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Paludisphaera borealis gen. nov., sp. nov., a hydrolytic planctomycete from northern wetlands, and the proposal of Isosphaeraceae fam. nov.
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Abstract:	Two isolates of aerobic, budding, pink-pigmented bacteria, designated strains PX4T and PT1, were isolated from a boreal Sphagnum peat bog and a forested tundra wetland. Cells of these strains were non-motile spheres that occurred singly or in short chains. Novel isolates were capable of growth at pH values between 3.5 and 6.5 (optimum at pH 5.0-5.5) and at temperatures between 6 and 30C (optimum at 15-25C). Most sugars and a number of polysaccharides including pectin, xylan, lichenan and Phytigel were used as growth substrates. The major fatty acids were C16:0, C18:1w9 and C18:0; the major polar lipids were phosphocholine and trimethylornithine. The quinone was MK-6, and the G+C content of the DNA was 66 mol%. Strains PX4T and PT1 were members of the order Planctomycetales and displayed 93-94% 16S rRNA gene sequence similarity to Aquisphaera giovannonii, 91-92% to Singulisphaera species and 90-91% to Isosphaera pallida. Two novel strains, however, differed from members of these genera by cell morphology, substrate utilization pattern and a number of physiological characteristics. Based on these data, the novel isolates should be considered as representing a novel genus and species of planctomycetes, for which the name Paludisphaera borealis gen. nov., sp. nov. is proposed. The type strain is PX4T (= DSM 28747T = VKM B-2904T). We also suggest to establish a novel family, the Isosphaeraceae fam. nov., to accommodate stalk-free planctomycetes with spherical cells, which can be assembled in short chains, long filaments or shapeless aggregates. This family includes the genera Isosphaera, Aquisphaera, Singulisphaera, and Paludisphaera.

1 ***Paludisphaera borealis* gen. nov., sp. nov., a hydrolytic planctomycete from**
2 **northern wetlands, and the proposal of *Isosphaeraceae* fam. nov.**

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22 The GenBank/EMBL/DDBJ accession number for the 16S rRNA gene sequences of

23 *Paludisphaera borealis* strains PX4^T and PT1 are KF467528 and KT372165, respectively.

24 **ABSTRACT**

25 **Two isolates of aerobic, budding, pink-pigmented bacteria, designated strains PX4^T and**
26 **PT1, were isolated from a boreal *Sphagnum* peat bog and a forested tundra wetland. Cells**
27 **of these strains were non-motile spheres that occurred singly or in short chains. Novel**
28 **isolates were capable of growth at pH values between 3.5 and 6.5 (optimum at pH 5.0-5.5)**
29 **and at temperatures between 6 and 30°C (optimum at 15-25°C). Most sugars and a number**
30 **of polysaccharides including pectin, xylan, lichenan and Phytigel were used as growth**
31 **substrates. The major fatty acids were C_{16:0}, C_{18:1ω9} and C_{18:0}; the major polar lipids were**
32 **phosphocholine and trimethylornithine. The quinone was MK-6, and the G+C content of**
33 **the DNA was 66 mol%. Strains PX4^T and PT1 were members of the order *Planctomycetales***
34 **and displayed 93-94% 16S rRNA gene sequence similarity to *Aquisphaera giovannonii*, 91-**
35 **92% to *Singulisphaera* species and 90-91% to *Isosphaera pallida*. The two novel strains,**
36 **however, differed from members of these genera by cell morphology, substrate utilization**
37 **pattern and a number of physiological characteristics. Based on these data, the novel**
38 **isolates should be considered as representing a novel genus and species of planctomycetes,**
39 **for which the name *Paludisphaera borealis* gen. nov., sp. nov, is proposed. The type strain is**
40 **PX4^T (= DSM 28747^T = VKM B-2904^T). We also suggest to establish a novel family, the**
41 ***Isosphaeraceae* fam. nov., to accommodate stalk-free planctomycetes with spherical cells,**
42 **which can be assembled in short chains, long filaments or shapeless aggregates. This family**
43 **includes the genera *Isosphaera*, *Aquisphaera*, *Singulisphaera*, and *Paludisphaera*.**

44

45 **Keywords:** the phylum *Planctomycetes*, *Paludisphaera borealis* gen. nov., sp. nov.,

46 *Isosphaeraceae* fam. nov.

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48

49 Planctomycetes of the phylogenetic lineage defined by the genus *Isosphaera* colonize a wide
50 range of terrestrial and aquatic environments with diverse conditions. The first taxonomically
51 described member of this lineage, i.e. the filamentous budding bacterium *Isosphaera pallida*,
52 was isolated from hot springs (Giovannoni *et al.*, 1987). This moderately thermophilic and
53 neutrophilic planctomycete displayed a number of unique features, such as gliding motility and
54 phototaxis. The second characterized genus in this phylogenetic group, *Singulisphaera*, was
55 distinctly different from *Isosphaera pallida* and was represented by non-filamentous, non-motile,
56 moderately acidophilic and cold-adapted spherical cells that occurred singly, in pairs or in
57 shapeless aggregates (Kulichevskaya *et al.*, 2008, 2012). The strains representing the two
58 described species of this genus, *Singulisphaera acidiphila* and *Singulisphaera rosea*, were
59 isolated from *Sphagnum*-dominated acidic wetlands. The third currently recognized genus in the
60 group of *Isosphaera*-like planctomycetes is *Aquisphaera*. It includes a single species of non-
61 filamentous and neutrophilic planctomycete, which was isolated from a freshwater aquarium, i.e.
62 *Aquisphaera giovannoni* (Bondoso *et al.*, 2011).

