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Manganese oxide biomineralization provides protection against nitrite toxicity in a cell density dependent manner.

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Author Contributions

CZ, JCO and OSS designed the study and the experiments. CZ performed the experiments and analyzed the data. All authors contributed to the writing of the manuscript and have given approval to the final version.

Conflict of Interest

The authors declare that there are no conflicts of interest.
Abstract

Manganese bio-mineralization is a widespread process among bacteria and fungi. To date there is no conclusive experimental evidence for, how and if this process impacts microbial fitness in the environment. Here we show how a model organism for manganese oxidation is growth-inhibited by nitrite, and that this inhibition is mitigated in presence of manganese. We show that such manganese-mediated mitigation of nitrite-inhibition is dependent on the culture inoculum size and that manganese oxide (MnO\textsubscript{X}) forms granular precipitates in the culture, rather than sheaths around individual cells. We provide evidence that MnO\textsubscript{X} protection involves both its ability to catalyze nitrite oxidation into (non-toxic) nitrate under physiological conditions, and its potential role in influencing processes involving reactive oxygen species (ROS). Taken together, these results demonstrate improved microbial fitness through MnO\textsubscript{X} deposition in an ecological setting, i.e. mitigation of nitrite toxicity, and point to a key role of MnO\textsubscript{X} in handling stresses arising from ROS.

Importance

We present here a direct fitness benefit (i.e. growth-advantage) for manganese oxide bio-mineralization activity in Roseobacter sp. AzwK-3b, a model organism used to study this process. We find that AzwK-3b in a laboratory culture experiment is growth-inhibited by nitrite in manganese free cultures, while the inhibition is considerably relieved by manganese supplementation and MnO\textsubscript{X} formation. We show that biogenic MnO\textsubscript{X} interacts directly with nitrite and possibly with reactive oxygen species, and find that its beneficial effects are established through formation of dispersed MnO\textsubscript{X} granules in a manner dependent on the population size. These experiments raise the possibility that manganese bio-mineralisation could confer protection against nitrite-toxicity to a population of cells. They open up new avenues of interrogating this process in other species, and provide possible routes to their biotechnological applications, including in metal recovery, biomaterials production, and in synthetic community engineering.
A large variety of biominerals based on different cations (e.g. iron, manganese, calcium) and anions (e.g. carbonates, oxides, phosphates) are deposited by different microorganisms (1). One of these is manganese oxide (2–5), which is deposited by the oxidation of soluble Mn$^{II}$. Microbial Mn$^{II}$ oxidation received attention with the discovery of polymetallic, manganese-rich biogenic deep sea nodules, which have been shown to harbor both manganese-oxidizing, and manganese-reducing organisms (6). While it is suggested that such nodules could potentially be mined for rare earth elements, and the associated metal-active organisms be used in biotechnology of metal recovery (2, 3, 5–8), it remains unclear in many cases why organisms carry out such metal-oxidizing and -reducing activities. In the case of metal-reducing organisms, it has been shown that metabolic energy can be gained under anaerobic conditions from using metal oxides (i.e. manganese, iron, or others) as an alternative terminal electron acceptor (9–11). Some metals can be oxidized by microbes and act as an inorganic energy source for so-called chemolithotrophic growth, as in the case of iron lithotrophy (12). While it has been suggested that manganese oxidation can also be used as a chemolithotrophic source of energy (2), little experimental evidence has been found. In most cases studied, Mn oxidation is not used as a lithotrophic source of energy and, hence, evolutionary advantages of this process are not well understood (2, 7, 8). Two running hypotheses for non-lithotrophic manganese oxidation are that the resulting manganese oxides (MnO$_X$) can i) increase accessibility of organic nutrients, or ii) protect microbes from potentially toxic compounds and superoxide stress (13, 14). The validity of the former hypothesis remains to be tested conclusively. MnO$_X$ has been shown to react with complex organic (i.e. humic) substances (15), but it is not clear if the resulting organic products form such reactions are utilized by microbes. It is suggested that certain fungi employ ligand-stabilized Mn$^{III}$ to oxidize recalcitrant litter (16), but these studies were not performed with single (defined) carbon/energy sources. The latter hypothesis regarding the protective potential of MnO$_X$ remains unproven to date for metal toxicity (2, 7). It is shown that MnO$_X$ can mediate a protection against superoxides in *Pseudomonas* species (14), but it is not clear how significant this benefit is given that these and other Mn-oxidising species also possess specific superoxide scavenging enzymes such as catalases and superoxide dismutases (17–19). It has been suggested that MnO$_X$ precipitates can act as strong sorbents of heavy metals,
hence mitigating the toxic effects of such metals on microorganisms, but this has yet to be tested in a biological context (2). Taken together, the biological significance of microbial manganese oxidation remains largely a paradox, as no clear benefits have been demonstrated.

In recent years, *Roseobacter sp.* AzwK-3b emerged as a model organism to study the generation of MnO$_X$ (20). AzwK-3b is a bacterium that shows significant manganese oxidizing activity in vitro when grown in a complex (rich) K-Medium (20) and defined (acetate-fed) J-medium (21). This activity was shown to be mediated by a secreted exoenzyme - a haem type oxidase - that can catalyze the in vitro generation of superoxides from NADH and oxygen (22) (this and later reactions shown in Figure 1), demonstrating the use of biological reductive energy equivalents. The resulting superoxide can in turn facilitate the Mn$^{	ext{II}}$ oxidation into Mn$^{	ext{III}}$, which undergoes further disproportionation to result in MnO$_2$ (22–26) – or more specifically mixed valence state MnO$_X$. While NADH was a suitable electron donor for the in vitro superoxide production by haem peroxidase, the natural reducing agent, and the way it is delivered, is not known. It has been suggested that the haem peroxidase might be loosely membrane-bound (27), which would mean that electrons could be shuttled from cytoplasmic reductive metabolites to the haem peroxidase e.g. via membrane proteins, though would imply that the natural site of superoxide production (and subsequent manganese oxidation) would be in the immediate proximity of the cell. Since haem peroxidases are also found in culture supernatants (22), an extracellular reaction would require that electron donor metabolites are secreted also, which would imply a considerable investment for AzwK-3b. Thus, these mechanistic findings strongly suggest that AzwK-3b is making a significant metabolic investment into production of MnO$_X$ in form of secreted enzymes and possibly also reductive energy donating metabolites. Furthermore, AzwK-3b’s cellular and excreted proteome is shown to be different when grown under the presence or absence of Mn, while it is notable that the haem peroxidase described above was not found to be differentially expressed (28). It is currently not clear how and if the metabolically costly process of extracellular Mn oxidation benefits individual cells and how it could have been maintained over evolutionary timescales.

