffering and control experiments in order to rule out this possibility. The sample pH was not varied in our experiments by the addition of NaHCO₃, and intentionally changing the pH failed to reproduce a comparable λ_{max} shift. It might be further speculated that HCO_3^- enhances the iron accessibility to PCA in the form of ferric bicarbonate, since the initial complex formation rate was accelerated by 46%.

Although the presence of HCO_3^{-} alone had no effect on the reduction of iron *in vitro*, the growth-promoting effect of CO_2/HCO_3^{-} correlated with increased intracellular Fe²⁺ levels reported *in vivo*. We conclude that the growth-promoting effect of CO_2/HCO_3^{-} requires the natural production of suitable reductants by *C. glutamicum*. Surprisingly, PCA production in *C. glutamicum* does not appear to be regulated in response to iron availability, unlike other organisms (16). The expression of *qsuB* encoding the dehydroshikimate dehydratase that catalyzes the conversion of 3-dehydroshikimic acid to PCA is controlled by QsuR in *C. glutamicum* and provides a shunt when chorismate accumulates (48, 49). The question of major iron reductants produced by *C. glutamicum* must be addressed in future research.

There is continuous debate about the redox reaction in Fe³⁺-catecholate complexes and its interpretation as antioxidant or prooxidant, since contradictory observations have been published (20, 50). It must be noted that the experimental setup is essentially decisive for the outcome. Many studies emphasized the antioxidant effect of PCA and other catecholates in the past (20). Fe²⁺ generation was monitored, e.g., by lipid oxidation experiments, which in contrast to BPS detection, do not exhibit high affinity for Fe²⁺. As a consequence, observations might differ from ours. The detection of Fe²⁺ with BPS in this study might have been beneficial for two reasons. (i) Fe²⁺ and BPS form a stable complex, thus preventing Fe^{2+} from reoxidation. (ii) Capturing Fe^{2+} in the complex constantly shifts the chemical equilibrium in favor of reduction. Thus, our in vitro screening system might be well suited for representing a biological context, considering the high Fe²⁺ affinity of iron processing enzymes, e.g., Escherichia coli IscA (the first enzyme of the iron sulfur cluster assembly machinery, $K_D = 3.3 \times 10^{-20}$ to 5 \times 10^{-20} M) (51, 52). Dissociation constants between iron and human transferrin (53) have been determined in a very similar range (2.1×10^{-21} and 4.2×10^{-20} M). However, the most obvious support of these hypothetical considerations is provided by the consistency of the results of in vitro and in vivo experiments in this study.

Given the predominance of Fe³⁺ in an oxygenic environment, the abiotic acceleration of iron reduction through HCO₃⁻ is of global importance. The two prerequisites the presence of functional aromatic compounds on one hand and high HCO3concentrations on the other hand-combine in a number of habitats. Aromatic compounds are ubiquitous in cellular metabolism, and plant-derived catecholic compounds can be released, e.g., in soils during biomass degradation or reach high concentrations in human plasma shortly after ingestion (20). A number of stress hormones and neurotransmitters, including dopamine, adrenaline, and norepinephrine, contain a functional catechol group and exhibit interesting features with regard to the iron homeostasis. The stimulating role of catecholamine on growth of a number of pathogenic bacteria in iron-restricted growth media containing serum, transferrin, or lactoferrin has been reported (54-58). Pathogens possess an outstanding role within the inhabitants of the human body as they encounter a constant battle for iron, the majority of which is sequestered in host transferrin and lactoferrin (35). Several studies demonstrated that catecholamines or dietary catechols can liberate iron from transferrin and lactoferrin (55, 59, 60) and might even reduce Fe³⁺ by promoting bacterial growth (61). The greater availability of liberated Fe³⁺ and Fe²⁺ resulting from this mechanism might be further increased at the high concentrations of CO₂ species typically found in calcareous soils, but also in the human body, where HCO_3^{-} concentrations can reach up to 140 mM (22). Considering the HCO_3^- interaction with iron reduction demonstrated in this study thus might be of particular interest in understanding pathogenicity. Of the currently most threatening human pathogens, Pseu-

domonas aeruginosa, Staphylococcus aureus, Klebsiella pneumoniae, Mycobacterium tuberculosis, and pathogenic Escherichia coli strains as well as several other pathogenic bacteria regulate toxin expression on the transcriptional level in response to iron availability (33, 62–64).

In summary, we show that high CO_2/HCO_3^- concentrations accelerate the chemical reduction of Fe³⁺ through various phenolic compounds. This increases the intracellular Fe²⁺ availability and stimulates growth of *C. glutamicum*.