63 Recently, representatives of the *Isosphaera*-like group were recognized as one of the
64 most abundant planctomycete populations in acidic northern wetlands (Ivanova, Dedysh, 2012;
65 Serkebaeva *et al.*, 2013; Moore *et al.*, 2015). Notably, their abundance peaked at the oxic/anoxic
66 interface, where the transition occurs from living vegetation to dead plant material. At the
67 oxic/anoxic interface of the boreal peat bog Obukhovskoye, Yaroslavl region, Russia (58° 14' N,
68 38° 12' E), 16S rRNA gene reads from *Isosphaera*-like planctomycetes comprised 53% of total
69 reads retrieved from the peat horizon (Moore *et al.*, 2015). One of the isolates described here,
70 strain PX4^T, was obtained from just above the oxic/anoxic interface (depth of 15–20 cm) of this
71 peat bog. The enrichment approach was designed in order to select for chitin-degrading
72 microorganisms capable of growth under micro-oxic or anoxic conditions. Freshly collected peat
73 (5 g) was used to inoculate 500-ml flasks containing 200 ml of liquid medium MM1 of the
74 following composition (g per liter distilled water): KH₂PO₄, 0.1; (NH₄)Cl, 0.2; MgCl₂, 0.1;

75 CaCl₂ × 2H₂O, 0.02; yeast extract, 0.05; chitin from crab shells (Sigma), 500; pH 5.5-5.8. The
76 flasks were then sealed with rubber septa, flushed with N₂ for 10 min and incubated in static
77 conditions at 20°C. After 1 month of incubation, the aliquots of peat suspension from these
78 enrichment cultures were screened by hybridization with two planctomycete-specific Cy3-
79 labelled probes PLA46 and PLA886 (Neef *et al.*, 1998). The probes hybridized to numerous
80 spherical cells that occurred in short chains or in shapeless aggregates, suggesting the presence
81 of *Isosphaera-Singulisphaera*-like planctomycetes. Further isolation strategy involved spread-
82 plating of cell suspensions from the enrichment cultures onto the same medium solidified with
83 10 g of Phytigel (Sigma–Aldrich), since this solidifying agent was shown to be highly useful for
84 isolation of diverse peat-inhabiting bacteria (Dedysh, 2011). One portion of plates was placed
85 into anaerobic jars with AnaeroGen anaerobic system envelopes (Oxoid), while another portion
86 of plates was kept in aerobic conditions. Examination of the plates after 1 month of incubation
87 revealed no growth under anoxic conditions. On plates kept under aerobic conditions, however,
88 we noticed development of numerous small (0.5-1 mm in diameter), circular, bright-pink
89 colonies that formed visible depressions in a Phytigel-solidified medium (Fig. 1a). These
90 colonies contained spherical cells, which reproduced by budding and occurred singly, in pairs or
91 in short chains containing up to 10 cells (Fig. 1b).

92 Another isolate with identical colony and cell morphologies, strain PT1, was obtained
93 from a forested tundra wetland, Nadym, Western Siberia, Russia (65°37' N, 72°43' E). In this
94 case, the cultivation approach was targeted at isolation of methanotrophic bacteria. Peat
95 suspensions were spread-plated onto the same medium MM1 with Phytigel but without yeast
96 extract and chitin, and were incubated at 20°C for 1 month in desiccators containing 30%
97 methane (v/v) in the gas phase. Small bright-pink colonies, which produced depressions in
98 Phytigel, formed on these plates among the colorless colonies produced by methanotrophic
99 bacteria.

100 The apparent ability of strains PX4^T and PT1 to hydrolyze Phytigel (a complex
101 heteropolysaccharide of microbial origin), as indicated by the depressions produced in
102 association with colonies, made them attractive objects for further studies since hydrolytic
103 capabilities of planctomycetes remain poorly characterized. Partial sequencing of the 16S rRNA
104 gene fragments (~500 bp) from these isolates showed that they affiliate with the *Isosphaera*-like
105 lineage of the family *Planctomycetaceae* but display only a distant relationship (90-94% 16S
106 rRNA gene similarity) to currently described members of this lineage, i.e. the genera *Isosphaera*,
107 *Singulisphaera*, and *Aquisphaera*. This study, therefore, was undertaken in order to characterize
108 strains PX4^T and PT1 and to describe them taxonomically.

109 Although both strains were isolated on medium MM1, they grew significantly better on
110 either agar- or Phytigel-solidified medium M31 (modification of medium 31 described by Staley
111 *et al.*, 1992) containing (g per liter distilled water): KH₂PO₄, 0.1; Hutner's basal salts, 20 ml; N-
112 acetylglucosamine, 0.5; ampicillin sodium salt, 0.2; yeast extract, 0.1; pH 5.8. Successive re-
113 streaking on agar medium M31 was used to purify strains PX4^T and PT1, which were then
114 routinely maintained on this medium without ampicillin and were subcultured at 1 month
115 intervals.

116 Morphological observations and cell-size measurements were made with a Zeiss
117 Axioplan 2 microscope and Axiovision 4.2 software (Zeiss). Negative staining was performed as
118 described for *Planctomicrobium piriforme* (Kulichevskaya *et al.*, 2015), with the only difference
119 that one additional round of staining with 1% aqueous solution of phosphotungstic acid (pH 7.0)
120 was made. The specimen samples were examined with a JEM-100C transmission electron
121 microscope.

