

1 **Geochemical and metagenomic characterization of Jinata Onsen, a Proterozoic-analog hot spring,**  
2 **reveals novel microbial diversity including iron-tolerant phototrophs and thermophilic lithotrophs**

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18

19 **Abstract**

20 Hydrothermal systems, including terrestrial hot springs, contain diverse geochemical  
21 conditions that vary over short spatial scales due to progressive interaction between the reducing  
22 hydrothermal fluids, the oxygenated atmosphere, and in some cases seawater. At Jinata Onsen,  
23 on Shikinejima Island, Japan, an intertidal, anoxic, iron-rich hot spring mixes with the  
24 oxygenated atmosphere and seawater over short spatial scales, creating a diversity of chemical  
25 potentials and redox pairs over a distance ~10 m. We characterized the geochemical conditions  
26 along the outflow of Jinata Onsen as well as the microbial communities present in biofilms,  
27 mats, and mineral crusts along its traverse via 16S rDNA amplicon and genome-resolved  
28 shotgun metagenomic sequencing. The microbial community changed significantly downstream  
29 as temperatures and dissolved iron concentrations decreased and dissolved oxygen increased.  
30 Near the spring source, biomass is limited relative to downstream, and primary productivity may  
31 be fueled by oxidation of ferrous iron and molecular hydrogen by members of the  
32 Zetaproteobacteria and Aquificae. Downstream, the microbial community is dominated by  
33 oxygenic Cyanobacteria. Cyanobacteria are abundant and active even at ferrous iron  
34 concentrations of ~150  $\mu\text{M}$ , which challenges the idea that iron toxicity limited cyanobacterial  
35 expansion in Precambrian oceans. Several novel lineages of Bacteria are also present at Jinata  
36 Onsen, including previously uncharacterized members of the Chloroflexi and Caldithrichaeota  
37 phyla, positioning Jinata Onsen as a valuable site for future characterization of these clades.

38 **Importance**

39 High temperatures and reducing conditions allow hot springs to support microbial  
40 communities that are very different from those found elsewhere on the surface of the Earth  
41 today; in some ways, these environments and the communities they support can be similar to  
42 environments that existed on the early Earth and that may exist on other planets. Here, we  
43 describe a novel hot spring system where hot, iron-rich but oxygen-poor water flows into the  
44 ocean, supporting a range of unique microbial communities. Metagenomic sequencing recovered  
45 many novel microbial lineages, including deep-branching and uniquely thermotolerant members

46 of known groups. Comparison of the biological communities in the upstream part of the hot  
47 spring, potentially supported by biological iron and hydrogen oxidizing metabolisms, to  
48 downstream microbial mats, supported by oxygenic photosynthesis, provides insight into the  
49 potential productivity of life during Proterozoic time and on other planets where oxygenic  
50 photosynthesis is not possible.

51

## 52 **Introduction**

53 A major theme of environmental microbiology has been the enumeration of microbial  
54 groups that are capable of exploiting diverse chemical potentials (i.e. chemical disequilibria) that  
55 occur in nature (e.g. 1-3). Hot springs, with their varied chemical compositions, provide  
56 reservoirs of novel microbial diversity, where environmental and geochemical conditions select  
57 for lineages and metabolisms distinct from other Earth-surface environments (e.g. 4-7). In  
58 addition to their value as sources of microbial diversity, hot springs also provide valuable test  
59 beds for understanding microbial community processes driven by different suites of metabolisms  
60 (e.g. 8)—this in turn allows these systems to serve as process analogs and to provide a window  
61 into biosphere function during early times in Earth history, for example when the O<sub>2</sub> content of  
62 surface waters was low or non-existent. In contrast to most surface ecosystems which are fueled  
63 almost entirely by oxygenic photosynthesis by plants, algae, and Cyanobacteria, hot spring  
64 microbial communities are commonly supported by lithotrophic or anoxygenic phototrophic  
65 organisms that derive energy and electrons for carbon fixation by oxidizing geologically sourced  
66 electron donors such as Fe<sup>2+</sup>, sulfide, and molecular hydrogen (e.g. 9-11). These communities  
67 may therefore provide insight into the function of microbial communities on the early Earth or  
68 other planets, in which oxygenic photosynthesis may be absent or less significant and  
69 anoxygenic photosynthetic or lithotrophic metabolisms may play a larger role, resulting in  
70 overall lower rates of primary productivity (e.g. 12-15).

71 Here, we present a geomicrobiological characterization of a novel Precambrian Earth  
72 process analog site: Jinata Onsen, on Shikinejima Island, Tokyo Prefecture, Japan. At Jinata hot  
73 spring, anoxic, iron-rich hydrothermal fluids feed a subaerial spring that flows into a small bay,  
74 and mixes with seawater over the course of a few meters. Over its course the waters transition  
75 from low-oxygen, iron-rich conditions analogous to some aspects of the early Proterozoic  
76 oceans, toward iron-poor and oxygen-rich conditions typical of modern coastal oceans. In  
77 upstream regions of the stream where oxygenic Cyanobacteria are absent, biomass is visibly  
78 sparse; however, downstream, biomass accumulates in the form of thick microbial mats  
79 containing abundant Cyanobacteria. Visible differences in accumulation and appearance of  
80 biomass across the temperature and redox gradient establish the hypothesis that microbial  
81 community composition, as well as magnitude and metabolic drivers of primary productivity,  
82 varies along the spring flow. To begin testing this hypothesis and to provide a baseline  
83 description of the geochemistry and microbiology of this site in support of future investigation,  
84 we performed geochemical measurements, 16S rDNA amplicon sequencing, and genome-  
85 resolved metagenomic sequencing to recover draft genomes of diverse novel microbial lineages  
86 that inhabit Jinata Onsen.

87

## 88 **Results**

### 89 **Site description**

90 The source water of Jinata Onsen emerges with dissolved oxygen concentrations below  
91 our limit of detection, is iron-rich, and gently bubbles gas from the spring source (Table 1,

