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Туре	Journal Article	
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26 Abstract

27 Carboxydotrophic 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 carboxydotrophic bacteria *Carboxydothermus*

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 *Carboxydothermus* spp. in various hot

35 springs to determine environmental factors that affect CO-oxidizers and to see whether

36 *Carboxydothermus* 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 *Carboxydothermus* spp. did not correlate with the

40 presence of potential microbial CO-oxidation activities; however, *Carboxydothermus*

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.



45

46

47 **One-sentence summary**

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5

- 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

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54 Keywords

55 carboxydotroph, 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 CO_2
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 CO ₂ , 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 *Carboxydothermus* is one of the most
96 studied CO-oxidizing anaerobes. Five species of the genus *Carboxydothermus* 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 Caboxydothermus 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 Carboxydothermus
109	pertinax was previously isolated (Yoneda et al., 2013a). Considering that the optimal
110	growth pH of Carboxydothermus spp. is moderately acidic or neutral, the abundance of
111	these bacteria is intriguingly high. This led us to further question whether
112	Carboxydothermus 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 carboxydotrophic 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 Carboxydothermus 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 °C until use. The other samples were refrigerated at
141	4 °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% $H_2/95\%$ N_2 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 themophilic 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 *Carboxydothermus* 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 *Carboxydothermus cooS-II* cooS2_1442F
- 189 (5'-TGATGCGTCACGGCTTTATGG-3') and cooS2-1606R

190	(5'-CTAAAGCTACTGCCCGGGAGT-3') were designed based on the <i>cooS-II</i> genes of
191	C. hydrogenoformans and C. pertinax (Yoneda et al., 2013a) because no other
192	sequences closely related to Carboxydothermus 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 Carboxydothermus
195	siderophilus, but not Carboxydothermus 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

204Results

205Sample description

At Unagi-onsen, samples designated UG-20, UG-55 and UG-90 were collected from 206

three points with different temperatures in a hot spring pool. These samples were acidic 207

208in 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
- 210site 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 $^{\circ}$ C and 65 $^{\circ}$ 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 carboxydotrophic 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

using environmental parameters as shown in Fig. 3. Sample YG-65 was excluded from
this analysis because an ORP value was not available. There were no apparent
similarities among samples where potential microbial CO-oxidation activities were
observed, though they were quite distinct from negative samples (Fig. 3). There was a
rough tendency of positive samples to be relatively rich in nutrients (TOC and TN) and
have higher ORP values and soil-water contents than negative samples. Among samples
associated with the environmental temperature of approximately 65–71 $^\circ$ C and
circum-neutral pH, the TOC and TN values of JI-65 and JI-70 (positive samples) were
more than double those of KS-65 and BZ-65 (negative samples). ORP values of positive
samples ranged from -49 mV to 487 mV and were higher than those of negative
samples (Table 1). The OPR values were in a range of 'reduced' to 'moderately reduced'
redox condition that appears too oxidative for sulphate reduction and methanogenesis
(DeLaune & Reddy, 2005; Klüpfel et al., 2014). One of the presumed sources of CO in
environments is methanogens and sulphate reducers because some of them produce CO
under laboratory conditions (Conrad & Thauer, 1983; Lupton et al., 1984; Techtmann et
al., 2009). However, in our case, it appeared that the metabolic activity of those
microorganisms did not necessarily correlate with the distribution of CO-oxidizing
anaerobes. Alternatively, CO is produced from volcanic gas and organic substances
(e.g., plant matter and humus). CO production from organic matter is enhanced in the
presence of oxygen, higher temperatures and higher water contents (Conrad & Seller
1985; Tarr et al., 1995; Shade et al., 1999; Hellebrand & Shade, 2008). Our result
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 (high279 ORP values, TOC, TN and soil-water contents).

280

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

- 282 Results of qPCR are shown in Fig. 4. The technical lower detection limit of qPCR for
- the standard bacterial 16S rRNA gene, archaeal 16S rRNA gene and cooS-II were
- 1.0×10^3 , 1.0×10^3 , and 5.0×10^1 copies/µL, respectively. In environmental samples,
- 285 *Carboxydothermus cooS-II* genes were detected in more than two replicates of samples
- of UN-55, UN-90 and YG-65. *Carboxydothermus cooS-II* was also detected in one or
- two replicates from UN-20, U-Lake, BZ-65, ME-90, JI-70 and JI-65. The highest copy
- number was 6.05×10^4 copies/g sediment in UN-55 corresponding to 7.95×10^{-4} % of
- 289 bacterial 16S rRNA. *Carboxydothermus cooS-II* copies were detected from samples
- 290 UN-90 and ME-90, where the environmental temperatures were above growth
- temperature (≥78 °C) of *Carboxydothermus* spp. (Yoneda *et al.*, 2012). On the other
- 292 hand, Carboxydothermus cooS-II was below detection level in sample KS-65 despite its
- 293 environmental temperature being optimal (65 °C) to most *Carboxydothermus* 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
- from the start time of the experiment. *Carboxydothermus cooS-II* was detected in more
- than two replicates in CO-incubated samples UN-20, UN-55, UN-90, KS-90 and KS-65.
- 298 Interestingly, *Carboxydothermus 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, Carboxydothermus cooS-II was not detected in CO-incubated JI-70

- 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
- 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
- remaining four sequences, all from UN-90, showed 94% similarity to both
- 315 Carboxydothermus 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 *Carboxydothermus* 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

and abundance of carboxydotrophic anaerobic thermophiles in various hydrothermal

328 environments in relation to environmental factors. Most carboxydotrophic anaerobic

329 thermophiles require strictly anaerobic, reduced conditions for carboxydotrophic growth.

However, most of them are not obligate carboxydotrophs and are able to utilize organic

331 substrate and electron acceptors such as ferric iron, AQDS

332 (anthraquione-2,6-disulfonate) and sulphur compounds (Sokolova *et al.*, 2009;

333 Sokolova & Lebedinsky, 2013). Those alternative metabolic pathways may enable

334 carboxydotrophic 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

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

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

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.,

344Moorella thermoacetica) and lactate (e.g., C. hydrogenoformans) (Fröstl et al., 1996; 345Henstra 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, 351suggesting that CO-oxidizers may have a niche in an intermediate zone of aerobic and 352anaerobic environments thanks to their utilization of electron acceptors such as nitrate, 353 Fe(III) and fumarate. 354The abundance of *Carboxydothermus* 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 Carboxydothermus spp. were detected (Fig. 2, Fig. 3 and Fig. 4a). Moreover, no significant changes in the Carboxdothermus spp. population 357358 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 *Carboxydothermus* 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 *Carboxydothermus* 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. Carboxydothermus 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 <i>cooS-I</i> , <i>cooS-II</i> and <i>cooS-IV</i> 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	
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- 505

1	Figure	legends
-	I Iguite	iegenas

2	Fig. 1. Sampling	sites in (a) the	Kagoshima Pref.	in southern Kyushu	Island and (b)
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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 sample	oles
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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 *Carboxydothermus 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. 2.

CO concentration (µmol/ mL in head space)







Fig. 4

