

Title	Detection of anaerobic carbon monoxide-oxidizing thermophiles in hydrothermal environments.
Author(s)	Yoneda, Yasuko; Kano, Sanae I; Yoshida, Takashi; Ikeda, Eitaro; Fukuyama, Yuto; Omae, Kimiho; Kimura-Sakai, Shigeiko; Daifuku, Takashi; Watanabe, Tetsuhiro; Sako, Yoshihiko
Citation	FEMS microbiology ecology (2015), 91(9)
Issue Date	2015-09
URL	http://hdl.handle.net/2433/207705
Right	<p>This is a pre-copyedited, author-produced PDF of an article accepted for publication in 'FEMS microbiology ecology' following peer review. The version of record [Yasuko Yoneda, Sanae I. Kano, Takashi Yoshida, Eitaro Ikeda, Yuto Fukuyama, Kimiho Omae, Shigeiko Kimura-Sakai, Takashi Daifuku, Tetsuhiro Watanabe, Yoshihiko Sako. Detection of anaerobic carbon monoxide-oxidizing thermophiles in hydrothermal environments. Volume 91, Issue 9, fiv093, September 2015] is available online at: http://femsec.oxfordjournals.org/content/91/9/fiv093.; The full-text file will be made open to the public on 28 July 2016 in accordance with publisher's 'Terms and Conditions for Self-Archiving'.; This is not the published version. Please cite only the published version. この論文は出版社版ではありません。引用の際には出版社版をご確認ご利用ください。</p>
Type	Journal Article
Textversion	author

1 **Title:** Detection of anaerobic carbon monoxide-oxidizing thermophiles in hydrothermal
2 environments

3

4 **Authors:** Yasuko Yoneda^{1,§}, Sanae I. Kano¹, Takashi Yoshida¹, Eitaro Ikeda¹, Yuto
5 Fukuyama¹, Kimiho Omae¹, Shigeko Kimura-Sakai^{1,¶}, Takashi Daifuku¹, Tetsuhiro
6 Watanabe² and Yoshihiko Sako^{1,*}.

7

8 **Authors Affiliations:**

9 ¹Laboratory of Marine Microbiology, Graduate School of Agriculture, Kyoto University,
10 Kitashirakawa Oiwake-cho, Sakyo-ku, Kyoto 606-8502, Japan.

11 ²Laboratory of Soil Science, Graduate School of Agriculture, Kyoto University,
12 Kitashirakawa Oiwake-cho, Sakyo-ku, Kyoto 606-8502, Japan.

13 [§]Current affiliation: Bioproduction Research Institute, National Institute of Advanced
14 Industrial Science and Technology (AIST), Central 6, Higashi 1-1-1, Tsukuba, Ibaraki
15 305-8566, Japan.

16 [¶]Current affiliation: School of Environmental Science, The University of Shiga
17 Prefecture, 2500 Hassaka-cho, Hikone city, Shiga 522-8533, Japan.

18

19 **Running title:** Anaerobic CO-oxidizing thermophiles in hydrothermal environments

20

21 ***Corresponding Author:** Yoshihiko Sako

22 **Phone:** +81 75 753 6217

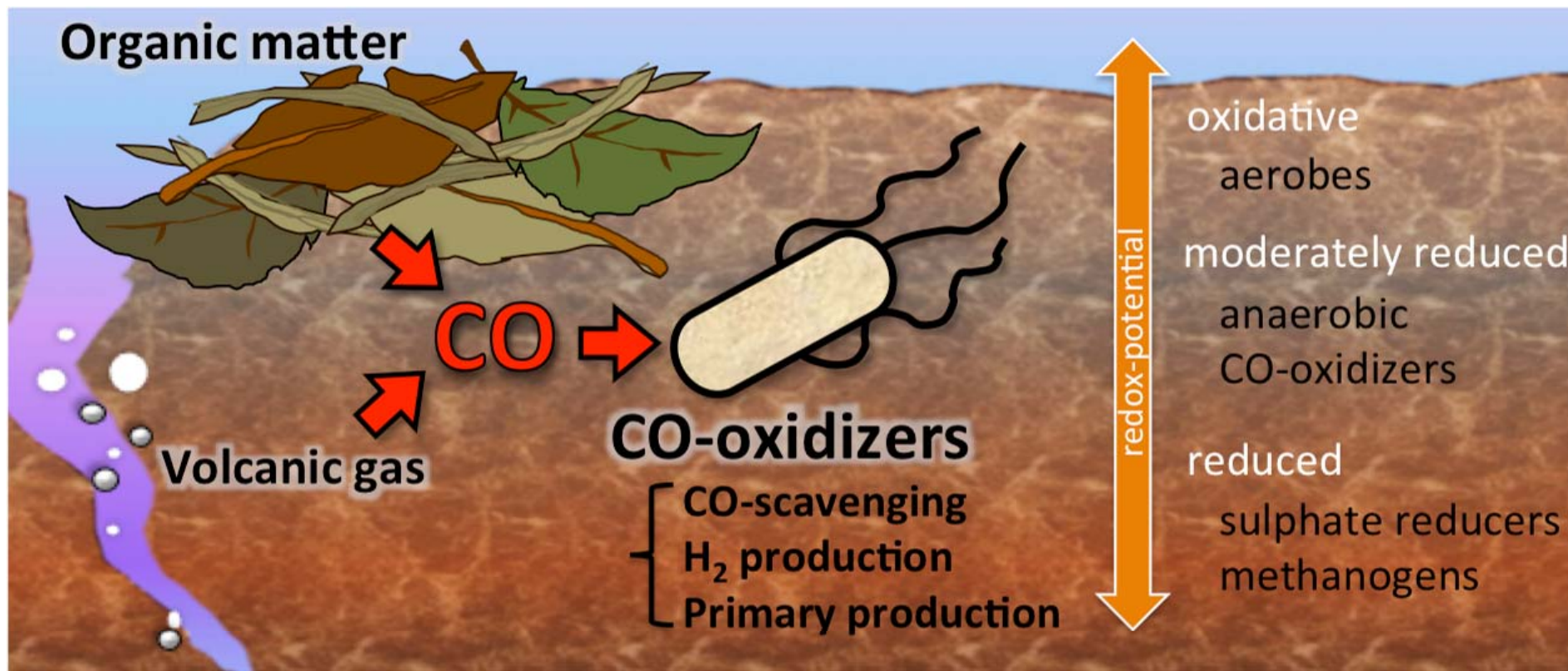
23 **Fax:** +81 75 753 6226

24 **E-mail:** sako@kais.kyoto-u.ac.jp

25

26 **Abstract**

27 Carboxydrotrophic anaerobic thermophiles have been isolated from various
28 hydrothermal environments and are considered to be important carbon monoxide (CO)
29 scavengers or primary producers. However, the ecological factors that influence the
30 distribution, abundance and CO-oxidizing activities of these bacteria are poorly
31 understood. A previous study detected the carboxydrotrophic bacteria *Carboxydotherrmus*
32 spp. in a hot spring sample and found that they constituted up to 10% of the total
33 bacterial cells. In this study, we investigated environmental features, potential microbial
34 CO-oxidation activities and the abundance of *Carboxydotherrmus* spp. in various hot
35 springs to determine environmental factors that affect CO-oxidizers and to see whether
36 *Carboxydotherrmus* spp. are common in those environments. We detected potential
37 microbial CO-oxidation activities in samples that showed relatively high values of total
38 organic carbon (TOC), total nitrogen (TN), oxidation–reduction potential (ORP) and
39 soil-water content. The abundance of *Carboxydotherrmus* spp. did not correlate with the
40 presence of potential microbial CO-oxidation activities; however, *Carboxydotherrmus*
41 spp. were detected in a wide range of environments, suggesting that these bacteria are
42 widely distributed in spite of the relatively low population size. This study implies
43 thermophilic CO-oxidizers occur in a wide range of environments and oxidize CO in
44 somewhat oxidative environments rich in organic matter.



