# Supplementary Information for Neuenschwander et al.

## Supplementary text

Additional genomic features of 'Ca. Nanopelagicales' Description of the proposed 'Candidatus' taxa

Additional references

- **Figure S1:** Microphotographs and cell volumes (µm<sup>3</sup>) of isolates. The scale bar at the bottom left applies to all pictures.
- **Figure S2:** Phylogenetic positioning of '*Ca*. Nanopelagicales' based on 16S rRNA genes. a, bootstrapped maximum likelihood tree of 16S rRNA genes; '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are shown on the nodes. b, 16S rRNA gene sequence similarity matrix. Species borders are marked with solid lines, genus borders with dashed lines. An asterisk indicates the positioning of the described mixed culture '*Ca*. P. limnetica'.
- **Figure S3:** Average nucleotide identity (ANI) matrix of '*Ca*. Nanopelagicales'. Species borders are marked with solid lines, genus borders with dashed lines.
- Figure S4: a, Average amino acid identity (AAI) matrix of '*Ca*. Nanopelagicales'. b, Protein similarity (>50% identity, >50% coverage) matrix of '*Ca*. Nanopelagicales'. Species borders are marked with solid lines, genus borders with dashed lines.
- **Figure S5:** Phylogenomic tree with complete genomes of '*Ca*. Nanopelagicales' only. 462 concatenated conserved proteins were used to generate a maximumlikelihood phylogenetic tree. The genomes of '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are indicated by black, grey, and white circles on the nodes, and a colour key is shown on the left.
- **Figure S6:** Phylogenomic tree with complete genomes of the phylum Actinobacteria. Forty-eight concatenated conserved proteins were used to generate a maximum-likelihood phylogenetic tree. The genomes of *Staphylococcus aureus* and *Listeria monocytogenes* were used as outgoup. Bootstrap values are indicated by black, grey or white circles on the nodes, and a colour key is shown on the left. The proposed novel order '*Ca*. Nanopelagicales' is highlighted in green.
- **Figure S7:** Genome streamlining in '*Ca*. Nanopelagicales'. Number of predicted CDS, number of predicted sigma factor homologs, median size of intergenic spacers, and coding density versus genome size for all complete published genomes of Actinobacteria (n=610; data taken from RefSeq). '*Ca*. Nanopelagicales' and *Rhodoluna lacicola* are marked in different colours.
- **Figure S8:** Bootstrapped phylogenetic tree of rhodopsin protein sequences of 'Ca. Nanopelagicales' and other Actinobacteria. Xanthorhodopsin sequences of *Salinibacter ruber* and *Thermus aquaticus* were used as outgroup. Bootstrap values are indicated by coloured circles on the nodes, and a colour key is shown on the left.

**Figure S9:** Bootstrapped maximum likelihood tree of 23S rRNA genes of '*Ca*. Nanopelagicales'. Target hits for the newly designed probes Pver-23S-1420 and Npel-23S-2669 are shown in brackets. '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are shown on the nodes.

Figure S10: Physico-chemical data from Lake Zurich.

- **Figure S11:** Redundancy analysis of environmental parameters explaining the variability in cell numbers of microbes affiliated to all '*Ca*. Nanopelagicales', '*Ca*. Nanopelagicus', and '*Ca*. P. vernalis' in Lake Zurich. temp, water temperature; picocyano, abundance of picocyanobacteria; irrad, irradiation; chloro, chlorophyll *a* associated with chlorophytes; diatom, chlorophyll *a* associated with diatoms; NH4, ammonium concentrations; O2, oxygen concentrations; PO4, phosphate concentrations; NO3, nitrate concentrations; depth, sampling depth.
- **Figure S12:** Metagenomic fragment recruitment of '*Ca*. Nanopelagicus' (a) and '*Ca*. Planktophila' (b) across diverse freshwater ecosystems (see Table S7 for details).
- Figure S13: Recruitment plots of time-series metagenomes from Lake Mendota, USA.
- **Figure S14:** Whole-genome alignment of all 13 '*Ca.* Planktophila' strains. The genomes have been linearized for simplicity and are arranged in the same order as in the phylogenomic tree in Fig. 1b. Synteny and different degrees of sequence similarity are indicated by vertical lines connecting the genomes.
- **Figure S15:** Whole-genome alignment of the three '*Ca.* Nanopelagicus' strains. The genomes have been linearized for simplicity and are arranged in the same order as in the phylogenomic tree in Fig. 1b. Synteny and different degrees of sequence similarity are indicated by vertical lines connecting the genomes. rRNA operons in the individual genomes are displayed as red arrows and tRNAs as short vertical lines. Genomic islands (GI) have been marked in different colours and numbered (see Table S10 for genes encoded in each island). Red: genes encoding mainly cell wall biosynthesis and modifications; Yellow: genes encoding mainly membrane transport and / or carbohydrate metabolism.
- Table S1: Details of the isolation experiments conducted in Lake Zurich, Switzerland.
- Table S2: Preprocessing and assembly parameters.
- Table S3: Details of the sequenced strains of planktonic 'Ca. Nanopelagicales'.
- Table S4: Cell size measurements of different strains and *in-situ* in Lake Zurich.
- **Table S5:** Cell sizes and genomic details of genome-streamlined microbes.
- Table S6: Details of the applied oligonucleotide probes.
- **Table S7:** Details of the publically available metagenomes that were used for recruitment in Figs. 3, S12, S13. Metagenomes are sorted according to habitat (rivers and lakes) and latitude (separately for North America and Europe). This table is provided as Excel-file (Tables\_S7-S10.xlsx).
- **Table S8**: Metabolic pathways in 'Ca. Nanopelagicales'. This table is provided as Excel-file (Tables\_S7-S10.xlsx).

- **Table S9:** Genomic islands in 11 '*Ca.* Planktophila sp.' strains. Locus tags and gene annotations are listed and a general function of the genes encoded in each island is given at the left. This table is provided as Excel-file (Tables\_S7-S10.xlsx).
- **Table S10:** Genomic islands in the three '*Ca*. Nanopelagicus sp.' strains. Locus tags and gene annotations are listed and a general function of the genes encoded in each island is given at the left. This table is provided as Excel-file (Tables\_S7-S10.xlsx).

# 1 Supplementary text

# 2 Additional genomic features of 'Ca. Nanopelagicales'

None of the strains encoded genes for flagella or chemotaxis, confirming the nonmotile lifestyle of '*Ca*. Nanopelagicales'. No CRISPR-Cas system was identified, and
the number of signal transduction genes was low with only two two-component
regulatory systems for phosphate starvation and osmotic stress response and 2-3
additional histidine kinases (Table S7). Nickel superoxide dismutases involved in
oxidative stress response were found in all strains, while catalase-peroxidases were

9 present in 'Ca. Planktophila', but absent in 'Ca. Nanopelagicus'.