92 Figure 1, Figure 2). Temperatures at the source are  $\sim 63^{\circ}\text{C}$ . Water emerges into the Source Pool,  
93 which has no visible microbial mats or biofilms (Figure 2D). Surfaces are instead coated with a  
94 fluffy red precipitate, likely a poorly ordered or short range-ordered ferric iron oxide phase such  
95 as ferrihydrite. Flow from the source is—at least in part—tidally charged, with the highest water  
96 levels and flow rates occurring at high tide. At low tide, flow rates drop and the water level of  
97 the Source Pool can drop by decimeters and portions of the Source Pool can drain during spring  
98 low tides. Downstream, the spring water collects into a series of pools (Pool 1-3) (Figure 2C,E-  
99 F), which cool sequentially (Figure 3, Supplemental Table 1). Pool 1 contains iron oxides like  
100 the Source Pool, but also develops macroscopic microbial streamers that are coated in iron  
101 oxides and thin veil-like layers of microorganisms overlaying iron oxide sediments—structures  
102 similar to those typically made by marine iron-oxidizing Zetaproteobacteria (e.g. 16). Streamers  
103 are very fine (mm-scale) and delicate (break apart on contact with forceps) but can reach several  
104 centimeters in length. Cyanobacteria in Pool 2 and Pool 3 display high levels of photosynthetic  
105 activity as revealed by high dissolved oxygen concentration ( $\sim 234\ \mu\text{M}$ ), low dissolved inorganic  
106 carbon concentrations, and the accumulation of visible  $\text{O}_2$  bubbles on the surface and within the  
107 fabric of the mat. Downstream pools (Pools 2 and 3) mix with seawater during high tide due to  
108 wave action, but this seawater influence does not appear to influence the Source Pool or Pool 1.  
109 Samples were collected and temperatures were measured at high tide, reflecting the lowest  
110 temperatures experienced by microbes in the pools—at low tide, hot spring input is dominant  
111 and temperatures rise (observed range at each site in Supplemental Table 1). Subaqueous  
112 surfaces in Pools 2 and 3 are covered in thick microbial mats. In Pool 2, the mat is coated in a  
113 layer of fluffy iron oxide similar to that in the Source Pool, with dense microbial mat below  
114 (Figure 2E). Pool 3 contains only patchy iron oxides, with mostly exposed microbial mats  
115 displaying a finger-like morphology. These “fingers” were 0.5-1 cm in diameter and up to  $\sim 5$  cm  
116 long and were closely packed and carpeting surfaces of Pool 3 below the high tide line,  
117 potentially related to turbulent mixing from wave action during high tide (Figure 2F). The  
118 Outflow is the outlet of a channel connecting Pool 2 to the bay. Its hydrology is dominantly  
119 marine with small admixtures of inflowing spring water (Figure 2G).

### 120 **Geochemistry**

121 Geochemical measurements along the flow path of Jinata Onsen revealed a significant  
122 shift from hot, low-oxygen, high-iron source water to cooler, more oxygen-rich water with less  
123 dissolved iron downstream. Geochemistry measurements of Jinata source water are summarized  
124 in Table 1 and Supplemental Table 1, while geochemical gradients along the stream outflow are  
125 summarized in Figure 3 and Supplemental Table 2. Source waters were slightly enriched in  
126 chloride relative to seawater ( $\sim 23.2$  g/L in Jinata source water versus  $\sim 19.4$  g/L in typical  
127 seawater), depleted in sulfate ( $\sim 1.6$  g/L in Jinata versus  $\sim 2.7$  g/L in seawater) but approached  
128 seawater concentrations downstream as mixing increased. Water emerging from the source was  
129  $63^{\circ}\text{C}$ , very low in dissolved oxygen ( $\sim 4.7\ \mu\text{M}$ ), at pH 5.4, and contained substantial  
130 concentrations of dissolved iron ( $\sim 250\ \mu\text{M Fe}^{2+}$ ). Dissolved organic carbon (DOC) in the source  
131 water was high ( $\sim 1.31$  mM). It is unknown whether this is produced *in situ* or whether the source  
132 water emerges with high DOC. Both DOC and DIC decrease along the outflow of the spring  
133 (Supplemental Table 1). After emerging from the source, the spring water exchanges gases with  
134 the air due to mixing associated with water flow and gas ebullition, and DO rose to  $39\ \mu\text{M}$  at the  
135 surface of the Source Pool. As water flows downstream from the Source Pool, it cools slightly,  
136 exchanges gases with the atmosphere, and intermittently mixes with seawater below Pool 1.

### 137 **16S rDNA and genome-resolved metagenomic sequencing**

138 16S rDNA and metagenomic sequencing of microbial communities at Jinata Onsen  
139 revealed a highly diverse community. In total, 16S rDNA amplicon sequencing recovered  
140 456,737 sequences from the 10 samples at Jinata (Supplemental Tables 3-5). Reads per sample  
141 ranged from 26,057 Source Pool Sample A to 97,445 for Pool 1 Sample A (median 43,331, mean  
142 45,673, and standard deviation 19,568). On average 74% of the microbial community was  
143 recovered from Jinata samples at the 99% OTU level based on the Good's Coverage statistic of  
144 the 16S rRNA gene (ranging from 54% coverage in the Outflow Sample A to 85% in the Pool 1  
145 Sample A) and 87% at the 97% OTU level (74% for the Outflow Sample A to 94.5% for the  
146 Pool 1 Sample B). The incomplete sampling—despite sequencing to relatively high depth  
147 (>26000 reads per sample)—probably reflects uneven diversity. Greater than 50% of the reads  
148 observed at most sites map to the 10 most abundant taxa (Supplemental Table 4). MDS analysis  
149 (Supplemental Figure 1) demonstrates that samples from the same site are highly similar, and  
150 adjacent sites (e.g. Source Pool and Pool 1, Outflow and Pool 3) show significant similarity.  
151 However, there is a significant transition in microbial community diversity between the most  
152 distant samples (e.g. Source Pool and Outflow).

153 Shotgun metagenomic sequencing of four samples from Jinata Onsen recovered 121 GB  
154 of data, forming a 1.48 Gb coassembly consisting of 1531443 contigs with an N50 of 1494 bp.  
155 Nucleotide composition and differential coverage-based binning of the coassembly via multiple  
156 methods followed by dereplication and refinement resulted in a final set of 161 medium- or high-  
157 quality metagenome-assembled genomes (MAGs) following current standards (i.e. completeness  
158 >50% and contamination <10%) (17). These MAGs are from diverse phyla of Bacteria and  
159 Archaea (Figure 4); metagenome and MAG statistics with tentative taxonomic assignments for  
160 recovered MAGs are available in Supplementary Table 5, while MAGs of particular interest are  
161 discussed in depth below and in the SI and shown in phylogenetic trees alongside reference  
162 strains in Figures 5-7.

163

## 164 Discussion

165 As Jinata spring water flows from source to ocean, it transitions from hot, low-oxygen,  
166 high-iron water to cooler, iron-depleted, oxygen-rich water in downstream regions (Figure 3).  
167 Following this geochemical transition is a major shift in the composition of the microbial  
168 community, from a high-temperature, putatively lithotrophic community which produces little  
169 visible biomass upstream, to a lower temperature, community with well-developed, thick  
170 microbial mats downstream. This shift in community composition is summarized in Figure 3,  
171 with complete diversity data in the Supplemental Information (including OTU counts per  
172 samples in Supplemental Table 4 and relative abundance binned at the class level in  
173 Supplemental Table 5). Below, we discuss the overall physiological and taxonomic trends across  
174 the spring sites as inferred from diversity and genomic analysis.