122 Mature cells of strains PX4^T and PT1 were spherical and varied in size from 1.5 to 2.5 μm.
123 Cells occurred singly, in pairs or in short chains (Fig. 1b) and reproduced by budding (Fig. 1c,
124 d). Examination of negatively stained cells using electron microscopy showed the presence of
125 crateriform pits on the cell surface (Fig. 1c). Negative staining revealed also that many cells

126 produce some extracellular material of a net-like structure (Fig. 1e). The nature of this excreted
127 material remains unclear. On agar-solidified M31 medium, strains PX4^T and PT1 formed small
128 (1-3 mm in diameter), bright-pink-pigmented, round colonies. Notably, no depressions were
129 formed, i.e. these isolates were incapable of hydrolyzing agar. Hydrolysis of Phytigel was
130 observed on medium MM1 or on medium M31 without N-acetylglucosamine. Liquid cultures
131 displayed light-pink turbidity.

132 Physiological tests were performed in liquid medium M31. Growth of strains PX4^T and PT1
133 was monitored by nephelometry at 600 nm in an Eppendorf BioPhotometer for 7–14 days under
134 a variety of conditions, including temperatures of 4–37 °C, pH 3.8–8.0 and NaCl concentrations
135 of 0–3.0% (w/v). Variations in the pH were achieved by using MES (pH 4.0–6.5) and MOPS
136 (pH 6.5–7.9) buffer systems. The pH range of 3-4 was achieved by adjusting medium pH with
137 0.5 M H₂SO₄. Strain PX4^T grew in the temperature range of 10-30°C, with an optimum at 22-
138 25°C. Strain PT1 developed in the temperature range of 6-30°C, with an optimum at 15-25°C.
139 The pH range for growth was 3.5-6.5, with an optimum at pH 5.0-5.5 (Supplementary Fig. S1).
140 Growth was completely inhibited at NaCl concentrations above 0.5% (w/v). The doubling time
141 of this bacterium under optimal growth conditions was about 32 h.

142 Carbon source utilization was determined using mineral medium MM supplemented with
143 respective carbon sources (0.05 %, w/v). Medium MM contained (g per liter distilled water):
144 KH₂PO₄, 0.1; (NH₄)₂SO₄, 0.1; MgSO₄ × 7H₂O, 0.1; yeast extract, 0.05; 1 ml metal salt solution
145 ‘44’ (Staley *et al.*, 1992), the pH being adjusted to 5.8. Cultivation was done in 100 ml flasks
146 containing 10 ml medium. Cultures were incubated at 24°C for 2–3 weeks on a shaker. The
147 capacity to degrade different biopolymers was examined by measuring the rate of CO₂
148 production in tightly closed 120 ml flasks containing 10 ml liquid medium MM with 0.05%
149 (w/v) of the corresponding polymer substrate for 1 month at 24°C. Control incubations were run
150 in parallel under the same conditions but without substrate. Nitrogen sources were tested using
151 liquid MM medium with 0.05% (w/v) glucose in which (NH₄)₂SO₄ was replaced with one of the

152 following compounds at a concentration of 0.01% (w/v): KNO₃, KNO₂, urea or one of the amino
153 acids listed in in the species description. Cultures were tested for growth under anaerobic
154 conditions in anaerobic jars by using AnaeroGen anaerobic system envelopes (Oxoid), which
155 absorb atmospheric oxygen with the simultaneous generation of CO₂ (up to 9-13%, vol/vol). For
156 cultivation in micro-oxic conditions, medium M31 was boiled for 10 min to remove oxygen.
157 After that, hermetically closed 500 ml flasks were filled with 450 ml medium M31, inoculated
158 with examined cultures and incubated under static conditions for 2 weeks. Dissolved O₂
159 concentration was 1.5 mg O₂ L⁻¹ (measured in cultivation flasks prior to inoculation by using
160 Dissolved Oxygen Meter sensION6, Hach, USA).

161 Strains PX4^T and PT1 grew best under aerobic conditions on media with carbohydrates or N-
162 acetylglucosamine. The carbon substrates tested in our study and the results are given in the
163 species description (see below). Most sugars tested, including N-acetylglucosamine, were the
164 preferred growth substrates. With the exception of pyruvate and succinate, organic acids were
165 not utilized. Strains PX4^T and PT1 were capable of hydrolyzing aesculin, gelatin, lichenan,
166 pectin, xylan and Phytigel, but not casein, chondroitin sulfate, laminarin, starch, pullulan,
167 protein hydrolysate or cellulose. Despite its isolation from the enrichment culture of chitin-
168 degrading microorganisms, no growth of strain PX4^T was detected in chitin-containing medium
169 MM. The same was true for strain PT1. Ammonia, nitrate, N-acetylglucosamine, Bacto peptone,
170 Bacto yeast extract, alanine, asparagine and valine were utilized as nitrogen sources. Neither of
171 the two isolates grew under anoxic conditions, although they showed stable albeit slow growth
172 ($\mu \sim 0.01 \text{ h}^{-1}$) under micro-oxic conditions.

173 Oxidative and fermentative utilization of carbohydrates and gelatin liquefaction were
174 determined by using an API 20NE kit (bioMérieux) and as described for the Hugh–Leifson test
175 (Gerhardt *et al.*, 1981). Enzymatic activities were examined by using an API ZYM kit
176 (bioMérieux). Catalase was tested by using the method 1 described by Gerhardt *et al.* (1981).
177 Oxidase was tested using a REF 55 635 Oxidase Reagent (bioMérieux). Strains PX4^T and PT1

178 were catalase- and cytochrome oxidase -positive, but urease -negative. Dissimilatory nitrate
179 reduction was negative. The following enzymatic activities (API ZYM) were detected in strains
180 PX4^T and PT1: acid phosphatase, esterase, esterase lipase, leucine and valine arylamidases,
181 phosphohydrolase, N-acetyl- β -glucosaminidase and β -galactosidase (API ZYM test). The
182 following enzyme activities were not detected: alkaline phosphatase, cystine arylamidase, lipase,
183 trypsin, chymotrypsin, α -galactosidase, β -glucuronidase, α -fucosidase and α -mannosidase.