10 We could confirm the presence of glycolysis via the Embden-Meyerhof pathway, pentose phosphate pathway, tricarboxylic acid (TCA) cycle and oxidative 11 12 phosphorylation in all genomes, as was proposed from SAGs and MAGs (Garcia et 13 al., 2013, Ghai et al., 2014, Ghylin et al., 2014). Nevertheless, some variations between the different strains were found: 'Ca. Planktophila' strains encoded a class I 14 15 fructose-bisphosphate aldolase, while the class II variant was annotated in all 'Ca. 16 Nanopelagicus' genomes. The former is in contrast to the 'Ca. Planktophila' genomes sequenced of Kang et al. (2017) which all encoded the class II variant. The non-17 18 oxidative branch of the pentose phosphate pathway was detected in all genomes, 19 whereas the oxidative branch was only present in 'Ca. P. dulcis' strains and 'Ca. P. 20 sulfonica' MMS-IA-56. This is in agreement with previous studies in which genes involved in the oxidative branch of the pentose phosphate pathway were only 21 22 detected in 'Ca. Planktophila' (i.e., acl-A / acl-A1) genomes (Ghylin et al., 2014, Kang 23 et al., 2017). Pyruvate dehydrogenase E2 components, involved in the conversion of 24 pyruvate to acetyl-CoA, were only detected in one of the 'Ca. P. versatilis' strains (MMS-IIB-142). All genomes did however contain genes coding for the 2-25 oxoglutarate dehydrogenase E2 component, a potential substitute for the latter, 26 27 which was also the case in the four acl genomes described by Kang et al. (2017). 28 Complete gluconeogenesis pathways were detected in 'Ca. N. limnes' and 'Ca. N. 29 hibericus' as well as in 'Ca. P. limnetica' and 'Ca. P. vernalis', whereas in the other 30 genomes either the first steps (pyruvate to PEP) and/or the last steps (fructose-1,6P2 31 to β-D-fructose-6P) were missing. Key enzymes involved in gluconeogenesis have 32 previously been detected in acl-A7, acl-C1 and acl-B1 genomes (Kang et al., 2017, 33 Ghylin et al., 2014) but not in other ac1-A1 genomes. The presence of carbonic 34 anhydrases and PEP carboxylases in all genomes suggests the ability to replenish 35 precursors needed for growth by anapleurotic CO<sub>2</sub> fixation. Previous studies 36 indicated a facultative aerobic lifestyle based on the presence of pathways for pyruvate fermentation (Garcia et al., 2013, Ghai et al., 2014, Ghylin et al., 2014) We 37 38 cannot confirm this for our genomes. However, all our strains were isolated from 5 m 39 depth from Lake Zurich, where oxic conditions prevail. Oxidative phosphorylation 40 pathways were present in all genomes with the anomaly of missing succinate dehydrogenase subunits SDHC and SDHD as previously described by Kang et al., 41 42 (2017).

Exopolyphosphatases, high affinity membrane transporters for inorganic
phosphate uptake (Pst system), inorganic pyrophosphatases, and two-component
regulatory systems for phosphate stress were annotated for all strains, in agreement
with predictions from SAGs (Ghylin *et al.*, 2014) and MAGs (Ghai *et al.*, 2014).
Phosphorus is usually the limiting element in freshwater systems (Vadstein 2000,

48 Wetzel 2001), and it seems that 'Ca. Nanopelagicales have efficient phosphate acquisition systems, although they did not show higher *in-situ* incorporation of 49 50 inorganic phosphate than other freshwater microbes (Rofner et al., 2016b) and were 51 underrepresented in dissolved organic phosphate uptake (Rofner et al., 2016a). All strains encoded genes for ammonium transport (Amt family), and 'Ca. P. sulfonica', 52 53 'Ca. P. versatilis' and 'Ca. P. lacus' strains had two copies. Transporters for other N-54 rich components like polyamines (i.e., spermidine/putrescine) and amino acids as 55 well as cyanophycinases were predicted for all strains and cyanate transporters were present in 'Ca. P. versatilis' and 'Ca. P. vernalis'. This high prevalence of genes 56 57 involved in the uptake of organic N-rich compounds appears to be a key 58 characteristic for 'Ca. Nanopelagicales' (Garcia et al., 2013; Ghai et al., 2014; Kang 59 et al., 2017). A high in-situ uptake of different amino acids was previously reported from MAR-FISH assays, as well as high glucose incorporation (Buck et al., 2009, 60 Pérez et al., 2015, Salcher et al., 2010, Salcher et al., 2013). Polyamine degradation 61 62 pathways were present in all genomes, however, not complete. With the exception of 63 'Ca. P. sulfonica' all strains were able to convert 4-aminobutyraldehyde to succinate; while enzymes catalysing the first three steps of the spermidine/putrescine 64 degradation pathway were not annotated. Most genomes, however, contained 65 66 acetylornithine aminotransferases which belong to the same superfamily as putrescine aminotransferases, and thus, might fulfil this function. Lysozymes were 67 predicted in all genomes, as well as putative chitinases (GH18 family, glycoside 68 69 hydrolase CAZy; (Garcia et al., 2013)) that include signal sequences for membrane transport, consistent with SAGs (Ghylin et al., 2014). Their low homology to 70 chitinases originating from other than acl SAGs or MAGs, however, makes 71 72 interpretation difficult, particularly since genes involved in amino-sugar degradation were only detected in 'Ca. P. dulcis' and 'Ca. P. sulfonica'. We could also not identify 73 74 transporters for amino sugars although 'Ca. Nanopelagicales' are known to take up 75 N-acetylglucosamine in situ (Beier and Bertilsson 2011, Eckert et al., 2012). Other transporters, especially for carbohydrates, as well as genes encoding carbohydrate 76 77 breakdown, were present in a highly variable fashion in the genomes: 'Ca. P. dulcis' 78 and 'Ca. P. vernalis' strains had the highest number of different carbohydrate transporters and enzymes involved in carbohydrate metabolism, while 'Ca. 79 80 Nanopelagicus' strains and 'Ca. P. limnetica' were less versatile in carbohydrate usage (Tables 2, S7). Membrane transporters for sulfonate and benzoate were 81 82 annotated in two strains only ('Ca. P. sulfonica' and 'Ca. P. limnetica', respectively). 83 Transporters for biotin (bioY) and energy-coupling factor transporters with preceding 84 AdoCbl riboswitches for cobalamin were annotated in all 'Ca. Planktophila' strains except for 'Ca. P. limnetica'. Although transporters for thiamine and riboflavin were 85 not found, additional NAD transporters (pnuC) preceded by riboswitches for thiamine 86 and/or riboflavin were annotated in all 'Ca. Nanopelagicus'. This is in accordance 87 with other genomes associated with 'Ca. Nanopelagicales' (Kang et al., 2017). As 88 89 Pnu transporters share high homologies to each other, it is likely that the putative 90 PnuC either function as thiamine (PnuT) or riboflavin (PnuX) transporters (Jaehme 91 and Slotboom Dirk 2015).