### 175 Potential for iron and hydrogen oxidation

176 The hot spring water emerging at the Source Pool at Jinata contains abundant electron  
177 donors including dissolved  $\text{Fe}^{2+}$  and likely  $\text{H}_2$  (though measurements of gas content varied, as  
178 discussed in the SI) (Table 1). Although rates of carbon fixation were not measured, the  
179 appearance of zetaproteobacterial veils and streamers and molecular evidence for  
180 lithoautotrophic microbes suggests that these electron donors may fuel productivity and  
181 determine the microbial community upstream at the Source Pool and Pool 1, where microbial  
182 mats are not well developed. The low accumulation of biomass in upstream regions of Jinata are  
183 similar to other microbial ecosystems fueled by iron oxidation (e.g. Oku-Okuhachikurou Onsen,

184 11, Fuschna Spring, 18, and Jackson Creek, 19), in which lithotrophic communities appear  
185 capable of accumulating less organic carbon than communities fueled by oxygenic  
186 photosynthesis (including those in downstream regions at Jinata).

187 Results of 16S rDNA sequencing indicate that the most abundant organisms in the Source  
188 Pool are members of the Aquificae family Hydrogenothermaceae (33% of reads in the Source  
189 Pool and 11.6% of reads in Pool 1). Members of this family of marine thermophilic lithotrophs  
190 are capable of iron and hydrogen oxidation, as well as heterotrophy (20) and may be utilizing  
191 either  $\text{Fe}^{2+}$  or  $\text{H}_2$  at Jinata. The seventh most abundant OTU in the Source Pool samples is a  
192 novel sequence 89% similar to a strain of *Persephonella* found in an alkaline hot spring in Papua  
193 New Guinea. *Persephonella* is a genus of thermophilic, microaerophilic hydrogen oxidizing  
194 bacteria within the Hydrogenothermaceae (21). Despite their abundance as assessed by 16S  
195 rDNA sequencing (Figure 3), only four partial Aquificae MAGs were recovered from Jinata of  
196 which only one (J026) was reasonably complete (~94%). Two Aquificae MAGs recovered  
197 Group 1 NiFe hydrogenase genes, which may be used in hydrogenotrophy; the absence of  
198 hydrogenases from the other MAGs may be related to their low completeness, or could reflect a  
199 utilization of iron or other electron donors and not  $\text{H}_2$  in these organisms.

200 The other most abundant organisms near the source are members of the  
201 Zetaproteobacteria—a group typified by the neutrophilic, aerobic iron-oxidizing genus  
202 *Mariprofundus* common in marine systems (22). Zetaproteobacteria accounted for 24.5% of 16S  
203 rDNA sequences in the Source Pool and 26.7% in Pool 1. All Zetaproteobacteria characterized to  
204 date are obligate iron- and/or hydrogen-oxidizing lithoautotrophs (23), suggesting that these  
205 organisms may play a substantial role in driving carbon fixation in the Source Pool and Pool 1.

206 Members of the Mariprofundaceae have been observed to have an upper temperature  
207 limit for growth of 30 °C (24), while Zetaproteobacteria are found at Jinata at temperatures up to  
208 63 °C. This currently represents a unique high-temperature environment for these organisms. In  
209 particular, the third most abundant OTU in the Source Pool and Pool 1 sample A is an unknown  
210 sequence that is 92% identical to a sequence from an uncultured zetaproteobacterium from a  
211 shallow hydrothermal vent in Papua New Guinea (25). This sequence likely marks a novel  
212 lineage of high-temperature iron-oxidizing Zetaproteobacteria. Four MAGs affiliated with the  
213 Zetaproteobacteria were recovered from Jinata with completeness estimates by CheckM ranging  
214 from ~80 to ~97% (J005, J009, J030, and J098). While these MAGs did not recover 16S rRNA  
215 genes, RpoB- and concatenated ribosomal protein-based phylogenies illustrated that members of  
216 this group at Jinata Onsen do not belong to the characterized genera *Mariprofundus* or *Ghiorsea*,  
217 but instead form separate basal lineages within the Zetaproteobacteria (Figure 5). Despite their  
218 phylogenetic distinctness, these MAGs largely recovered genes associated with aerobic iron  
219 oxidation as expected given the physiology of other Zetaproteobacteria, including *Cyc2*  
220 cytochrome and C-family heme copper oxidoreductase genes (SI). Notable exceptions are J005  
221 and J030 which did not recover genes for carbon fixation via the Calvin cycle (such as the large  
222 and small subunits of rubisco, phosphoribulose kinase, or carboxysome proteins); the high  
223 completeness of these MAGs (~94-97%) makes it unlikely that these genes would all fail to be  
224 recovered (MetaPOAP False Negative estimates  $10^{-5}$ - $10^{-7}$ ); the absence of carbon fixation  
225 pathways from these genomes together with the availability of abundant dissolved organic  
226 carbon in Pool 1 (~1.3 mM) suggest that these organisms may be heterotrophic, a lifestyle not  
227 previously observed for members of the Zetaproteobacteria.

228 Seven MAGs were recovered from the enigmatic bacterial phylum Calditrichaeota (J004,  
229 J008, J042, and J075) (Figure 6). While few members of Calditrichaeota have been isolated or  
230 sequenced, the best known of these is *Caldithrix abyssi* (26); this taxon was characterized as an  
231 anaerobic thermophile capable of lithoheterotrophic H<sub>2</sub> oxidation coupled to denitrification and  
232 organoheterotrophic fermentation (27, 28). The Calditrichaeota MAGs reported here are up to  
233 97% complete (J004) and contain members with variable putative metabolic capabilities,  
234 potentially including aerobic hydrogen- or iron-oxidizing lithoautotrophy (SI). Unlike previously  
235 described Calditrichaeota which are all heterotrophic (28), most of the Calditrichaeota MAGs  
236 reported here possess a putative capacity for carbon fixation via the Calvin cycle. J004 is closely  
237 related to *Caldithrix abyssi*, while the other MAGs form two distinct but related clades (Figure  
238 6).

### 239 **Oxygenic photosynthesis**

240 Cyanobacteria are nearly absent from near the Source Pool, but are observed in low  
241 numbers in Pool 1 and become dominant starting in Pool 2. The most abundant Cyanobacteria  
242 present are predominantly members of Subsection III, Family I. This group includes  
243 *Leptolyngbya*, a genus of filamentous non-heterocystous Cyanobacteria that is present in other  
244 hot springs of similar temperatures (e.g. 11, 29, 30). Diverse cyanobacterial MAGs were  
245 recovered, including members of the orders Pleurocapsales (J083), Chroococcales (J003 and  
246 J149), and Oscillatoriales (J007, J055, and J069). In the Outflow samples, chloroplast sequences  
247 are abundant, most closely related to the diatom *Melosira*.