In an attempt to better understand any ecologically relevant ‘fitness’ impacts of manganese oxidation, we have studied the physiology of *Roseobacter sp.* AzwK-3b in more
detail. While we did not find any significant difference in growth rate between manganese-
free and -supplemented media, we found that the manganese oxidizing activity of
*Roseobacter* sp. AzwK-3b supports growth of the bacterium at nitrite concentrations that fully
prevent growth in a manganese-free culture. MnO$_X$ formed as granules dispersed among
cells, and its nitrite-inhibition mitigation effects showed a significant population size effect,
suggesting a ‘community commodity’ nature of this compound. Mechanistically, we show
that biogenic MnO$_X$ was able to catalyze nitrite oxidation into nitrate under physiological
conditions (according to reaction (4) in Figure 1), and that the mitigation of nitrite-inhibition
was also affected by NADH. These results suggest that the ability of MnO$_X$ to alleviate nitrite
toxicity relates to providing catalytic scavenging of reactive oxygen species (ROS) within the
environment, whose effect can be leveraged by nitrite.

**Results**

To study the role of manganese oxidation on microbial fitness we have focused here
on *Roseobacter* sp. AzwK-3b, which has recently emerged as a model organism for this
process (2, 8). We refer to the oxidation product as MnO$_X$, since biogenic manganese oxides
are usually precipitates with mixed manganese oxidation states, particularly Mn$^{II}$, Mn$^{III}$ and
Mn$^{IV}$ (2, 29). AzwK-3b has been shown to oxidize manganese to MnO$_X$ by means of an
exoenzyme and reductive energy (e.g. NADH in vitro), and potentially involving an elaborate
redox reaction path (22–26). We first attempted to identify fully-defined growth conditions
for this bacterium, which has been to date studied in complex and lean K-medium (20), both
of which contained undefined complex ingredients such as peptone or vitamin mixtures and
yeast extract (20, 30) or standard vitamin supplements (21). Through systematic analysis of
media composition, we have created a minimal defined medium that supports AzwK-3b
growth (from now on referred to as modified artificial seawater medium, ASW$_m$) (Table 1),
and that has revealed the requirement for five specific vitamin supplements for growth
(Figure S1). Given this defined culture medium, we were then able to interrogate the impact
of manganese on the growth of AzwK-3b.
Manganese oxidation has negligible impact on growth rate. Despite potentially significant costs associated with exoenzyme secretion and the investment of reductive energy equivalents for superoxide generation, we did not find any substantial difference in growth rates and steady state population sizes with increasing MnII concentration for cultures grown with 25 mM acetate (Figure 2). A slightly slower growth at the highest manganese concentration (500 μM) was observed, but it was difficult to ascertain this effect, as both MnO_x particles and cells co-aggregating with those particles could have interfered with the absorbance measurements. The slightly reduced growth rate at 200 μM Mn^{II}Cl_2 is in line with an earlier report on AzwK-3b, where 100 μM Mn^{II} was found to decrease the growth rate in (complex) K-medium (20). Other manganese-oxidizing bacteria, such as Erythrobacter sp. SD-21 (31, 32) and a marine Bacillus strain (33), were reported to grow better when cultured with Mn^{II}-supplement. In light of these different findings and possible difficulties with growth rate measurements in the presence of manganese precipitation, we cannot be fully conclusive about the growth effects associated with manganese oxidation based on the presented results, however, they are suggestive of a low or no-impact on growth rate.

Manganese oxidation mitigates nitrite growth inhibition. With growth effects being limited, a possible alternative explanation for a positive role of manganese oxidation is a protective effect against inhibitors or stresses (2, 13). Here, we evaluated this hypothesis for nitrite. Nitrite is commonly found in the environment, where it results from the reduction of nitrate, a key terminal electron acceptor utilized by many microbes (34). We found nitrite inhibited the growth of AzwK-3b in manganese-free cultures, where already as little as 0.25 mM nitrite prevented growth of AzwK-3b (Figure 3A). To rule out a salinity effect, different concentrations of sodium chloride were tested (200 mM (default in ASWm) to 428 mM NaCl (default in original ASW medium (35)), and AzwK-3b grew in all tested conditions (Figure S2) with the addition of 200 μM Mn^{II}, we found that AzwK-3b is able to grow in the presence of up to 1 mM nitrite (Figure 3B). Increasing the nitrite concentration still affected both the growth rate and maximal culture density (based on A_{600}), but this effect was much lower compared to the manganese-free cultures (Figure 3). To overcome any potential confounding effects of MnO_x precipitation on spectroscopic culture density measurements, we additionally quantified acetate consumption by ion chromatography as a proxy for...
growth. As expected, manganese-free cultures with 0.25 mM (or higher) nitrite showed only insignificant decrease in acetate, while the Mn\textsuperscript{II} supplemented cultures showed acetate consumption in accordance with the $A_{600}$ measurements (see Figure S3). These findings confirm that Mn\textsuperscript{II} supplementation allows AzwK-3b to withstand nitrite inhibition.