MATERIALS AND METHODS

Genetic manipulation, cloning, and strain construction. Standard techniques of molecular biology were applied in accordance with the literature (65). Kits for purification of plasmids and PCR products and for isolation of genomic DNA were used according to Lange et al. (66). Cloning of plasmids and construction of *C. glutamicum* integration and deletion mutants have been performed as described in detail previously (66). All cloned fragments were verified by Sanger sequencing (GATC, Constance, Germany).

C. glutamicum FEM3 was composed of the chromosomally integrated sensor part and a replicative plasmid carrying the reporter gene. Expression of *lacl* was placed under the control of the DtxR-regulated promoter of the *ripA* gene (P_{ipA}), followed by the strong *rrnB* terminator (T_{rrnB}). Chromosomal integration of the cassette in *C. glutamicum* was targeted to CgLP13 (between *cg3344* and *cg3345*) by two 500-bp homologous sequences (66). *C. glutamicum* endogenous elements (500-bp flanks, P_{ripA}) were amplified from genomic DNA using the primer pairs 5'(cg3344)-1 plus 5'(cg3344)-2, 3'(cg3344)-1 plus 3'(cg3344)-2, and PripA-1 plus PripA-2, respectively, and exogenous elements (*lacl*, T_{rrnB}) were amplified from ploE7706.1 (67) with lacl-1 plus lacl-2 and TrrnB-1 plus TrrnB-2, respectively. All PCR amplification products were simultaneously inserted into BamH1- and Nhel-linearized pK19*mobsacB* (68) by isothermal assembling (69). The *egfp* gene under the control of the strong hybrid promoter P_{tac} and T_{rrnB} was amplified from pJOE7706.1 with primer pairs Ptac-1 plus Ptac-2 and egfp-1 plus egfp-2, respectively, and inserted into Xbal- and Nucl-linearized pL4 plasmid (70) in the same way.

For markerless deletions of the genes *pup* (*cg1689*), *ftn* (*cg2782*), and *dps* (*cg3327*) in *C. glutamicum*, 500-bp homologous sequences flanking the target gene were amplified with the respective primer pair (see Table S1 in the supplemental material) and simultaneously inserted in BamHI- and HindIII-linearized pK19*mobsacB* as described before.

For purification of the DtxR regulator protein, the *dtxR* gene was amplified from the *C. glutamicum* chromosomal DNA via PCR using the primer pair dtxR-1 and dtxR-2 and cloned into pJOE6089.4 by restriction and ligation using the Ndel and BamHI restriction sites. Competent *E. coli* DH5 α was transformed with the resulting plasmid pJOE6089-*dtxR* carrying a C-terminal *Strep*-tag II fused to the *dtxR* gene by electroporation (66).

Bacterial strains and cultivation conditions. An overview of all bacterial strains and plasmids used in this work is given in Table S2. Permanent cultures were maintained at -70° C in 30% (vol/vol) glycerol. *C. glutamicum* strains were cultivated at 30°C on a rotary shaker in baffled shaking flasks (500 ml filled with 50 ml medium, 120 rpm). Kanamycin was added at a working concentration of 50 μ g ml⁻¹ when appropriate. Permanent cultures were streaked on 2× yeast tryptone (2× YT) (65) agar (18 g liter⁻¹) plates and grown for 2 days. Liquid cultures (5 ml of 2× YT) were inoculated from the plate, grown overnight (O/N), and used to inoculate a 50-ml 2× YT culture, which was incubated for another 6 to 8 h. An appropriate volume of the complex preculture was harvested and resuspended in 2 ml of 0.9% (wt/vol) NaCl to inoculate a CgXII preculture to a starting optical density at 600 nm (OD₆₀₀) of 1. The main cultures were inoculated in the same way from the CgXII O/N culture and incubated for at least 25 h.

Shaking flask cultivation. *C. glutamicum* cultivations were performed in CgXII medium (pH 7.4) containing 20 g glucose liter⁻¹ supplemented with protocatechuic acid (PCA) to a final concentration of 195 μ M (10) or with 30 mM NaHCO₃ or not supplemented (71) as indicated. Since slight variations occurred in the past, the exact composition of the CgXII medium used in this study is outlined in Lange et al. (66). The functionality of the reporter strain *C. glutamicum* FEM3 was validated under iron starvation (1 μ M FeSO₄) and iron excess (100 μ M FeSO₄) conditions in the presence of 195 μ M PCA and starting from identical precultures as described before (32). Iron-depleted growth medium was prepared with a trace element solution lacking the iron source (FeSO₄), and the ferrous iron chelator 2,2-dipyridyl was added at a final concentration of 250 μ M immediately before inoculation to remove any residual iron from the medium (42). Precultures were performed in iron excess conditions (100 μ M FeSO₄).