Graphical Abstract Figure

45

46

47 **One-sentence summary**

48 Carbon-monoxide-oxidizing thermophiles are widely distributed in a range of
49 environments, and high potential microbial CO-oxidation activity is associated with the
50 levels of organic matter, oxidation–reduction potential and soil-water content of
51 environments.

52

53

54 **Keywords**

55 carboxidotroph, carbon monoxide, carbon monoxide dehydrogenase, thermophile, hot
56 spring

57

58 **Introduction**

59 It is believed that approximately 2,500–2,600 Tg of carbon monoxide (CO) is produced
60 annually and emitted to environments via the combustion of fossil fuels, oxidation or
61 photochemical degradation of organic compounds, volcanic gas, and metabolism in
62 animals, plants and microbes (King & Weber, 2007). CO concentrations in
63 environments usually occur in trace amounts; e.g., 2–15 nM in seawater (Moran &
64 Miller, 2007), 20–33 nM in hot springs (Kochetkova *et al.*, 2011) and approximately 92
65 ppb global mean mole in the atmosphere (WMO WDCGG, 2014). There is generally
66 equilibrium between CO production and consumption; approximately 90% of
67 atmospheric CO is consumed by reaction with hydroxyl radicals, and CO-oxidizing
68 microbes in soil are considered to be a CO sink accounting for 10% of the total CO
69 consumption (King, 2007).

70 Studies on microbiological CO consumption have been conducted under
71 aerobic conditions using soil, freshwater and seawater (Conrad & Seiler, 1980; Conrad
72 & Seiler, 1982). Aerobic CO-oxidizers utilize molybdenum-containing carbon
73 monoxide dehydrogenases (Mo-CODHs), which can be detected widely in
74 environments such as soils (plant root), sediments, volcanic deposits and the ocean floor
75 (King, 2003; King, 2007; Dunfield & King, 2004; Martin-Cuadrado *et al.*, 2009). In
76 anaerobes, CO oxidation is catalysed by nickel-containing CO dehydrogenases
77 (Ni-CODHs) (Ragsdale, 2004). Ni-CODHs can be divided into the following two major
78 groups: the monofunctional CODH group and the bifunctional CODH/acetyl CoA
79 synthase (ACS) group. Bifunctional CODH/ACSs comprise CODH subdomains and

80 ACS subdomains and are known to be key enzymes in the Wood–Ljungdahl pathway
81 (reductive acetyl CoA pathway) (Ragsdale, 2004). The CODH subdomains reduce CO₂
82 to CO, after which the ACS subdomains incorporate CO as a carboxyl group of
83 acetyl-CoA. Monofunctional CODHs catalyse oxidation of CO to CO₂, thereby deriving
84 low potential electrons whose energy can be converted to the transmembrane potential
85 and ATP when they are transferred to some oxidants (Techtmann *et al.*, 2009). In a
86 previous study, monofunctional and bifunctional CODH genes were predicted from
87 their genomic context. Monofunctional Ni-CODH genes are widely found in the
88 prokaryotic genomes of hydrogenogens, methanogens, sulphate reducers, acetogens and
89 gut microbes in animals (Techtmann *et al.*, 2012). CODH genes are found in diverse
90 microbes in various environments; however, only a few studies have reported anaerobic
91 microbial CO oxidation in environmental samples from soil, wetland and hot springs
92 (Conrad & Seiler, 1980; Rich & King, 1999; King, 2007; Kochetkova *et al.*, 2011), and
93 the environmental and physiological significance of anaerobic CO-oxidizers is less
94 understood.

95 The thermophilic bacterial genus *Carboxydotherrmus* is one of the most
96 studied CO-oxidizing anaerobes. Five species of the genus *Carboxydotherrmus* have
97 been isolated from hot springs and described (Yoneda *et al.*, 2012; Sokolova *et al.*,
98 2013). Four of the species produce hydrogen via CO-oxidization under 100% CO
99 atmospheric conditions. Because CO is a toxic gas and hydrogen is an important energy
100 source for many microbes in an anoxic environment, CO-oxidizers in the environment
101 are assumed to be important ‘CO scavengers’ or primary producers (Sokolova *et al.*,

102 2009; Techtmann *et al.*, 2009).

103 The genome of *C. hydrogenoformans* has four genes encoding
104 monofunctional CODH (*cooS-I*, *cooS-II*, *cooS-IV* and *cooS-V*), which have different
105 predicted functions and sequences (Wu *et al.*, 2005). Previously, the quantitative
106 detection of *Carboxydothemus* spp. by targeting the CODH-II gene (*cooS-II*) using
107 real-time PCR showed that these microorganisms can constitute up to approximately
108 10% of the total bacterial cells in acidic hot spring sediment where *Carboxydothemus*
109 *pertinax* was previously isolated (Yoneda *et al.*, 2013a). Considering that the optimal
110 growth pH of *Carboxydothemus* spp. is moderately acidic or neutral, the abundance of
111 these bacteria is intriguingly high. This led us to further question whether
112 *Carboxydothemus* spp. are common in other hot springs, where the environment is
113 more suitable for the growth of this genus, and are responsible for CO-oxidation *in situ*.
114 The physiological group of carboxydrotrophic hydrogenogens has been isolated from
115 various environments, such as hot springs, deep-sea hydrothermal vents, lake sediments
116 and bioreactors (Sokolova & Lebedinsky, 2013). Unfortunately, the range of
117 environments that offer a niche for these microorganisms is still unknown. The aim of
118 the current study was to determine the answer to the above question. In the current study,
119 we measured potential microbial CO-oxidation activities under anoxic conditions and
120 the abundance of *Carboxydothemus* spp. in various hot spring samples.

121

122 **Materials and methods**

123 **Sample collection**

124 Sample sediments were collected from hot springs located in the Kagoshima Pref.
125 (southern Kyushu Island) and Shizuoka Pref., Japan (Fig. 1). The hot springs were
126 Unagi-onsen in the Kagoshima Pref. and Atagawa-onsen (Benzainoyu and Izu
127 Atagawaso), Shimogamo-onsen (Jiunji temple), Mine-onsen (Mine Onsen Fountain
128 Park), Yatsu-onsen (Yakushinoyu) and Yugawara-onsen (Daikan-Soh) in Shizuoka Pref.
129 As references, samples were collected from the Unagi Lake near Unagi-onsen, a water
130 drain in Shimogamo-onsen and the Yamagawa coastal hydrothermal field that had high
131 salinity (Table 1). The Unagi Lake and a water drain near Shimogamo-onsen were
132 non-hydrothermal environments. The surface layer (above a 3.0 cm depth from the
133 water-sediment interface) of the sediments was sampled using a plastic core tube (7.0
134 cm diameter) or using a metal dipper. Samples were thoroughly mixed using metal
135 spoons and then subsampled into plastic tubes. For measurement of potential microbial
136 CO oxidation activities, samples were placed in a plastic bag with Aneropouch–Anaero
137 (Mitsubishi Gas Chemical, Tokyo, Japan) and sealed immediately. All sample tubes
138 were packed with ice and transported to the laboratory. Potential microbial CO
139 oxidation was measured as soon as possible after return to the laboratory. Samples for
140 DNA extraction were stored at $-80\text{ }^{\circ}\text{C}$ until use. The other samples were refrigerated at
141 $4\text{ }^{\circ}\text{C}$ until use.