Nitrite-inhibition relief is a community function that depends on culture size and that is mediated by dispersed, granular MnO\textsubscript{X} precipitates. It has been shown that MnO\textsubscript{X} precipitation by AzwK-3b is mediated by secreted exoenzymes (22). It is not known, however, whether the process of MnO\textsubscript{X} precipitation occurs primarily on individual cell surfaces, or whether it is a population level process with the secreted enzymes conferring to the notion of a “community commodity” (36–39). We hypothesized that these two different scenarios could be distinguished by analyzing population size effects on MnO\textsubscript{X} mediated mitigation of nitrite-inhibition. In particular, we designed an experiment in which cultures pre-grown without Mn\textsuperscript{II} are subsequently sub-cultured into media with Mn\textsuperscript{II} and nitrite, using different inoculum size. We argue that in the case of MnO\textsubscript{X}-based protection being a process confined to individual cells, there should be no effect of inoculation size.

We found that MnO\textsubscript{X}-based protection against nitrite inhibition was dependent on inoculum size (Figure 4). A pre-culture was grown without nitrite and manganese, and from this, inocula were generated at two different time points within the first third of the exponential phase (labelled IT1 and IT2 in Figure S4). When these inocula were subjected to nitrite in the main-culture, the earlier, low-density inoculum IT1 was inhibited by nitrite regardless of the presence or absence of Mn\textsuperscript{II} (Figure 4 A,B), while manganese-mediated mitigation of nitrite inhibition was clearly evident for the larger, high-density inoculum IT2 (Figure 4 C,D). In the IT1 cultures half of the acetate was unused at 0.25 mM nitrite, and gradually more acetate resided with increasing nitrite concentration (Figure S5). In the IT2 cultures with Mn\textsuperscript{II} supplementation, however, acetate was completely removed at all nitrite levels below 2.5 mM and only 25 – 50 % of acetate remained at 5 – 10 mM nitrite. In the control samples (no inoculation) there was no change in acetate concentration ruling out any cross-activity with manganese.

Rather than a true population size effect, these observed inocula effects could be due to cells from the Mn-free, early-phase pre-cultures not having ‘turned on’ expression of
exoenzymes required for MnO\(_X\) precipitation. To rule out this possibility, we performed an additional experiment, where the pre-cultures were already grown with 200 μM Mn\(^{II}\). Using this pre-adapted culture, inocula were again prepared by sampling at different growth time points (IT 1 – 4 in Figure S6, A). Cultures grown from these different inocula displayed much weaker inhibition by increasing nitrite concentrations up to 10 mM (Figure S6, B) and were able to consume acetate (Figure S6, C), yet there were still inoculum size effects on overcoming nitrite inhibition (Figure 5, green). Interestingly, the extent of this effect seems similar to that observed with inocula originating from pre-cultures grown without Mn\(^{II}\) but supplied with Mn\(^{II}\) after subculturing into nitrite containing media (Figure 5, blue). In particular, at 5 and 10 mM nitrite, maximum growth rate (and final density) data from these two treatments can all be fitted on to a single (sigmoidal) curve that describes the relation between this data and initial inocula density (Figure 5, black line). This shows that the presence of Mn\(^{II}\) in the pre-culture does not impact the dynamics of the process, but rather allows the main-culture populations to grow to a higher density under a given nitrite level. In the absence of Mn\(^{II}\) in both pre- and main-cultures, a much denser inoculum was required to achieve growth at a given nitrite level and even then both growth rate and final growth density were lower compared to the case in the presence of Mn\(^{II}\). Under this condition, there was still a density dependence of the nitrite effect.

These results suggest that MnO\(_X\) precipitation is a community level function. To further collaborate on this result, we explored the micro-structure of the AzwK-3b cultures in the presence of Mn\(^{II}\). Analysis of cultures using electron microscopy revealed that MnO\(_X\) precipitates as granules dispersed within the culture, and attaching to clusters of cells, rather than forming sheaths around individual cells (as seen in some other cases of metal oxide precipitations (40)) (Figure 6, left). Employing electron dispersive X-ray spectroscopy, we confirmed that these granular structures contained manganese, while no manganese was detected in locations with cells only (i.e. without granular structures, see Figure 6, right).

MnO\(_X\) mediated nitrite protection involves redox reactions and oxygen radicals. After establishing the community level functionality of biogenic MnOX as a protective agent against nitrite, we next wanted to evaluate the mechanistic basis of this function in the context of nitrite toxicity. While multiple mechanisms of nitrite-toxicity are reported (41, 42), two key
reactive species are usually implicated, i.e. free nitrous acid (43) and peroxynitrite. The former forms through protonation of nitrite, while the latter forms from the reaction of nitrite with hydrogen peroxide (44–46). Thus, two non-exclusive, possible mechanisms of MnO\textsubscript{X} relief on nitrite toxicity are: (i) MnO\textsubscript{X} catalyzed oxidation of nitrite to nitrate (a reaction that has been shown to be feasible chemically under low pH (47)) and thereby avoiding formation of either free nitrous acid or peroxynitrite; or (ii) MnO\textsubscript{X} catalyzed degradation of hydrogen peroxide and thereby avoiding the reaction of this compound with nitrite to form peroxynitrite.