## 92 Description of the proposed 'Candidatus' taxa

93 Based on our analysis we suggest that our strains represent a novel '*Candidatus*'

94 order and 'Candidatus' family in the phylum Actinobacteria with two new genera and

95 nine new species.

96 'Candidatus Nanopelagicales' [Na.no.pe.la.gi.ca'les. N.L. masc. n. 97 Nanopelagicus type genus of the order; suff. -ales, ending to denote an order; N.L. 98 fem. pl. n. Nanopelagicales, the order of the genus Nanopelagicus]. Aerobic 99 chemoheterotrophs. Cells are tiny, non-motile, and inhabit the plankton of 100 freshwaters. 'Ca. Nanopelagicales' can be recognized by the presence of the 101 diagnostic oligonucleotide sequence 5'-AATGCGTTAGCTGCGTCGCA-3' in the 16S 102 rRNA gene (positions 852-872, E. coli numbering). A member of the phylum Actinobacteria. Contains a single family, 'Candidatus Nanopelagicus', and two 103 104 genera 'Candidatus Nanopelagicus' and 'Candidatus Planktophila'. Basis of the 105 assignment is a phylogenetic tree of 48 conserved concatenated proteins of >100 complete genomes of all orders of Actinobacteria (Fig. 2, S6). 106

107 'Candidatus Nanopelagicaceae' [Na.no.pe.la.gi.ca.ce'ae. N.L. masc. n.
 108 Nanopelagicus type genus of the family; suff. -aceae, ending to denote a family; N.L.
 109 fem. pl. n. Nanopelagicaceae, the order of genus Nanopelagicus.]

110 'Candidatus Nanopelagicus limnes' [Na.no.pe.la.gi.cus. N.L. masc. n. nano very small; L. masc. adj. *pelagicus* belonging to the pelagial; N.L. masc. n. *Nanopelagicus* 111 112 very small pelagic; referring to the small cell and genome size and the pelagic 113 habitat; lim'nes. L. gen. n. limnes of a lake]. Represented by strain MMS-21-122, which was isolated from Lake Zurich, Switzerland. Curved rods with lengths of 114 115 0.45±0.09 µm and diameters of 0.25±0.03 µm (Table S3, Figure S1). The initial pure culture was lost after a few propagations to fresh medium; no growing culture is 116 117 available. The initial culture grew well in sterile lake water amended with minimal 118 carbon medium, vitamins and amino acids. 'Ca. N. limnes' MMS-21-122 has a 119 genome size of 1.24 Mbp and a genomic GC content of 41.5 %. It is auxotrophic for 120 reduced sulfur sources, several amino acids (proline, ornithine, histidine, betaine) and several vitamins (B1, B2, B5, B7, B12) and possesses rhodopsins (Table S7). 121 122 Members of the genus 'Ca. N. abundans' can be recognized by the presence of the diagnostic oligonucleotide sequence 5'-ACAAGAGGTTCGTCCGTCC-3' in the 23S 123 124 rRNA gene (positions 2669-2688, E. coli numbering). A complete genome of 'Ca. N. 125 limnes' MMS-21-122 is available at Genbank (CP016768). Phylogenetic analyses of 126 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and rhodpdopsin genes, as well as average nucleotide identities and protein similarities indicated that 127 128 the Candidatus taxon belongs to a novel genus ('Ca. Nanopelagicus') of the family 129 'Ca. Nanopelagicaceae' and the novel order 'Ca. Nanopelagicales'.

130 'Candidatus Nanopelagicus hibericus' [hi.be'ri.cus. L. masc. adj. hibericus 131 Spanish; referring to a high abundance in two Spanish reservoirs]. Represented by strain MMS-21-160, which was isolated from Lake Zurich, Switzerland. Rods with 132 133 lengths of 0.33±0.07 µm and diameters of 0.24±0.03 µm (Table S3, Figure S1). The 134 initial pure culture was lost after a few propagations to fresh medium; no growing 135 culture is available. The initial culture grew well in sterile lake water amended with 136 minimal carbon medium, vitamins and amino acids. 'Ca. N. hibericus' MMS-21-160 137 has a genome size of 1.22 Mbp and a genomic GC content of 42.4 %. It is 138 auxotrophic for reduced sulfur sources, several amino acids (ornithine, histidine, betaine) and several vitamins (B1, B2, B5, B7, B12) and possesses rhodopsins 139 140 (Table S7). A complete genome is available at Genbank (CP016771). Phylogenetic 141 analyses of 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and 142 rhodpdopsin genes, as well as average nucleotide identities and protein similarities 143 indicated the affiliation of the Candidatus taxon to the novel genus 'Ca.

144 Nanopelagicus' of the family 'Ca. Nanopelagicaceae' and the novel order 'Ca.

145 Nanopelagicales'.

146 'Candidatus Nanopelagicus abundans' [a.bun'dans. L. pres. part. abundans 147 abundant; referring to high global abundances]. Represented by strain MMS-IIB-91, 148 which was isolated from Lake Zurich, Switzerland. Rods with lengths of 0.46±0.47 µm 149 and diameters of 0.26±0.20 µm (Table S3, Figure S1). The initial pure culture was 150 lost after a few propagations to fresh medium; no growing culture is available. The 151 initial culture grew well in sterile lake water amended with inorganic basal medium, minimal carbon medium, vitamins and amino acids. 'Ca. N. abundans' MMS-IIB-91 152 153 has a genome size of 1.16 Mbp and a genomic GC content of 40.2 %. It is 154 auxotrophic for reduced sulfur sources, several amino acids (methionine, lysine, 155 ornithine, histidine, betaine), several vitamins (B1, B5, B7, B12), and possesses 156 rhodopsins (Table S7). A complete genome is available at Genbank (CP016779). Phylogenetic analyses of 476 concatenated protein sequences, 16S rRNA, 23S 157 158 rRNA, and rhodpdopsin genes, as well as average nucleotide identities and protein similarities indicated the affiliation of the Candidatus taxon to the novel genus 'Ca. 159 Nanopelagicus' of the family 'Ca. Nanopelagicaceae' and the novel order 'Ca. 160

161 Nanopelagicales'.

162 'Candidatus Planktophila limnetica' [Plank.to'phi.la. N.L.neut. n. plankton 163 plankton; N.L. fem. adj. phila friendly to; N.L. fem. n. Planktophila the friend (fem.) of plankton; lim.ne'ti.ca. N.L. fem. adj. limnetica pertaining to lakes]. According to the 164 165 previous description of a mixed culture (MWH-EgelM2-3.acl)(Jezbera et al., 2009) 166 that has a 100% identical 16S rRNA gene sequences with strain MMS-VB-114, we 167 propose the name 'Ca. P. limnetica' for strain MMS-VB-114 and make an amended description based on a complete genome. Represented by strain MMS-VB-114, 168 which was isolated from Lake Zurich, Switzerland. Curved rods with lengths of 169 170 0.38±0.10 µm and diameters of 0.25±0.06 µm (Table S3, Figure S1). The initial pure 171 culture was lost after a few propagations to fresh medium; no growing culture is 172 available. The initial culture grew well in sterile lake water amended with pyruvate, 173 urea, vitamins, and amino acids. 'Ca. P. limnetica' MMS-VB-114 has a genome size 174 of 1.33 Mbp and a genomic GC content of 45.0 %. It is auxotrophic for reduced sulfur sources and vitamins (B1, B5, B7), and possesses rhodopsins (Table S7). A 175 176 complete genome is available at Genbank (CP016782). Phylogenetic analyses of 177 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and rhodopsin genes, as well as average nucleotide identities and protein similarities indicated the affiliation 178 179 of the Candidatus taxon to the family 'Ca. Nanopelagicaceae' and the novel order 180 'Ca. Nanopelagicales'.

