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1 ***Hydrogenimonas urashimensis* sp. nov., a hydrogen-oxidizing**
2 **chemolithoautotroph isolated from a deep-sea hydrothermal vent in the**
3 **Southern Mariana Trough**

4
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19 **Keywords:** “*Campylobacteria*”; *Epsilonproteobacteria*; *Hydrogenimonas*; Deep-sea hydrothermal
20 vent, Thermophile

21

22 **Abbreviations**

23 ANI, average nucleotide identity; *in silico* DDH, *in silico* DNA-DNA hybridization; AAI, average
24 amino acid identity; SGC, single-copy core gene

25

26 **Nucleotide sequence data**

27 The 16S rRNA gene sequence and the genome sequence of strain SSM-sur55^T were deposited in
28 DDBJ/EMBL/GenBank under LC469141 and AP023212, respectively.

29

30 **Strain repositories**

31 JCM 19825 = KCTC 15926

32 **Abstract**

33 A novel thermophilic bacterium, strain SSM-sur55^T, was isolated from a chimney structure
34 at the Urashima site on the Southern Mariana Trough in the Pacific Ocean. Growth was observed at
35 temperatures between 25 and 60°C (optimum, 55°C; 180 min doubling time), at pH values between
36 5.3 and 7.2 (optimum, pH 5.9) and in the presence of between 1.6 and 5.6% (w/v) NaCl (optimum,
37 3.2%). The isolate used molecular hydrogen as its sole energy source, carbon dioxide as its sole
38 carbon source, ammonium as its sole nitrogen source, and elemental sulfur as its sole sulfur source.
39 Thiosulfate, molecular oxygen (0.1%, v/v) or elemental sulfur was utilized as its sole electron
40 acceptor. Phylogenetic analysis based on 16S rRNA gene sequences indicated that strain SSM-sur55^T
41 belonged to the genus *Hydrogenimonas* of the class “*Campylobacteria*”, and its closest relative was
42 *Hydrogenimonas thermophila* EP1-55-1%^T (94.9%). On the basis of the phylogenetic, physiological
43 and molecular characteristics, strain SSM-sur55^T represents a novel species within the genus
44 *Hydrogenimonas*, for which the name *Hydrogenimonas urashimensis* sp. nov. is proposed, with the
45 type strain SSM-sur55^T (JCM 19825 = KCTC 15926).

46 **Introduction**

47 The class “*Campylobacteria*” is one of the ecologically important chemolithoautotrophic
48 primary producers in deep-sea hydrothermal systems, where they are often found to be dominant in
49 bacterial communities [1, 2]. Species of the class “*Campylobacteria*” from hydrothermal
50 environments with validly published name fall into two mesophilic (*Thiovulaceae* and *Sulfurovaceae*)
51 and three thermophilic (*Hydrogenimonadaceae*, *Nitratiruptoraceae*, and *Nautiliaceae*) families [3, 4].
52 *Hydrogenimonadaceae* is presently a monogeneric family comprised solely of a validly described
53 hydrogen-oxidizing thermophile, *Hydrogenimonas thermophila*, which was isolated from a deep-sea
54 hydrothermal vent in the Central Indian Ridge [5]. Although sequences affiliated to the genus
55 *Hydrogenimonas* have been detected from worldwide hydrothermal systems [6-9], physiological

56 diversity and the ecological role of this group in hydrothermal ecosystems have not been fully
57 elucidated yet due to the lack of additional cultivated strains within this genus.

58 The Southern Mariana Trough is a spreading back-arc basin, where geochemically diverse
59 hydrothermal sites have been discovered [10]. The geochemical diversity would provide diverse
60 habitats and a great metabolic potential of microbial communities [11]. In some hydrothermal sites,
61 sulfur- and/or hydrogen-oxidizing mesophilic and thermophilic “*Campylobacteria*” such as the
62 genera *Sulfurovum* and *Lebetimonas* have been detected as dominant in bacterial lineages [11, 12]. A
63 recent study successfully has enriched a member of the genus *Hydrogenimonas* that possessed the
64 potential of global warming mitigation [13]. Although these previous studies imply that deep-sea
65 hydrothermal environments in the Southern Mariana Trough are expected to be good candidates for
66 obtaining new microbial species, “*Campylobacteria*” species from this area has never been validly
67 described. Here, we succeeded to isolate a novel strain SSM-sur55^T from the hydrothermal field of
68 the Urashima site in the Southern Mariana Trough, and determined its physiological and phylogenetic
69 characteristics.

70 **Methods**

71 **Sample collection, enrichment and purification**

72 Sample collection and subsampling procedures were carried out as described previously [13].
73 The chimney structure was taken with *R/V Yokosuka* and *DSV Shinkai 6500* from the Baltan chimney
74 at the Urashima site (12°55.3’N, 143°38.9’E) in the Southern Mariana Trough in 2010. The physical
75 and chemical characterization of the vent emissions has been described elsewhere [10]. After retrieval
76 on board, the sample was anaerobically suspended in 20 ml sterilized MJ synthetic sea water [14, 15]
77 containing 0.05% (w/v) sodium sulfide in a 100 ml glass bottle (Schott Glaswerke) tightly sealed with
78 a butyl-rubber cap under a gas phase of 100% N₂ (0.1 MPa). The suspended slurry was used to
79 inoculate a series of media, including MMJHS medium [16], under a gas phase of H₂/CO₂/O₂
80 (80:19:1; 0.3 MPa), and the cultures were incubated at 55°C in a dry oven. Growth of thermophiles

81 was observed in MMJHS medium after 2 days incubation at 55°C. Enrichment cultures at 55°C
82 contained rod-shaped cells. A pure culture was obtained by using the dilution-to-extinction technique
83 at 55°C with the same medium as that used for the enrichment [17]. The culture in the tube showing
84 growth at the highest dilution was designated strain SSM-sur55^T. Purity was confirmed with a routine
85 microscopic examination and by repeated partial sequencing of the 16S rRNA gene using several
86 PCR primers.