184 Susceptibility of strains PX4^T and PT1 to antibiotics was determined on M31 agar plates
185 using discs containing the following antibiotics: ampicillin (10 mg), gentamicin (10 mg),
186 kanamycin (30 mg), neomycin (10 mg), novobiocin (30 mg), streptomycin (10 mg),
187 chloramphenicol (30 mg) and lincomycin (10 mg) (Oxoid). The isolates were resistant to
188 ampicillin, streptomycin, chloramphenicol, lincomycin, and novobiocin, but sensitive to
189 kanamycin, neomycin and gentamicin.

190 For the analysis of fatty acids, intact polar lipids (IPLs), neutral lipids and quinones, cells
191 of the novel isolates were grown on liquid medium M31 and harvested in the late exponential
192 growth phase. Lipids were analyzed after acid hydrolysis of whole cells following the procedure
193 described by Sinninghe Damsté *et al.* (2011). The major fatty acids (FA) detected in strain PX4^T
194 and PT1 were $nC_{16:0}$, $nC_{18:1\omega9}$ and $nC_{18:0}$ (Table 1). A number of hydroxy FA, including the (ω -
195 1)-OH $nC_{28:0}$ hydroxy FA, and the n-C31 : 9 hydrocarbon were also detected. The latter seems
196 to be very common in most planctomycetes. As shown in Table 2, the major polar lipids were
197 phosphocholine (PC) and trimethylornithine (TMO). Phosphoglycerol (PG) and 1-acyl-glycero-
198 3-phosphocholine were also present in minor amounts. As reported by Moore *et al.* (2015), the
199 ratio of TMO/(PC +PG) in cells grown in micro-oxic conditions was higher than that in fully
200 aerobic conditions, which suggests that TMOs could be synthesized as a response to changing
201 redox conditions in the oxic/anoxic interface.

202 Isoprenoid quinones of strains PX4^T and PT1 were analyzed as described for
203 *Planctomicrobium piriforme* (Kulichevskaya *et al.*, 2015). Similar to other members of the order

204 *Planctomycetales* ([Ward, 2010](#)), our isolates contained menaquinone-6 (MK-6) as the only
205 isoprenoid quinone. Based on genome sequence analysis, the DNA G+C content of strain PX4^T
206 is 66 mol % (Ivanova *et al.*, unpublished data), which is higher than that in *Isosphaera* and
207 *Singulisphaera* but lower than that in *Aquisphaera* (Table 3).

208 PCR-mediated amplification of the 16S rRNA gene from DNA of strains PX4^T and PT1 was
209 performed using primers 9f and 1492r and reaction conditions described by Weisburg *et al.*
210 (1991). 16S rRNA gene amplicons were sequenced on an ABI 377A DNA sequencer using
211 BigDye terminator chemistry, as specified by the manufacturer (PE Applied Biosystems).
212 Phylogenetic analysis was carried out using the ARB program package (Ludwig *et al.*, 2004).
213 The significance levels of interior branch points obtained in the neighbour-joining analysis were
214 determined by bootstrap analysis (based on 1000 data resamplings) using PHYLIP (Felsenstein,
215 1989). The comparative analysis based on nearly full-length 16S rRNA gene sequences
216 confirmed that strains PX4^T and PT1 belong to the order *Planctomycetales* and are members of
217 the coherent phylogenetic cluster defined by the genus *Isosphaera* (Fig. 2). The minimum
218 sequence identity within this cluster is about 90%, which is close to the taxonomic threshold
219 defined for the family level (Yarza *et al.*, 2014). This cluster is strongly supported by all
220 algorithms used for the tree construction and accommodates morphologically similar, stalk-free
221 planctomycetes with spherical cells, which can be assembled in short chains, long filaments or
222 shapeless aggregates. Notably, daughter cells of these budding bacteria are non-motile. The 16S
223 rRNA gene sequence identity between members of the *Isosphaera*-like cluster and other
224 taxonomically described organisms within the order *Planctomycetales* is in the range of 76.8-
225 81.9%. Based on this phylogenetic divergence (Fig. 2) and the morphological similarity between
226 members of the *Isosphaera*-like cluster, the latter should be given the status of a family, i.e.
227 *Isosphaeraceae* fam. nov. Strains PX4^T and PT1 are members of the *Isosphaeraceae*, but are
228 phylogenetically divergent, morphologically distinct and phenotypically different from other
229 characterized representatives of this family. These non-filamentous, non-motile, acidophilic and

230 cold-tolerant planctomycetes could clearly be differentiated from filamentous, gliding,
231 thermophilic and neutrophilic *Isosphaera pallida*. Abilities to grow at temperatures below 10°C,
232 pH ≤ 6.0 as well as to develop under micro-oxic conditions differentiate the novel isolates and
233 *Aquisphaera giovannonii*. Finally, strains PX4^T and PT1 can be distinguished from members of
234 the genus *Singulisphaera* by the formation of cell chains, the ability to hydrolyze Phytigel, the
235 absence of C18:2ω6c,12c fatty acid, and the lower DNA G+C content (Table 3). In addition,
236 cells of novel isolates are smaller than cells of *Singulisphaera* spp.