181 'Candidatus Planktophila dulcis' [dul'cis. L. fem. adj. dulcis sweet; referring to a 182 high diversity of sugar transporters and metabolism]. Represented by strain MMS-183 IIA-65, which was isolated from Lake Zurich, Switzerland. Curved rods with lengths of 184 0.44±0.09 µm and diameters of 0.26±0.03 µm (Table S3, Figure S1). The initial pure 185 culture was lost after a few propagations to fresh medium; no growing culture is 186 available. The initial culture grew well in sterile lake water amended with pyruvate, 187 urea, minimal carbon medium, vitamins, and amino acids. 'Ca. P. dulcis' MMS-IIA-65 188 has a genome size of 1.35 Mbp and a genomic GC content of 48.0 %. It is auxotrophic for reduced sulfur sources, betaine, and several vitamins (B1, B5, B7, 189 190 B12), and possesses rhodopsins (Table S7). It has a high number of different 191 carbohydrate transporters. A complete genome is available at Genbank (CP016777).

192 Phylogenetic analyses of 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and rhodopsin genes, as well as average nucleotide identities and protein 193 194 similarities indicated the affiliation of the Candidatus taxon to the genus 'Ca. 195 Planktophila' of the family 'Ca. Nanopelagicaceae' and the novel order 'Ca. 196 Nanopelagicales'. Two more strains (MMS-IA-53 and MMS-21-155) isolated from 197 Lake Zurich share high similarities with strain MMS-IIA-65 and are thus affiliated to 198 the same *Candidatus* species. This assignment is based on similar cell and genome 199 sizes, similar metabolism, phylogenetic analyses of 476 concatenated protein 200 sequences, 16S rRNA (100% identical), 23S rRNA, and rhodopsin genes, as well as 201 average nucleotide identities (<97%) and protein similarities (<94%). Complete genomes of 'Ca. P. dulcis' MMS-IA-53 and MMS-21-155 are available at Genbank 202 203 (CP016772, CP016770).

204 'Candidatus Planktophila sulfonica' [sul.fo'ni.ca. N.L. fem. adj. sulfonica pertaining to sulfonate; referring to sulfonate transporters]. Represented by strain 205 206 MMS-IA-56, which was isolated from Lake Zurich, Switzerland. Curved rods with 207 lengths of 0.50±0.14 µm and diameters of 0.28±0.06 µm (Table S3, Figure S1). The 208 initial pure culture was lost after a few propagations to fresh medium; no growing 209 culture is available. The initial culture grew well in sterile lake water amended with 210 inorganic basal medium, minimal carbon medium, vitamins, and amino acids. It is so far the only member of the order 'Ca. Nanopelagicales' that possesses membrane 211 212 transporters for sulfonates. 'Ca. P. sulfonica' MMS-IA-56 has a genome size of 1.34 Mbp and a genomic GC content of 48.6 %. It is auxotrophic for reduced sulfur 213 214 sources, betaine, and several vitamins (B1, B5, B7, B12), and possesses rhodopsins 215 (Table S7). A complete genome is available at Genbank (CP016773). Phylogenetic 216 analyses of 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and rhodopsin genes, as well as average nucleotide identities and protein similarities 217 218 indicated the affiliation of the Candidatus taxon to the genus 'Ca. Planktophila' of the 219 family 'Ca. Nanopelagicaceae' and the novel order 'Ca. Nanopelagicales'.

220 'Candidatus Planktophila versatilis' [ver.sa'ti.lis. L. fem. adj. versatilis versatile; 221 referring to high metabolic versatility]. Represented by strain MMS-IA-79, which was 222 isolated from Lake Zurich, Switzerland. Curved rods with lengths of 0.45±0.10 µm and diameters of 0.27±0.04 µm (Table S3, Figure S1). The initial pure culture was 223 224 lost after a few propagations to fresh medium; no growing culture is available. The 225 initial culture grew well in sterile lake water amended with inorganic basal medium, minimal carbon medium, vitamins, and amino acids. 'Ca. P. versatilis' MMS-IA-79 226 227 has a genome size of 1.33 Mbp and a genomic GC content of 48.2 %. It is 228 auxotrophic for reduced sulfur sources, betaine, and several vitamins (B1, B5, B7, 229 B12), and possesses rhodopsins (Table S7). A complete genome is available at 230 Genbank (CP016778). Phylogenetic analyses of 476 concatenated protein sequences, 16S rRNA, 23S rRNA, and rhodopsin genes, as well as average 231 nucleotide identities and protein similarities indicated the affiliation of the Candidatus 232 233 taxon to the genus 'Ca. Planktophila' of the family 'Ca. Nanopelagicaceae' and the 234 novel order 'Ca. Nanopelagicales'. Three additional strains (MMS-IIB-79, MMS-IA-235 105, and MMS-IIB-142) isolated from Lake Zurich share high similarities with strain 236 MMS-IA-79 and are thus affiliated to the same *Candidatus* species. This assignment 237 is based on similar cell and genome sizes, similar metabolism, phylogenetic analyses 238 of 476 concatenated protein sequences, 16S rRNA (100% identical), 23S rRNA, and 239 rhodopsin genes, as well as average nucleotide identities (<95%) and protein similarities (<90%). Complete genomes of 'Ca. P. versatilis' MMS-IIB-79, MMS-IA-240

- 105, and MMS-IIB-142 are available at Genbank (CP016774, CP016775,
- 242 CP016781).