87 **Growth characteristics**

88 Growth of strain SSM-sur55^T was measured by direct cell counting after staining with 4',6-
89 diamidino-2-phenylindole using the ZEISS Axiophot microscope (Carl Zeiss Co., Oberkochen,
90 Germany). To determine optimum temperature, pH and NaCl concentrations, cultures were prepared
91 in a 3 ml MMJHS medium under various conditions.

92 To determine combinations of a single electron donor and acceptor, the isolate was tested
93 using MJ synthetic seawater containing 0.1% (w/v) NaHCO₃ as the basal medium. For testing the
94 growth on hydrogen as an electron donor, H₂/CO₂ (80:20) was used as the gas phase. To examine the
95 growth on thiosulfate (0.1%, w/v) or elemental sulfur (S⁰) (1%, w/v) as an electron donor, N₂/CO₂
96 (80:20) was used as the gas phase. Nitrate (0.1%, w/v), thiosulfate (0.1%, w/v), nitrite (0.1 and 0.01%,
97 w/v) sulfite (0.01%, w/v), elemental sulfur (1%, w/v) or molecular oxygen (0.1 and 1%, v/v) were
98 tested as sole electron acceptors. The presence or absence of the cell growth was determined by
99 microscopic observation.

100 For testing heterotrophic growth of strain SSM-sur55^T, experiments were conducted using
101 MMJHS medium without NaHCO₃ under a gas phase of 100% H₂ (0.3 MPa). Each of the following
102 potential carbon sources was tested: yeast extract, peptone, tryptone, casamino acids, D(+)-glucose,
103 galactose, sucrose, fructose, lactose, maltose, starch (all 0.2%, w/v), formate, acetate, glycerol, citrate,
104 tartrate, malate, succinate, propionate, lactate, oxalate, pyruvate (all 10 mM), methanol (0.05%, v/v),
105 ethanol (0.1%, v/v) and 2-propanol (0.2%, v/v).

106 Potential nitrogen and sulfur sources required for the growth of the isolate were tested. To

107 determine the nitrogen sources for growth of strain SSM-sur55^T, NH₄Cl (0.025%, w/v), NaNO₃
108 (0.1%, w/v) and NaNO₂ (0.1 and 0.01%, w/v) were added in MMJHS medium lacking all nitrogen
109 sources, under a H₂/CO₂ (80:20) gas phase (0.3 MPa). In addition, utilization of N₂ was examined
110 under a H₂/N₂/CO₂ (60:20:20) gas phase.

111 To examine the sulfur sources for the growth of the isolate, sulfate (0.42%, w/v), thiosulfate
112 (0.1%, w/v), sulfite (0.01%, w/v) and elemental sulfur (1%, w/v) were examined in MMJHS medium
113 in which sulfur compounds were removed and replaced with the chloride salts under an H₂/CO₂
114 (80:20) gas phase (0.3 MPa).

115 **Morphological observation**

116 The cell morphology of strain SSM-sur55^T was observed by phase-contrast microscopy with
117 a Zeiss Axiophot microscope and a JEM-1011 transmission electron microscope (JEOL). Cells grown
118 in MMJHS medium at 55°C in the mid-exponential phase of growth were used for microscopic
119 observation. Cells were fixed with 2.5% glutaraldehyde, followed by several washes with 10mM
120 PIPES buffer before negative staining with EM Stainer (Nisshin EM, Tokyo, Japan).

121 **Genome sequencing and assembly**

122 Genomic DNA of strain SSM-sur55^T was extracted from the cells grown in MMJHS medium
123 with Wizard genomic DNA purification kit (Promega, Madison, Wisconsin, USA) according to the
124 protocol provided by the manufacturer. The genome was sequenced using both Oxford Nanopore
125 Technology (ONT) and Illumina platforms. For the ONT sequencing, library was prepared using the
126 Rapid Barcoding Sequence kit (Oxford Nanopore Technologies, Oxford, UK) according to the
127 standard protocol provided by the manufacturer. The constructed library was loaded into the FlowCell
128 (FLO-MIN106) on a MinION device and a 48-hour sequencing run with MinKNOW 1.15.4 software
129 was performed. After basecalling ONT reads with Guppy v1.1 (Oxford Nanopore Technologies) with
130 following settings: --qscore_filtering and --calib_detect, basecalled reads were binned with
131 Deepbinner [18]. For the Illumina sequencing, paired-end libraries were generated using Nextera

132 library preparation methods. Genome sequencing was then performed on an Illumina MiSeq platform
133 (2x300 bp paired-end). ONT and Illumina reads were then assembled using Unicycler version 0.4.7
134 [19]. The genome was annotated using DFAST [20]. KEGG pathway annotation and mapping were
135 performed with BlastKOALA [21] and KEGG Mapper [22], respectively.

