

1 **Novel insights into the taxonomic diversity and molecular mechanisms of bacterial Mn(III)**
2 **reduction**

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15 **Running title:** Novel undecaheme in Betaproteobacteria

16 **Originality-Significance Statement:** Recent observations suggest that Mn(III)-ligand
17 complexes are geochemically important in diverse aquatic environments. Thus far, microbially-
18 driven Mn(III) reduction has only been associated with *Gammaproteobacteria* encoding three-
19 component outer-membrane porin-cytochrome c conduits. Here, we demonstrate that
20 *Betaproteobacteria* dominate in abundance and with respect to protein expression during
21 biologically-mediated Mn(III) reduction in an enrichment culture from an anoxic lacustrine
22 system. Using metaproteomics, we detect for the first time that *Betaproteobacteria* express a
23 two-component porin-cytochrome c conduit, and an uncharacterized extracellular undecaheme
24 (11-heme) c-type cytochrome. Although this is the first definitive report of an undecaheme
25 within the *Betaproteobacteria*, we find evidence that they are widespread in uncultivated strains.
26 These results widen the phylogenetic diversity of Mn(III)-reducing bacteria, and provide new
27 insights into potential molecular mechanisms for soluble Mn(III) reduction.

28 **Summary:** Soluble ligand-bound Mn(III) can support anaerobic microbial respiration in diverse
29 aquatic environments. Thus far, Mn(III) reduction has only been associated with certain
30 *Gammaproteobacteria*. Here, we characterized microbial communities enriched from Mn-replete
31 sediments of Lake Matano, Indonesia. Our results provide the first evidence for biological
32 reduction of soluble Mn(III) outside the *Gammaproteobacteria*. Metagenome assembly and
33 binning revealed a novel betaproteobacterium, which we designate “*Candidatus* Dechloromonas
34 *occultata*.” This organism dominated the enrichment and expressed a porin-cytochrome c
35 complex typically associated with iron-oxidizing *Betaproteobacteria* and a novel cytochrome c-
36 rich protein cluster (Occ), including an undecaheme putatively involved in extracellular electron
37 transfer. This *occ* gene cluster was also detected in diverse aquatic bacteria, including
38 uncultivated *Betaproteobacteria* from the deep subsurface. These observations provide new
39 insight into the taxonomic and functional diversity of microbially-driven Mn(III) reduction in
40 natural environments.

41 **Introduction.** Manganese(III) is a strong oxidant with a reduction potential close to molecular
42 oxygen (Kostka et al., 1995). Mn(III) is short-lived and unstable, but its stability is greatly
43 increased when bound to ligands (Luther III et al., 2015). Ligand-bound Mn(III) is often the most
44 abundant dissolved Mn species in sediment porewaters (Madison et al., 2013; Oldham et al.,
45 2019) and soils (Heintze and Mann, 1947), with the potential to facilitate one-electron redox
46 reactions in a variety of biogeochemical cycles (Luther III et al., 2015). Microbes accelerate the
47 oxidation and reduction of Mn by orders of magnitude compared to abiotic mechanisms (Hem,
48 1963; Diem and Stumm, 1984; Morgan, 2005; Tebo et al., 2005; Learman et al., 2011; Luther et
49 al., 2018; Jung et al., 2020). Yet, despite clear evidence for the environmental importance of
50 Mn(III), knowledge about microbial Mn(III) cycling pathways remains fragmented.

51 To date, only *Shewanella* spp. (*Gammaproteobacteria*) have been confirmed to respire
52 soluble Mn(III) (Kostka et al., 1995; Szeinbaum et al., 2014). *Shewanella* respire Mn(III) using
53 the Mtr pathway (Szeinbaum et al., 2017), a porin-cytochrome (PCC) conduit that transports
54 electrons across the periplasm for extracellular respiration of Mn(III/IV), Fe(III), and other
55 metals (Richardson et al., 2012; Shi et al., 2016). Many Fe(II)-oxidizing *Betaproteobacteria* also
56 contain PCCs (MtoAB, generally lacking the C subunit), which are proposed to oxidize Fe(II) to
57 Fe(III) by running the PCC in reverse (Emerson et al., 2013; Kato et al., 2015; He et al., 2017).
58 In some metal-reducing *Gammaproteobacteria* and *Deltaproteobacteria*, extracellular
59 undecaheme (11-heme) UndA is thought to play a key functional role in soluble Fe(III) reduction
60 (Fredrickson et al., 2008; Shi et al., 2011; Smith et al., 2013; Yang et al., 2013). UndA's crystal
61 structure shows a surface-exposed heme surrounded by positive charges, which may bind
62 negatively-charged soluble iron chelates (Edwards et al., 2012). Environmental omics suggest
63 that metal reduction by *Betaproteobacteria* may be widespread in the deep subsurface

64 (Anantharaman et al., 2016; HERNSDORF et al., 2017). However, only a few Fe(III)-reducing
65 *Betaproteobacteria* isolates have been characterized to date (Cummings et al., 1999; Finneran et
66 al., 2003), and little is known about metal reduction pathways in *Betaproteobacteria*.