248 Cyanobacteria are sometimes underrepresented in iTag datasets as a result of poor DNA  
249 yield or amplification biases (e.g. 31, 32), but the low abundance of Cyanobacteria near the  
250 Source Pool was confirmed by fluorescent microscopy, in which cells displaying cyanobacterial  
251 autofluorescence were observed abundantly in samples from the downstream samples but not in  
252 the Source Pool (Supplemental Figure 2). Thick microbial mats first appear in Pool 2 when  
253 Cyanobacteria become abundant, suggesting that oxygenic photosynthesis fuels more net carbon  
254 fixation than lithotrophy in these environments. Previously, it has been suggested that high  
255 ferrous iron concentrations are toxic to Cyanobacteria, and that this would have greatly reduced  
256 their productivity under ferruginous ocean conditions such as those that may have persisted  
257 through much of the Archean era (33). The abundant Cyanobacteria observed to be active at  
258 Jinata under high iron concentrations suggest that Cyanobacteria can adapt to ferruginous  
259 conditions, and therefore iron toxicity might not inhibit Cyanobacteria over geological  
260 timescales. Indeed, the soluble iron concentrations observed at Jinata are higher (150-250  $\mu\text{M}$ )  
261 than predicted for the Archean oceans (<120  $\mu\text{M}$ , 34) or observed at other iron-rich hot springs  
262 (~100-200  $\mu\text{M}$ , 11, 35), making Jinata an excellent test case for determining the ability of  
263 Cyanobacteria to adapt to high iron concentrations. Culture-based physiological experiments  
264 may be useful to determine whether Jinata Cyanobacteria utilize similar strategies to other iron-  
265 tolerant strains (e.g. by those in Chocolate Pots Hot Spring, 35, or the ferric iron-tolerant  
266 *Leptolyngbya*-relative *Marsacia ferruginosa*, 36) or whether Jinata strains possess unique  
267 adaptations that allow them to grow at higher iron concentrations than known for other  
268 environmental Cyanobacteria strains. This will in turn provide insight into whether iron tolerance  
269 is due to evolutionarily conserved strategies or whether this is a trait that has evolved  
270 convergently multiple times.

### 271 **Diverse novel Chloroflexi from Jinata Onsen**

272 In addition to the primary phototrophic and lithotrophic carbon fixers at Jinata, 16S  
273 rDNA and metagenomic data sets revealed diverse novel lineages within the Chloroflexi phylum.

274 A total of 23 Chloroflexi MAGs were recovered, introducing substantial genetic and metabolic  
275 diversity that expands our understanding of this group. While the best known members of this  
276 phylum are Type 2 Reaction Center-containing lineages such as *Chloroflexus* and *Roseiflexus*  
277 within the class Chloroflexia (e.g. 37), phototrophy is not a synapomorphy of the Chloroflexi  
278 phylum or even the Chloroflexia class (e.g. 38) and most of the diversity of the phylum belongs  
279 to several other classes made up primarily of nonphototrophic lineages (7). The bulk of  
280 Chloroflexi diversity recovered from Jinata belongs to “subphylum I”, a broad group of  
281 predominantly nonphototrophic lineages that was originally described based on the classes  
282 Anaerolineae and Caldilineae (39), but also encompasses the related classes Ardenticatenia,  
283 Thermoflexia, and *Candidatus* Thermofonsia (7, 40, 41).

284 16S rDNA analysis indicates that members of the Chloroflexi class Anaerolineae are  
285 common throughout Jinata with the exception of the Outflow (average 3.5% relative abundance).  
286 However, it is likely that a large fraction of 16S rDNA sequences annotated as Anaerolineae at  
287 Jinata Onsen belong to the sister class *Candidatus* Thermofonsia (7). Three MAGs recovered  
288 from Jinata (J082, J097, and J130) are associated with the Anaerolineae class as determined by  
289 RpoB and concatenated ribosomal protein phylogenies, as compared to seven associated with  
290 *Ca.* Thermofonsia (J027, J033, J036, J038, J039, J064, and J076). Particularly notable among  
291 these MAGs is J036, a close relative of the phototrophic *Ca.* Roseilinea gracile (42-44). J036  
292 contains a 16S rRNA gene that is 96% similar to that of *Ca.* Roseilinea gracile, and two-way  
293 AAI estimates (Rodriguez and Konstantinidis 2014) showed 73.6% similarity between the two  
294 strains, indicating these strains are probably best classified as distinct species within the same  
295 genus. Unlike other phototrophs in the Chloroflexi phylum that are capable of photoautotrophy  
296 via the 3-hydroxypropionate bicycle or the Calvin Cycle (45, 46), J036 and *Ca.* Roseilinea  
297 gracile do not encode carbon fixation and are likely photoheterotrophic.

298 Members of the Chloroflexi class Caldilineae were present at up to ~1% abundance at  
299 Jinata in the 16S rDNA dataset. Three MAGs were recovered that form a deeply branching  
300 lineage with the Caldilineae class (J095, J111, and J123), sister to the previously characterized  
301 genera *Caldilinea* and *Litorilinea*. Like other members of the Caldilineae, these strains encode  
302 aerobic respiration via A-family heme copper oxidoreductases and both a *bc* complex III and an  
303 alternative complex III, and are therefore likely at least facultatively aerobic. J095 also encodes  
304 carbon fixation via the Calvin cycle as well as a Group 1f NiFe hydrogenase, suggesting a  
305 potential capability for lithoautotrophy by hydrogen oxidation, expanding the known metabolic  
306 diversity of this class and the Chloroflexi phylum as a whole. MAG J114 branches at the base of  
307 subphylum I of the Chloroflexi, potentially the first member of a novel class-level lineage. The  
308 divergence between Anaerolineae and Caldilineae has been estimated to have occurred on the  
309 order of 1.7 billion years ago (46). The phylogenetic placement of J114 suggests that it diverged  
310 from other members of subphylum I even earlier, and it may be a good target for future  
311 investigation to assess aspects of the early evolution of the Chloroflexi phylum. J114 encodes  
312 genes that suggest the capacity for aerobic hydrogen-oxidizing autotrophy (including those  
313 encoding an A-family heme copper oxidoreductase, a Group 1f NiFe hydrogenase, and rubisco,  
314 SI) —a lifestyle not previously described for members of the Chloroflexi.