136 **Phylogeny based on 16S rRNA gene sequences**

137 Phylogenetic trees were constructed by using the almost full-length sequences of the 16S
138 rRNA genes. The 16S rRNA gene of strain SSM-sur55^T was amplified by PCR using primers Eubac
139 27F and 1492R [23]. The nearly complete 16S rRNA gene sequence (1,422 bp) was obtained by direct
140 sequencing of both strands. The 16S rRNA gene sequence was analyzed using BLAST search
141 algorithm [24]. To conduct the phylogenetic analysis of the strain, the other “*Campylobacteria*”
142 sequences were retrieved and aligned using Silva database [25] and Silva Incremental Aligner v1.2.11
143 [26], respectively. A phylogenetic tree was constructed by neighbor-joining algorithm [27] with the
144 MEGA 7.0.21 software [28] using 1,273 base pairs. Bootstrap analysis was done using 1,000
145 replications to provide confidence estimates for the phylogenetic tree topologies.

146 **Pangenomic, phylogenomic, and genome sequence similarities analyses**

147 Phylogenomic tree construction and pangenomic analysis were carried out with 16
148 thermophilic campylobacterial genomes using the anvi’o v5.5 [29]. The phylogenomics workflow
149 (<http://merenlab.org/2017/06/07/phylogenomics/>) was followed to infer evolutionary associations
150 between genomes. Briefly, the fasta files containing nucleotide sequences of genomes was used for
151 generating the database of each genome (anvi-script-FASTA-to-contigs-db). We then identified an
152 HMM profiles (anvi-get-sequences-for-hmm-hits) and extracted 139 single-copy core genes (SCGs)
153 proposed by Campbell et al. [30]. The amino acid sequences of 139 SCGs were concatenated in a
154 fasta file (anvi-get-sequences-for-hmm-hits) and a phylogenomic tree was constructed (anvi-gen-
155 phylogenomic-tree) using FastTree [31]. For pangenome analysis, we generated a storage database
156 (anvi-gen-genomes-storage) from the genomes of thermophilic relatives and then computed the

157 pangenome (anvi-pan-genome). COG-annotated strains-specific gene clusters were exported from
158 pangenome (anvi-summarize) for the gene functional analysis. Average nucleotide identity (ANI)
159 between genomes was calculated with PyANI (anvi-compute-ani)
160 (<https://github.com/widdowquinn/pyani/releases/tag/v0.1.2>). The pangenome was visualized with
161 anvi-display-pan. Average amino acid identity (AAI) and *in silico* DNA-DNA hybridization values
162 were calculated using aai.rb script within the enveomics collection [32] and Genome-to-Genome
163 Distance Calculator 2.1 [33], respectively.

164 **Results and Discussion**

165 **Growth characteristics**

166 With MMJHS medium, strain SSM-sur55^T grew at temperature between 25°C and 60°C,
167 showing optimum growth at 55°C. No growth was observed below 20°C or above 65°C. Growth
168 occurred between pH 5.3 and 7.2, with optimum growth at pH 5.9. No growth was observed below
169 pH 4.9 or above pH 7.7. Growth was observed between 1.6 and 5.6% (w/v) NaCl concentrations,
170 with optimum growth at 3.2%. No growth was detected below 0.8% or above 7.2% NaCl
171 concentration levels (Fig. S1). These growth characteristics of strain SSM-sur55^T were similar to
172 those of *Hydrogenimonas thermophila* EP1-55-1%^T [5] (Table 1).

173 Strain SSM-sur55^T was unable to utilize any electron donors other than H₂. Thiosulfate
174 (0.1%, w/v), elemental sulfur (1%, w/v) and molecular oxygen (0.1%, v/v) were able to serve as its
175 sole electron acceptors. None of the organic compounds sustained growth of strain SSM-sur55^T as
176 energy or carbon sources. These results indicated that strain SSM-sur55^T was a strictly hydrogen-
177 oxidizing thermophilic chemolithoautotroph. Strain SSM-sur55^T utilized ammonium as its sole
178 nitrogen source and did not utilize nitrate, nitrite and molecular nitrogen. Strain SSM-sur55^T utilized
179 elemental sulfur as its sole sulfur source. Utilization of thiosulfate, sulfate and sulfite were not
180 observed.

181 **Cell morphology**

182 Cells of strain SSM-sur55^T were Gram-stain negative and rod-shaped (Fig. S2). Flagella
183 were not observed although motility was confirmed under the light microscopic observation (Fig. S2).
184 A small percentage of cells grown in MMJHS medium exhibited spherical shape, as reported for *H.*
185 *thermophila* EP1-55-1%^T [5]

186 **Phylogenetic analysis based on 16S rRNA gene sequences**

187 With a nearly full length 16S rRNA gene sequence of strain SSM-sur55^T as a query in
188 BLAST search, the highest similarity with *Hydrogenimonas* sp. RS_Sur55-1 (96.2%),
189 *Hydrogenimonas* sp. MAG (95.9%), and *H. thermophila* EP1-55-1%^T (94.9%) were obtained. More
190 closely related sequence was retrieved from an environmental clone from the detritus from tubeworm
191 in the East Pacific Rise (LF8GH2b132, 96.7% 16S rRNA gene sequence similarity). The phylogenetic
192 tree based on 16S rRNA gene sequences showed that strain SSM-sur55^T formed a clade with *H.*
193 *thermophila* (Fig. 1). These results indicated that strain SSM-sur55^T is a novel species belonging to
194 the genus *Hydrogenimonas*.

195 **Genome properties and genome relatedness**

196 The complete genome of strain SSM-sur55^T was reconstructed. The genome size was
197 2,297,889 bp with an average G + C content was 52.8% (Table 1). In total, 2,270 coding sequence
198 regions, 48 tRNA genes, and 3 set of rRNA genes were respectively predicted. The G + C content of
199 strain SSM-sur55^T was higher than that of previously sequenced genome of *H. thermophila* (33.5%)
200 and comparable with that of *Hydrogenimonas* sp. MAG (50.2%) [13].