237 Strains PX4^T and PT1 displayed 93-94% 16S rRNA gene sequence similarity to *Aquisphaera*
238 *giovannonii*, 91-92% to *Singulisphaera* species and 90-91% to *Isosphaera pallida*. The overall
239 similarities between the genome of strain PX4^T (Ivanova *et al.*, unpublished data) and the
240 genomes of *Isosphaera pallida* IS1B^T and *Singulisphaera acidiphila* DSM 18658^T estimated
241 using formula 2 of the Genome-to-Genome-Distance-Calculator (Auch *et al.*, 2010) is 19.8 ± 2.3
242 and 20.0 ± 2.3, respectively. These DDH values are similar to those calculated for members of
243 different genera (Scheuner *et al.*, 2014). Average nucleotide identity values generated by
244 comparing the genome of strain PX4^T and the genomes of *Isosphaera pallida* IS1B^T and
245 *Singulisphaera acidiphila* DSM 18658^T are also very low, i.e. 75 and 77%, respectively. We,
246 therefore, propose to classify strains PX4^T and PT1 as representing a novel genus and species,
247 *Paludisphaera borealis* gen. nov., sp. nov. The characteristics that differentiate the genus
248 *Paludisphaera* from the genera *Isosphaera*, *Singulisphaera* and *Aquisphaera* are summarized in
249 Table 3.

250

251 **Description of the genus *Paludisphaera*.**

252 *Paludisphaera* (Pa.lu.di.sphae'ra. L. n. *palus* –*udis* a swamp, marsh; L. fem. n. *sphaera* a ball,
253 globe sphere; N.L. fem. n. *Paludisphaera* a spherical cell from wetland).

254 Cells are Gram-stain-negative, non-motile spheres that occur singly, in pairs or in short chains.

255 Reproduce by budding. Daughter cells are non-motile. Crateriform pits are scattered all over cell

256 surface. Stalk-like structures are absent. Colonies are opaque and pink colored. Chemo-
257 organotrophic aerobes. Capable of growth under micro-oxic conditions. Possess hydrolytic
258 capabilities. Dissimilatory nitrate reduction is negative. Moderately acidophilic and mesophilic.
259 Sensitive to NaCl. The major quinone is MK-6. The major fatty acids are *n*C16:0, *n*C18:1 ω 9 and
260 *n*C18:0. The major polar lipids are phosphocholine and trimethylornithine. The genus is a
261 member of the phylum *Planctomycetes*, order *Planctomycetales*, family *Isosphaeraceae*. The
262 type species is *Paludisphaera borealis*.

263

264 **Description of *Paludisphaera borealis* sp. nov.**

265 *Paludisphaera borealis* (bo.re.a'lis. L. fem. adj. *borealis* pertaining to the north, boreal).

266 The description is as for the genus but with the following additional traits. Spherical cells with
267 diameter of 1.5–2.5 μ m. Carbon sources (0.05 %, w/v) include glucose, fructose, galactose,
268 lactose, cellobiose, maltose, mannose, melibiose, rhamnose, ribose, trehalose, sucrose, xylose,
269 N-acetylglucosamine, salicin, pyruvate, and succinate. Cannot utilize leucrose, raffinose,
270 sorbose, melezitose, fucose, glycerol, methanol, ethanol, starch, glucuronic acid, benzoate,
271 caproate, citrate, formate, formaldehyde, fumarate, glutarate, lactate, malate, propionate,
272 mannitol, tartrate, alanine, arginine, asparagine, aspartate, cysteine, cystine, glutamine, glycine,
273 histidine, isoleucine, leucine, lysine, methionine, norleucine, ornithine, phenylalanine, proline,
274 serine, threonine, tryptophan, tyrosine or valine. Capable of hydrolyzing aesculin, gelatin,
275 lichenan, pectin, xylan and Phytigel. Cannot hydrolyze casein, chondroitin sulfate, laminarin,
276 starch, peptone, pullulan, protein hydrolysate, fucoidan, chitin or cellulose. Catalase-,
277 cytochrome oxidase-positive, but urease-negative. Nitrogen sources (0.05 %, w/v) include
278 ammonia, nitrate, N-acetylglucosamine, Bacto peptone, Bacto yeast extract, alanine, asparagine
279 and valine. Aspartate, arginine, glutamine, glycine, isoleucine, lysine, threonine, tryptophan,
280 phenylalanine, proline and urea are not utilized. Growth factors are required. Possess the
281 following enzyme activities: acid phosphatases, esterase, esterase lipase, leucine arylamidase,

282 valine arylamidase, phosphohydrolase, N-acetyl- β -glucosaminidase and β -galactosidase (API
283 ZYM test). The following enzyme activities are not present: alkaline phosphatase, cystine
284 arylamidase, lipase, trypsin, chymotrypsin, α -galactosidase, β -glucuronidase, α -fucosidase and α -
285 mannosidase. Resistant to ampicillin, streptomycin, chloramphenicol, lincomycin, kanamycin,
286 but sensitive to neomycin, novobiocin and gentamicin. Growth occurs at pH 3.5-6.5 (optimum at
287 pH 5.0-5.5) and at temperatures between 6°C and 30°C (optimum at 15-25°C). Growth is
288 inhibited at NaCl concentrations above 0.5% (w/v). The DNA G+C content is 66 mol%.
289 Wetlands are the main habitat. The type strain, PX4^T (= DSM 28747^T = VKM B-2904^T), was
290 isolated from the peat bog Obukhovskoye, Yaroslavl region, Russia.

291

292 **Description of *Isosphaeraceae* fam. nov.**

293 *Isosphaeraceae* (I.so.sphae.ra.ce´ae. N.L. fem. n. *Isosphaera* type genus of the family; -aceae
294 ending to denote a family; N.L. fem. pl. n. *Isosphaeraceae* the *Isosphaera* family).