## 315 316 **Conclusions**

317 The diversity of iron oxidizing bacteria at Jinata is different than in other Fe<sup>2+</sup>-rich  
318 springs and environments. For example, in freshwater systems such as Oku-Okuhachikurou  
319 Onsen in Akita Prefecture, Japan (11), and Budo Pond in Hiroshima, Japan (47), iron oxidation

320 is driven primarily by the activity of chemoautotrophs such as members of the Gallionellaceae.  
321 In contrast, at Chocolate Pots hot spring in Yellowstone National Park, USA, iron oxidation is  
322 primarily abiotic, driven by O<sub>2</sub> produced by Cyanobacteria, with only a small contribution from  
323 iron oxidizing bacteria (48, 49). The distinct iron-oxidizing community at Jinata Onsen may be  
324 related to the salinity of the spring water, or biogeographically by access to the ocean, as  
325 Zetaproteobacteria are typically found in marine settings, particularly in deep ocean basins  
326 associated with hydrothermal iron sources (24). Despite the taxonomically distinct iron oxidizer  
327 communities between Jinata and Oku-Okuhachikuro Onsen, both communities support only  
328 limited biomass in regions dominated by iron oxidizers (11), perhaps reflecting the shared  
329 biochemical and bioenergetic challenges iron oxidation incurred by diverse iron oxidizing  
330 bacteria including Gallionellaceae and Zetaproteobacteria (11, 24, 50). Future work focused on  
331 isolation and physiological characterization of microbes, quantification of rates and  
332 determination of microbial drivers of carbon fixation and aerobic and anaerobic heterotrophy,  
333 and carbon isotope profiling of organic and inorganic species along the flow path of the hot  
334 spring will be necessary to fully characterize the activity of microbes at Jinata and to fully  
335 compare this system to other areas with high dissolved ferrous iron concentrations (e.g. Oku-  
336 Okuhachikuro Onsen, 11, Fuschna Spring, 18, Jackson Creek, 19, and Chocolate Pots Hot  
337 Spring, 48, 49).

338 Throughout Earth history, the metabolic opportunities available to life, and the resulting  
339 organisms and metabolisms responsible for driving primary productivity, have been shaped by  
340 the geochemical conditions of the atmosphere and oceans. The modern, sulfate-rich, well-  
341 oxygenated oceans we see today reflect a relatively recent state—one typical of only the last few  
342 hundred million years (e.g. 51). For the first half of Earth history, until ~2.3 billion years ago  
343 (Ga), the atmosphere and oceans were anoxic (52), and the oceans were largely rich in dissolved  
344 iron but poor in sulfur (53). At this time, productivity was low and fueled by metabolisms such  
345 as methanogenesis and anoxygenic photosynthesis (12, 13, 15). Following the expansion of  
346 oxygenic photosynthesis by Cyanobacteria and higher primary productivity around the Great  
347 Oxygenation Event ~2.3 Ga (54-56), the atmosphere and surface ocean accumulated some  
348 oxygen, and the ocean transitioned into a state with oxygenated surface waters but often anoxic  
349 deeper waters, rich in either dissolved iron or sulfide (57-60). At Jinata Onsen, this range of  
350 geochemical conditions is recapitulated over just a few meters, providing a useful test case for  
351 probing the shifts of microbial productivity over the course of Earth history as conditions vary  
352 over short spatial scales. In particular, the concomitant increase in visible biomass at Jinata as the  
353 community shifts from lithotrophy toward water-oxidizing phototrophy (i.e. oxygenic  
354 photosynthesis) is consistent with estimates for greatly increased primary production following  
355 the evolution and expansion of Cyanobacteria around the GOE (14, 15, 55, 56).

356 The dynamic abundances of redox-active compounds including oxygen, iron, and  
357 hydrogen at Jinata may not only be analogous to conditions on the early Earth, but may have  
358 relevance for potentially habitable environments on Mars as well. Early Mars is thought to have  
359 supported environments with metabolic opportunities provided by the redox gradient between  
360 the oxidizing atmosphere and abundant electron donors such as ferrous iron and molecular  
361 hydrogen sourced from water/rock interactions (e.g. 61), and production of these substrates may  
362 continue today (62, 63). Understanding the potential productivity of microbial communities  
363 fueled by lithotrophic metabolisms is critical for setting expectations of the presence and size of  
364 potential biospheres on other worlds and early in Earth history (e.g. 11, 15). Uncovering the  
365 range of microbial metabolisms present under the environmental conditions at Jinata, and their

366 relative contributions to primary productivity, may therefore find application to predicting  
367 environments on Mars most able to support productive microbial communities.

368

## 369 **Materials and Methods:**

### 370 **Geological context and sedimentology of Jinata:**

371 Jinata Onsen is located at 34.318 N, 139.216 E on the island of Shikinejima, Tokyo  
372 Prefecture, Japan. Shikinejima is part of the Izu Islands, a chain of volcanic islands that formed  
373 in the last few million years along the northern edge of the Izu-Bonin-Mariana Arc (64).  
374 Shikinejima is formed of Late Paleopleistocene- to-Holocene non-alkaline felsic volcanics and  
375 Late-Miocene to Pleistocene non-alkaline pyroclastic volcanic flows, with Jinata Onsen located  
376 on a small bay on the southern side of the island (Figure 1).

### 377 **Sample collections:**

378 Five sites were sampled at Jinata Onsen: the Source Pool, Pool 1, Pool 2, Pool 3, and the  
379 Outflow (Figure 1, Figure 2). During the first sampling trip in January 2016, two whole  
380 community DNA samples were collected from each site for 16S rDNA amplicon sequencing.  
381 During the second sampling trip, additional DNA was collected from the Source Pool and Pool 2  
382 for shotgun metagenomic sequencing along with gas samples for qualitative analysis. Samples  
383 for quantitative gas analysis were collected in October 2017 and April 2018.

384 Samples were collected as mineral scrapings of loosely attached, fluffy iron oxide coating  
385 from surfaces and clasts upstream (Source Pool and Pool 1) and as samples of microbial mat  
386 downstream (Pools 2 and 3, and Outflow) using sterile forceps and spatulas (~0.25 cm<sup>3</sup> of  
387 material). Immediately after sampling, cells were lysed and DNA preserved with a Zymo  
388 Terralyzer BashingBead Matrix and Xpedition Lysis Buffer. Lysis was achieved by attaching  
389 tubes to the blade of a cordless reciprocating saw (Black & Decker, Towson, MD) and operating  
390 for 1 minute. Aqueous geochemistry samples consisted of water collected with sterile syringes  
391 and filtered through a 0.2 µm filter. Gas samples were collected near sites of ebullition emerging  
392 from the bottom of the Source Pool; collection was done into serum vials by water substitution,  
393 and then sealed underwater to prevent contamination by air.