201 Genome analysis of the strain SSM-sur55^T supported its phenotypic characteristics. The H₂-
202 uptake hydrogenase gene cluster for H₂ oxidation was present in strain SSM-sur55^T. The ability to
203 utilize oxygen as a sole electron acceptor was ensured by the presence of four genes encoding the
204 CcoNOQP proteins, which constitutes a *cbb*₃-type terminal cytochrome oxidase. A gene encoding
205 putative thiosulfate sulfurtransferase was found in its genome, supporting the ability to utilize
206 thiosulfate as a sole electron acceptor. Although strain SSM-sur55^T showed no ability to use nitrate

207 and N₂O as a sole electron acceptor under hydrogen-oxidizing condition, the complete set of
208 denitrification genes such as *nap*, *nir*, *nor*, and *nos* was found on its genome, that might allow to the
209 strain to contribute to the nitrogen cycle as a denitrifier in deep-sea hydrothermal vent environments.

210 The ANI and *in silico* DDH values between strain SSM-sur55^T and *H. thermophila* EP1-55-
211 1%^T were found to be 72.7% and 17.4%, respectively, well below the species threshold (95-96% and
212 70%, respectively) [34]. Strain SSM-sur55^T shared AAI values of 71.4% with *H. thermophila* EP1-
213 55-1%^T that considerably above the recently proposed AAI genus threshold among
214 “*Campylobacterota*” (60-62%) (35). These results also support the proposal that the isolate is a novel
215 species within the genus *Hydrogenimonas*.

216

217

Table 1. Comparison of physiological characteristics of SSM-sur55^T with species of “*Campylobacteria*” from deep-sea vents.

Characteristics	<i>Hydrogenimonas</i> sp. SSM-sur55 ^T (This study)	<i>Hydrogenimonas</i> <i>thermophila</i> EP1-55-1% ^T [5]	<i>Nitratiruptor</i> <i>tergarcus</i> MI55-1 ^T [36]	“ <i>Nitrosophilus labii</i> ” HRV44 ^T [35, 37]	“ <i>Nitrosophilus alvini</i> ” EPR55-1 ^T [35]
Origin	Southern Mariana Trough	Central Indian Ridge	Mid-Okinawa Trough	Mid-Okinawa Trough	East Pacific Rise
Temperature range (°C)	25-60	35-65	40-55	45-60	50-60
Temperature optimum (°C)	55	55	55	53	60
pH range	5.3-7.2	4.9-7.2	5.4-6.9	5.4-6.4	5.4-8.6
pH optimum	5.9	5.9	6.4	6.0	6.6
NaCl range (% w/v)	1.6-5.6	1.6-5.6	1.5-4.0	2.0-4.0	2.4-3.2
NaCl optimum (% w/v)	3.2	3.2	2.5	2.5	2.4
Electron donors	H ₂	H ₂	H ₂	H ₂	H ₂
Electron acceptors	S ₂ O ₃ ²⁻ , O ₂ , S ⁰	NO ₃ ⁻ , O ₂ , S ⁰	NO ₃ ⁻ , O ₂ , S ^{0†}	NO ₃ ⁻ , N ₂ O, O ₂ , S ⁰	NO ₃ ⁻ , N ₂ O, S ₂ O ₃ ²⁻ , O ₂ , S ⁰
Carbon sources other than CO ₂	-	-	-	-	-
Nitrogen sources	NH ₄ ⁺	NO ₃ ⁻ , NH ₄ ⁺	NO ₃ ⁻ , NH ₄ ⁺	NO ₃ ⁻ , NH ₄ ⁺	NH ₄ ⁺
DNA G+C content (%)	52.8	33.5	36.9	33.4	37.7

-, negative; ND, not determined.

†S⁰ did not serve as a sole electron acceptor to support growth.

220 Pangenomic and phylogenomic analyses

221 The pangenome of thermophilic campylobacteria revealed a total of 8,272 gene clusters
222 comprising 35,198 genes (Fig. 2). 557 gene clusters were represented core genome, and 468 of those
223 were SCGs. 577 gene clusters were found to be unique to the strain SSM-sur55^T. Although we
224 searched gene clusters enriched in *Hydrogenimonas* genomes, the statistically significant trends were
225 not obtained (corrected $p \geq 0.25$), possibly due to the small number of genomes in each genus. The
226 COG functional annotation of the strain-specific gene clusters showed that relatively abundance of
227 genes related to energy production and conversion (C) in SSM-sur55^T is higher (9.6% in without
228 poorly characterized categories) than those in *H. thermophila* EP1-55-1%^T and *Hydrogenimonas* sp.
229 BAL40 [13] (Fig. S3). Further pangenomic analyses with additional genome sequences could provide
230 us a better understanding on genus-specific genomic traits of thermophilic “*Campylobacteria*”.

231 Conclusion

232 On the basis of physiological and molecular characteristics of strain SSM-sur55^T, the strain
233 is considered to represent a novel species in the genus *Hydrogenimonas*, for which the name
234 *Hydrogenimonas urashimensis* sp. nov. is proposed. Its Protologue description is listed in Table 2.