142

143 **Physiochemical analyses of the environment**

144 The temperature, pH and oxidation–reduction potential (ORP) of the samples were
145 measured before sample collection using a portable meter HM-31P (DKK-TOA, Tokyo,

146 Japan). Soil-water content was calculated on the basis of the weight of the sediments
147 before and after overnight oven-drying at 105 °C. Pore water was extracted by filtration
148 using filter paper (Advantec quantitative ashless filter paper grade no. 5C, Toyo Roshi
149 Kaisha, Tokyo). The total organic carbon (TOC) and total nitrogen (TN) contents of
150 pore water were analysed using TOC-L (Shimadzu, Kyoto, Japan) equipped with a TN
151 meter unit, TNM-L (Shimadzu). The salinity of pore water was measured using a digital
152 salinity probe SS-31A (Sekisui Chemical, Osaka, Japan) in the laboratory at room
153 temperature.

154 A principal component analysis (PCA) was performed using R 3.0.2 (R
155 Development Core Team, 2008) to compare the environmental features of the samples.

156

157 **Measurement of potential microbial CO oxidation**

158 Samples were manipulated in an anaerobic glove box (Coy Laboratory Products, Grass
159 Lake, MI, USA) under a 5% H₂/ 95% N₂ atmosphere. Approximately 5.0 mL of
160 sediment was placed in each glass vial (63.8 mL), which was then sealed with a rubber
161 stopper. Glutaraldehyde solution (25% v/v), which is reported to effectively inhibit
162 thermophilic microbial CO-oxidation (Slepova *et al.*, 2007), was added to the control
163 samples (5% v/v at final concentration). All of the following manipulations were
164 conducted outside of the anaerobic glove box. Glass vials were vigorously vortexed and
165 pre-incubated at experimental temperatures for 1 h. The experimental temperature for
166 each sample was set as close to the temperature of its origin as possible. Five mL of
167 pure CO gas (CO PURE, Sumitomo Seika Chemicals, Osaka, Japan) was introduced in

168 each glass vial using a syringe (under atmospheric pressure at room temperature). From
169 the start time at CO injection, 1.0 mL of gas phase was subsampled from each vial using
170 a gas-tight syringe at approximately 24 h intervals over 5 days. CO concentrations in
171 subsamples were measured by gas chromatography as previously described (Yoneda *et*
172 *al.*, 2012). Potential microbial CO-oxidation in each sample was calculated based on the
173 difference in the average CO concentrations between the samples and controls. The
174 potential microbial CO-oxidation rate was evaluated by regression analysis and
175 regarded as positive when the slope of microbial CO-oxidation increased over the time
176 course ($p < 0.05$). Measurements of CO concentration in samples and controls were
177 performed in triplicate and duplicate, respectively.

178

179 **DNA extraction and real-time PCR**

180 DNA extraction was conducted using the commercial kit Extrap Soil DNA Kit Plus ver.
181 2 (NIPPON STEEL and SUMIKIN eco-tech, Tokyo, Japan) following the
182 manufacturer's protocols. Bead-beating steps were carried out using a Beads Crusher
183 μ T-12 (Taitec, Koshigawa, Japan) at a speed of 3,200 r/min for 60 s.

184 Quantitative PCR (qPCR) was performed for the *Carboxydotherrmus* CODH-II gene
185 (*cooS-II*), the bacterial 16S rRNA gene and the archaeal 16S rRNA gene as previously
186 described (Yoneda *et al.*, 2013a). Real-time PCR primers should be designed against
187 conserved regions of closely related strain sequences to cover the target organisms.

188 However, the primer set for *Carboxydotherrmus cooS-II* cooS2_1442F

189 (5'-TGATGCGTCACGGCTTTATGG-3') and cooS2-1606R

190 (5'-CTAAAGCTACTGCCCGGGAGT-3') were designed based on the *cooS-II* genes of
191 *C. hydrogenoformans* and *C. pertinax* (Yoneda *et al.*, 2013a) because no other
192 sequences closely related to *Carboxydothemus* species were available. No sequence
193 matches to the primers were found in NCBI nr database using BLASTn search. This
194 primer set can also amplify *cooS-II* from the genomic DNA of *Carboxydothemus*
195 *siderophilus*, but not *Carboxydothemus ferrireducens*. Therefore, the abundance of
196 CODH copy numbers observed in this study might be underestimated. Bacterial and
197 archaeal 16S rRNA gene primer sets were described previously (Einen *et al.*, 2008). For
198 *cooS-II*, qPCR was conducted in 4 to 7 replicates. The abundance of bacterial and
199 archaeal 16S rRNA genes was detected in triplicate. Some representative samples of
200 *cooS-II* qPCR products were purified, cloned and sequenced as previously described
201 (Yoneda *et al.*, 2013b). The obtained sequences were analysed using the MEGA 6
202 program (Tamura, 2013).

203

204 **Results**

205 **Sample description**

206 At Unagi-onsen, samples designated UG-20, UG-55 and UG-90 were collected from
207 three points with different temperatures in a hot spring pool. These samples were acidic
208 in the range of pH 2.8-5.3 (Table 1). As a reference, sediments from the Unagi Lake, a
209 freshwater maar lake located approximately 300 m downhill away from the sampling
210 site at Unagi-onsen, were also collected and designated by the label U-Lake. Sample
211 YG-65 was taken from beach sands under hot water flow at Yamagawa coastal

212 hydrothermal field, where general physicochemical and microbial characteristics have
213 been previously studied (Kawaichi *et al.*, 2013). The hot water of Yamagawa had a high
214 salinity of up to 2.2% (w/v). Hot springs in the Shizuoka Pref. were neutral to slightly
215 alkaline (pH 7.1–8.4) (Table 1). At Atagawa-onsen, deposits from open-air pools of the
216 hot spring well at Benzainoyu and Izu Atagawaso were designated by the labels BZ-65
217 and AT-90, respectively. At Yakushinoyu in Yatsu-onsen, two samples (samples KS-90
218 and KS-65) of hot spring deposits and mud were collected from open-air hot pools
219 created by continuous hot water inflow. A sample from Mine-onsen designated by the
220 label ME-90 was composed of white sinters scratched out from a pipe that was used to
221 upwell underground hot water. Sediment samples from Shimogamo-onsen were
222 collected from a trench drain of hot spring water in the grounds of the Jiunji Temple
223 (Samples JI-70 and JI-65). Salinity was as high as 0.7% (w/v) in samples JI-70 and
224 JI-65. Sediment samples designated by the label J-Drain were also collected from a
225 water drain that was next to, but not connected to, the hot water drain in the grounds of
226 the Jiunji Temple. Sample DA-80 was collected from a small hot water container owned
227 by Daikan-soh inn in Yugawara-onsen. White and orange-coloured hot spring deposits
228 were collected from the container, which is usually closed with a cover.