243 'Candidatus Planktophila lacus' [la'cus. L. gen. masc. n. lacus of a lake]. Represented by strain MMS-21-148, which was isolated from Lake Zurich, 244 245 Switzerland. Rods with lengths of 0.41±0.10 µm and diameters of 0.30±0.06 µm 246 (Table S3, Figure S1). The initial pure culture was lost after a few propagations to 247 fresh medium; no growing culture is available. The initial culture grew well in sterile 248 lake water amended with minimal carbon medium, vitamins, and amino acids. 'Ca. P. 249 lacus' MMS-21-148 has a genome size of 1.46 Mbp and a genomic GC content of 250 47.5 %. It is auxotrophic for reduced sulfur sources, betaine, and several vitamins 251 (B1, B5, B7, B12), and possesses rhodopsins (Table S7). A complete genome is 252 available at Genbank (CP016769). Phylogenetic analyses of 476 concatenated 253 protein sequences, 16S rRNA, 23S rRNA, and rhodopsin genes, as well as average 254 nucleotide identities and protein similarities indicated the affiliation of the Candidatus 255 taxon to the genus 'Ca. Planktophila' of the family 'Ca. Nanopelagicaceae' and the 256 novel order 'Ca. Nanopelagicales'. Two additional strains (MMS-IIB-106 and MMS-257 IIB-60) isolated from Lake Zurich share high similarities with strain MMS-21-148 and 258 are thus affiliated to the same Candidatus species. This assignment is based on 259 similar cell and genome sizes, similar metabolism, phylogenetic analyses of 476 concatenated protein sequences, 16S rRNA (100% identical), 23S rRNA, and 260 261 rhodopsin genes, as well as average nucleotide identities (<94%) and protein similarities (<95%). Complete genomes of 'Ca. P. lacus' MMS-IIB-60 and MMS-IIB-262 106 are available at Genbank (CP016780, CP016783). 263

264 'Candidatus Planktophila vernalis' [ver.na'lis. L. fem. adj. vernalis belonging to spring; referring to high abundances in spring]. Represented by strain MMS-IIA-15, 265 which was isolated from Lake Zurich, Switzerland. Curved rods with lengths of 266 267 0.43±0.12 µm and diameters of 0.26±0.05 µm (Table S3, Figure S1). The initial pure 268 culture was lost after a few propagations to fresh medium; no growing culture is 269 available. The initial culture grew well in sterile lake water amended with pyruvate, 270 urea, minimal carbon medium, vitamins, and amino acids. 'Ca. P. vernalis' MMS-IIA-271 15 has a genome size of 1.36 Mbp and a genomic GC content of 45.7 %. It is 272 auxotrophic for reduced sulfur sources, serine, and several vitamins (B1, B5, B7, 273 B12), and possesses rhodopsins (Table S7). 'Ca. P. vernalis' MMS-IIA-15 can be recognized by the presence of the diagnostic oligonucleotide sequence 5'-274 275 AACTACTACCACACCGGTTCG-3' in the 23S rRNA gene (positions 1420-1441, E. 276 coli numbering). A complete genome is available at Genbank (CP016776). 277 Phylogenetic analyses of 476 concatenated protein sequences, 16S rRNA, 23S 278 rRNA, and rhodopsin genes, as well as average nucleotide identities and protein 279 similarities indicated the affiliation of the Candidatus taxon to the genus 'Ca. 280 Planktophila' of the family 'Ca. Nanopelagicaceae' and the novel order 'Ca. 281 Nanopelagicales'.

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**Figure S1:** Microphotographs and cell volumes ( $\mu$ m<sup>3</sup>) of isolates. The scale bar at the bottom left applies to all pictures.



**Figure S2:** Phylogenetic positioning of '*Ca*. Nanopelagicales' based on 16S rRNA genes. a, bootstrapped maximum likelihood tree of 16S rRNA genes; '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are shown on the nodes. b, 16S rRNA gene sequence similarity matrix. Species borders are marked with solid lines, genus borders with dashed lines. An asterisk indicates the positioning of the described mixed culture '*Ca*. P. limnetica'.



**Figure S3:** Average nucleotide identity (ANI) matrix of '*Ca*. Nanopelagicales'. Species borders are marked with solid lines, genus borders with dashed lines.



**Figure S4:** a, Average amino acid identity (AAI) matrix of '*Ca*. Nanopelagicales'. b, Protein similarity (>50% identity, >50% coverage) matrix of '*Ca*. Nanopelagicales'. Species borders are marked with solid lines, genus borders with dashed lines.



**Figure S5:** Phylogenomic tree with complete genomes of '*Ca*. Nanopelagicales' only. 462 concatenated conserved proteins were used to generate a maximum-likelihood phylogenetic tree. The genomes of '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are indicated by black, grey, and white circles on the nodes, and a colour key is shown on the left.



**Figure S6:** Phylogenomic tree with complete genomes of the phylum Actinobacteria. Forty-eight concatenated conserved proteins were used to generate a maximum-likelihood phylogenetic tree. The genomes of *Staphylococcus aureus* and *Listeria monocytogenes* were used as outgoup. Bootstrap values are indicated by black, grey or white circles on the nodes, and a colour key is shown on the left. The proposed novel order '*Ca*. Nanopelagicales' is highlighted in green.



**Figure S7:** Genome streamlining in '*Ca*. Nanopelagicales'. Number of predicted CDS, number of predicted sigma factor homologs, median size of intergenic spacers, and coding density versus genome size for all complete published genomes of Actinobacteria (n=610; data taken from RefSeq). '*Ca*. Nanopelagicales' and *Rhodoluna lacicola* are marked in different colours.



**Figure S8:** a, Bootstrapped phylogenetic tree of rhodopsin protein sequences of '*Ca*. Nanopelagicales' and other Actinobacteria. Xanthorhodopsin sequences of *Salinibacter ruber* and *Thermus aquaticus* were used as outgroup. Bootstrap values are indicated by coloured circles on the nodes, and a colour key is shown on the left. b, Arrangement of actinorhodopsin (AR) and carotenoid synthesis genes, i.e., beta carotene (*crtE, crtI, crtB, crtY*) and retinal (*blh*) biosynthesis genes. Genomes are arranged according to the phylogenomic tree in Fig. S5.



**Figure S9:** Bootstrapped maximum likelihood tree of 23S rRNA genes of '*Ca*. Nanopelagicales'. Target hits for the newly designed probes Pver-23S-1420 and Npel-23S-2669 are shown in brackets. '*Ca*. Aquiluna sp.' and *Rhodoluna lacicola* were used as outgroup. Bootstrap values are shown on the nodes.





Figure S10: Physico-chemical data from Lake Zurich

**Figure S11:** Redundancy analysis of environmental parameters explaining the variability in cell numbers of microbes affiliated to all '*Ca*. Nanopelagicales', '*Ca*. Nanopelagicus', and '*Ca*. P. vernalis' in Lake Zurich. temp, water temperature; picocyano, abundance of picocyanobacteria; irrad, irradiation; chloro, chlorophyll *a* associated with chlorophytes; diatom, chlorophyll *a* associated with diatoms; NH4, ammonium concentrations; O2, oxygen concentrations; PO4, phosphate concentrations; NO3, nitrate concentrations; depth, sampling depth.









Figure S13: Recruitment plots of time-series metagenomes from Lake Mendota, USA.