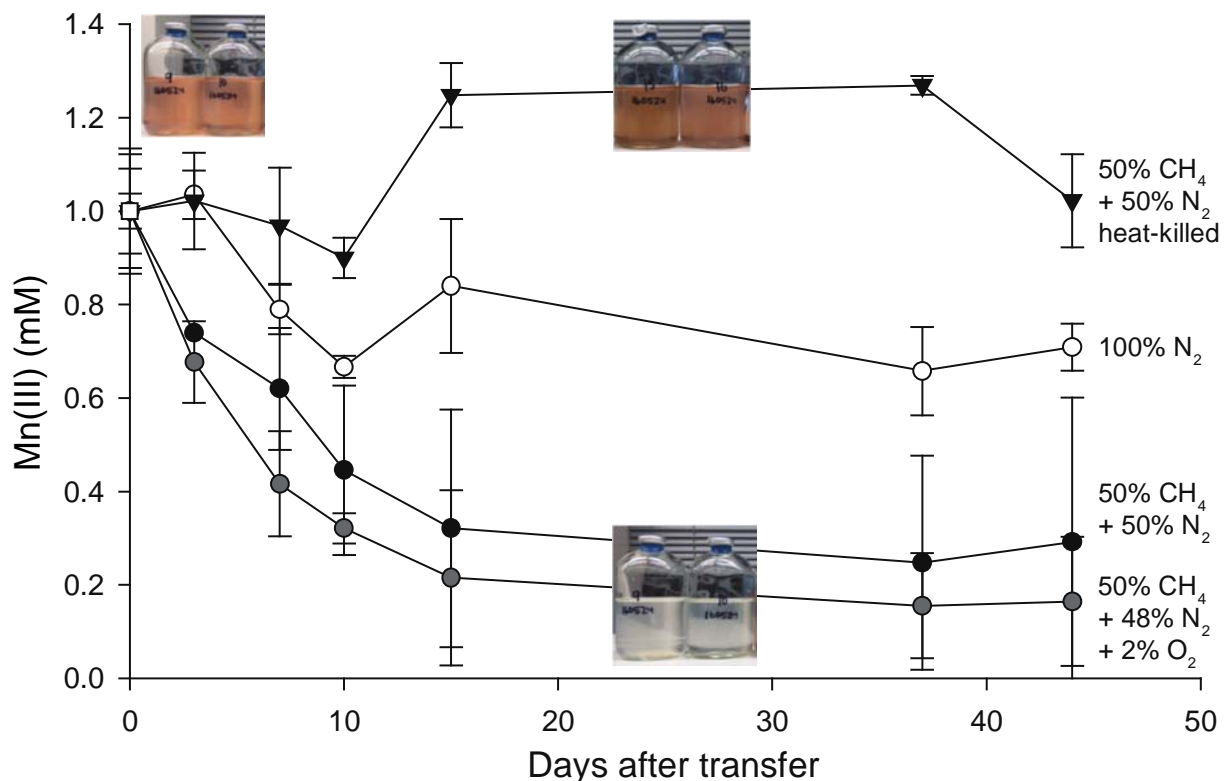
67 Manganese reduction coupled to methane (CH₄) oxidation is a novel metabolism only
68 recently discovered in cultures enriched in Archaea (Ettwig et al., 2016; Leu et al., 2020).
69 Biological and geochemical evidence suggest that this metabolism may be found in a variety of
70 environments (Beal et al., 2009; Crowe et al., 2011; Riedinger et al., 2014), including Fe-rich
71 Lake Matano, Indonesia. In an attempt to explore whether CH₄ can fuel microbial Mn(III)
72 reduction in enrichments inoculated with sediments from Lake Matano, Indonesia, which has
73 active and pronounced microbial Mn and CH₄ cycles (Jones et al., 2011), we uncovered a novel
74 betaproteobacterium as the most dominant and active member of our Mn(III)-reducing
75 enrichment culture. Our results provide the first evidence for biological reduction of soluble
76 Mn(III) outside *Gammaproteobacteria* and provide evidence for a new biochemical pathway
77 involved in extracellular electron transfer.

78

79 **Results and discussion**

80 ***Enrichment of Mn(III)-reducing populations.*** Lake Matano, Indonesia, is a permanently
81 stratified ultraoligotrophic lake (Crowe et al., 2008). Below its oxic surface waters, Lake
82 Matano's permanently anoxic and stratified waters are highly enriched in iron and manganese,
83 and support the activity of Mn cycling organisms with organic carbon and CH₄ as potential
84 sources of electrons (Crowe et al., 2011; Jones et al., 2011; Kuntz et al., 2015; Sturm et al.,
85 2019). We designed an enrichment strategy to select for microbes capable of anaerobic CH₄
86 oxidation coupled to soluble Mn(III) reduction by incubating anoxic Lake Matano sediment

87 communities with soluble Mn(III)-pyrophosphate as the electron acceptor (with 2% O₂ in a
88 subset of bottles), and CH₄ as the sole electron donor and carbon source after pre-incubation to
89 deplete endogenous organic carbon (see **Supporting Information** for enrichment details).
90 Enrichment cultures were transferred into fresh media after Mn(III) was completely reduced to
91 Mn(II), for a total of five transfers over 395 days. By the fourth transfer, cultures with CH₄
92 headspace (with or without 2% O₂) reduced ~80% of soluble Mn(III) compared to ~30% with N₂
93 headspace (**Fig. 1**). 16S rRNA gene sequences were dominated by *Betaproteobacteria*
94 (*Rhodocyclales*; 8-35%) and *Deltaproteobacteria* (*Desulfuromonadales*; 13-26%; **Fig. S1**).
95 ¹³CH₄ oxidation to ¹³CO₂ was undetectable (**Fig. S2**).