229

230 **Potential microbial CO-oxidation and environmental factors**

231 Incubation temperature settings for each sample are shown in Table 2. Potential
232 microbial CO-oxidation was observed in UN-55, UN-90, JI-70, JI-65 and J-Drain (Fig.
233 2). In samples that were positive for potential microbial CO-oxidation activities, ORP,

234 TOC and TN ranged from -49 mV to 487 mV, 14.7 mg/L to 147.1 mg/L and 3.7 mg/L
235 to 10.8 mg/L, respectively (Table 1). The highest amount of potential microbial CO
236 consumption was observed in JI-65, where as much as 63.6% of initial CO was
237 consumed during the experimental period of 5 days (Table 2). Relatively high CO
238 consumptions were also observed in J-Drain (51.2%), UN-55 (46.1%) and JI-70
239 (30.4%). Among the hot spring samples from Unagi-onsen (UN-20, UN-55 and UN-90)
240 and Shimogamo-onsen (JI-65 and JI-70), potential microbial CO-oxidation activities
241 were the highest at temperatures of 55 °C and 65 °C, respectively. These results were
242 expected as most of CO-oxidizing thermophilic isolates from terrestrial hot springs
243 show optimal growth at temperatures of approximately 55–65 °C (Sokolova *et al.*,
244 2009; Sokolova & Lebedinsky, 2013 and references therein). So far, only a few
245 hyperthermophilic carboxydrotrophic isolates that can grow at temperature >80 °C have
246 been reported, and they are archaeal species of the genera *Thermococcus* and
247 *Archeoglobus* from marine hydrothermal environments (Sokolova *et al.*, 2004; Henstra
248 *et al.*, 2007; Lee *et al.*, 2008). We observed potential microbial CO-oxidation in sample
249 UN-90 incubated at 90 °C. Previously, microbial CO oxidation has been reported in hot
250 springs as high as 80 °C and 90 °C (Kochetkova *et al.*, 2011). These results suggest the
251 existence of unknown hyperthermophilic CO-oxidizers in terrestrial hot springs.

252 Significant potential microbial CO-oxidation activities were not observed in
253 the other hot spring samples, such as KS-65 and BZ-65, in which temperature and pH
254 conditions seemed suitable for CO-oxidizers. To investigate the differences between
255 positive and negative samples, principal component analysis (PCA) was carried out

256 using environmental parameters as shown in Fig. 3. Sample YG-65 was excluded from
257 this analysis because an ORP value was not available. There were no apparent
258 similarities among samples where potential microbial CO-oxidation activities were
259 observed, though they were quite distinct from negative samples (Fig. 3). There was a
260 rough tendency of positive samples to be relatively rich in nutrients (TOC and TN) and
261 have higher ORP values and soil-water contents than negative samples. Among samples
262 associated with the environmental temperature of approximately 65–71 °C and
263 circum-neutral pH, the TOC and TN values of JI-65 and JI-70 (positive samples) were
264 more than double those of KS-65 and BZ-65 (negative samples). ORP values of positive
265 samples ranged from –49 mV to 487 mV and were higher than those of negative
266 samples (Table 1). The OPR values were in a range of ‘reduced’ to ‘moderately reduced’
267 redox condition that appears too oxidative for sulphate reduction and methanogenesis
268 (DeLaune & Reddy, 2005; Klüpfel *et al.*, 2014). One of the presumed sources of CO in
269 environments is methanogens and sulphate reducers because some of them produce CO
270 under laboratory conditions (Conrad & Thauer, 1983; Lupton *et al.*, 1984; Techtmann *et*
271 *al.*, 2009). However, in our case, it appeared that the metabolic activity of those
272 microorganisms did not necessarily correlate with the distribution of CO-oxidizing
273 anaerobes. Alternatively, CO is produced from volcanic gas and organic substances
274 (e.g., plant matter and humus). CO production from organic matter is enhanced in the
275 presence of oxygen, higher temperatures and higher water contents (Conrad & Seller
276 1985; Tarr *et al.*, 1995; Shade *et al.*, 1999; Hellebrand & Shade, 2008). Our result
277 implies that the CO source is likely from organic substances and suggests that

278 CO-oxidizers prefer environments where high CO productivity can be expected (high
279 ORP values, TOC, TN and soil-water contents).

280

281 **Abundance of *Carboxydotherrmus* CODH-II genes and 16S rRNA genes**

282 Results of qPCR are shown in Fig. 4. The technical lower detection limit of qPCR for
283 the standard bacterial 16S rRNA gene, archaeal 16S rRNA gene and *cooS-II* were
284 1.0×10^3 , 1.0×10^3 , and 5.0×10^1 copies/ μ L, respectively. In environmental samples,
285 *Carboxydotherrmus cooS-II* genes were detected in more than two replicates of samples
286 of UN-55, UN-90 and YG-65. *Carboxydotherrmus cooS-II* was also detected in one or
287 two replicates from UN-20, U-Lake, BZ-65, ME-90, JI-70 and JI-65. The highest copy
288 number was 6.05×10^4 copies/g sediment in UN-55 corresponding to 7.95×10^{-4} % of
289 bacterial 16S rRNA. *Carboxydotherrmus cooS-II* copies were detected from samples
290 UN-90 and ME-90, where the environmental temperatures were above growth
291 temperature (≥ 78 °C) of *Carboxydotherrmus* spp. (Yoneda *et al.*, 2012). On the other
292 hand, *Carboxydotherrmus cooS-II* was below detection level in sample KS-65 despite its
293 environmental temperature being optimal (65 °C) to most *Carboxydotherrmus* spp.

294 The qPCR series was conducted on CO-incubated samples used for
295 measurement of potential microbial CO-oxidation activities after incubation for a week
296 from the start time of the experiment. *Carboxydotherrmus cooS-II* was detected in more
297 than two replicates in CO-incubated samples UN-20, UN-55, UN-90, KS-90 and KS-65.
298 Interestingly, *Carboxydotherrmus cooS-II* was detected in CO-incubated KS-90 and
299 KS-65 samples where its corresponding environmental samples produced negative

300 results. In contrast, *Carboxydotherrmus cooS-II* was not detected in CO-incubated JI-70
301 and JI-65 samples. Abundances of *cooS-II* in environments and in CO incubated
302 samples from UN-55 and UN-90 were compared by t-test. There were no significant
303 differences in *cooS-II* copy numbers in either sample (Fig. 4).

304 Some of the representatives of qPCR products from environmental samples and CO
305 incubated sediments were sequenced and the inner region of the *cooS-II* primers (123
306 bp) were compared to *cooS-II* of *C. hydrogenoformans* (CP000141), *C. ferrireducens*
307 (IMG accession no. 2510417609) and *C. pertinax* (AB723512) obtained from
308 DDBJ/EMBL/GenBank or Integrated Microbial Genome (IMG) databases (data not
309 shown). A total of 58 sequences were obtained from the environmental samples. Of
310 these, 31 sequences showed 97–100% similarity to *C. pertinax cooS-II*: UN-20 (4),
311 UN-55 (8), UN-90 (6), U-Lake (3), YG-65 (3), BZ-65 (3), MI-90 (1) and JI-65 (3). A
312 total of 23 sequences showed 98–100% similarity to *C. hydrogenoformans cooS-II* from
313 samples UN-90 (7), U-Lake (1), YG-65 (2), BZ-65 (1), MI-90 (7), JI-70 (4). The
314 remaining four sequences, all from UN-90, showed 94% similarity to both
315 *Carboxydotherrmus ferrireducens* and *C. pertinax cooS-II*. A total of 66 sequences were
316 obtained from CO incubated sediment samples. Of these, 39 sequences were most
317 related to *C. pertinax cooS-II* with 97–100% sequence similarity: U-Lake (5), UN-20
318 (6), UN-55 (10), UN-90 (5), YG-65 (1), BZ-65 (2), KS-65 (5) and KS-90 (5). The other
319 27 sequences were most related to *C. hydrogenoformans* with 97–99% similarity:
320 UN-20 (2), UN-55 (2), UN-90 (3), YG-65 (3), BZ-65 (5), KS-65 (8) and KS-90(4). The
321 divergent sequences found in our study sites showed there are variations in

322 *Carboxydothemus* spp. and suggested that species-level diverse CO-oxidizers co-exist
323 in certain environments.