235

236 **Table 2. Protologue description of *Hydrogenimonas urashimensis* sp. nov.**

Genus name	<i>Hydrogenimonas</i>
Species name	<i>Hydrogenimonas urashimensis</i>
Specific epithet	<i>urashimensis</i>
Species status	sp. nov.
Species etymology	u.ra'shi men'sis. N.L. fem. adj. <i>urashimensis</i> pertaining to the Urashima deep-sea hydrothermal site in the Southern Mariana Trough
Description of the new taxon and diagnostic traits	Cells are gram-negative, motile, and rod-shaped. The temperature range for growth is 25-60°C (optimum, 55°C; 180 min doubling time). The pH range for growth is 5.3-7.2 (optimum, pH5.9). NaCl in the concentration range 1.6-5.6% (w/v) is an absolute growth requirement; optimum growth occurs at 3.2%. Strain SSM-sur55 ^T is hydrogen-oxidizing, facultatively anaerobic and chemolithoautotroph with molecular hydrogen as its sole electron donor and with thiosulfate, molecular

	oxygen or elemental sulfur as its sole electron acceptors. Ammonium is utilized as its sole nitrogen source. Elemental sulfur is utilized as its sole sulfur source.
Country of origin	USA
Region of origin	The Southern Mariana Trough
Date of isolation (dd/mm/yyyy)	17/12/2010
Source of isolation	deep-sea hydrothermal vent
Sampling date (dd/mm/yyyy)	23/08/2010
Latitude (xx°xx'xx"N/S)	12°55'18"N
Longitude (xx°xx'xx"E/W)	143°38'54"E
Altitude (meters above sea level)	-2,922 m
16S rRNA gene accession nr.	LC469141
Genome accession number [RefSeq; EMBL; ...]	GenBank = AP023212
Genome status	Complete
Genome size	2,297 kbp
GC mol%	52.8 (based on the complete genome sequence)
Number of strains in study	1
Source of isolation of non-type strains	not applicable
Information related to the Nagoya Protocol	not applicable
Designation of the Type Strain	SSM-sur55 ^T
Strain Collection Numbers	JCM 19825 ^T = KCTC 15926 ^T

237

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354 **Figure Legends**

355 **Fig. 1. Phylogenetic tree based on 16S rRNA gene sequences.** Phylogenetic tree of the members of
356 thermophilic “*Campylobacteria*”, inferred by the neighbor-joining algorithm using 1,229 homologous
357 sequence positions. Numbers at branches are bootstrap values (%) based on 1,000 replicates.

358

359 **Fig. 2. Pangenome of thermophilic “*Campylobacteria*”.** Pangenomic analysis of 16 sequenced
360 thermophilic “*Campylobacteria*” revealing 468 SCGs among 8,272 total gene clusters along with
361 their distribution and ANI.

362

363 **Supplementary**

364 **Fig. S1. Growth rates of strain SSM-sur55^T.** Growth rates of temperature (a), pH (b) and NaCl
365 concentration (c) in MMJHS medium.

366

367 **Fig. S2. Transmission electron micrograph of a rod-shaped cell of strain SSM-sur55^T.** Bar, 1
368 μm .

369

370 **Fig. S3. The COG category annotation of strain-specific gene clusters.** All COGs (a) and without poorly
371 characterized categories, “R”, “S”, and “-”.