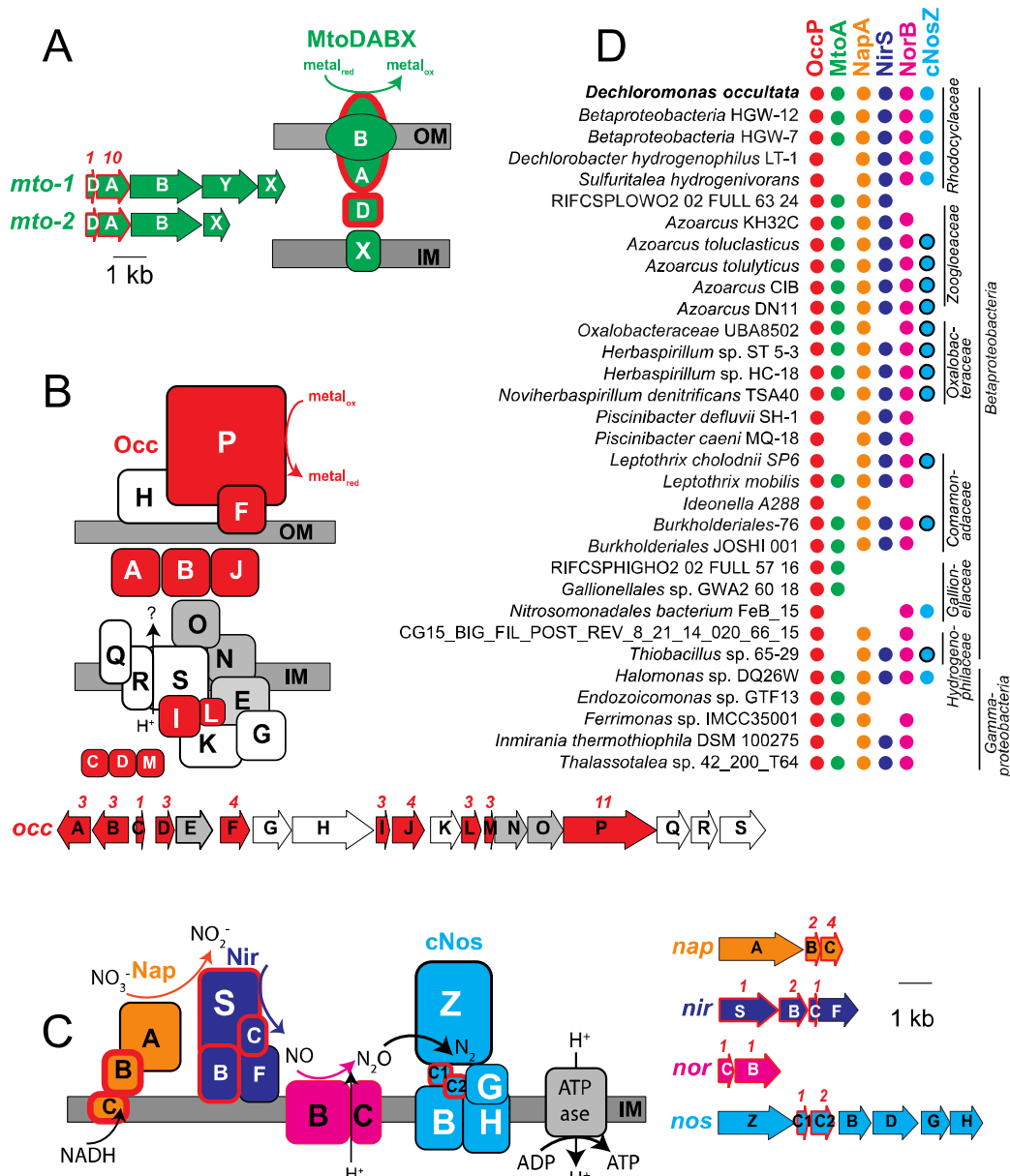
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97 **Figure 1. Consumption of Mn(III) in Lake Matano enrichments in the presence and absence of methane.**
98 Sediment-free cultures (transfer 4), from 335 days after the initial enrichment, were incubated for 45 days with 1
99 mM Mn(III) pyrophosphate as the sole electron acceptor. One set was incubated with Mn(III) and 2% O₂. Initial
100 bottle headspace contained 50% CH₄ + 50% N₂ (black circles), 50% CH₄+48% N₂+2% O₂ (gray circles), 100% N₂
101 (white circles), and 50% CH₄+50% N₂ heat killed controls (black triangles). Error bars are standard deviations from
102 duplicate experiments. Color change from red to clear indicates Mn(III) reduction.
103

104 Samples for metagenomic and metaproteomic analysis were harvested from the fifth
105 transfer (**Fig. 1; Fig. S1**). Out of 2,952 proteins identified in the proteome, 90% were assigned to
106 *Betaproteobacteria*; of those, 72% mapped to a 99.5% complete metagenome-assembled genome
107 (MAG; *Rhodocyclales* bacterium GT-UBC; NCBI accession QXPY01000000) with 81-82%
108 average nucleotide identity (ANI) and phylogenetic affiliation to *Dechloromonas* spp. (**Table**
109 **S1; Fig. S3**). This MAG is named here “*Candidatus Dechloromonas occultata*” sp. nov.;
110 etymology: *occultata*; (L. fem. adj. ‘hidden’). The remaining 10% of proteins mapped to
111 *Deltaproteobacteria*; of those, 70% mapped to a nearly complete MAG (*Desulfuromonadales*
112 bacterium GT-UBC; NCBI accession RHLS01000000) with 80% ANI to *Geobacter*
113 *sulfurreducens*. This MAG is named here “*Candidatus Geobacter occultata*”.