324

325 **Discussion**

326 To the best of our knowledge, our work is the first attempt to discover the distribution
327 and abundance of carboxydrotrophic anaerobic thermophiles in various hydrothermal
328 environments in relation to environmental factors. Most carboxydrotrophic anaerobic
329 thermophiles require strictly anaerobic, reduced conditions for carboxydrotrophic growth.
330 However, most of them are not obligate carboxydrotrophs and are able to utilize organic
331 substrate and electron acceptors such as ferric iron, AQDS
332 (anthraquinone-2,6-disulfonate) and sulphur compounds (Sokolova *et al.*, 2009;
333 Sokolova & Lebedinsky, 2013). Those alternative metabolic pathways may enable
334 carboxydrotrophic bacteria to survive in moderately reduced conditions. We observed
335 potential microbial CO-oxidation in samples where ORP ranged from -49 mV to 487
336 mV. Within this ORP range, respiration of iron, fumarate and humic substances can
337 take place (Klüpfel *et al.*, 2014), and CO-oxidation may take place as a subsidiary
338 reaction. Our results suggested that anaerobic CO-oxidizing activity is associated with
339 redox conditions, soil-water content, TOC and TN. TOC can be presumed as a CO
340 source where a positive association is conceivable between CO-oxidizing activity and
341 TOC value. Fig. 3 shows that TN is also a putative factor that influences potential
342 microbial CO-oxidation activity. Some CO-oxidizing thermophiles can reduce nitrates
343 as electron acceptors when grown on organic compounds such as yeast extract (e.g.,

344 *Moorella thermoacetica*) and lactate (e.g., *C. hydrogenoformans*) (Fröstl *et al.*, 1996;
345 Henstra *et al.*, 2004). Further studies are needed to determine which nitrogen
346 compounds are associated with the distribution or activity of CO-oxidizers. In
347 environments rich in organic matter, the ability to withstand and utilize CO may be an
348 advantage, allowing microbes to compete with fast-growing aerobes, as CO generally
349 inhibits cytochrome oxidase within aerobic respiration (Cooper & Brown, 2008).
350 Potential microbial CO-oxidation activities were detected in relatively high ORP sites,
351 suggesting that CO-oxidizers may have a niche in an intermediate zone of aerobic and
352 anaerobic environments thanks to their utilization of electron acceptors such as nitrate,
353 Fe(III) and fumarate.

354 The abundance of *Carboxydotherrmus* spp. did not correlate with potential
355 CO-oxidation activities and there were no apparent similarities in the environmental
356 features among samples in which *Carboxydotherrmus* spp. were detected (Fig. 2, Fig. 3
357 and Fig. 4a). Moreover, no significant changes in the *Carboxdotherrmus* spp. population
358 were observed between environmental samples and CO-incubated samples from UN-55
359 and UN-90 (Fig. 4a and 4b). In samples JI-65 and JI-70, abundance of
360 *Carboxydotherrmus* spp. dropped below the threshold for detection despite the presence
361 of potential microbial CO oxidation activity. These results indicate that there were other
362 unknown CO-oxidizers responsible for CO-oxidation in some of our samples.
363 Interestingly, the abundance of *Carboxydotherrmus* spp. was maintained or even
364 increased to a detectable level when incubated at a high temperature of 90 °C (in sample
365 UN-90 and KS-90), although sharp decreases in bacterial population were observed. *C.*

366 *hydrogenoformans* can produce spores (Wu *et al.*, 2005) that may allow this species to
367 maintain its population in inhospitable environments. *Carboxydotherrmus* spp. can be
368 regarded as rare and robust bacteria that are distributed in a wide range of environments
369 including extremely thermophilic hot springs, non-hydrothermal environments, and
370 coastal environments.

371 Anaerobic CODH genes are widely distributed in physiologically diverse bacteria and
372 archaea. Their phylogenetic relationships are often incongruent with taxonomy based on
373 16S rRNA genes, especially clade F including *cooS-I*, *cooS-II* and *cooS-IV* of *C.*
374 *hydrogenoformans* (Techtmann *et al.* 2012). For example, a gene cluster which include
375 *cooS-I* in *C. hydrogenoformans* is closely related with that of *Thermosinus*
376 *carboxydivorans* indicating horizontal gene transfer of the cluster (Techtmann *et al.*
377 2012). Thus, we could not exclude the possibility of unintended detection of the
378 recipient microorganisms using our *cooS-II* primer set.

379 Because CO is a common trace chemical in environments, further study on diversity,
380 competition or habitat segregation would be of interest. In addition, further research
381 should include exploration of new habitats (e.g., extremely hot or acidic terrestrial hot
382 springs) for CO-oxidizing anaerobes.

383

384 **Acknowledgement**

385 We would like to thank Makoto Shibata for support in analysing the chemical properties
386 of the samples. We would like thank Dr. Stephen M. Techtmann for kindly providing
387 CODH gene sequence information that greatly improved our knowledge regarding

388 CODH and anaerobic carboxydrotrophic microbes. We are also grateful to the local
389 people and the owners of the hot springs for willingly supporting our sampling.

390 This work was supported by a Grant-in-Aid for Scientific Research (A) (No.
391 20248023) and (A) (25252038) from The Ministry of Education, Culture, Sports,
392 Science and Technology (MEXT) and a Grant-in-Aid for JSPS Fellows (No. 244441)
393 from the Japan Society for the Promotion of Science (JSPS).

394 **References**

- 395 **Conrad R & Seiler W (1980)** Role of microorganisms in the consumption and
396 production of atmospheric carbon monoxide by soil. *Appl Environ Microbiol* **40**:
397 437-445.
- 398 **Conrad R & Seiler W (1982)** Utilization of traces of carbon monoxide by aerobic
399 oligotrophic microorganisms in ocean, lake and soil. *Arch Microbiol* **132**: 41-46.
- 400 **Conrad R & Thauer RK (1983)** Carbon monoxide production by *Methanobacterium*
401 *thermoautotrophicum*. *FEMS Microbiol Lett* **20**: 229-232.
- 402 **Conrad R & Seiler W (1985)** Characteristics of abiological carbon monoxide
403 formation from soil organic matter, humic acids, and phenolic compounds. *Environ Sci*
404 *Technol* **19**: 1165-1169.
- 405 **Cooper CE & Brown GC (2008)** The inhibition of mitochondrial cytochrome oxidase
406 by the gases carbon monoxide, nitric oxide, hydrogen cyanide and hydrogen sulfide:
407 chemical mechanism and physiological significance. *J Bioenerget Biomem* **40**: 533-539.
- 408 **DeLaune RD & Reddy KR (2005)** Redox potential. In *Encyclopedia of Soils in the*
409 *Environment* (pp 366-371) Hillel D (Ed.) Elsevier, Amsterdam, Netherlands.
- 410 **Dunfield KE & King GM (2004)** Molecular analysis of carbon monoxide-oxidizing
411 bacteria associated with recent Hawaiian volcanic deposits. *Appl Environ Microbiol* **70**:
412 4242-4248.

413 **Einen J, Thorseth IH & Øvreås L (2008)** Enumeration of *Archaea* and *Bacteria* in
414 seafloor basalt using real-time PCR quantitative PCR and fluorescence microscopy.
415 *FEMS Microbiol Lett* **282**: 182-187.