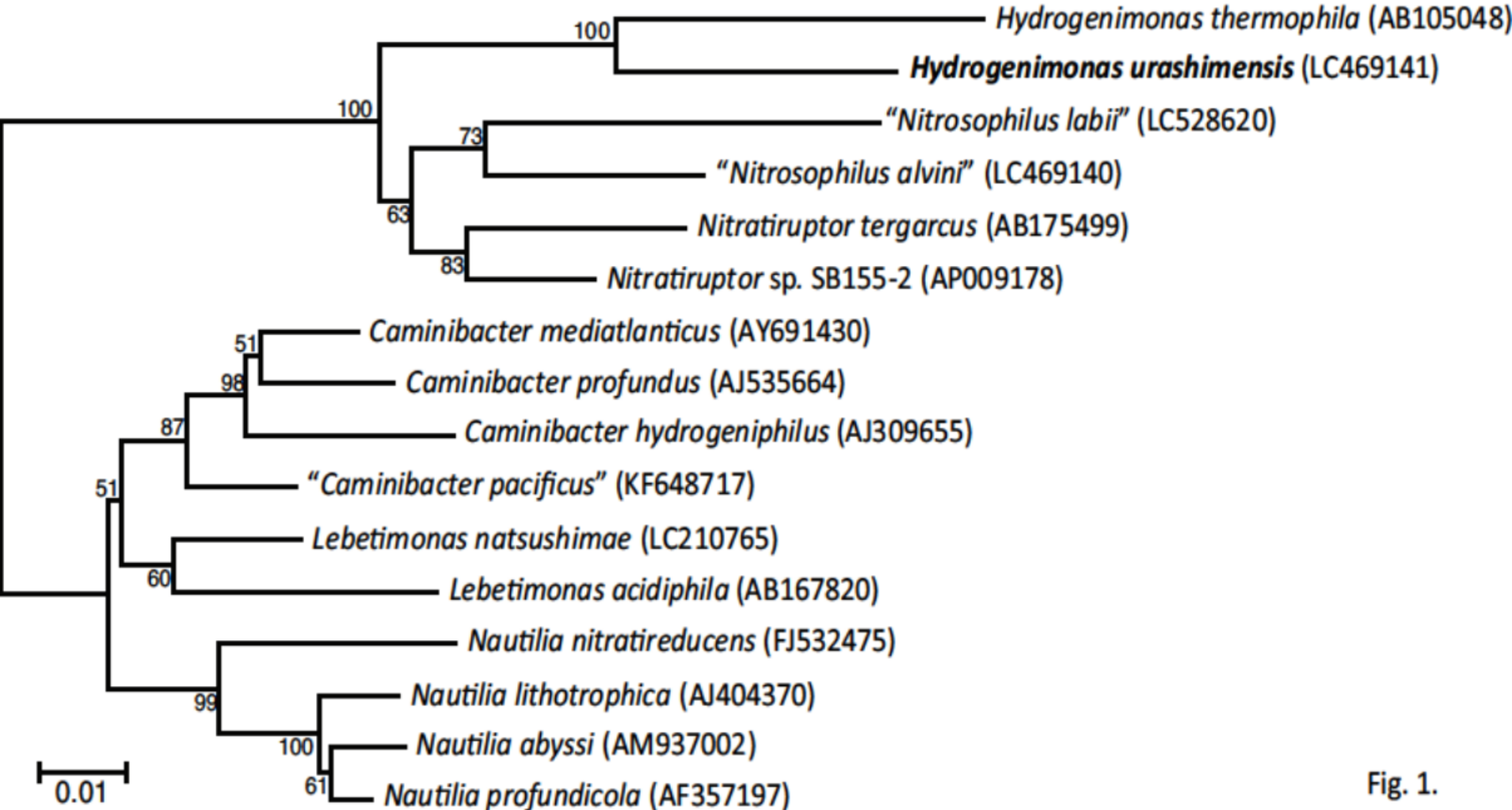


Fig. 1.

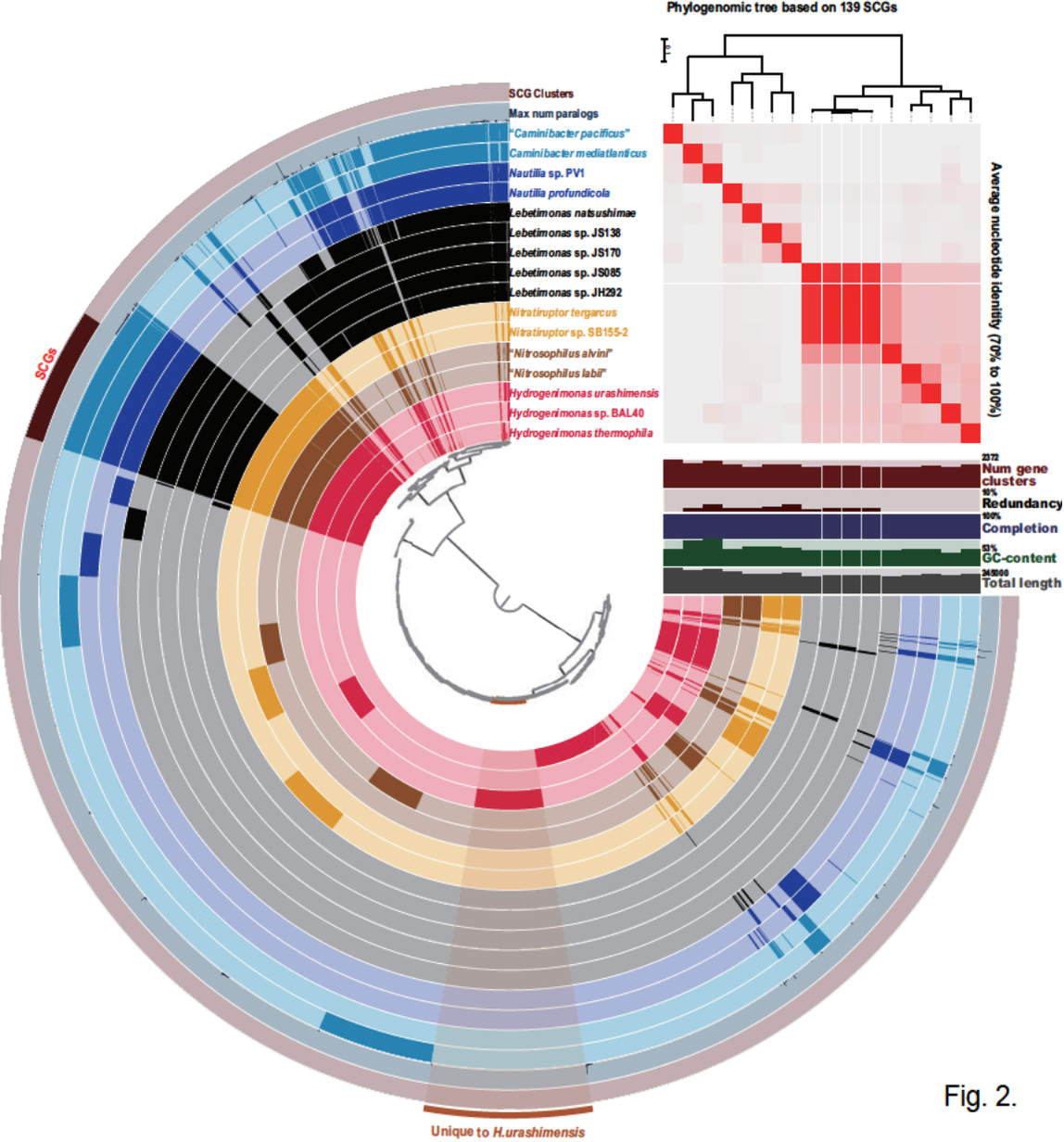


Fig. 2.