114 ***Cytochrome expression during Mn(III) reduction.*** Cytochromes containing multiple *c*-
115 type hemes are key for electron transport during microbial metal transformations, and therefore
116 also expected to play a role in Mn(III) reduction. Numerous mono-, di-, and multi (>3)-heme
117 cytochromes (MHCs) were expressed by “*Ca. D. occultata*” in Mn(III)-reducing cultures. Nine
118 out of 15 MHCs encoded by the “*Ca. D. occultata*” MAG were expressed, including two
119 decahemes similar to MtoA in Fe(II)-oxidizing *Betaproteobacteria* (**Tables 1, S2, S3; Figs. 2A,**
120 **S4**). Several highly expressed MHCs were encoded on a previously unreported 19-gene cluster
121 with 10 cytochrome-*c* proteins, hereafter *occA-S* (**Table 1; Figs. 2B, S5, S6**). OccP was
122 predicted to be an extracellular undecaheme protein of ~100 kDa (922 amino acids). “*Ca.*
123 *Dechloromonas occultata*” may reduce Mn(III) using the novel extracellular undecaheme OccP
124 as the terminal Mn(III) reductase. Experimental verification of the function of the putative Occ
125 complex is currently limited by the scarcity of genetically tractable *Betaproteobacteria*.

126 Proteins with 40-60% identity to the expressed “*Ca. D. occultata*” OccP protein were
 127 widely distributed in *Betaproteobacteria* from diverse freshwaters and deep subsurface
 128 groundwaters, as well as in several *Gammaproteobacteria* and one alphaproteobacterium (**Fig.**
 129 **2D; Table S3**). Most *occP*-containing bacteria also possessed *mtoA* and denitrification genes
 130 (**Fig. 2D; Figs. S7, S8**). These results widen the phylogenetic diversity of candidate extracellular
 131 MHCs that may be involved in microbial Mn(III) reduction.



132 **Figure 2. Gene arrangement, predicted protein location, and taxonomic distribution of major expressed**
 133 **respiratory complexes in “*Ca. D. occultata*”.** A: MtoDAB(Y)X porin-cytochrome c electron conduit; B: OccA-S;
 134

135 **C:** denitrification complexes (Nap, Nir, Nor and cNos); **D:** Occurrence of key marker genes in *Betaproteobacteria*
136 and *Gammaproteobacteria* with >95% complete genomes that encode OccP. Protein sequences from “*Ca. D.*
137 *occultata*” were used as query against a genome database and searched using PSI BLAST. Matches with identities
138 >40%, query coverage >80% and E values <10⁻⁵ were considered positive. Red fill around genes and proteins
139 indicate cytochrome-*c* proteins. Black outlines around blue circles in D indicate type I nitrous oxide reductase to
140 distinguish from blue dots (type II/cytochrome-nitrous oxide reductase). Gray-shaded genes on the *occ* gene cluster
141 indicate 6-NHL repeat proteins. Protein locations shown are based on P-sort predictions. Numbers above genes
142 indicate number of CxxCH motifs predicted to bind cytochrome *c*. IM: inner membrane; OM: outer membrane. For
143 more details, see **Table 1** and **Table S3**.

144 ***Heme-copper oxidases in “Ca. D. occultata”.*** “*Ca. D. occultata*” expressed high-affinity
145 *cbb*₃-type cytochrome *c* oxidase (CcoNOQP) associated with microaerobic respiration (**Table**
146 **S4**). Features of the “*Ca. D. occultata*” *occS* gene product, including conserved histidine residues
147 (H-94, H-411, and H-413) that bind hemes *a* and *a*₃, as well as the H-276 residue that binds Cu_B
148 (**Fig. S6**), suggest that OccS may function similarly to CcoN, the terminal heme-copper oxidase
149 proton pump in aerobic respiration. All identified OccS amino acid sequences lack Cu_B ligands
150 Y-280 and H-403, and most lack Cu_B ligands H-325 and H-326. OccS sequences also lack polar
151 and ionizable amino acids that comprise the well-studied D and K channels involved in proton
152 translocation in characterized cytochrome *c* oxidases (Blomberg and Siegbahn, 2014), but
153 contain conserved H, C, E, D, and Y residues that may serve in alternate proton translocation
154 pathways, similar to those recently discovered in qNOR (Gonska et al., 2018). OccS homologs
155 were also found in *Azoarcus* spp. and deep subsurface *Betaproteobacteria* (**Fig. S6**).

156 ***Expression of denitrification proteins and possible sources of oxidized nitrogen species.***

157 Periplasmic nitrate reductase (NapA), cytochrome nitrite reductase (NirS), and type II atypical
158 nitrous oxide reductase (cNosZ; **Fig. S7**) were highly expressed by “*Ca. D. occultata*” (**Table 1**).
159 Expression of the denitrification pathway was not expected because oxidized nitrogen species
160 were not added to the medium, to which the only nitrogen supplied was 0.2 mM NH₄Cl (along
161 with headspace N₂). Oxidized nitrogen species could result from the oxidation of NH₄Cl, but we
162 did not find any of the canonical genes for aerobic nor anaerobic ammonia oxidation, nor did we

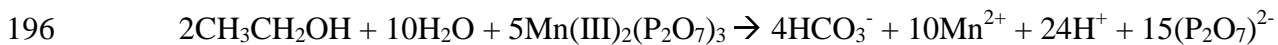
163 measure any ammonium oxidation in experimental bottles from the transfer used to make **Figure**
164 **1**.

165 The expression of denitrification genes is controlled by a diverse array of transcriptional
166 regulators that depend on different signals including low levels of oxygen, even in the absence of
167 nitrate (Spiro, 2012; Lin et al., 2018). The close redox potential of Mn³⁺-pyrophosphate (~0.8 V;
168 Yamaguchi and Sawyer, 1985) to oxidized nitrogen species (0.35-0.75 V) at circumneutral pH
169 and the lack of oxygen in the media could have induced the expression of denitrification genes
170 simultaneously with Mn(III)-reduction genes. *Gammaproteobacteria*, for example, reduce
171 Mn(III) even in the presence of nitrate (Kostka et al., 1995), and there is precedent for microbial
172 use of multiple electron acceptors, e.g. “co-respiration” of oxygen and nitrate during aerobic
173 denitrification (Chen and Strous, 2013; Ji et al., 2015).