To see if AzwK-3b generated MnO\textsubscript{X} can catalyze nitrite oxidation under physiological conditions, we collected it from culture supernatants and evaluated its reactivity with nitrite in our ASW\textsubscript{m}-medium at pH = 8.0. Over 27 days, we found nitrite oxidation by biogenic MnO\textsubscript{X} in a dose dependent manner, while neither synthetic MnO\textsubscript{2} powder nor the MnO\textsubscript{X}-free solution showed any significant nitrite oxidation (Figure 7A). The trend of nitrite oxidation matched with nitrate production (Figure 7B), thus confirming the assumed reaction pathway of nitrite-oxidation into nitrate (47). Taking into account the difficulties of accurately determining the amount of precipitated MnO\textsubscript{X} that were added into the nitrite assay, we can still estimate that the condition with highest MnO\textsubscript{X} levels contained at least 1-2 mM (with respect to Mn). This presents a stoichiometric minimum 2-fold excess over nitrite (at 0.5 mM), hence enough for complete nitrite oxidation. The fact that this reaction didn't proceed further than an oxidation of ~0.18 mM nitrite (i.e. ~35 %) indicates that either the biogenic MnO\textsubscript{X} was only partially reactive or that its reactivity reduced over time (as known to be the case for synthetic manganese oxides (2, 13)). Sample pH remained relatively stable with the biogenic MnO\textsubscript{X}, while samples without manganese and with synthetic MnO\textsubscript{2} reached a pH of 6.9 and 6.8, respectively at the end of the experiment (from an initial pH of 8.2 of the medium). This acidification of the control samples might be due to carbon dioxide dissolution, which might have been buffered in the samples with biogenic MnO\textsubscript{X} due to proton consumption during nitrite oxidation, or due to co-precipitated organic solutes (polymers, proteins) from the cell-free supernatant.

These findings confirmed that the biogenic MnO\textsubscript{X} were capable to oxidize nitrite at physiological conditions, and prompted us to test MnO\textsubscript{X} mediated nitrite oxidation directly in AzwK-3b cultures. We found some evidence for decreasing nitrite concentration in different cultures tested, but this was not significant (Figure S7), and some decrease was also seen in
the manganese free cultures (indicating possible measurement effects in the solution). If nitrite oxidation was the main mechanism of MnO$_X$ mediated protection in vivo, these cultures would have been expected to oxidize most of the nitrite present in the media. Thus, we conclude that under our experiment conditions nitrite-oxidation was only a potential contributing factor.

A plausible alternative mechanism of MnO$_X$ mediated nitrite-inhibition relief could be related to formation of reactive peroxynitrite, which is shown to be highly toxic to bacteria (45, 46, 48, 49), and which can form (particularly) at low pH from the reaction of hydrogen peroxide with nitrite (44). If peroxynitrite is the main species underpinning nitrite toxicity, then MnO$_X$ protection against nitrite could be due to its ability to degrade hydrogen peroxide and thereby reducing the rate of peroxynitrite formation. The reactivity of MnO$_X$ towards hydrogen peroxide has been demonstrated chemically (44, 50–57), but never shown or tested in a biological context. Here, we hypothesized that if these types of redox reactions were involved in MnO$_X$ mediated mitigation of nitrite-inhibition, the process dynamics can be modulated with the introduction of additional hydrogen peroxide or NADH (which can help increase the rate of MnO$_X$ formation (23), but which can also be directly involved in hydrogen peroxide reduction through peroxidase-catalysed reactions (17–19, 58, 59)). To test this hypothesis, we again grew pre-cultures of AzwK-3b without Mn$^{ll}$ and sub-cultured these in medium containing Mn$^{ll}$ and nitrite, but at the same time also spiking in hydrogen peroxide or NADH. Hydrogen peroxide spiking did not show any effect on nitrite inhibition or its release by Mn$^{ll}$ supplementation (Figure S8), possibly due to spiked hydrogen peroxide being cleared primarily through additional peroxidases rather than impacting MnO$_X$ mediated process dynamics. In line with this hypothesis, spiking NADH resulted in full mitigation of nitrite inhibitory effect (even without Mn$^{ll}$) (Figure 8). This suggests that nitrite toxicity relates to peroxynitrite formation via hydrogen peroxide, which can be decomposed by MnO$_X$ (as shown before (44, 50–57)) or NADH-utilizing peroxidases (that are shown to be present in Roseobacter species including AzwK-3b (22, 27) (see also Table S1)).

Discussion

Manganese bio-mineralization into MnO$_X$ is widespread among bacteria, but there is no clarity about its possible functional roles. Here, we developed a defined growth media for
the manganese oxidizing model organism *Roseobacter sp.* AzwK-3b and demonstrated that in a laboratory setting this organism’s strong growth-inhibition by nitrite is mitigated through its ability to precipitate biogenic MnO\(_X\). While our experiments were undertaken in an artificial lab environment, these findings provide a direct evidence for the impact of MnO\(_X\) on an organism’s growth, thus raising the possibility of a positive fitness effect and a possible ecological/evolutionary explanation to the costly process of MnO\(_X\) oxidation.

Interestingly, we also show that the MnO\(_X\)-mediated mitigation of nitrite toxicity is dependent on population size, and that MnO\(_X\) forms dispersed granules that are attached to clusters of cells in the population. These observations, combined with the established role of exoenzymes in the formation of MnO\(_X\) precipitates, suggests that these provide a community function to AzwK-3b and allows cultures grown to sufficient density in the presence of manganese to become resistant to the inhibitory effects of nitrite. Our attempts to elucidate the mechanistic basis of this functionality showed that biogenic MnO\(_X\) can oxidise nitrite to nitrate (under conditions that synthetic MnO\(_2\) cannot). Together with the known ability of MnO\(_X\) to degrade hydrogen peroxide (44, 50–57), these findings show that biogenic MnO\(_X\) can inhibit the two key routes to the formation of reactive nitrite species.

While mitigation of nitrite inhibition might not be the only evolutionary advantage of MnO\(_X\) oxidation in AzwK-3b or other manganese oxidizing species, it is definitely an ecologically relevant function. Nitrite is a known inhibitor in the environment (41, 42, 60), including in wastewater treatment applications (43). In soil, reported nitrite concentrations are in the range up to low micromole per kg or litre, respectively (61, 62), though can peak to higher than 0.5 millimole per kg by agricultural nitrogen fertilization (61). In biofilms, where diffusion is inhibited, oxygen is shown to rapidly diminish (63–65), which can favor anaerobic metabolism including nitrate respiration to nitrite (66). Furthermore, biofilms are shown to preferentially select for and accumulate ions such as phosphate and nitrite (67–69). For example, in freshwater lake biofilms, the annual variation range for nitrite was found to be from µM to mM range (i.e. 1,000-fold) in biofilms (68). In the case of AzwK-3b, these physical and ecological processes can be highly relevant, as this species was isolated from an "agriculturally impacted, shallow salt marsh" (20) where nitrite (among other nitrogen species) can occur due to microbial conversion of nitrogen fertilizers (61, 70–72). It is also interesting to note that oceanic manganese-rich modules are found to contain both
manganese oxidizing and reducing bacteria (6), with current-day representatives of the latter group, such as *Shewanella oneidensis* (9), also being nitrate-reducers (73–75). Thus, these nodules can or have harbored also high levels of nitrite, creating environments that select for manganese oxidation.