### 394 **Geochemical analysis:**

395 Dissolved oxygen (DO), pH, and temperature measurements were performed *in situ* using  
396 an Extech DO700 8-in-1 Portable Dissolved Oxygen Meter (FLIR Commercial Systems, Inc.,  
397 Nashua, NH). Iron concentrations were measured using the ferrozine assay (65) following  
398 acidification with 40 mM sulfamic acid to inhibit iron oxidation by O<sub>2</sub> or oxidized nitrogen  
399 species (66). Ammonia/ammonium concentrations were measured using a TetraTest NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup>  
400 Kit (TetraPond, Blacksburg, VA) following manufacturer's instructions but with colorimetry of  
401 samples and NH<sub>4</sub>Cl standards quantified with a Thermo Scientific Nanodrop 2000c  
402 spectrophotometer (Thermo Fisher Scientific, Waltham, MA) at 700 nm to improve sensitivity  
403 and accuracy. Anion concentrations were measured via ion chromatography on a Shimadzu Ion  
404 Chromatograph (Shimadzu Corp., Kyoto, JP) equipped with a Shodex SI-90 4E anion column  
405 (Showa Denko, Tokyo, JP).

406 Presence of H<sub>2</sub> and CH<sub>4</sub> in gas samples was qualitatively determined with a Shimadzu  
407 GC-14A gas chromatograph within 12 hours of collection to minimize oxidation of reduced  
408 gases. Quantitative gas composition was measured following methods described in Suda et al.  
409 2017 (67) and in the SI.

### 410 **16S rDNA and metagenomic sequencing and analysis:**

411 Sequencing of 16S rDNA followed methods described previously (11) and detailed in the  
412 SI. Following initial characterization via 16S rDNA sequencing, four samples were selected for  
413 shotgun metagenomic sequencing: JP1-A and JP3-A from the first sampling trip, and JP1L-1 and  
414 JP2-1 from the second sampling trip. Purified DNA was submitted to SeqMatic LLC (Fremont,  
415 CA) for library preparation and 2x100bp paired-end sequencing via Illumina HiSeq 4000  
416 technology. Samples JP1-A and JP3-A shared a single lane with two samples from another  
417 project (7), while JP1L-1 and JP2-1 shared a lane with one sample from another project.

418 Raw sequence reads from all four samples were co-assembled with MegaHit v. 1.02 (68)  
419 and genome bins constructed based on nucleotide composition and differential coverage using  
420 MetaBAT (69), MaxBin (70), and CONCOCT (71) prior to dereplication and refinement with  
421 DAS Tool (72) to produce the final bin set. Genome bins were assessed for completeness,  
422 contamination, and strain-level heterogeneity using CheckM (73), tRNA sequences found with  
423 Aragorn (74), and presence of metabolic pathways of interest predicted with MetaPOAP (75).  
424 Coverage was extracted using bbmap (76) and samtools (77). Genes of interest (e.g. coding for  
425 ribosomal, photosynthesis, iron oxidation, and electron transport proteins) were identified from  
426 assembled metagenomic data locally with BLAST+ (78) and were screened against outlier (e.g.  
427 likely contaminant) contigs as determined by CheckM using tetranucleotide, GC, and coding  
428 density content. Translated protein sequences of genes of interest were aligned with MUSCLE  
429 (79), and alignments manually curated in Jalview (80). Phylogenetic trees were calculated using  
430 RAxML (81) on the Cipres science gateway (82). Node support for phylogenies was calculated  
431 with transfer bootstraps by BOOSTER (83). Trees were visualized with the Interactive Tree of  
432 Life viewer (84). Because sequencing depth of each sample in the full metagenome was uneven,  
433 relative abundance of genes of interest between metagenomic datasets was normalized to the  
434 coverage of *rpoB* genes in each raw dataset as mapped onto the coassembly. Like the 16S rRNA  
435 gene, *rpoB* is a highly conserved, vertically-inherited gene useful for taxonomic identification of  
436 organisms but has the added advantage that it is only known to occur as a single copy per  
437 genome (85) and is more readily assembled in metagenomic datasets (e.g. 7). Presence and  
438 classification of hydrogenase genes was determined with HydDB (86). Taxonomic assignment of  
439 MAGs was made based on placement in a reference phylogeny built with concatenated  
440 ribosomal protein sequences following (87) and confirmed using GTDB-Tk (88).

441

#### 442 **Data availability:**

443 Raw 16S rDNA, raw metagenomic sequence data, and MAGs have been uploaded and made  
444 publicly available on NCBI under Project Number PRJNA392119 (genome accession numbers  
445 can be found in Supplemental Table 6).

446

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770

771 **Figure 1:**

772 Location of Jinata Onsen on Shikinejima Island, Japan, and inset overview sketch of field site  
773 with sampling localities marked.

774

775 **Figure 2:**  
776 Representative photos of Jinata. A) Panorama of field site, with Source Pool on the left (Pool 1  
777 below), Pool 2 and 3 in the center, and Out Flow to the bay on the right. B) Undistorted view  
778 north up the canyon. C) Undistorted view south toward the bay, overlooking Pool 2. D) Source  
779 Pool, coated in flocculent iron oxides and bubbling with gas mixture containing H<sub>2</sub>, CO<sub>2</sub>, and CH<sub>4</sub>.  
780 E) Pool 2, with mixture of red iron oxides and green from Cyanobacteria-rich microbial mats. F)  
781 Close up of textured microbial mats in Pool 3. G) Close up of Out Flow, where hot spring water  
782 mixes with ocean water.

783

784 **Table 1:** Geochemistry of source water at Jinata Onsen.

<b>T</b>	63°C
<b>pH</b>	5.4
<b>DO</b>	4.7 μM
<b>Fe<sup>2+</sup></b>	261 μM
<b>NH<sub>3</sub>/NH<sub>4</sub><sup>+</sup></b>	87 μM
<b>Cl<sup>-</sup></b>	654 mM
<b>SO<sub>4</sub><sup>2-</sup></b>	17 mM
<b>NO<sub>3</sub><sup>-</sup></b>	<1.6 μM
<b>NO<sub>2</sub><sup>-</sup></b>	<2.2 μM
<b>HPO<sub>4</sub><sup>-</sup></b>	<1 μM

785

786 **Figure 3:**  
787 Summary of geochemical and microbiological trends along the flow path of Jinata Onsen. Top:  
788 panoramic view of Jinata Onsen, with Source Pool at left and flow of spring water toward the  
789 bay at right, with sampling locations indicated. Middle: geochemical transect across the spring,  
790 showing temperature (°C, left axis) and dissolved Fe(II) and O<sub>2</sub> (μM, right axis). Bottom:  
791 stacked bar chart of relative community abundance of dominant microbial phyla as determined  
792 by 16S rDNA amplicon sequencing. Sequence data were binned at the phylum level averaged at  
793 each sample site. Reads that could not be assigned to a phylum were discarded; all phyla that do  
794 not make up more than 2% of the community at any one site have been collapsed to “Other”.  
795 Near the source, the community is predominantly made up of iron- and/or hydrogen-oxidizing  
796 organisms in the Proteobacteria and Aquificae phyla. As the hot spring water flows downstream,  
797 it equilibrates with the atmosphere and eventually mixes with seawater, resulting in downstream  
798 cooling, accumulation of oxygen, and loss of dissolved iron due to biological and abiotic  
799 processes. Oxygenic Cyanobacteria become progressively more abundant downstream  
800 Hydrogen- and iron-oxidizing lithotrophs dominate near the source, but phototrophic  
801 Cyanobacteria come to dominate downstream. Additional community diversity is found in  
802 Supplemental Table 4.