Bioreactor cultivation. Bioreactor cultivations were performed in batch mode in 1.5-liter stainless steel bioreactors (30) with a starting volume of 800 ml CgXII medium containing 40 g glucose liter⁻¹ as the sole carbon and energy source but lacking 3-(*N*-morpholino)propanesulfonic acid (MOPS) buffer and urea. The cultivation pH was maintained at pH 7.4 by the addition of 25% ammonium hydroxide. Two parallel fermentations were always inoculated from identical precultures to a starting OD₆₀₀ of 2. Both reactors were operated at a total pressure of 1.5 bar and contained two six-blade Rushton-type impellers. The stirrer speed started at 300 rpm and was gradually increased to maintain the dissolved oxygen concentration above 35%. Reactor 1 was aerated with 0.5 vvm (volume of gas per volume of liquid per minute) pressurized air (0.04% CO₂), and reactor 2 was aerated with synthetically mixed gas containing 20% CO₂, 21% O₂, and 59% N₂ (20% CO₂).

Biomass formation. Cell density was determined by measuring the optical density of a culture at 600 nm (OD₆₀₀). Biomass concentration (cell weight [dry weight] [CDW] in grams liter⁻¹) was calculated from it using the correlation coefficient CDW = $0.21 \times OD_{600}$ (72) specific for the spectrophotometer (Ultrospec 10 cell density meter; GE Healthcare, Little Chalfont, UK).

Fluorescence readings. Fluorescence (relative fluorescence units [RFU]) was detected in $100-\mu$ l samples of the pure culture using 96-well microtiter plates in the Synergy 2 device (BioTek Instruments, Bad Friedrichshall, Germany) at 37°C (excitation wavelength/bandwidth, 485/20 nm; emission wavelength/bandwidth 528/20 nm) as described before (73). The culture background (after removing cells by centrifugation, 13,300 rpm, 1 min) was subtracted from the sample value and normalized with respect to the biomass concentration (RFU CDW⁻¹).

LC-MS-QTOF. PCA degradation was detected by a relative decrease of the respective peak area via liquid chromatography-quadrupole time of flight mass spectrometry (LC-MS-QTOF). Samples (1-ml samples) containing 0.5 mM FeCl_a (from a stock in 10 mM HCl) were prepared in 1.5-ml Eppendorf cups. Samples were neutralized with 50 mM NaOH, and 19.5 μ M PCA, 6.5 mM BPS, and 50 mM NaHCO₃ were added when appropriate. At the indicated time points, $200-\mu$ l samples were transferred to fresh 1.5-ml Eppendorf cups and stored frozen at -20°C until further processing. After thawing, samples or respective standards were prepared in 60% (vol/vol) acetonitrile, 0.75 mM EDTA, and 10 mM ammonium acetate buffer (adjusted to pH 9.2) and analyzed by an Agilent 1260 bio-inert high-performance liquid chromatography (HPLC) system (Agilent Technologies, Waldbronn, Germany) coupled to an Agilent 6540 accurate-mass QTOF (Agilent Technologies, Santa Clara, CA, USA). Alkaline hydrophilic interaction liquid chromatography (HILIC) (74) and QTOF-MS (75) parameters were set as previously described, and analysis was conducted in the negative mode with a fragmentor voltage of 100 V in the MS mode. The extraction of chromatograms and integration were done in MassHunter Qualitative Analysis with the "Find by Formula" algorithm. Potential oxidation products were identified via the molecular formula calculator feature based on the accurate mass, isotope abundances, and isotope spacing (B.07.00, Agilent Technologies, Santa Clara, CA, USA).

Iron reduction assays. The kinetics of Fe³⁺ reduction was monitored by an increase of the absorbance at 534 nm due to complex formation with the Fe²⁺-specific chelator bathophenanthroline disulfonic acid (BPS) (76, 77) in 96-well microtiter plates containing 100 μ l of sample. The iron reduction assay was performed in 200 mM MOPS buffer (pH 7.4), 0.5 mM FeCl₃ was added from a stock in 10 mM HCl neutralized with 50 mM NaOH immediately before use. 19.5 μ M PCA, other benzoic acid derivatives, or 50 mM NaHCO₃ was applied as indicated. The reaction was started by the addition of 6.5 mM BPS and measured every 10 min.