Our study opens up additional investigations into the mechanism of nitrite toxicity and the role of MnO\textsubscript{X} oxidation in it. Multiple mechanisms of nitrite-inhibition have been reported (41, 42), and a key role for free nitrous acid (i.e. protonated nitrite) (43) and peroxynitrite, from nitrite and hydrogen peroxide (44–46), is proposed. Both molecules can prevent chemiosmotic coupling, and are primarily formed at low pH (nitrite is often found to inhibit bacterial survival at pH < 7 (45, 46)). Indeed, low pH can arise within the cellular microenvironment: energy metabolism coupled to chemiosmosis generates a proton motive force that can enrich the proton concentration at the charged membrane surface (values down to pH 5.5-6.5 are discussed). This local low pH environment can be further stabilized and inhibited from equilibration with the bulk due to an electrostatic barrier imposed by water layering (76, 77). Additionally, respiratory activity can increase hydrogen peroxide in the same cellular microenvironment (18, 48, 49, 58, 78–85), which can facilitate nitrite conversion to peroxynitrite. Interestingly, these local conditions could be avoided through the presence of MnO\textsubscript{X}, which can degrade hydrogen peroxide and catalyse the oxidation of nitrite to nitrate, which is a proton consuming process with increased rate at low pH (47). The latter mechanism is confirmed here under physiological conditions, as we show that biogenic MnO\textsubscript{X} can catalyze nitrite oxidation also at pH 8.

The MnO\textsubscript{X}-mediated hydrogen peroxide degradation as a mechanism to prevent peroxynitrite formation remains to be fully confirmed. Our experiments with spikes of hydrogen peroxide did not alter the gross dynamics of MnO\textsubscript{X} mediated nitrite-inhibition relief, but this could be due to the design of these experiments with hydrogen peroxide delivered in single doses rather than being delivered in a controlled manner in the vicinity of the cells. A single dose could have been readily dealt with by additional peroxidases, without altering MnO\textsubscript{X} mediated effects. On the other hand, our observation that the nitrite-stress is fully mitigated in NADH- supplemented cultures (even in the absence of MnO\textsubscript{X}) lends support to the idea that nitrite stress is mediated primarily through formation of peroxynitrite. In that case, the reductive power of NADH could be employed by peroxidases, as well as MnO\textsubscript{X}, to
reduce hydrogen peroxide (17, 58, 59) and thereby stopping the formation of peroxynitrite, explaining the observed mitigation effect of NADH.

These possible mechanistic scenarios of nitrite toxicity and roles of NADH, peroxidases, and MnOₓ in mitigating it, can shed light on why and if manganese oxidation is a functional, actively evolved trait or not. In particular, it is not clear why cells that already have several peroxidases, such as AzwK-3b (22, 27) (see also Table S1), might invest additional energy in the formation of MnOₓ precipitates. One possibility is that the the formation of MnOₓ is a mere side effect arising from the microbially generated superoxide (which appears widespread among bacteria) (27) reacting with the manganese (Mn²⁺), and the exoenzymes of AzwK-3b simply removing the resulting hydrogen peroxide that would otherwise lead to subsequent reduction of the oxidized manganese (24). An alternative possibility is that manganese oxidation is actively selected for due to the exact reaction mechanisms of ROS scavenging. It has been suggested, for example, that different ROS scavenging enzymes have different substrate affinities and efficiencies (18). In this context MnOₓ-mediated scavenging could be preferred under certain ROS concentrations and modes of production. In addition, and unlike peroxidases that require stoichiometric equivalents of reductants as e.g. NADH/NADPH for hydrogen peroxide reduction (18, 19), MnOₓ at its different oxidation states (II, III, IV) can, once formed, directly catalyze degradation of hydrogen peroxide without NADH involvement (23, 24, 26, 44, 50–57). The fact that some peroxidases, as well as the AzwK-3b enzyme catalyzing MnOₓ formation, are exoenzymes (22, 86) could be also highly relevant. The expression of such exoenzymes is a ‘social trait’, that can be exploited by cheating cells that do not invest the costs but reap the benefits (36–39). The presented finding that MnOₓ forms dispersed granules in the (agitated) liquid culture of AzwK-3b shows that, in this case, the ultimate functional effects arising from exoenzyme activity are localized. This kind of localization is a known strategy to stabilize a social trait in the face of evolution of cheating, as seen in exoenzymes with localized actions involved in sugar degradation (87) and metal scavenging (88). Thus, the reductive energy investment into the formation of MnOₓ mediated protection might be a metabolically less costly strategy that is also socially more stable, compared to for example exoenzymes that are freely diffusing.

Within a wider context, our findings are relevant to understand the different forms of metal mineralization observed in different microorganisms and under different ecological
contexts. Given the abundance of microorganisms being involved in reactions of the nitrogen cycle, there is indeed potential transient accumulation of nitrite in different environments. It is also possible that MnO₂ (or other minerals) can provide more broad protection against ROS chemistry. For example, manganese oxidation is also observed in spore-forming bacteria (89, 90), fungi and other microorganisms (as reviewed and shown in (2, 40)), where a role for nitrite stress remains to be elucidated. Our findings will facilitate such further studies of biomineralizing organisms and their different functional motives and social strategies.