174 Because solid-phase Mn(III) is known to chemically oxidize NH₄⁺ (Aigle et al., 2017;
175 Boumaiza et al., 2018), we tested for abiotic NH₄⁺ oxidation by soluble Mn(III) (1 mM).
176 Ammonium concentrations remained unchanged, and no N₂O or NO_x⁻ production was observed
177 (**Fig. S8**), likely because our experiments lacked solid surfaces to mediate electron transfer.
178 Similarly, N₂O levels in the headspace of our experimental bottles with Mn(III)-reducing
179 cultures were near or below the detection limit (data not shown). These findings are consistent
180 with lack of detectable ammonium oxidation by Mn(III) pyrophosphate in estuarine sediments
181 (Crowe et al., 2012).

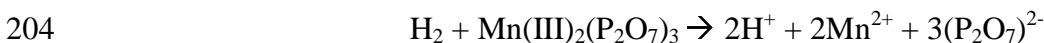
182 ***Electron donors.*** Methane was the only electron donor added intentionally to the
183 enrichment cultures, to select for organisms that oxidize methane anaerobically. Yet, we did not
184 detect ¹³CO₂ after addition of ¹³CH₄ (**Fig. S2**). One explanation is that ¹³CO₂ was produced, but
185 was subsequently assimilated by other members of the microbial community such as abundant

186 Deltaproteobacteria (**Fig. S1**), as observed in previous studies (Wegener et al., 2008). A filtration
187 step included in our protocol to measure $^{13}\text{CO}_2$ would have excluded ^{13}C -enriched biomass from
188 our analyses. Alternatively, we considered other electron donors that might have been
189 unintentionally present in trace amounts, but sufficiently abundant to drive the observed ~300-
190 600 μM Mn(III) reduction (**Fig. 1**). The ethanol catabolism pathway (PQQ-dependent
191 methanol/ethanol dehydrogenase (RIX45050), quinoprotein alcohol dehydrogenase (RIX45053),
192 and an NAD^+ -dependent aldehyde dehydrogenase-II (RIX45061)) were all highly expressed in
193 “*Ca. D. occultata*” (**Table 1**). Ethanol could have been introduced to the bottles during culture
194 preparation during sterilization of bottle stoppers. Based on the stoichiometry of ethanol
195 oxidation coupled to Mn(III) reduction:



197 150 μM ethanol would be required to reduce 600 μM of Mn(III), which equates to ~1 μL of 70%
198 ethanol (12 M) into 100 mL culture medium. We conclude that trace contamination of ethanol
199 was likely the major electron donor to our cultures.

200 It is also possible that other substrates, such as H_2 from fermentation by other microbes in
201 the enrichment or from impurities in the headspace gas, could have supplied another source of
202 electrons. Indeed, an NAD-reducing hydrogenase (RIX44099-100) was expressed by “*Ca. D.*
203 *occultata*” (**Table 1**). Based on the stoichiometry of H_2 oxidation coupled to Mn(III) reduction:

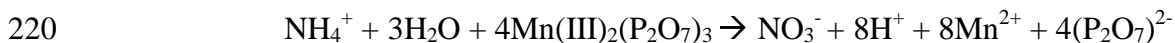


205 600 μM H_2 would be required to reduce 600 μM of Mn(III). Thus, H_2 may have contributed
206 electrons to Mn(III) reduction, but is not likely sole electron donor. A combination of ethanol,
207 H_2 and other trace contaminants would likely have been necessary to provide enough electrons
208 for the additional reduction of Mn(III) observed in the $^{13}\text{CH}_4$ -amended cultures compared to the

209 controls lacking $^{13}\text{CH}_4$. There is precedent for other metal-reducers simultaneously using H_2 and
210 an organic electron donor (Brown et al., 2005).

211 Another trace source of organics to our cultures could have been leaching from the rubber
212 stoppers, which were black bromobutyl and pre-boiled in 0.1 N NaOH. A previous study
213 reported that organics leaked an array of n-alkanes ($\text{C}_{16}\text{--}\text{C}_{34}$) and unidentified organic
214 contaminants in black bromobutyl stoppers (Niemann et al., 2015). It is also conceivable that
215 trace organic was introduced as impurities in solid Mn(III) oxide powder (99% purity) used to
216 synthesize Mn(III)-pyrophosphate.

217 Finally, we considered the possibility that 0.2 mM NH_4^+ , added to the cultures as a
218 nitrogen source, could have provided the electron donor, via an unknown pathway. Based on the
219 stoichiometry of NH_4^+ oxidation coupled to Mn(III) reduction:



221 0.2 mM NH_4^+ would supply 1.6 mM electron equivalents, which is more than enough to account
222 for the observed reduction of 600 μM of Mn(III). This process could operate cryptically if the
223 oxidized products were reduced to N_2 via denitrification enzymes, such as nitrous oxide
224 reductase (cNosZ), which was one of the most abundant proteins expressed in Mn(III)-reducing
225 cultures (**Fig. 2c, Table 1**).