416 **Fröstl, JM, Seifritz C & Drake HL (1996)** Effect of nitrate on the autotrophic
417 metabolism of the acetogens *Clostridium thermoautotrophicum* and *Clostridium*
418 *thermoaceticum*. *J Bacteriol* **178**: 4597-4603.

419 **Hellebrand HJ & Schade GW (2008)** Carbon monoxide from composting due to
420 thermal oxidation of biomass. *J Environ Qual* **37**: 592-598.

421 **Henstra AM & Stams AJM (2004)** Novel Physiological Features of
422 *Carboxythermus hydrogenoformans* and *Thermoterrabacterium ferrireducens*. *Appl*
423 *Environ Microbiol* **70**: 7236-7240.

424 **Henstra AM, Dijkema C & Stams AJ (2007)** *Archaeoglobus fulgidus* couples CO
425 oxidation to sulfate reduction and acetogenesis with transient formate accumulation.
426 *Environ Microbiol* **9**: 1836-1841.

427 **Kawaichi S, Ito N, Yoshida T & Sako Y. (2013)** Bacterial and Archaeal Diversity in
428 an Iron-Rich Coastal Hydrothermal Field in Yamagawa, Kagoshima, Japan. *Microbes*
429 *Environ* **28**: 405-413.

430 **King GM (2003)** Molecular and culture-based analyses of aerobic carbon monoxide
431 oxidizer diversity. *Appl Environ Microbiol* **69**: 7257-7265.

432 **King GM (2007)** Microbial carbon monoxide consumption in salt marsh sediments.
433 *FEMS Microbiol Ecol* **59**: 2-9.

434 **King GM & Weber CF (2007)** Distribution, diversity and ecology of aerobic
435 CO-oxidizing bacteria. *Nature Rev Microbiol* **5**: 107-118.

436 **Klüpfel L, Piepenbrock A, Kappler A & Sander M (2014)** Humic substances as fully
437 regenerable electron acceptors in recurrently anoxic environments. *Nature Geosci* **7**:
438 195-200.

439 **Kochetkova TV, Rusanov II, Pimenov NV, Kolganova TV, Lebedinsky AV,**
440 **Bonch-Osmolovskaya EA & Sokolova TG (2011)** Anaerobic transformation of carbon
441 monoxide by microbial communities of Kamchatka hot springs. *Extremophiles* **15**:
442 319-325.

443 **Lee HS, Kang SG, Bae SS, Lim JK, Cho Y, Kim YJ, Jeong JH, Cha S-S, Kwon KK,**
444 **Kim H-T, Park C-J, Lee H-W, Kim SI, Chun J, Colwell RR, Kim S-J & Lee J-H**
445 **(2008)** The complete genome sequence of *Thermococcus onnurineus* NA1 reveals a
446 mixed heterotrophic and carboxydrotrophic metabolism. *J Bacteriol* **190**: 7491-7499.

447 **Lupton FS, Conrad R & Zeikus JG (1984)** CO metabolism of *Desulfovibrio vulgaris*
448 strain Madison: physiological function in the absence or presence of exogeneous
449 substrates. *FEMS Microbiol Lett* **23**: 263-268.

450 **Martin-Cuadrado A.-B, Ghai R, Gonzaga A & Rodriguez-Valera F (2009)** CO
451 dehydrogenase genes found in metagenomic fosmid clones from the deep
452 Mediterranean Sea. *Appl Environ Microbiol* **75**: 7436-7444.

453 **Moran MA & Miller WL (2007)** Resourceful heterotrophs make the most of light in
454 the coastal ocean. *Nature Rev Microbiol* **5**: 792-800.

455 **R Development Core Team (2008)** R: a language and environment for statistical

456 computing. ISBN 3-900051-07-0. URL <http://www.R-project.org>.

457 **Ragsdale SW (2004)** Life with carbon monoxide. *Crit Rev Biochem Mol Biol* **39**:

458 165-195.

459 **Rich JJ & King GM (1999)** Carbon monoxide consumption and production by wetland

460 peats. *FEMS microbiol Ecol* **28**: 215-224.

461 **Schade GW, Hofmann RM & Crutzen PJ (1999)** CO emissions from degrading plant

462 matter. *Tellus B* **51**: 889-908.

463 **Slepova, TV, Rusanov II, Sokolova TG, Bonch-Osmolovskaya EA & Pimenov NV**

464 **(2007)** Radioisotopic tracing of carbon monoxide conversion by anaerobic thermophilic

465 prokaryotes. *Microbiology* **76**: 523-529.

466 **Sokolova TG, Jeanthon C, Kostrikina NA, Chernyh NA, Lebedinsky AV,**

467 **Stackebrandt E & Bonch-Osmolovskaya EA (2004)** The first evidence of anaerobic

468 CO oxidation coupled with H₂ production by a hyperthermophilic archaeon isolated

469 from a deep-sea hydrothermal vent. *Extremophiles* **8**: 317-323.

470 **Sokolova TG, Henstra AM, Sipma J, Parshina SN, Stams AJ & Lebedinsky AV**

471 **(2009)** Diversity and ecophysiological features of thermophilic carboxydrotrophic

472 anaerobes. *FEMS Microbial Ecol* **68**: 131-141.

473 **Sokolova T & Lebedinsky A (2013)** CO-Oxidizing Anaerobic Thermophilic

474 Prokaryotes. In *Thermophilic Microbes in Environ Indust Biotechnol* 2nd ed. (pp.

475 203-231). Satyanarayana T, Littlechild J & Kawarabayashi Y (Eds.) Springer,

476 Amsterdam, Netherlands.

477 **Tamura K, Stecher G, Peterson D, Filipowski A & Kumar S (2013)** MEGA6: Molecular
478 Evolutionary Genetics Analysis Version 6.0. *Mol Biol Evol* **30**: 2725-2729.

479 **Tarr MA, Miller WL & Zepp RG (1995)** Direct carbon monoxide photoproduction
480 from plant matter. *J Geophys Res* **100**: 11403-11413.

481 **Techtmann SM, Colman AS & Robb FT (2009)** ‘That which does not kill us only
482 makes us stronger’: the role of carbon monoxide in thermophilic microbial consortia.
483 *Environ Microbiol* **11**: 1027-1037.

484 **Techtmann SM, Lebedinsky AV, Colman AS, Sokolova TG, Woyke T, Goodwin L &
485 Robb FT (2012)** Evidence for horizontal gene transfer of anaerobic carbon monoxide
486 dehydrogenases. *Front Microbiol* **3**: 132.

487 **WMO WDCGG (2014)** Data Summary No. 38. Japan Meteorological Agency. World
488 Meteorological Organization, Tokyo, Japan, 2014.

489 **Wu M, Ren Q, Durkin AS, Daugherty SC, Brinkac LM, Dodson RJ, Madupu R,
490 Sullivan SA, Kolonay JF, Nelson WC, Tallon LJ, Jones KM, Ulrich LE, Gonzalez
491 JM, Zhulin IB, Robb FT & Eisen JA (2005)** Life in hot carbon monoxide: the
492 complete genome sequence of *Carboxydothemus hydrogenoformans* Z-2901. *PLoS*
493 *Genet* **1**: e65.

494 **Yoneda Y, Yoshida T, Kawaichi S, Daifuku T, Takabe K & Sako Y. (2012)**
495 *Carboxydothemus pertinax* sp. nov., a thermophilic, hydrogenogenic, Fe (III)-reducing,
496 sulfur-reducing carboxydrotrophic bacterium from an acidic hot spring. *Int J Syst Evol*
497 *Microbiol* **62**: 1692-1697.