**Materials and Methods**

**Bacterial Strain and Culture Conditions.** *Roseobacter sp.* AzwK-3b was obtained from Colleen Hansel (Woods Hole Oceanographic Institution, Falmouth, MA/USA), who isolated the strain (20). Cultures were grown in a defined medium, which was established by modifying the pre-defined artificial seawater (ASW) medium (35). This media is referred to as ASW<sub>m</sub> from now on, and its composition is shown in Table 1. ASW<sub>m</sub> contained sodium acetate as the sole carbon source (at concentrations specified per experiment), 200mM sodium chloride (instead of 428 mM, as in ASW), ammonium as nitrogen source (instead of nitrate, as in ASW), and five vitamins that were added as supplement. In manganese-supplemented ASW<sub>m</sub>, manganese chloride (MnCl₂) was added to 200 μM. Cultures were grown at 30 °C in appropriate (100 ml) Erlenmeyer flasks (shaking at 150 strokes per minute) or 96 well polystyrene plates (Corning Inc.) closed with lid and parafilm (shaking at 300 strokes per minute). For flask cultures, a MaxQ 4000 shaking incubator (Thermo Fisher Scientific) was used. Plates were incubated in a CLARIOstar plate reader (BMG labtech) and absorbance measurements were done at 600 nm (A<sub>600</sub>) and with path length-correction, so to present absorbance per 1 cm.

**Electron microscopy (EM) and Energy Dispersive X-ray spectroscopy (EDS) analysis of AzwK-3b cultures.** A culture of AzwK-3b (40 ml in 100 ml Erlenmeyer flasks) was inoculated in ASW<sub>m</sub> without manganese and nitrite, and containing 50 mM acetate. After 3 days at 150 strokes per minute shaking and 30 °C (by which time the culture reached the stationary phase), dilutions (25x – 200x) were made for a second passage of culture in the same medium, supplemented with 200 μM manganese. After further 2 days of culturing, samples for EM
were prepared as follows: Cells from 2.5 ml culture were harvested by centrifugation (5 min at 5,000 g), and the supernatant was discarded. From here, several washing and dehydration steps were conducted by re-suspending the pellet in different solutions and subsequently centrifuging for 5 min at 5,000 g (supernatant discarded): (1) first, pellets were twice re-suspended in ASW\textsubscript{m} medium basis (no manganese, no acetate, no ammonium, no nitrite, no trace metals); (2) afterwards, samples were re-suspended in 200 μl 70 % ethanol, incubated for 1 min, and pelleted by centrifugation; (3) for a washing-dehydration step, pellets were twice re-suspended in 200 μl 100 % ethanol and harvested by centrifugation; (4) finally, samples were re-suspended in 100 μl of 100 % ethanol. This suspension was then applied to Transmission Electron Microscopy (TEM) grids (Lacey carbon film coated copper grids (Agar Scientific)) by pipetting, in 1 μl portions (allowed to dry in between), until a total of 2 or 5 μl was accumulated (on different grids). After letting dry on the bench overnight, grids were analysed by EM.

EM analysis was done on a Gemini SEM 500 (Zeiss) equipped with EDS X-Max detector (Oxford Instruments). Data analysis was done on the associated AZtec software, which contained the spectral information to identify individual elements. Electron micrographs had the best quality in scanning transmission EM mode (STEM) with a high angle annular dark field detector (HAADF). For EDS, the sample needed to be moved, and the HAADF detector had to be withdrawn, so the location of analysis after changing the setup was confirmed by additional scanning EM (SEM) recording. The HAADF recording presented in Figure 6 was recorded at 25 kV and 4.3 mm working distance, with a 50,000 x magnification. The EDS was recorded at 25 kV, and spectra were accumulated for the same time (40 seconds for the two locations compared in Figure 6).

**Large inocula preparation for nitrite-assays.** AzwK-3b was grown in Erlenmeyer flasks (usually 40 ml culture volume in 100 ml Erlenmeyer flasks) in ASW\textsubscript{m} with 25 mM acetate. The culture absorbance \(A_{600}\) was recorded regularly on a Spectronic 200 spectrophotometer (Thermo Fisher) with 1 cm path length polystyrene cuvettes, and inocula were sampled at various stages of the growth curve (e.g. see Figures S4, S6, S8). This culture was used to inoculate into 96 well plates, which were supplemented by 1:1 dilution with fresh medium supplemented with manganese and/or nitrite and other additives, as described for the
particular results shown (see legends of Figures 4, 6, S6, S8). Where noted (see respective
figure captions), the fresh medium used for dilution was also supplemented with NADH or
hydrogen peroxide at different concentrations. NADH or hydrogen peroxide were added as
last additives (to prevent reaction e.g. between hydrogen peroxide and Mn^{II} before
inoculation) and the completed fresh medium was used immediately.

Growth curve fitting and analysis. Growth curves were analyzed using the R-package Grofit
(91) applying the Gompertz growth model (91, 92). Plate reader data (measurements every
10 minutes) were de-noised by averaging over 6 measurements (i.e. hourly averages). The
maximum $A_{600}$ reached was read directly from the data. For curve fitting, all data later than
the maximum $A_{600}$, i.e. decaying growth phase, were removed. Then, the data was read
backwards in time to find the first reading that was below 5 % of the maximum $A_{600}$. This data-
trimming was done to facilitate the fitting of the Gompertz growth model without bias from
different lag-phases (which were ignored), or different lengths and scales of decaying phases
recorded. From the resulting model, the maximum growth rate $\mu$ (in $A_{600}$ nm(a.u.) per hour)
was recorded.