226 **Carbon metabolism.** “*Ca. D. occultata*” appeared to be growing mixotrophically. “*Ca. D.*
227 *occultata*” encoded several central metabolic pathways, including a complete TCA cycle with a
228 glyoxylate bypass, an incomplete (acetate-dependent) 3-hydroxypropionate bicycle (3-HP), a
229 modified Calvin-Benson-Bassham (CBB) pathway, and a pathway for synthesis of
230 polyhydroxybutyrate (**Fig. S11**). In addition, “*Ca. D. occultata*” encoded genes for organic
231 carbon transport, and lactate, acetate, and propionate utilization (**Fig. S11**). Like *D. agitata* and

232 *D. denitrificans*, the CBB pathway of “*Ca. D. occultata*” did not encode RuBisCO and
233 sedoheptulose-1,7-bisphosphatase (SHbisPase; **Fig. S10**); SHbisPase may be replaced by 6-
234 phosphofructokinase and an energy-generating pyrophosphatase (RIX41248; Kleiner et al.,
235 2012; Zorz et al., 2018). The presence of incomplete carbon fixation pathways and for organic
236 carbon utilization pathways suggests that “*Ca. D. occultata*” relies on organic carbon to fix
237 inorganic carbon mixotrophically. The source of this organic carbon could have been ethanol,
238 which is converted to acetate via the pathway discussed in the previous section.

239 ***Effect of methane.*** Although we did not measure appreciable $^{13}\text{CH}_4$ oxidation to $^{13}\text{CO}_2$,
240 CH_4 stimulated Mn(III) reduction and cytochrome expression in “*Ca. D. occultata*” enrichment
241 cultures. While the specific role of CH_4 in Mn(III) reduction remains unknown, the addition of
242 CH_4 appeared to significantly stimulate expression of many cytochrome *c* proteins, including
243 OccABGJK, MtoD-2, and cytochrome-*c4* and -*c5* proteins associated with anaerobic respiration
244 ($p < 0.05$; **Table 1**; **Fig. 2C**). Expression of several “*Ca. D. occultata*” proteins involved in outer
245 membrane structure and composition - including an extracellular DUF4214 protein located next
246 to an S-layer protein similar to those involved in manganese binding and deposition (Wang et al.,
247 2009), a serine protease possibly involved in Fe(III) particle attachment (Burns et al., 2009), an
248 extracellular PEP-CTERM sorting protein for protein export (Haft et al., 2006), and a Tol-Pal
249 system for outer membrane integrity - was higher in the presence of CH_4 (**Table 1**).

250 ***Transporters and sensors.*** Numerous transporters were present in the “*Ca. D. occultata*”
251 genome, including 26 TonB-dependent siderophore transporters, 13 TRAP transporters for
252 dicarboxylate transport, as well as ABC transporters for branched-chained amino acids and
253 dipeptides and polypeptides (**Table S4**). “*Ca. D. occultata*” also contained a large number of
254 environmental sensing genes: 52 bacterial hemoglobins with PAS-PAC sensors, 8 TonB-

255 dependent receptors, and 8 NO responsive regulators (Dnr: Crp/fr family; **Table S4**). Uniquely
256 in “*Ca. D. occultata*”, PAC-PAS sensors flanked accessory genes *nosFLY* on the *c-nosZ* operon
257 (**Fig. S7**). Comparison of these flanking PAC-PAS sensors in “*Ca. D. occultata*” with O₂-binding
258 sensors revealed that an arginine ~20 aa upstream from the conserved histidine as the distal
259 pocket ligand for O₂-binding is not present in either sensor (**Fig. S11**), suggesting that the sensor
260 may bind a different ligand, possibly NO, consistent with the placement of these genes next to
261 *cNosZ* (Shimizu et al., 2015).

262 **Nutrient storage.** Active synthesis of storage polymers suggested that “*Ca. D. occultata*”
263 was experiencing electron acceptor starvation at the time of harvesting, consistent with Mn(III)
264 depletion in the bottles (Liu et al., 2015; Guanghuan et al., 2018). Polyphosphate-related
265 proteins, including phosphate transporters, polyphosphate kinase, polyphosphatase, and poly-3-
266 hydroxybutyrate synthesis machinery were detected in the proteome (**Table S4**). Polyphosphate-
267 accumulating organisms store polyphosphates with energy generated from organic carbon
268 oxidation during aerobic respiration or denitrification. These stored compounds are later
269 hydrolyzed when respiratory electron acceptors for ATP production are limiting. Cyanophycin
270 was actively synthesized for nitrogen storage.

271 **Geobacter.** “*Ca. Geobacter occultata*” expressed proteins in the TCA cycle at moderate
272 abundance. “*Ca. G. occultata*” contained 17 multiheme c-type cytochromes, none of which were
273 detected in the proteome. The lack of expression of electron transport and metal-reducing
274 pathways makes it unlikely that “*Ca. G. occultata*” was solely responsible for Mn(III) reduction
275 observed in the incubations. A periplasmic group I Ni-Fe hydrogenase (RNC64340; 91% identity
276 to a protein (RLB64899) from *Geobacter* MAG from terrestrial hot spring sediment) and a type
277 IV pilin (RNC67631; 10% aromatics, 87% identity to *Geobacter pickeringii* (Holmes et al.,

278 2016)) were significantly more expressed in the presence of CH₄ than N₂ in the “*Ca. G.*
279 *occultata*” proteome ($p < 0.05$; **Table 1**). It is possible that “*Ca. G. occultata*” transferred
280 electrons to “*Ca. D. occultata*” via e-pilins (e.g. direct interspecies electron transfer),
281 contributing to the higher rates of Mn(III) reduction in the presence of CH₄ vs. N₂. The possible
282 involvement of *Geobacter* e-pilins in Mn(III) reduction remains an open question, due to the lack
283 of studies examining the possibility of Mn(III) reduction in *Deltaproteobacteria*.