498 **Yoneda Y, Yoshida T, Daifuku T, Kitamura T, Inoue T, Kano S & Sako Y. (2013a)**

499 Quantitative detection of carboxydophilic bacteria *Carboxydothemus* in a hot aquatic
500 environment. *Fundam Appl Limnol* **182**: 161-170.

501 **Yoneda Y, Yoshida T, Yasuda H, Imada C & Sako Y. (2013b)** A novel thermophilic,
502 hydrogenogenic, and carboxydophilic bacterium *Calderihabitans maritimus* gen. nov.,
503 sp. nov. from a marine sediment core of an undersea caldera. *Int J Syst Evol Microbiol*
504 **63**: 3602-3608.

505

1 Figure legends

2 Fig. 1. Sampling sites in (a) the Kagoshima Pref. in southern Kyushu Island and (b)
3 Shizuoka Pref.

4 Original maps were obtained from a digital map provided by The Geospatial
5 Information Authority of Japan (GSI) (<http://www.gsi.go.jp/>) and have been edited by
6 the authors.

7

8 Fig. 2. Potential microbial CO-oxidation in environmental samples

9 Each result of regression analysis is shown outside each graph. Lines showing potential
10 microbial CO-oxidation are dotted when negative in the regression analysis.

11

12 Fig. 3. Principal component analysis of environmental features

13 Closed symbols show positive samples for potential microbial CO-oxidation.

14

15 Fig. 4. Abundance of the *Carboxydotherrmus cooS-II* and 16S rRNA genes in (a)
16 environmental samples and (b) in CO-incubated samples

17 *CooS-II* bars are average values of positive replicates (negative replicates are not
18 included in the analysis). Error bars are shown when the genes were detected in more
19 than two replicates (i.e., detected at least in triplicate).

20

Fig. 1

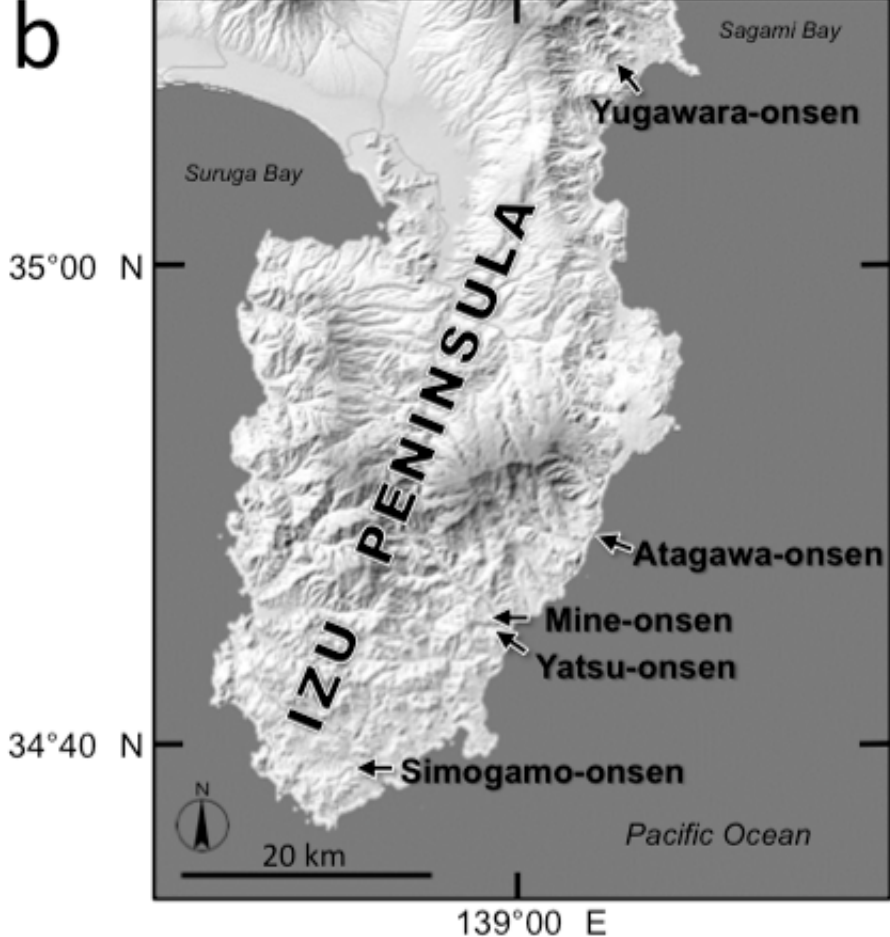
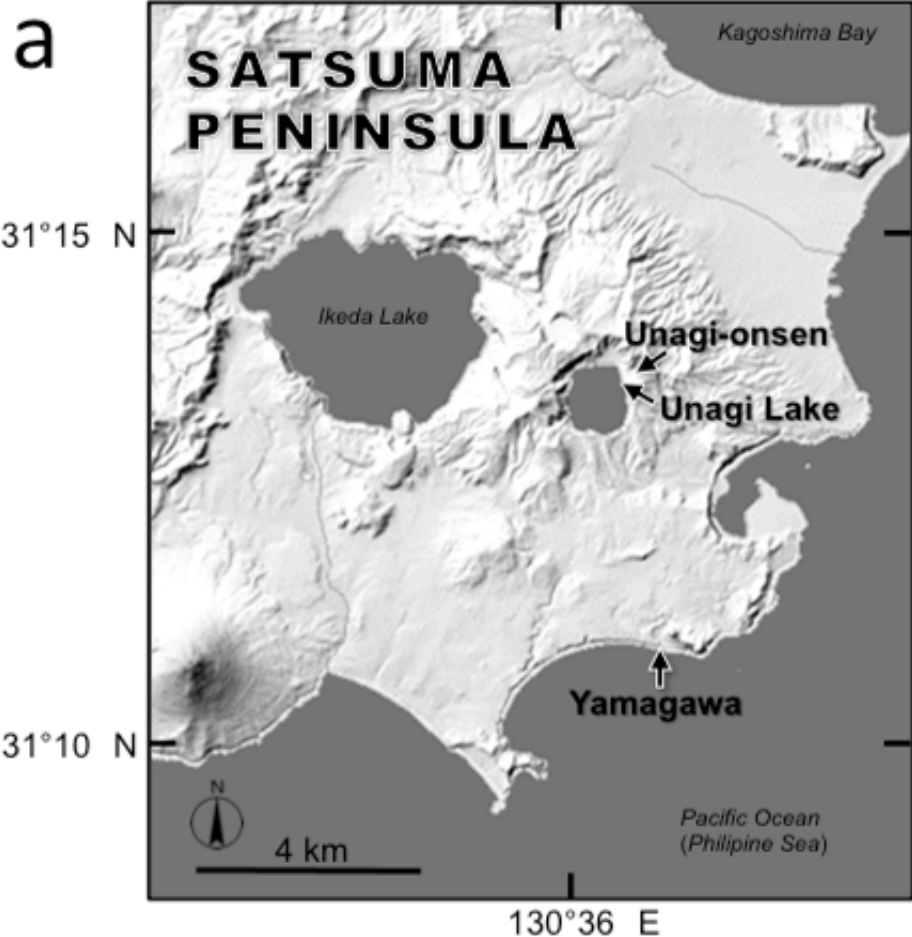