Preparation of cell-free bio-manganese oxide. The procedure was adapted from previous
publications using the cell free supernatant of Roseobacter sp. AzwK-3b grown in complex
medium (20, 22–24). AzwK-3b was grown in ASW$_m$ supplemented with 50 mM sodium acetate
for nine days, using individual 50 or 100 ml cultures in 100 or 200 ml Erlenmeyer flasks,
respectively, at 30 °C with shaking (150 strokes per minute). In total, 2 liters of culture was
prepared, cells were removed by centrifugation (5 minutes at 10,000 g) and the supernatants
were combined. From this (cell-free) supernatant, individual samples of 100 or 200 ml were
prepared and supplemented with 200 μM manganese chloride, MnCl$_2$. Manganese oxidation
was allowed to proceed for five days at 30 °C with shaking (150 strokes per minute), after
which the manganese oxide was harvested by centrifugation (5 minutes at 10,000 g) from
each 50/100 ml sample. These were combined and washed by suspending in 25 ml acetate-
free ASW$_m$ medium and re-sedimented by centrifugation. The pellet was brown in
appearance and had considerable volume, indicating co-precipitation of organic material (e.g.
secreted proteins) from the cell-culture supernatant. To estimate the amount of manganese
precipitated in the assay, the supernatants from centrifugation and the washing steps were combined, and the residual manganese determined by the 3,3',5,5'-tetramethylbenzidine (TMB)-assay (93) for soluble manganese. Note that this was not a precise quantification, but was conclusive enough to allow conservative stoichiometric relations to be inferred. In particular, we inferred that ca. 75 % of the 200 μM manganese chloride had been removed from the solution and this value was used for downstream calculations. The MnO$_X$ precipitate was suspended in an appropriate volume of the acetate-free medium to produce a “10 mM” suspension of manganese oxide, and this value is used in the manuscript as indicator for manganese oxide concentration. The pH was 8.2, which is well in line with the pH 8.0 of the ASW$_m$ medium, showing that the suspended manganese oxide did not alter the pH.

Quantification of nitrite, nitrate and acetate. Quantification was done by Ion Chromatography (IC) on a DIONEX ICS-5000+ (ThermoFisher, UK) equipped with conductivity detector, potassium hydroxide (KOH) eluent generator, appropriate suppressor, and a DIONEX IonPac AS11-HC-4μm (2 x 250 mm ThermoFisher, UK) anion separation column with appropriate guard column. Culture samples were filtered (0.22 μm polyamide spin filter Costar Spin-X, Corning, NY/USA) and 10-fold diluted with MilliQ-water (resistance R > 18.2 MΩ), of which 2.5 μl were injected for IC separation. The IC was run as continuous gradient as follows (flow rate 0.38 ml/min, column temperature 30 °C, conductivity detector cell temperature of 35 °C): -7-0 min – 1.5 mM KOH (equilibration), 0-8 min – 1.5 mM KOH, 8-18 increase to 15 mM KOH, 18-23 min – increase to 24 mM KOH, 23-24 min – increase to 60 mM KOH, 24-30 min – stay at 60 mM KOH. Reference samples with known concentrations were run for calibration. During the course of the experiments (see below) evaporation of the samples was noted (indicated by the increase in the peak area of chloride, which is expected to be unaltered by any biologic means and therefore should have displayed no concentration change). To correct for this evaporation effect, the concentrations of the analytes of interest were corrected by the same ratio as that obtained from the chloride peak area (from the beginning and end point samples of a particular time-course experiment).
Acknowledgments

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Conflict of Interest

The authors declare that there are no conflicts of interest.

Supplementary Information

Supplementary information is available at journal’s website.
References


Impact of agriculture and land use on nitrate contamination in groundwater and running waters in central-west Poland. Environ Monit Assess 188:172.


Table 1. Detailed composition of the defined AzwK-3b growth medium, ASW_m. The medium was developed starting out from artificial seawater (ASW) (35) with extra trace metals taken from (9, 94) and a 5-vitamin solution identified starting out from Wolfe’s vitamin mixture (95).

<table>
<thead>
<tr>
<th>Compound</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Base salts (1 x AzwK-3b medium)</strong></td>
<td></td>
</tr>
<tr>
<td>Sodium chloride (NaCl)</td>
<td>200 mM</td>
</tr>
<tr>
<td>Ammonium chloride (NH₄Cl)</td>
<td>8.82 mM</td>
</tr>
<tr>
<td>Potassium chloride (KCl)</td>
<td>6.71 mM</td>
</tr>
<tr>
<td>Di-potassium hydrogenphosphate (KH₂PO₄)</td>
<td>131 µM</td>
</tr>
<tr>
<td>Magnesium sulphate (MgSO₄)</td>
<td>14.2 mM</td>
</tr>
<tr>
<td>Magnesium chloride (MgCl₂)</td>
<td>9.84 mM</td>
</tr>
<tr>
<td>Calcium chloride (CaCl₂)</td>
<td>3 mM</td>
</tr>
<tr>
<td>Tris(hydroxymethyl)aminomethane (TRIS)</td>
<td>1.1 mM</td>
</tr>
<tr>
<td><strong>pH of the medium</strong></td>
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</tr>
<tr>
<td><strong>Trace metal solution (1,000 x)</strong></td>
<td></td>
</tr>
<tr>
<td>Copper chloride (CuCl₂)</td>
<td>32 µM</td>
</tr>
<tr>
<td>Zink sulphate (ZnSO₄)</td>
<td>765 µM</td>
</tr>
<tr>
<td>Cobalt chloride (CoCl₂)</td>
<td>169 µM</td>
</tr>
<tr>
<td>Sodium molybdate (Na₂MoO₄)</td>
<td>1.65 mM</td>
</tr>
<tr>
<td>Boric acid (H₃BO₃)</td>
<td>46.3 mM</td>
</tr>
<tr>
<td>Nickel chloride (NiCl₂)</td>
<td>4.2 mM</td>
</tr>
<tr>
<td>Sodium tungstate (Na₂WoO₄)</td>
<td>243 µM</td>
</tr>
<tr>
<td>Sodium selenite (Na₂SeO₃)</td>
<td>228 µM</td>
</tr>
<tr>
<td><strong>Additional (1,000 x) supplement solutions</strong></td>
<td></td>
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<tr>
<td>Iron chloride (FeCl₃; prepared in 10 mM HCl, balanced with extra 10 mM NaOH solution)</td>
<td>10.4 mM</td>
</tr>
<tr>
<td>Ethylenediaminetetraacetate (EDTA, pH 8.0; sodium salt)</td>
<td>1.34 mM</td>
</tr>
<tr>
<td>Manganese chloride (MnCl₂, only added where desired)</td>
<td>200 mM</td>
</tr>
<tr>
<td><strong>Vitamin supplement (1,000 x)</strong></td>
<td></td>
</tr>
<tr>
<td>Biotin</td>
<td>82 µM</td>
</tr>
<tr>
<td>Pyridoxine hydrochloride</td>
<td>484 µM</td>
</tr>
<tr>
<td>Thiamine hydrochloride</td>
<td>148 µM</td>
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<tr>
<td>Riboflavin</td>
<td>133 µM</td>
</tr>
<tr>
<td>Nicotinic acid</td>
<td>406 µM</td>
</tr>
</tbody>
</table>
Figure 1. Biological oxidation of manganese via superoxide, and nitrite oxidation by the product manganese oxide. These reactions are taken from references (24) (manganese oxidation) and (47) (nitrite oxidation). Note that only representative reactions are presented. For instance, the text refers to a mixed oxide (MnOₓ), while this reaction scheme simplyfies to MnO₂. The cellular reductant (cell. reductant) which serves as electron donor for superoxide production is not unambiguously identified.