803

804 **Figure 4:**  
805 Phylogeny of Bacteria and Archaea based on concatenated ribosomal proteins. Numbers in  
806 parentheses next to phylum labels refer to number of MAGs recovered from Jinata Onsen.  
807 Labels for phyla with two or fewer MAGs recovered from Jinata omitted for clarity. Reference  
808 alignment modified from (87). Full list of MAGs recovered available in Supplemental Table 6.

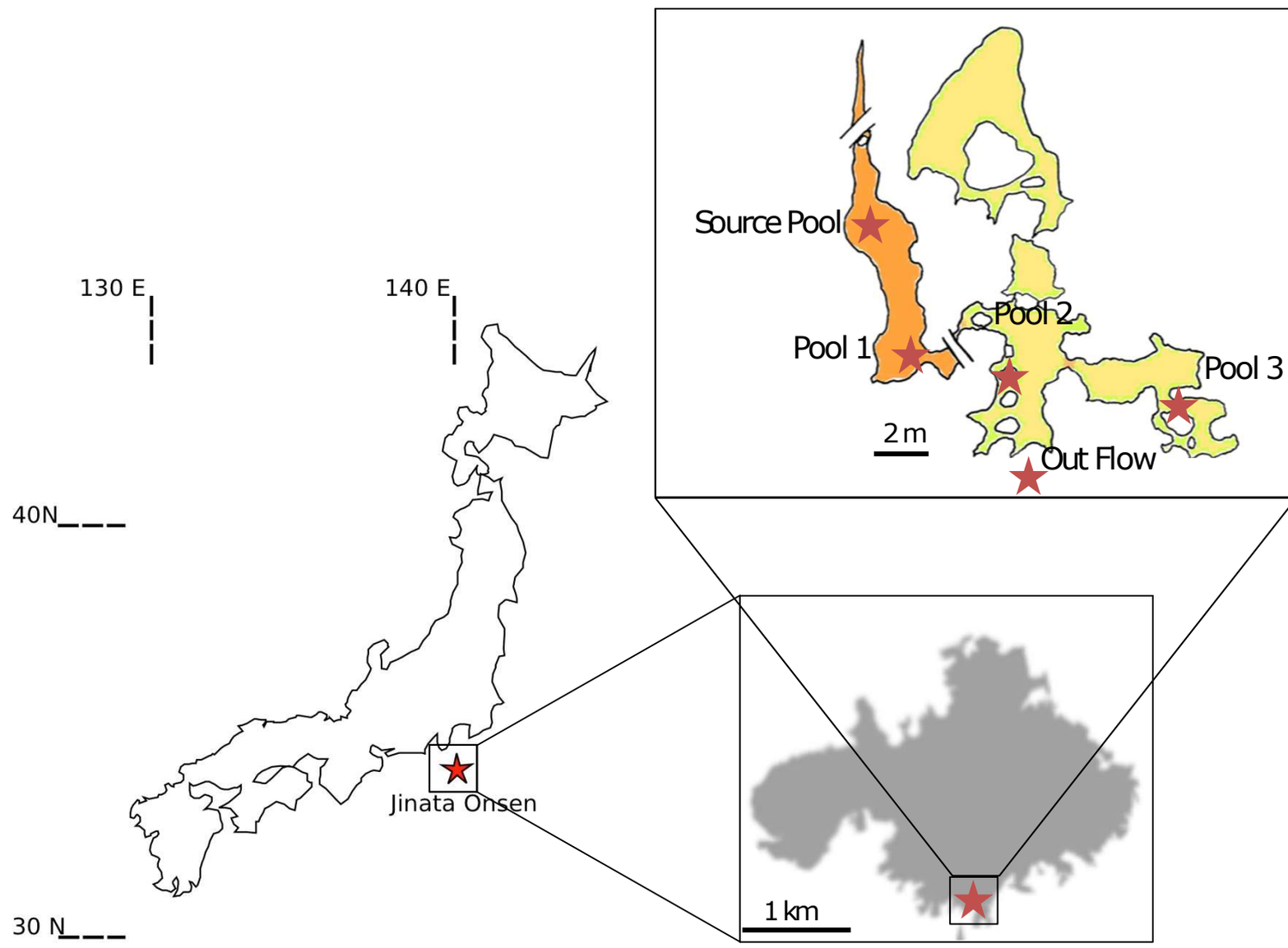
809

810 **Figure 5:** Phylogeny of the Zetaproteobacteria, rooted with Alphaproteobacteria, built with  
811 concatenated ribosomal protein sequences. Data from references 23, 89, 90, and other draft  
812 genomes available on Genbank. All nodes recovered TBE support values greater than 0.7. In  
813 cases where reference genomes have a unique strain name or identifier, this is included;  
814 otherwise Genbank WGS genome prefixes are used.