Fig. 3

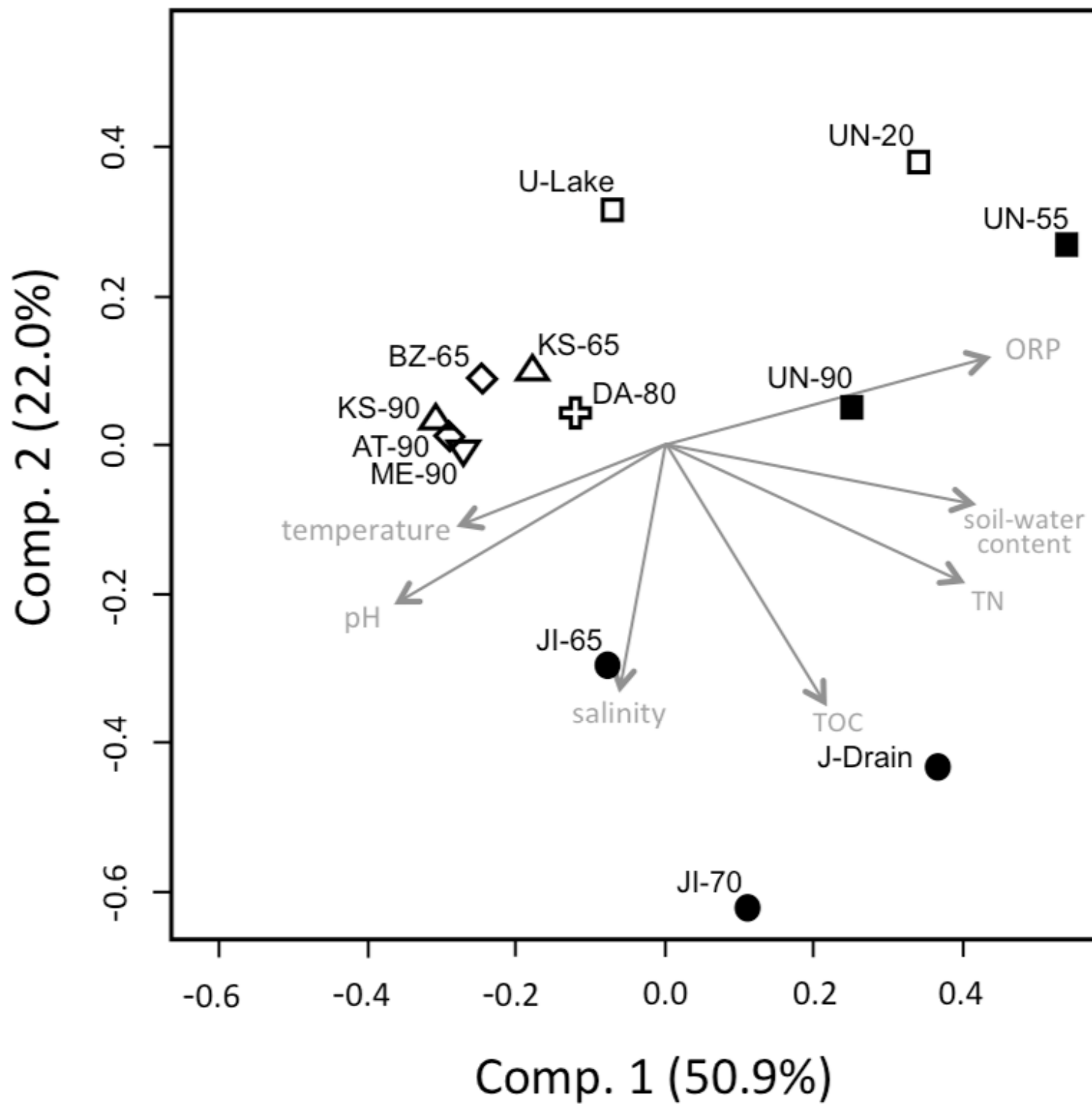
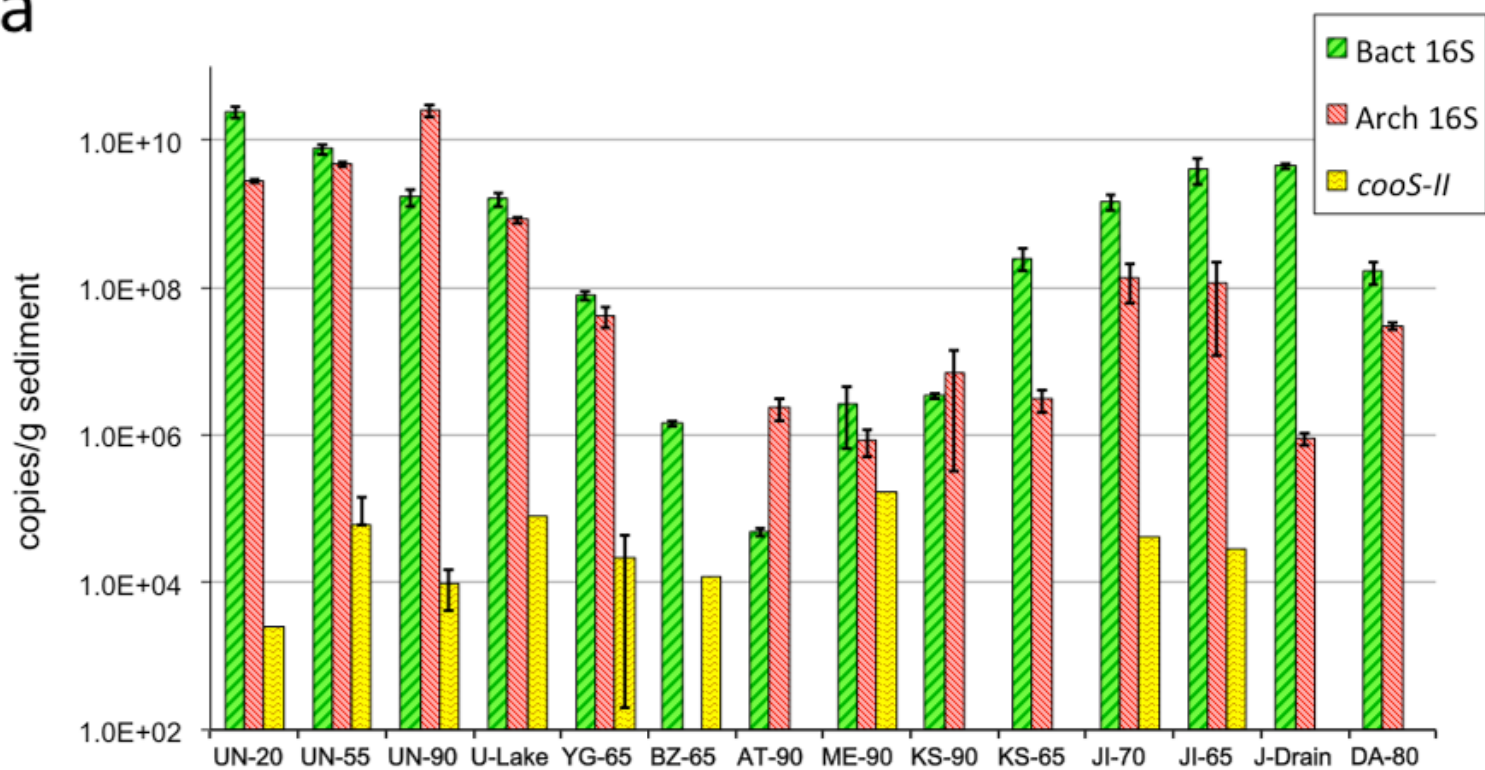


Fig. 4

a



b