Figure 2. Effect of Mn²⁺ on the growth of Roseobacter sp. AzwK-3b in the defined growth medium (see Table 1). The concentrations of manganese were 0 μM (black), 200 μM (red) and 500 μM (dark green), with no growth (zero line) in the respective non-inoculated controls (blue, magenta, light blue). Cultures were grown in a 96 well plate (200 μl culture) with shaking and absorbance measurement every 10 minutes (see Methods).
**Figure 3.** Growth of *Roseobacter* sp. AzwK-3b in the defined growth medium supplemented with sodium nitrite. Media were prepared without (Figure A) or with (Figure B) 200 μM manganese chloride, MnCl$_2$. Nitrite-concentrations were 0 mM (black), 0.25 mM (red), 0.5 mM (green), 1 mM (dark blue) and 2.5 mM (light blue). All conditions were tested in triplicates, and the growth curves represent averages and their standard deviations (see Methods).
Figure 4. Larger AzwK-3b inocula are less inhibited by nitrite. A pre-culture without manganese or nitrite was grown and sampled in the exponential growth phase (Figure S4) to prepare inocula from a very early time point in the exponential phase (IT 1, Figures A and B), and from a later time point (IT 2, Figures C and D; both sampled in first third of exponential phase). These inocula were 1:1 diluted with fresh medium, and tested for growth at different nitrite concentrations (see below for colour code) without (A, C) or with (B, D) 200 μM Mn$^{2+}$Cl$_2$ supplement. The nitrite concentrations were: Black – control no nitrite. Red – 0.25 mM nitrite. Green – 0.5 mM nitrite. Blue – 1 mM nitrite. Yellow – 2 mM nitrite. Magenta – 5 mM nitrite. Light blue – 7.5 mM nitrite. Dark red – 10 mM nitrite. Growth curves show the averages and standard deviations over a triplicate analysis (see Methods).
Figure 5. Inoculum-size effect on MnO\textsubscript{x} mediated mitigation of nitrite-inhibition. Data from different AzwK-3b growth experiments of similar type (“Large inocula”, see Methods) were analyzed for the maximum A\textsubscript{600} (bottom row) and growth rate (top row) by fitting the growth curves. Each condition was done in three technical replicates (note that error bars are not visible in some cases due to only small differences). Nitrite-concentrations of the main-cultures are indicated as headings of the figure-columns. The x-axes show the calculated A\textsubscript{600} of the initial cultures after diluting them 1:1 from the pre-cultures, while the y-axes show the maximum A\textsubscript{600} and maximum growth rate as calculated with the Gompertz model (91, 92) (see Methods). The colours represent different conditions: Red: Neither pre-, nor main-culture contained manganese; Blue: Pre-culture without, main-culture with manganese; Green: both pre- and main-culture with manganese. The black curve is a sigmoidal fit (logistic model) from the Grofit R-package (91), for the results of the combined blue and green dataset where the nitrite-exposed main-cultures all contained manganese.
Figure 6. Scanning transmission electron micrograph (left figure, high angle annular dark field) of (granular) manganese-containing precipitate (center) surrounded by AzwK-3b cells, and associated energy dispersive X-ray spectroscopic analysis (right figure) in this location. Only the energy range containing the manganese-specific X-ray energies at 5.90 keV ($K_{\alpha 1}$) and 6.49 keV ($K_{\beta 1}$) is shown, and the manganese transitions are indicated by vertical gray dashed lines.
**Figure 7.** Oxidation of nitrite by biogenic manganese oxide (MnO$_x$) produced in cell-free culture supernatant of AzwK-3b. The figures show the concentration of nitrite (A) and nitrate (B), determined by ion chromatography, over time (note that concentrations were corrected for the IC-peak from chloride, to account for evaporation during the experiment). As controls, samples without MnO$_x$ (green), or with MnO$_2$ powder (orange) were included in the experiment (see Methods). The samples with AzwK-3b cell-free manganese oxide contained (from grey to black) 0.2, 0.5, 1 and 2 mM manganese oxide equivalent (see Methods).
Figure 8. Reductive power (NADH) mitigates the growth inhibitory effects of nitrite in AzwK-3b. Cultures (pre- and main-culture without manganese) were grown in the absence (A) and presence (B) of 5 mM nitrite and supplement of 0, 50, 100 and 200 μM NADH (black, red, green and blue) at the start of the culture.