295 Gram-stain negative, budding bacteria with spherical cells, which occur singly, in pair or are
296 assembled in short chains, long filaments or shapeless aggregates. Crateriform pits are scattered
297 all over cell surface. Stalk-like structures are absent. Do not form rosettes. Daughter cells are
298 non-motile. Chemo-organotrophic aerobes. Some representatives are capable of growth in micro-
299 oxic conditions. The family belongs to the class *Planctomycetacia*, order *Planctomycetales*. The
300 type genus is *Isosphaera*. Other genera in this family are *Singulisphaera*, *Aquisphaera*, and
301 *Paludisphaera*.

302

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307

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352 *Rubinisphaera* gen. nov. and emended descriptions of the order *Planctomycetales* and the family
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376 **Table 1.** Relative abundance (% of total) of lipids extracted after acid hydrolysis of cell material
 377 of strain PX4^T and PT1. (Major components are given in bold).
 378

	PX4^T	PT1
Fatty acids		
<i>n</i> C14:0	3.4	3.2
<i>n</i> C16:1 ω 9	1.8	1.7
<i>n</i> C16:1 ω 7	0.3	0.6
<i>n</i> C16:0	17.4	16.5
<i>n</i> C17:1 ω 8	0.4	-
<i>n</i> C18:1 ω 9	47.9	50.9
<i>n</i> C18:0	12.4	7.4
Hydroxy fatty acids		
β -OH <i>n</i> C14:0	-	1.0
β -OH <i>n</i> C18:0	0.8	2.4
α - OH <i>n</i> C25:0	0.4	0.3
α - OH <i>n</i> C27:1	-	0.5
α - OH <i>n</i> C27:0	4.7	4.4
α - OH <i>n</i> C29:1	0.8	1.7
α - OH <i>n</i> C29:0	0.5	0.5
α ,(ω -1)-diOH <i>n</i> C29:0	0.7	0.5
(ω -1)-OH <i>n</i> C28:0	1.2	1.3
(ω -1)-OH <i>n</i> C30:1	0.6	1.3
α ,(ω -1)-diOH <i>n</i> C31:0	1.9	1.6
C ₃₃ α -OH hopanoic acid	2.4	2.9
Other lipids		
<i>n</i> C31:9 hydrocarbon	2.4	1.2

379

380 **Table 2.** Relative abundances and fatty acid composition of IPLs of strain PX4^T and PT1.

381

IPL	Strain PX4 ^T	Strain PT1	Fatty acid composition
PG	+	+	(C36:2, C34:1, C32:1)
PC	+++	+++	(C36:2, C34:1, C32:1)
TMO	++	+	(C18:1, β OH C18)
lyso-PC	tr	+	(C18:1, C16:0)

382

383 The abundance is relative to the major peak in the LC/MS base peak chromatogram: +++=base
 384 peak, ++=50-100% of base peak, +=10-50% of base peak, tr=<10% of base peak. Note that the
 385 mass spectral response factor for different IPL groups can be quite different. The predominant
 386 fatty acid composition is reported as the total number of the acyl moieties and the number of
 387 double bonds (except for TMO and lyso-PC where individual acyl moieties are given).

388 PG=phosphoglycerol, PC=phosphocholine, TMO= trimethylornithine , lyso-PC=1-acyl-glycero-
 389 3-phosphocholine

390

391 **Table 3.** Major characteristics that distinguish the genus *Paludisphaera* gen. nov. and the genera
 392 *Isosphaera*, *Singulisphaera*, and *Aquisphaera*

Characteristic	<i>Paludisphaera</i>	<i>Isosphaera</i> *	<i>Singulisphaera</i> **	<i>Aquisphaera</i> ***
Arrangement of cells	Single, in pairs or short chains	Filaments	Single or in pairs	Single or aggregates
Cell size, μm	1.5-2.5	2.5-3.0	1.6-3.5	1.6-2.0
Gliding motility	-	+	-	-
Photo-taxis	-	+	-	-
Habitat	Wetlands	Hot springs	Wetlands	Freshwater
Colony colour	Pink	Pink	Colourless or pink	Pink
Respiration	Aerobic or microaerophilic	Strictly aerobic	Aerobic or microaerophilic	Strictly aerobic
Hydrolysis of Phytigel aesculin starch xylan	+ + - +	ND ND ND ND	- + - +	ND - + -
Temperature growth range ($^{\circ}\text{C}$) Optimal temperature ($^{\circ}\text{C}$)	6-30 15-25	34-55 40-50	4-33 15-28	10-35 30-35
pH growth range pH optimum	3.5-6.5 5.0-5.5	ND 7.8-8.8	4.2-7.5 5.0-6.2	6.5-9.5 7.5-8.5
Vitamin requirement	None	ND	None	B ₁₂
Presence of C18 : 2 ω 6c,12c fatty acids	-	ND	+	-
G+C content (mol%****)	66	62	62	70

393 *Data from Giovannoni *et al.* (1987) and Göker *et al.* (2011).

394 **Data from Kulichevskaya *et al.* (2008, 2012) and Scheuner *et al.* (2014).

395 ***Data from Bondoso *et al.* (2011).

396 ****Values given for *Paludisphaera*, *Isosphaera*, and *Singulisphaera* are based on genome
 397 sequence analyses.