284 **Conclusions.** To our knowledge, this study provides the first evidence for biological
285 reduction of soluble Mn(III) by a bacterium outside of the *Gammaproteobacteria* class. The
286 dominant bacterium in Mn(III)-reducing enrichment cultures was “*Ca. D. occultata*”, a member
287 of the *Rhodocyclales* order of *Betaproteobacteria*. “*Ca. D. occultata*” expressed decahemes
288 similar to the Mto pathway, and *occ* genes, including a novel extracellular undecaheme (OccP),
289 which are predicted to encode a new respiratory electron transport pathway. The novel *occ*
290 operon was found to be widespread in *Betaproteobacteria* from the deep subsurface, where metal
291 cycling can fuel microbial metabolism. We also found highly expressed peptides from various
292 central metabolic cycles and organic substrate utilization pathways, suggesting that “*Ca. D.*
293 *occultata*” may have been using multiple pathways simultaneously for energy generation and
294 carbon assimilation during Mn(III) reduction.

295 Puzzles remain about whether “*Ca. D. occultata*” can transform two potent greenhouse
296 gases: methane and nitrous oxide. Although “*Ca. D. occultata*” was enriched with CH₄ as the
297 sole electron donor and cultures reduced Mn(III) more rapidly in the presence of CH₄, no CH₄
298 oxidation activity was measured in Mn(III)-reducing cultures, and proteomic data suggested that
299 “*Ca. D. occultata*” was growing mixotrophically rather than assimilating CH₄. Further, although
300 we did not add oxidized nitrogen compounds to our media, and Mn(III) did not chemically

301 oxidize NH_4^+ under our culture conditions, type II nitrous oxide reductase (cNosZ) was one of
302 the most abundant proteins expressed in Mn(III)-reducing cultures. The role of cNosZ and other
303 denitrification enzymes in “*Ca. D. occultata*” metabolism, and their possible connection to
304 Mn(III) reduction, remain to be investigated.

305

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314

315 **Competing Interests:** The authors declare no competing interests.

316 **Table 1. Expression levels for select “*Ca. Dechloromonas occultata*” and “*Ca. Geobacter occultata*” proteins**
317 **in the presence of CH₄ and N₂.** Gray boxes indicate membrane proteins¹. Bold proteins indicate proteins that were
318 significantly more expressed with CH₄ than N₂ (CH₄/N₂>1; p<0.05). P values indicate significance of abundance
319 difference between CH₄ and N₂ treatments.

¹SP: signal peptide (Y:present/N:absent); TMH: numbers of transmembrane helices; CxxCH: number of heme-binding motifs; P-sort: predicted cellular location based on Psortb v.3.0. MCP: methyl-accepting chemotaxis protein; PPIase: Peptidyl-proline isomerase; P: periplasm, C: cytoplasm; OM: outer membrane; IM: inner membrane, E: extracellular; U: unknown. MtoX and MtoY were predicted to be an inner membrane cytochrome-b protein and a methyl-accepting chemotaxis protein, respectively. Membrane proteins may be underrepresented by mass spectrometry-based metaproteomic analyses, which inherently favor soluble over insoluble membrane-bound or hydrophobic proteins.