Fe³⁺-PCA complex formation. Complex formation between Fe³⁺ and PCA was monitored in a setup identical to the setup used in the iron reduction assay but in the absence of Fe²⁺-specific BPS. The Fe³⁺-PCA complex formation assay was started by the addition of 1.5 mM PCA to provide it in a 3:1 stoichiometry with iron. Fe³⁺-catecholate complexes show characteristic λ_{max} values between 561 and 586 nm (20), and for Fe³⁺-PCA, 575 nm was reported as λ_{max} (19). Wavelength scans of the complexes were performed with 100- μ l samples in 96-well plates (1-nm step size) after 5 h of incubation. Since the absorbance peaked around 560 nm in the standard setup (see Fig. S5 in the supplemental material), kinetic measurements of complex formation was performed in the following experiments every 5 min at 560 nm in the same setup.

Purification of DtxR. For the production of DtxR protein, a cryogenic culture of *E. coli* DH5 α (pJOE6089-*dtxR*) was streaked on a 2× YT agar plate containing 100 μ g ampicillin (Amp¹⁰⁰) ml⁻¹ and incubated for 24 h at 37°C. A single cell was used to inoculate 5 ml of 2× YT (Amp¹⁰⁰) which was incubated overnight (O/N) at 37°C on a rotary shaker at 120 rpm before 1.5 ml of this culture was transferred to fresh 150 ml of 2× YT (Amp¹⁰⁰) medium in a 1-liter baffled shaking flask. This culture was incubated at 37°C on a rotary shaker at 120 rpm until an OD₆₀₀ of 0.84 was reached. The expression of *dtxR* was induced by the addition of L-rhamnose to a final concentration of 2 g liter⁻¹, and incubation continued for 6 h at 30°C before the cells were harvested by centrifugation (10 min, 5,000 rpm at 4°C in an Eppendorf centrifuge 5804 R). The cell pellets were stored frozen at -20°C until further processing.

Strep-tagged protein was purified with the Strep-tagged protein purification kit (Qiagen, Hilden, Germany) after lysing cells by sonication (five cycles, Sonopuls HD2200, Bandelin, Berlin, Germany; tip type MS73, 30 s, power 40%).

Thermal shift assay. Differential scanning fluorimetry experiments were performed in a final sample volume of 50 μ l in 96-well plates using the Mastercycler EP Realplex² (epgradient S; Eppendorf, Hamburg, Germany). Briefly, at a gradually increasing temperature, protein denaturation is monitored by an increase of the fluorescence which is caused by the binding of a dye to successively presented hydrophobic areas of the protein (78). Ligand binding might increase the protein stability. Thermal protein denaturation was monitored by an increase of the absorbance at 550 nm due to binding of SYPRO Orange (Thermo Fisher Scientific Inc., Bremen, Germany).

Protein samples (50 μ g ml⁻¹) were prepared in 100 mM potassium phosphate buffer (pH 7.4) containing 600 mM NaCl and 5 mM dithiothreitol (DTT). Divalent metal ions were applied at various concentrations (0.1 μ M to 1 mM NiCl₂ or 0.1 μ M to 5 mM MnCl₂). When 10 to 200 mM NaHCO₃ was added to the sample, osmolarity was maintained at an identical level by adjusting the NaCl concentration. One microliter of SYPRO Orange was added to each sample immediately before starting the assay. Then, the 96-well microtiter plate was sealed with adhesive foil and incubated in the mastercycler (20°C for 15 s, 20°C to 90°C in 30 min, 90°C for 15 s). The range of the greatest absorbance increase over temperature was fitted by a fourth order polynomial (best fit), and the melting temperature (T_m) was determined at the inflection point. The T_m at a given ligand concentration constants (K_D) were calculated for the inflection point of the T_m on the ligand concentration curve (79).

Statistical analysis. All growth experiments were performed as independent biological triplicates on different days. (Bio)chemical assays were replicated 4 to 6 times, and LC-MS-QTOF analysis was performed on the three biological replicate samples. All values represent mean values with error bars indicating standard deviations. When appropriate, statistically significant differences of sample means were tested with a two-sample *t* test.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. FIG S1, EPS file, 0.2 MB. FIG S2, EPS file, 0.2 MB. FIG S3, EPS file, 0.2 MB. FIG S4, EPS file, 0.2 MB. FIG S5, EPS file, 0.2 MB. TABLE S1, DOCX file, 0.01 MB. TABLE S2, DOCX file, 0.02 MB.

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F.M., R.T., and B.B. conceived and designed the experiments. F.M., J.R., A.-L.H., and A.F. performed the experiments. F.M., J.R., A.-L.H., A.F., and B.B. analyzed the data. F.M. and B.B. wrote the manuscript.

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