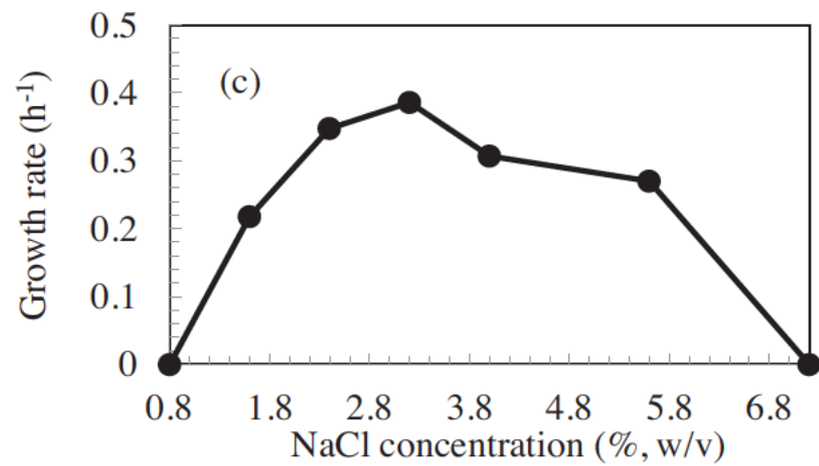
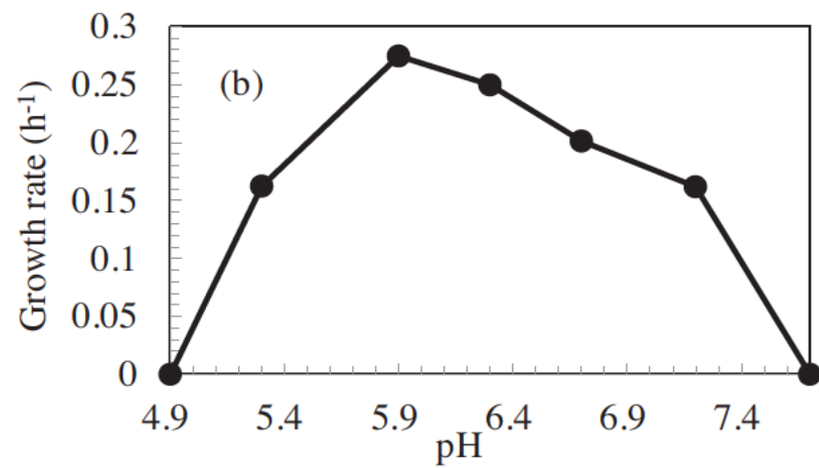
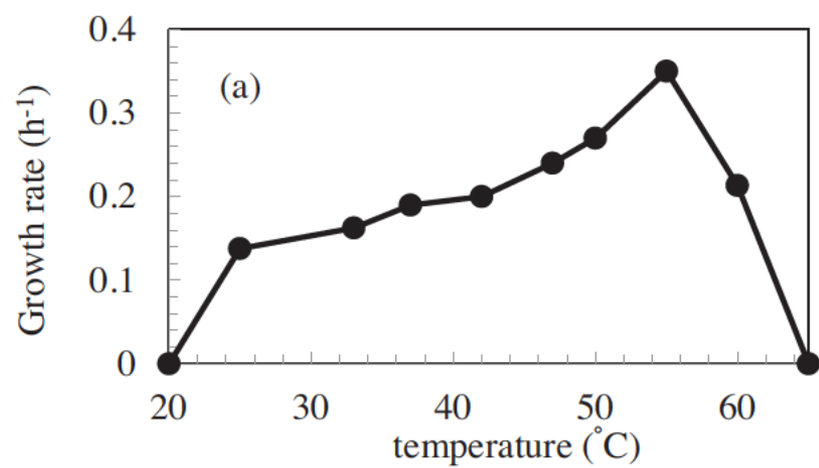


Fig. S1.