**Figure S14:** Whole-genome alignment of all 13 '*Ca*. Planktophila' strains. The genomes have been linearized for simplicity and are arranged in the same order as in the phylogenomic tree in Fig. 1b. Synteny and different degrees of sequence similarity are indicated by vertical lines connecting the genomes.



**Figure S15:** Whole-genome alignment of the three '*Ca*. Nanopelagicus' strains. The genomes have been linearized for simplicity and are arranged in the same order as in the phylogenomic tree in Fig. 1b. Synteny and different degrees of sequence similarity are indicated by vertical lines connecting the genomes. rRNA operons in the individual genomes are displayed as red arrows and tRNAs as short vertical lines. Genomic islands (GI) have been marked in different colours and numbered (see Table S10 for genes encoded in each island), except for '*Ca*. N. limnes' MMS-21-122, which displayed a large inversion in the genome. Red: genes encoding mainly cell wall biosynthesis and modifications; Yellow: genes encoding mainly membrane transport and / or carbohydrate metabolism.



Date	water temp	med	# wells inoc	# cells /well	# + wells	# pure cult	# acl	label	affiliation
25/05/2012	13.5	1	168	2.0	18	17	4	MMS-21	Pdul, Plac, Nlim, Nhib
22/05/2013	13.2	2	120	1.6	20	11	4	MMS-IA	Pdul, Psul, Pvers
23/05/2013	13.2	3	168	2.0	128	95	2	MMS-IIA	Pdul, Pvern
23/05/2013	13.2	2	168	2.0	92	49	5	MMS-IIB	Pvers, Plac, Nabu
15/07/2013	20.1	4	144	1.8	51	30	1	MMS-VB	Plim
SUM			768		309	202	16		

**Table S1:** Details of the isolation experiments conducted in Lake Zurich, Switzerland.

Abbreviations: water temp, water temperature; med, medium used for dilution-to-extinction isolation; # wells inoc, number of wells inoculated; # cells/well, number of cells inoculated per well; # + wells, number of wells with dense cultures; # pure cult, number of pure cultures; # acl, number of cultures affiliated to acl Actinobacteria; label, labeling suffix for pure cultures; Nhib, '*Ca.* Nanopelagicus hibericus'; Nlim, '*Ca.* Nanopelagicus limnes'; Nabu, '*Ca.* Nanopelagicus abundans'; Pdul, '*Ca.* Planktophila dulcis'; Plim, '*Ca.* Planktophila limnetica'; Pvern, '*Ca.* Planktophila vernalis; Psul, '*Ca.* Planktophila sulfonica', Pvers, '*Ca.* Planktophila versatilis', Plac, '*Ca.* Planktophila lacus'.

### Media:

- 1: filtered and autoclaved lake water (LW) + 10x MC#2 + vitamins (V) + amino acids (AA)
- 2: LW + MC#2 + MC#3 + inorganic basal medium (IBM) + stock4 + V + 10xAA
- 3: LW + pyruvate (50 µM) + urea (0.5 µM) + MC#4 + V + 10xAA
- 4: LW + pyruvate (50 µM) + urea (0.5 µM) + V + 10xAA

Minimal carbon medium MC#2: 1  $\mu$ M NH<sub>4</sub>Cl, 0.1  $\mu$ M K<sub>2</sub>HPO<sub>4</sub>, 55.5  $\mu$ M D-glucose, 66.6  $\mu$ M D-ribose, 217.2  $\mu$ M formate, 217.2  $\mu$ M ethanol, 84.7  $\mu$ M succinate, 131.5  $\mu$ M glycolate, 108.6  $\mu$ M glycerol, 45.2  $\mu$ M N-acetylglucosamine

MC#3: 62.1 µM putrescine, 39.3 µM spermidine, 66.6 µM D-xylose, 66.6 µM arabinose

MC#4: 30 µM NH<sub>4</sub>Cl, 50 µM oxaloacetate, 50 µM taurine, 1 µM betaine, 40 µM CaCl<sub>2</sub>

Amino acids (AA): 0.5  $\mu$ M isoleucine, leucine, lysine, methionine, alanine, phenylalanine, threonine, tryptophane, valine, arginine, histidine, asparagine, aspartate, cysteine, proline, serine, tyrosine, 2  $\mu$ M glutamine, glutamate, glycine

Vitamins (V): 0.593  $\mu$ M thiamine, 0.08  $\mu$ M niacin, 0.000074  $\mu$ M cobalamine, 0.005  $\mu$ M paraamino benzoic acid, 0.074  $\mu$ M pyridoxine, 0.081  $\mu$ M pantothenic acid, 0.004  $\mu$ M biotin, 0.004  $\mu$ M folic acid, 0.555  $\mu$ M myo-inositol

Inorganic basal medium (IBM, Hahn et al. 2003): 304  $\mu$ M MgSO<sub>4</sub>, 182  $\mu$ M Ca(NO<sub>3</sub>)<sub>2</sub>, 190  $\mu$ M NaHCO<sub>3</sub>, 20  $\mu$ M KCI, 16  $\mu$ M K<sub>2</sub>HP<sub>4</sub>, 17  $\mu$ M Na<sub>2</sub>EDTA-Fe, 0.1 ml TES

TES (trace element solution): 12  $\mu$ M FeCl<sub>3</sub>, 16  $\mu$ M H<sub>3</sub>BO<sub>3</sub>, 1  $\mu$ M MnCl<sub>2</sub>, 0.1  $\mu$ M ZnSO<sub>4</sub>, 0.04  $\mu$ M CuSO<sub>4</sub>, 0.04  $\mu$ M CoCl<sub>2</sub>, 0.03  $\mu$ M NaMoO<sub>4</sub>, 0.4  $\mu$ M NiCl<sub>2</sub>