398 ND, not determined

399

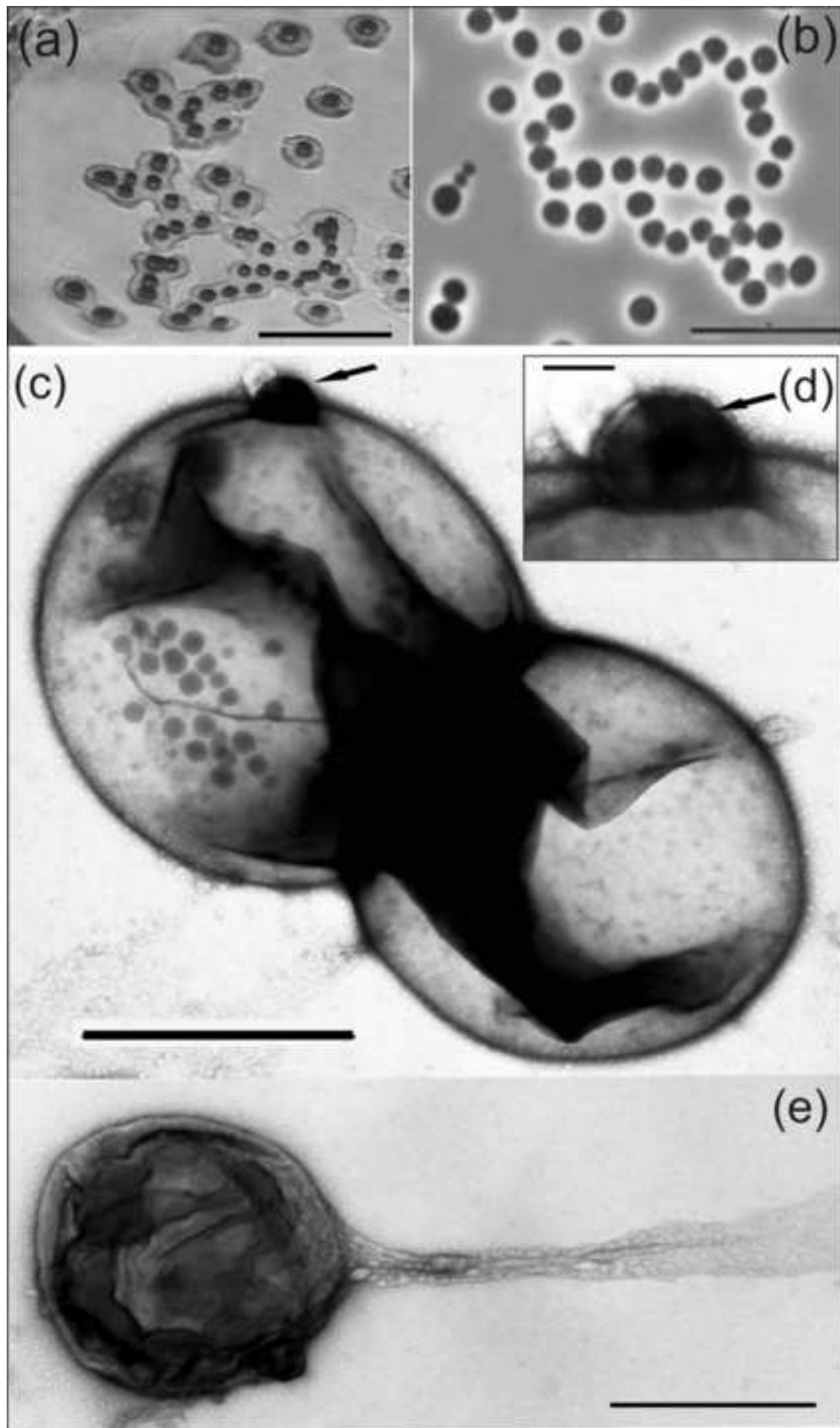
400 **FIGURE CAPTIONS**

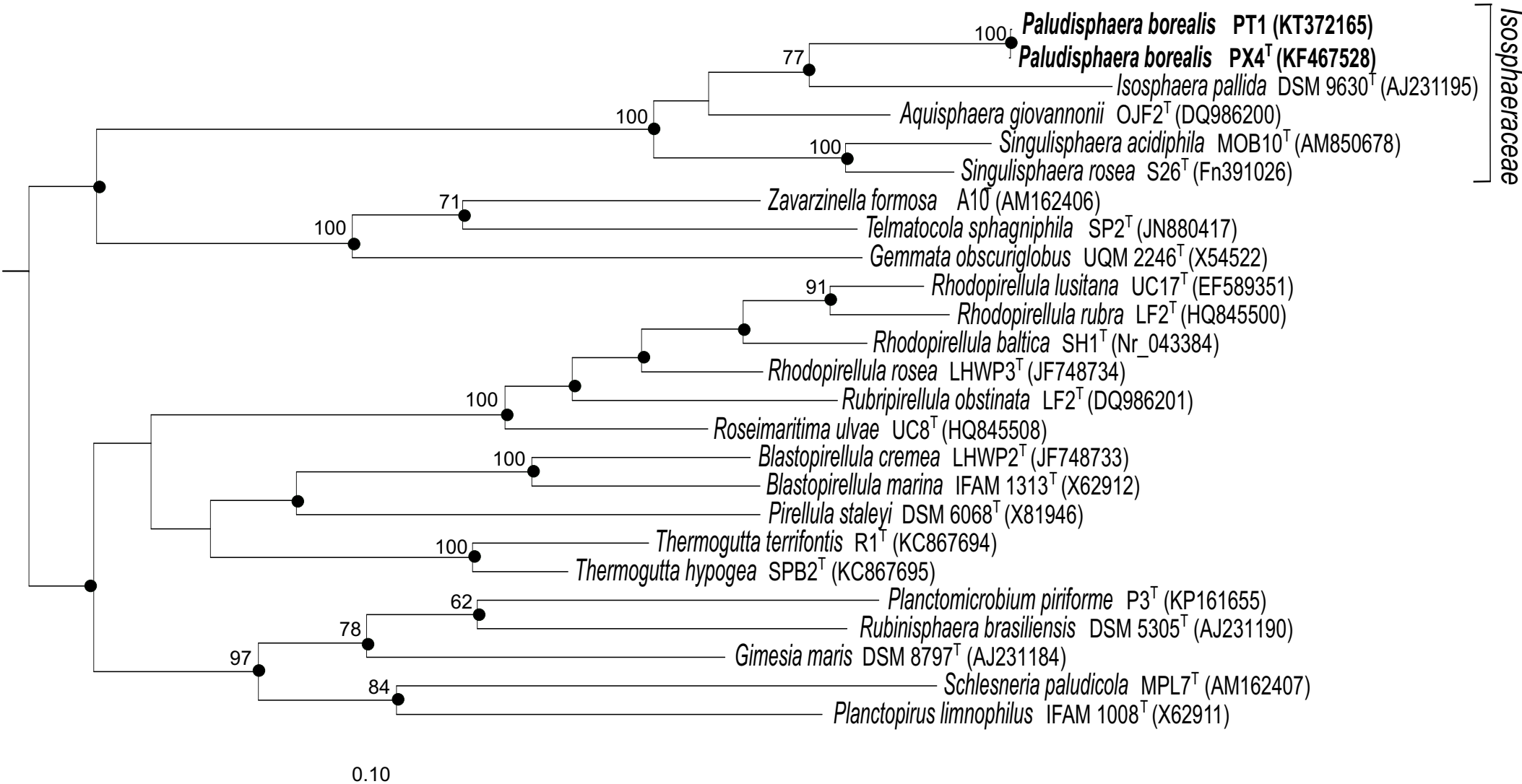
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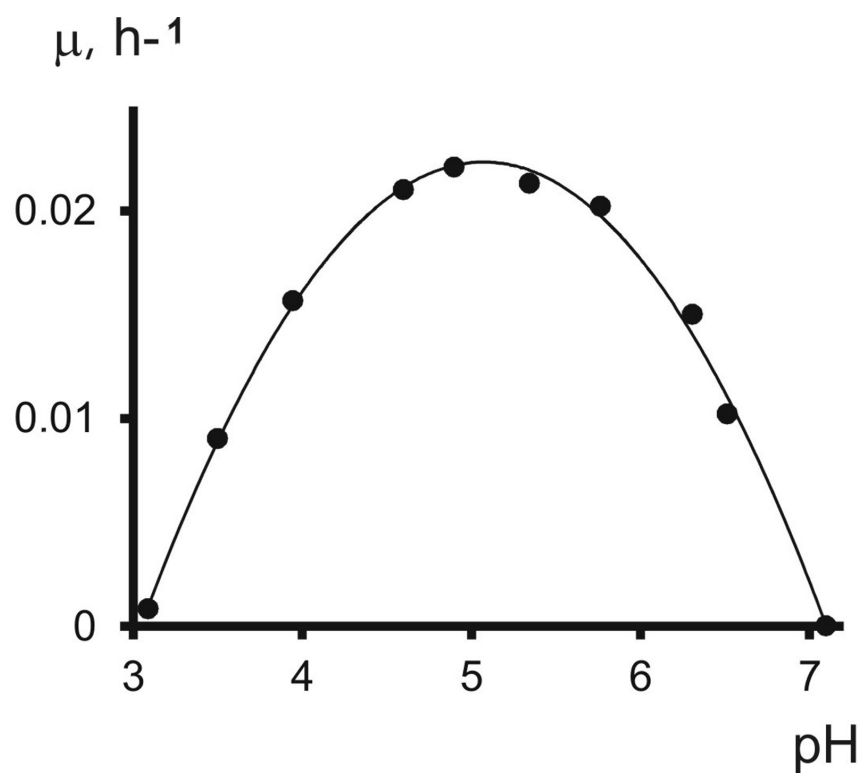
402 **Figure 1.** (a) Development of depressions in Phytigel-solidified medium during colony growth
403 of strain PX4^T. Bar, 10 mm. (b) Phase-contrast image of cells of strain PX4^T in 10-day-old
404 culture. Bar, 10µm. (c -d) Electron micrographs of negatively stained cells of strain PX4^T
405 displaying crateriform pits scattered all over cell surface (c), a newly formed bud (indicated by
406 black arrow in c and d) and an extracellular material of a net-like structure excreted by the cell
407 (e). Bars, 1 µm (c, e) and 0.2 µm (d).

408

409 **Figure 2.** 16S rRNA gene-based neighbour-joining tree (Jukes-Cantor correction) showing the
410 phylogenetic relationship of strain PX4^T and PT1 to representative members of the order
411 *Planctomycetales*. The bracket on the right indicates boundaries proposed for the family
412 *Isosphaeraceae*. The significance levels of interior branch points obtained in neighbor-joining
413 analysis were determined by bootstrap analysis (1000 data re-samplings) using PHYLIP
414 (Felsenstein, 1989). Bootstrap values (1000 data resamplings) of >50% are shown. Black circles
415 indicate that the corresponding nodes were also recovered in the maximum-likelihood and
416 maximum-parsimony trees. The root (not shown) was composed of five 16S rRNA gene
417 sequences from anammox planctomycetes (AF375994, AF375995, AY254883, AY257181,
418 AY254882). Bar, 0.1 substitutions per nucleotide position.







Supplementary Fig. S1. Influence of medium pH on the growth of strain PX4^T.



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