Enzyme Complex/Category	Function	Protein Sequence Predictions				Normalized Peptide Abundance							
		Proteins	NCBI ID	Motifs			By Treatment			Differential Peptide			
				SP	TMH	CxxCH	P-sort	CH ₂	SD	N ₂	SD	Avg	SD
Ca. Dechloromonas occulta													
Mtc-1	Outer membrane porin-cytochrome c electron conduit	MtoX-1 (cyt-b)	RIX49676	N	5	0	IM						
		MtoY-1 (MCP)	RIX49677	N	2	1	IM	2.7	0.5	3.6	0.2	0.8	0.2
		MtoB-1 (porin)	RIX49678	Y	0	0	OM	10	2	15	2	0.6	0.1
		MtoA-1	RIX49874	Y	1	10	P	5	1	2.5	0.1	1.9	0.4
Mtc-2	Outer membrane porin-cytochrome c electron conduit	MtoX-2 (cyt-b)	RIX48942	N	4	0	IM						
		MtoB-2 (porin)	RIX48943	Y	0	0	OM	8	1	16	0.2	0.5	0.1
		MtoA-2	RIX48944	Y	1	10	P	7.3	0.8	4	2	2.1	1.3
		MtoD-2	RIX48945	Y	1	1	U	2.6	0.3	0.7	0.3	4.0	1.4
Occ	Membrane-spanning electron transport cytochromes	OccA	RIX49688	Y	1	3	P	4	0.5	0.7	0.6	7.8	5.7
		OccB	RIX49689	Y	0	3	U	41	4	19	2	2.2	0.0
		OccC	RIX49877	N	0	1	U						
		OccD	RIX49878	N	0	3	U						
		OccE (6-NHL)	RIX49690	N	1	0	U	22	2.1	20.5	0.2	1.1	0.1
		OccF	RIX49691	Y	2	4	E	13	0.7	10.1	0.1	1.3	0.1
		OccG (P/Phase)	RIX49692	N	0	0	U	14	1	3.3	0.5	4.2	0.3
		OccH	RIX49693	N	0	0	OM/E	6.0	0.2	7.7	0.6	0.8	0.1
		OccI	RIX49694	N	1	3	U	7	2.5	2.3	0.0	2.9	1.1
		OccJ	RIX49879	Y	0	4	U	44	0.2	19	3	2.4	0.4
		OccK	RIX49880	N	0	0	C	39	6	13	1	3.0	0.2
		OccL	RIX49695	N	1	3	U						
		OccM	RIX49881	N	0	3	U						
		OccN (6 NHL)	RIX49696	N	2	0	U	5.7	0.3	6	1	0.9	0.1
		OccO (6 NHL)	RIX49882	N	0	0	U	12	0.8	4.2	0.4	0.3	0.2
		OccP	RIX49697	N	0	11	E	34	2	12	3	1.2	0.5
		OccQ	RIX49698	Y	4	0	IM						
OccR	RIX49883	N	8	0	IM								
OccS	RIX49699	N	12	0	IM								
Cyt c	Mono- and dihemetic cytochromes involved in electron transfer	Cyt c5	RIX47670	N	1	1	U	27	2	9	3	3.2	0.8
		Cyt c5	RIX49884	Y	1	2	P	19	2	6	1	3.3	1.0
		Cyt c'/C ₂	RIX44710	Y	1	1	P	17	5	3.6	0.8	4.8	2.3
		Cyt c'/C ₂	RIX49630	Y	1	1	P	7	1	1.2	0.9	8.2	6.6
		Cyt c551/c552	RIX49807	Y	0	1	P	13	3	2.8	0.0	4.8	1.1
		Cyt c4	RIX48804	Y	0	2	P	16	0.8	9.8	0.8	1.6	0.2
		Cyt c4	RIX44782	Y	0	2	P	4	2	1.7	0.7	2.6	0.1
		Cyt c4	RIX45018	Y	0	2	P	7	0.6	2.2	0.2	3.0	0.0
Nap	Periplasmic nitrate reductase	NapA	RIX41011	Y	0	0	P	76	2	67	3	1.1	0.1
		NapB	RIX41010	Y	1	2	P	15	1	5	2	3.2	0.9
		NapC	RIX41009	N	1	4	IM	12	3	13	1	1.0	0.2
Nir	Nitrite reductase	NirS	RIX44719	Y	0	1	P	58	2	44	4	1.3	0.2
		NirB	RIX44720	Y	1	2	P	14	3	10	2	1.5	0.6
		NirC	RIX44788	N	0	1	P						
		NirF	RIX44721	Y	1	0	Per C	2	1	7	1	0.3	0.1
Nor	Nitric oxide reductase	NorC	RIX45182	N	1	1	IM	3.5	0.7	3.2	0.7	1.1	0.0
		NorB	RIX45183	N	12	1	IM						
cNos	Type II nitrous oxide reductase	cNosZ	RIX42539	Y	0	0	P	77	17	66	8	1.2	0.3
		cNosC1	RIX42538	Y	1	1	P	16	2	4	2	4.9	3.3
		cNosC2	RIX42537	Y	1	2	P	10	0.1	3.9	0.3	2.6	0.1
		cNosB	RIX42536	N	6	0	IM						
		cNosD	RIX42535	N	0	0	P						
		cNosG	RIX42534	N	1	0	C						
Qcr	Menquinone-cytochrome c reductase complex	QcrA	RIX41976	N	9	0	CM						
		QcrB	RIX41977	N	9	0	CM						
		QcrC	RIX41978	N	1	0	CM						
Proteases	Serine protease		RIX49468	N	0	0	P	27	2	1.0	0.3	29.0	9.9
			RIX48818	N	1	0	CM	18.5	0.8	8.0	0.9	2.3	0.1
Membrane/Extracellular	DUF4214 protein 5-layer protein PEP-CTERM sorting protein Tol-Pal system protein Peptidoglycan-associated lipoprotein Tol-Pal system protein Pilus assembly protein		RIX44180	N	0	0	OM/E	146	25	43	6	3.4	0.5
			RIX44181	N	0	0	U	8	0.5	10	0.6	0.8	0.1
			RIX45463	Y	1	0	E	68	6	33	10	2.1	0.5
		TolB	RIX44015	Y	0	0	P	20	2	12	1	1.7	0.0
		Pal	RIX44016	N	0	0	OM	27.3	0.2	10	3	2.7	0.7
		YbgF	RIX44017	Y	0	0	U	10.8	0.4	4	2	3.7	2.2
			RIX46961	N	0	0	U	54	5	30	5	1.8	0.1
Other	Ethanol/methanol dehydrogenase Alcohol dehydrogenase Aldehyde dehydrogenase Phasin family granule-associated protein Phasin family granule-associated protein High potential iron-sulfur protein Electron transfer flavoprotein NAD-reducing hydroge nase		RIX45050	Y	0	0	P	37	4	17	1	2.2	0.1
			RIX45053	Y	0	0	P	12.4	1.4	14.2	1.7	0.9	0.0
			RIX45061	Y	0	0	P	125	31	221	75	0.6	0.1
			RIX40682	N	0	0	U	49	2	22	1	2.2	0.2
			RIX40683	Y	0	0	U	34	4	16	1	2.1	0.0
			RIX49681	Y	0	0	U	10.79	0.01	6.5	0.4	1.7	0.1
		FixA	RIX43544	N	0	0	C	16	3	10	2	1.7	0.0
HoxH	RIX46736	N	0	0	C	22.9	0.7	37	3	0.6	0.1		
Ca. Geobacter occulta													
Hydrogenase	[Ni/Fe] hydrogenase, group 1, small subunit [Ni/Fe] hydrogenase, group 1, large subunit	HyaA	RNC64339	Y	0	0	P	11.1	0.4	3	1	5	2
		HyaB	RNC64340	N	0	0	P	32	0	11	5	3	1
E-pilus	Type IV pilin	PilA	RNC67631	N	1	0	E	93	3	18	3	5.6	0.6

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