#### Reference:

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			MiSoa	road	
sample	strain	library prep	reagents	filtering*	assembler
dampio	'Ca Planktonhila	TruSeg PCR-	1000000000000000000000000000000000000	Intering	
MMS-IIA-65	dulcis'	free	cycle	tnk	SPAdes-370
	'Ca Planktonhila	TruSeg PCR-	$v_{2}$ 500	ι, ρ, κ	
MMS-IA-53	dulcis'	free	cycle	tnk	SPAdes-362
	'Ca Planktonhila	TruSeg PCR-	v2 500	ι, ρ, κ	
MMS-21-155	dulcis'	free	cycle	tnk	a5 mised linux 20150522
11110 21 100	'Ca Planktophila	TruSeg PCR-	$v_{2}$ 500	ι, <b>ρ</b> , κ	
MMS-IA-56	sulfonica'	free	vcle	tnk	SPAdes-362
	'Ca Planktonhila	TruSeg PCR-	$v_{2}$ 500	ι, ρ, κ	01 Adds 0.0.2
MMS-IIB-76	versatilis'	free	cycle	tnk	SPAdes-362
	'Ca Planktonhila	TruSeg PCR-	$v_{2}$ 500	ι, ρ, κ	017/000 0.0.2
MMS-IA-79	versatilis'	free	cycle	tnk	SPAdes-370
	'Ca Planktophila	TruSeg PCR-	$v_{2}^{2}$ 300	ι, <b>ρ</b> , κ	
MMS-IA-105	versatilis'	free	cycle	tnk	a5 misea linux 20140604
	'Ca Planktophila	TruSea PCR-	$v_{2}$ 500	ι, <b>ρ</b> , κ	
MMS-IIB-142	versatilis'	free	cvcle	t. p. k	a5 miseg linux 20150522
	'Ca. Planktophila	TruSea PCR-	v2. 500	-, [- ,	···_·····
MMS-IIB-106	lacus'	free	cvcle	t. p. k	SPAdes-3.6.2
	'Ca, Planktophila	TruSea PCR-	v2. 500	217	
MMS-IIB-60	lacus'	free	cvcle	t. p. k	SPAdes-3.6.2
	'Ca. Planktophila	TruSea PCR-	v2. 500	-, [- ,	
MMS-21-148	lacus'	free	cvcle	-	SPAdes-3.7.0
	'Ca, Planktophila	TruSea PCR-	v2. 500		
MMS-VB-114	limnetica'	free	cycle	t, p	SPAdes-3.7.0
	'Ca. Planktophila	TruSeg PCR-	v2, 500	<i>i</i> <b>i</b>	
MMS-IIA-15	vernalis'	free	cycle	t, p, k	SPAdes-3.6.2
	'Ca. Nanopelagicus	TruSeg PCR-	v2, 500	· • •	
MMS-21-122	limnes'	free	cycle	-	SPAdes-3.7.0
	'Ca. Nanopelagicus	TruSeq PCR-	v2, 500		
MMS-21-160	hibericus'	free	cycle	-	a5 miseg linux 20150522
	'Ca. Nanopelagicus	TruSeq PCR-	v2, 300		
MMS-IIB-91	abundans'	free	cycle	t, p, k	a5_miseq_linux_20140604

<u>MMS-IIB-91</u> abundans' tree cycle t, p, k a5\_miseq\_linux\_20140604 \*t: trimmomatic-0.32.jar (MMS-IA-105, MMS-IA-105), trimmomatic-0.35.jar (others), parameters: PE ILLUMINACLIP:TruSeq3-PE-2.fa:2:30:10; p: prinseq-lite-0.20.4, parameters: -out\_format 3 range\_len 60-160 / 60-260 (MMS-IA-105, MMS-IA-105 / others) -range\_gc 30-80 -min\_qual\_mean 20 -ns\_max\_n 2 -derep 14 -derep\_min 2 -noniupac -trim\_qual\_right 18 -trim\_qual\_left 18 -lc\_method entropy -lc\_threshold 15 -trim\_tail\_left 3 -trim\_tail\_right 3 -trim\_right 1; k: kmernorm-1.0.5, parameters: -k 16 -c 2 -t 30.

The REPLI-g single cell kit (Qiagen, Venlo, Netherlands) was used for multiple displacement amplification (MDA). All pre-amplification steps were performed in a particle free environment dedicated to MDA. Fresh PCR-clean pipet tips were used for each MDA session and reaction tubes and PCR plates were UV-treated before usage. MDA was conducted according to the manufacturer's protocol with the following modifications: Reaction volumes were reduced to 12.5  $\mu$ l and lysates for 6 to 8 replicate MDA reactions were produced in a 0.5 ml reaction tube (8  $\mu$ l sample containing 2000 to 20000 cells, 6  $\mu$ l each of the reagents D2 and stop) and subsequently distributed among wells of 96 well plates each containing 10  $\mu$ l MDA reaction mix and SYBR I green (0.2x final concentration). Each MDA-reaction contained 2'000 - 20'000 cells.

taxonomy	strain	isolation date	genome size (Mbp)	eGC content (%)	# CDS	# tRNA	spacer length (bp)	coding density (%)	# σ factors
'Ca. Planktophila dulcis'	MMS-IIA-65	23.05.2013	1.35	48.0	1344	40	12	95.7	3
' <i>Ca</i> . Planktophila dulcis'	MMS-IA-53	22.05.2013	1.37	48.0	1356	40	12	95.7	3
' <i>Ca</i> . Planktophila dulcis'	MMS-21-155	25.05.2012	1.36	47.9	1361	40	11	95.7	3
' <i>Ca</i> . Planktophila sulfonica'	MMS-IA-56	22.05.2013	1.34	48.6	1336	37	12	96.0	4
'Ca. Planktophila versatilis'	MMS-IIB-76	23.05.2013	1.33	48.2	1318	40	14	95.3	4
'Ca. Planktophila versatilis'	MMS-IA-79	22.05.2013	1.33	48.3	1327	38	14	95.2	4
'Ca. Planktophila versatilis'	MMS-IA-105	22.05.2013	1.33	48.2	1329	40	14	95.4	4
'Ca. Planktophila versatilis'	MMS-IIB-142	23.05.2013	1.27	48.3	1258	39	14	95.5	4
' <i>Ca</i> . Planktophila lacus'	MMS-IIB-106	23.05.2013	1.38	47.8	1368	40	15	95.6	4
' <i>Ca</i> . Planktophila lacus'	MMS-IIB-60	23.05.2013	1.41	47.8	1389	40	16	95.5	5
' <i>Ca</i> . Planktophila lacus'	MMS-21-148	25.05.2012	1.46	47.5	1438	41	15	95.3	4
' <i>Ca</i> . Planktophila limnetica'	MMS-VB-114	15.07.2013	1.33	45.0	1333	41	10	96.0	4
' <i>Ca</i> . Planktophila vernalis'	MMS-IIA-15	23.05.2013	1.36	45.7	1355	39	10	95.7	4
' <i>Ca</i> . Nanopelagicus limnes'	MMS-21-122	25.05.2012	1.24	41.5	1216	38	11	95.7	3
<i>'Ca</i> . Nanopelagicus hibericus'	MMS-21-160	25.05.2012	1.22	42.4	1211	38	13	95.4	4
' <i>Ca</i> . Nanopelagicus abundans'	MMS-IIB-91	23.05.2013	1.16	40.2	1150	39	14	95.3	2

Table S3: Details of the sequenced strains of planktonic 'Ca. Nanopelagicales'.