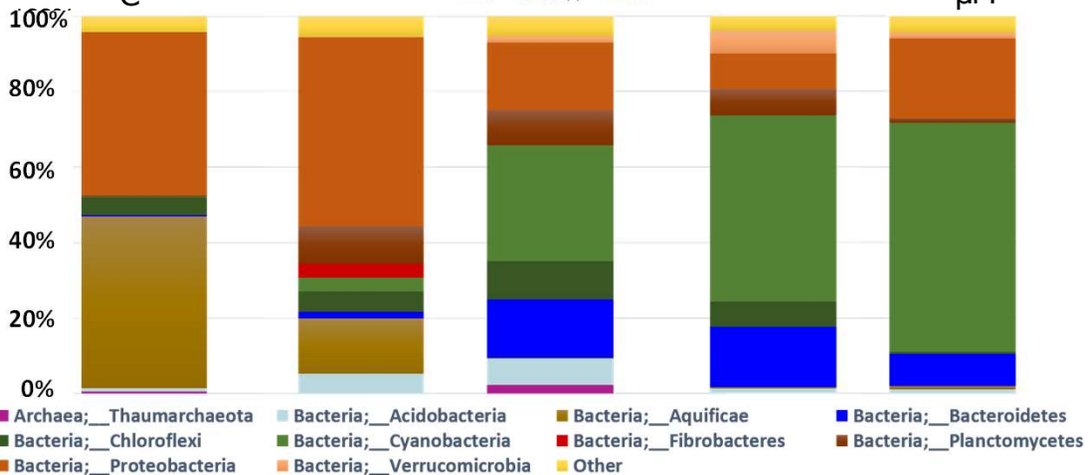
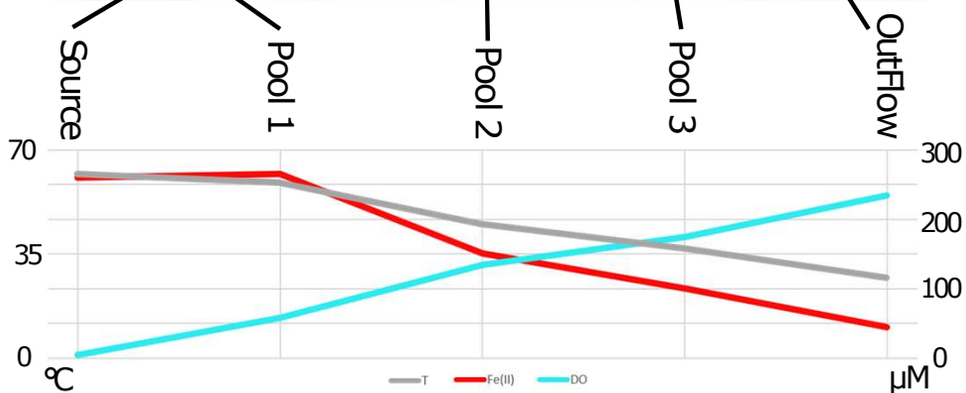
815  
816 **Figure 6:** Phylogeny of the Calditrarchaeota, rooted with Bacteroidetes, built with concatenated  
817 ribosomal protein sequences. Data from reference 91 and other draft genomes available on  
818 genomes have a unique strain name or identifier, this is included; otherwise Genbank WGS  
819 genome prefixes are used.

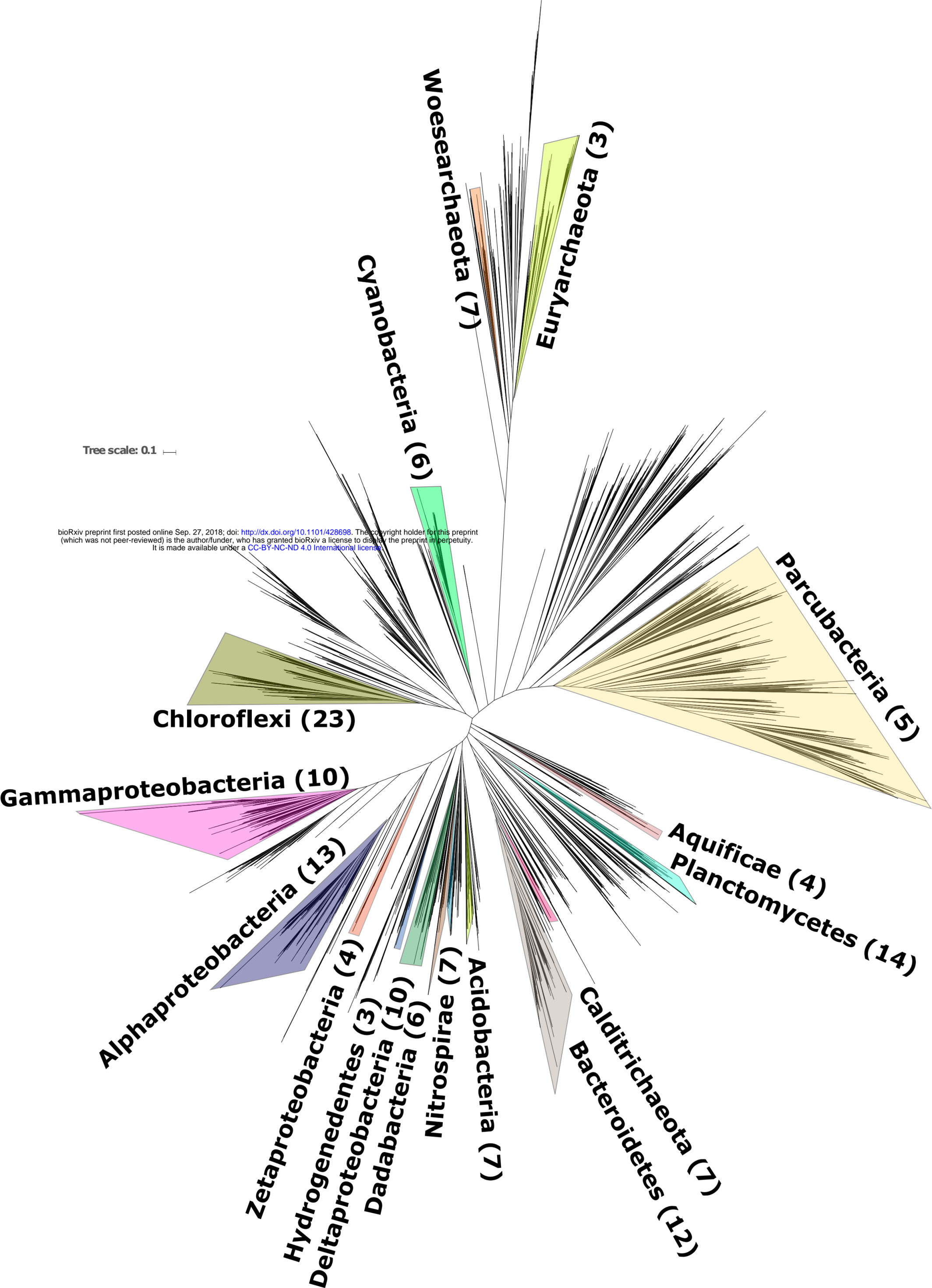
820  
821 **Figure 7:** Detailed phylogeny of the Chloroflexi phylum, with class-level clades highlighted in  
822 gray, built with concatenated ribosomal protein sequences. The large basal class  
823 Dehalococcoidia, which was not observed in 16S rDNA or metagenome data from Jinata, is  
824 omitted for clarity. Contains MAGs reported here, members of the Chloroflexi phylum  
825 previously described (7, 38, 41, 92-104), and members of the closely related phylum  
826 Armatimonadetes as an outgroup (105, 106). MAGs described here highlighted in green, MAGs  
827 previously reported from Jinata Onsen highlighted in pink. All nodes recovered TBE support  
828 values greater than 0.7. In cases where reference genomes have a unique strain name or  
829 identifier, this is included; otherwise Genbank WGS genome prefixes are used.

830









Tree scale: 0.1