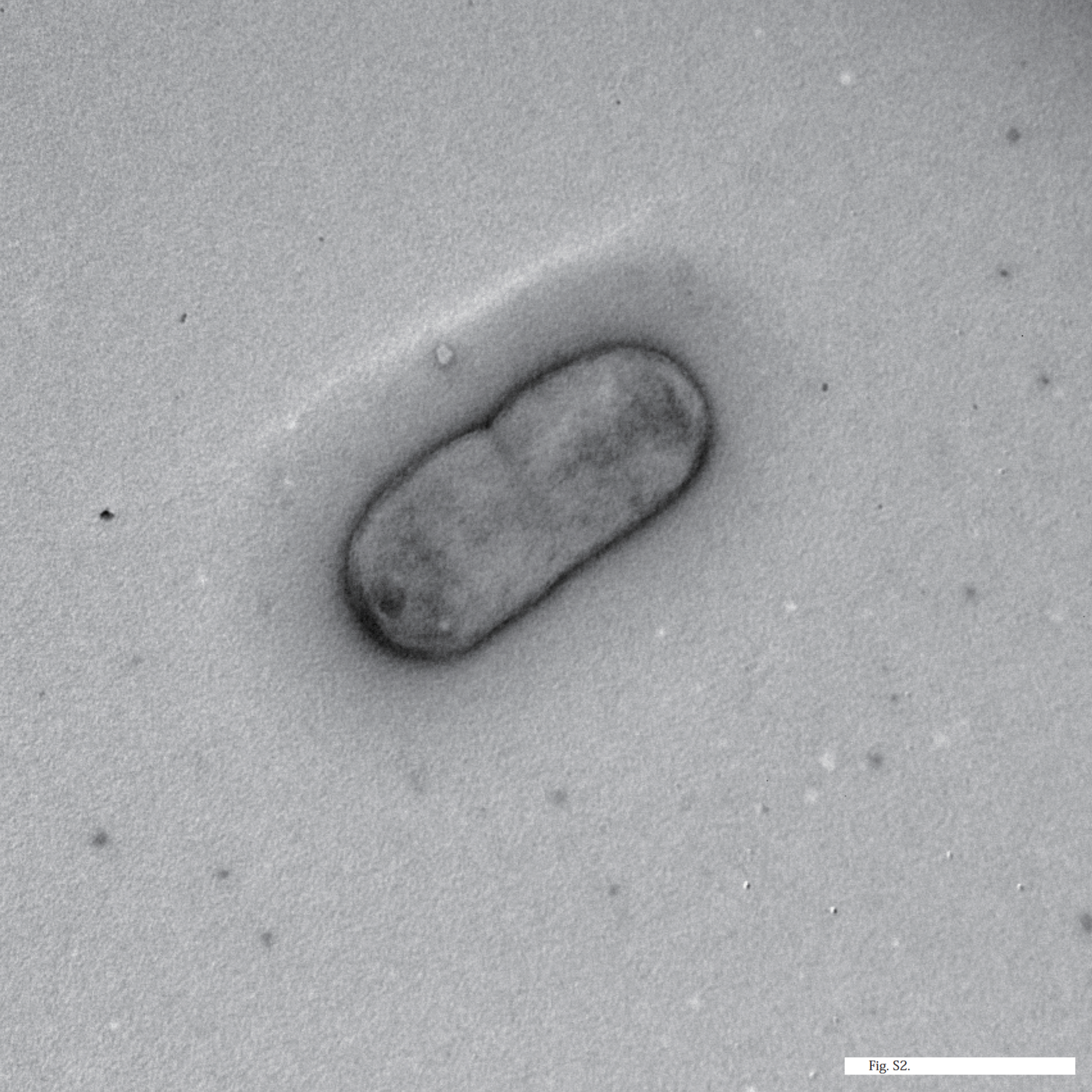


Fig. S2.

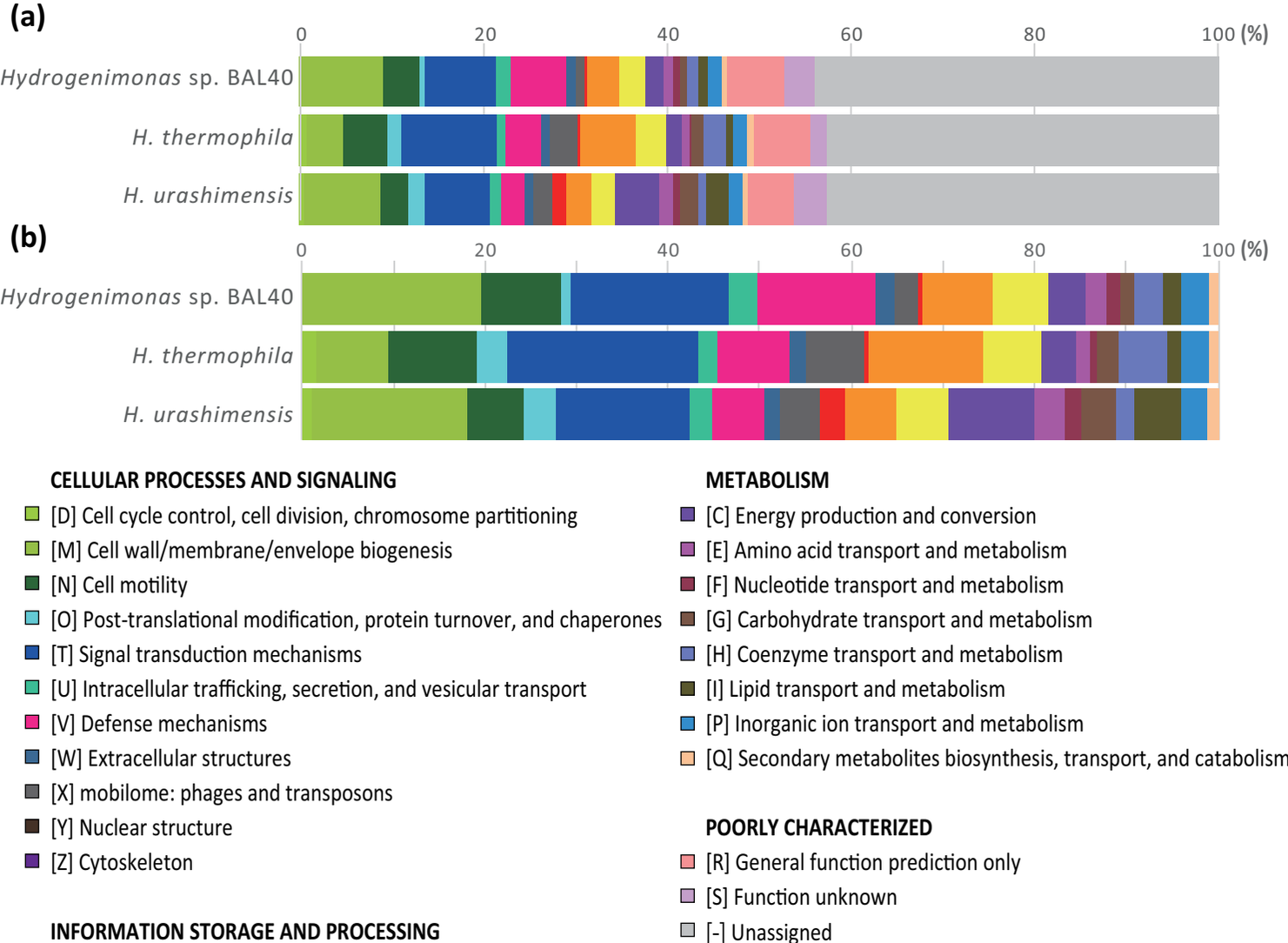


Fig. S3.