	length		width		volume		CC (fg		
strain	(µm)	s.d.	(µm)	s.d.	(µm³)	s.d.	C cell <sup>-1</sup> )	s.d.	n
MMS-IIA-65	0.44	±0.09	0.26	±0.03	0.019	±0.007	7.1	±2.3	71
MMS-IA-53	0.44	±0.14	0.28	±0.04	0.023	±0.012	8.5	±3.7	35
MMS-21-155	0.43	±0.12	0.25	±0.02	0.018	±0.007	6.8	±2.4	77
MMS-IA-56	0.50	±0.14	0.28	±0.06	0.026	±0.021	9.3	±6.4	23
MMS-IIB-76	0.49	±0.12	0.25	±0.03	0.021	±0.009	7.6	±2.8	18
MMS-IA-79	0.45	±0.10	0.27	±0.04	0.022	±0.010	8.1	±3.1	51
MMS-IA-105	0.47	±0.15	0.29	±0.05	0.027	±0.014	9.6	±4.4	64
MMS-IIB-142	0.50	±0.47	0.28	±0.29	0.026	±0.024	9.3	±8.8	23
MMS-IIB-106*	0.50	±0.14	0.30	±0.04	0.029	±0.015	10.3	±4.5	10
MMS-IIB-60	0.47	±0.08	0.30	±0.07	0.028	±0.013	10.0	±4.0	18
MMS-21-148	0.41	±0.10	0.30	±0.06	0.025	±0.014	8.8	±4.4	128
MMS-VB-114	0.38	±0.10	0.25	±0.06	0.016	±0.009	6.0	±3.0	52
MMS-IIA-15	0.43	±0.12	0.26	±0.05	0.020	±0.012	7.3	±3.9	52
MMS-21-122	0.45	±0.09	0.25	±0.03	0.018	±0.009	6.9	±2.9	85
MMS-21-160	0.33	±0.07	0.24	±0.03	0.012	±0.005	4.8	±1.7	49
MMS-IIB-91*	0.46	±0.47	0.26	±0.26	0.020	±0.020	7.5	±7.4	5

**Table S4:** Cell sizes of different strains and *in-situ* in Lake Zurich. Asterisks indicate values based on <10 measured cells and should be treated with caution.

probe (samp- ling date)	length (µm)	s.d.	width (µm)	s.d.	volume (µm³)	s.d.	CC (fg C cell <sup>-1</sup> )	s.d.	n
Npel-2669 (18/07/2012)	0.35	±0.09	0.25	±0.04	0.014	±0.008	5.4	±2.6	97
Pver-1420 (18/07/2012)	0.37	±0.12	0.24	±0.04	0.015	±0.011	5.7	±3.4	91
Pver-1420 (15/05/2013)	0.35	±0.09	0.23	±0.03	0.012	±0.005	4.8	±1.8	87

s.d., standard deviation; CC, carbon content; n, number of cells measured

organism	habita	cell volume t (μm³)	source	genome size (Mbp	)GC (%)	coding density (%)	rho	. ref.
'Ca. Nanopelagicus'	fw	0.012-0.020	cultures	1.16-1.24	40.2-42.4	95.3-95.7	х	this study
'Ca. Planktophila'	fw	0.016-0.029	cultures	1.27-1.46	45.0-48.6	95.2-96.0	х	this study
'Ca. Nanopelagicales'	fw	N.A.	SAGs	1.10-1.66	41.4-47.6	95.8-96.3	х	1, 2
'Ca. Nanopelagicus'	fw	N.A.	MAGs	1.38-2.2	40.4-42.1	96.1-96.4	х	3
' <i>Ca</i> . Actinomarina minuta'	mar	0.013	MAG	0.8-1.03	33.4		х	4
' <i>Ca</i> . Pelagibacter' (SAR11)	mar	0.025-0.045	cultures	1.24-1.48	28.6-32.3	93.6-97	х	5, 6, 7
LD12	fw	0.017	SAGs	1.03-1.39	29-30	95.5-96.5	х	8, 9
<i>'Ca</i> . Methylopumilus planktonicus'	fw	0.030-0.075	cultures	1.28-1.35	37.0-37.3	94.9-95.3	х	10, unpubl.
OM43	mar	0.023	cultures	1.30-1.37	35.4-37.9	95.0-96.7	х	11
Rhodoluna lacicola	fw	0.053	cultures	1.43	51.5	93.7	х	12
' <i>Ca</i> . Aquiluna sp.'	fw	N.A.	cultures	1.36	51.7	93.5	х	13

Table S5: Cell sizes and genomic details of genome-streamlined microbes.

fw, freshwater; mar, marine; rho., rhodopsins; ref., references; N.A., data not available

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probe	specificity	sequence (5' to 3')	target (rRNA, 5' position)	FA%
Npel-23S-2669	' <i>Ca</i> . Nanopelagicus' (acl-B1)	ACA AGA GGT TCG TCC GTC C	23S, 2669	60
Npel-C1	competitor 1 for Npel- 23S-2669	AC <u>Y</u> AGA GGT TCG TCC GTC C		
Npel-C2	competitor 2 for Npel- 23S-2669	ACA AGA GGT TCG TCC <u>A</u> TC C		
Npel-C3	competitor 3 for Npel- 23S-2669	AC <u>Y</u> AGA GGT TCG TCC <u>A</u> TC C		
Npel-H1	helper 1 for Npel-23S- 2669	CGG TCC TCT CGT ACT AGG GAC AGC		
Npel-H2	helper 2 for Npel-23S- 2669	GTG CTY CTG GCG RAA CAA CCG ACA C		
Npel-H3	helper 3 for Npel-23S- 2669	CYT TCC RAA CGT TGC WAA TCG GCC		
Pver-23S-1420	'Ca. Planktophila	AAC TAC TAC CAC ACC GGT TCG	23S, 1420	55
	vernalis' (acl-A7)			
Pver-C1	competitor 1 for Pver- 23S-1420	AAC TAC TAC CAC ACC GGT TCA		
Pver-C2	competitor 2 for Pver- 23S-1420	AAC TAC TAC CAC ACC GG <u>G</u> TCG		
Pver-C3	competitor 3 for Pver- 23S-1420	AAC TAC TAC <u>A</u> AC ACC GGT TCG		
Pver-C4	competitor 4 for Pver- 23S-1420	AAC TAC TAC <u>A</u> AC ACC GG <u>G</u> TC <u>A</u>		
Pver-H1	helper 1 for Pver-23S- 1420	CAT TAG TGG RTT CGT YAT GGG CGA ATT A		
Pver-H2	helper 2 for Pver-23S- 1420	AGC CAT CCA CCC ACG CRG CTT		
Pver-H3	helper 3 for Pver-23S- 1420	CTG TGT CAC ACC ATT GCT T		
Acl-852 <sup>1</sup>	<i>'Ca</i> . Nanopelagicales' (acl lineage)	AAT GCG TTA GCT GCG TCG CA	16S, 852	55
Acl-852-H1 <sup>1</sup>	Helper for Acl-852	AAA CCG TGG AAG GTY CSC ACA ACT AG		
Acl-852-H2 <sup>1</sup>	Helper for Acl-852	TCC CCA GGC GGG GCR CTT		

Abbreviation: FA%, formamide concentration required for the CARD-FISH hybridization buffer.

Reference:

[1] Warnecke F, Sommaruga R, Sekar R, Hofer JS, Pernthaler J (2005). Abundances, identity, and growth state of Actinobacteria in mountain lakes of different UV transparency. Appl Environ Microbiol 71: 5551-5559