



Microbial network, phylogenetic diversity and community membership in the active layer across a permafrost thaw gradient

Journal:	<i>Environmental Microbiology and Environmental Microbiology Reports</i>
Manuscript ID	Draft
Journal:	Environmental Microbiology
Manuscript Type:	EMI - Research article
Date Submitted by the Author:	n/a
Complete List of Authors:	Mondav, Rhiannon; Uppsala University, Ecology and Genetics, Limnology; University of Queensland, Chemistry and Molecular Biosciences McCalley, Carmody; Rochester Institute of Technology, Thomas H. Gosnell School of Life Sciences; University of New Hampshire, Earth Systems Research Center Hodgkins, Suzanne ; Florida State University , Department of Earth Ocean and Atmospheric Science Frolking, Steve ; University of New Hampshire, Institute for the Study of Earth, Oceans, and Space Saleska, Scott; University of Arizona, Ecology and Evolutionary Biology Rich, Virginia; University of Arizona, Department of Soil, Water and Environmental Science; Ohio State University, Microbiology Department Chanton, Jeff; Florida State University , Department of Earth Ocean and Atmospheric Science Crill, Patrick; Stockholm University, Department of Geological Sciences
Keywords:	microbial ecology, microbially-influenced global change, environmental signal/stress responses, microbial communities

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Manuscripts

1 **Microbial network, phylogenetic diversity and community membership in the**
2 **active layer across a permafrost thaw gradient**

3 Authors:

4 Rhiannon Mondav^{1,2*}, Carmody K McCalley^{3,4,+}, Suzanne B Hodgkins⁵, Steve
5 Frolking⁴, Scott R Saleska³, Virginia I Rich^{6,^}, Jeff P Chanton⁵, Patrick M Crill⁷

6 Affiliations:

7 ¹ Department of Ecology and Genetics, Limnology, Uppsala University, Uppsala
8 75236, Sweden

9 ² School of Chemistry and Molecular Biosciences, University of Queensland,
10 Brisbane 4072, Australia

11 ³ Department of Ecology and Evolutionary Biology, University of Arizona, Tucson
12 85721, USA

13 ⁴ Institute for the Study of Earth, Oceans, and Space, University of New Hampshire,
14 Durham NH 03824, USA

15 ⁵ Department of Earth Ocean and Atmospheric Science, Florida State University,
16 Tallahassee 32306-4320, USA

17 ⁶ Department of Soil, Water and Environmental Science, University of Arizona,
18 Tucson 85721, USA

19 ⁷ Department of Geology and Geochemistry, Stockholm University, Stockholm 10691,
20 Sweden

21 *Corresponding author:

22 RMondav, Norbyvagen, Uppsala University, Uppsala, SE75236 Sweden, +46 (0)7
23 2731 7286, rhiannon.mondav@ebc.uu.se

24 Running Title: Microbial community across a permafrost thaw gradient

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26 Current addresses: ⁺ Thomas H. Gosnell School of Life Sciences, Rochester Institute
27 of Technology, Rochester, New York 14623, USA; [^] Department of Microbiology, The
28 Ohio State University, Columbus 43210, USA.

29 **Summary**

30 Biogenic production and release of methane (CH₄) from thawing permafrost has the
31 potential to be a strong source of radiative forcing. We investigated changes in the
32 active layer microbial community of three sites representative of distinct permafrost
33 thaw stages at a palsa mire in northern Sweden. The palsa sites with intact
34 permafrost, and low radiative forcing signature had a phylogenetically clustered
35 community dominated by *Acidobacteria* and *Proteobacteria*. The bog with thawing
36 permafrost and low radiative forcing signature was dominated by hydrogenotrophic
37 methanogens and *Acidobacteria*, had lower alpha diversity, and midrange
38 phylogenetic clustering, characteristic of ecosystem disturbance affecting habitat
39 filtering, shifting from palsa-like to fen-like at the waterline. The fen had no underlying
40 permafrost, and the highest alpha, beta and phylogenetic diversity, was dominated
41 by *Proteobacteria* and *Euryarchaeota*, and was significantly enriched in
42 methanogens. The mire microbial network was modular with module cores
43 consisting of clusters of *Acidobacteria*, *Euryarchaeota*, or *Xanthomonadales*. Loss of
44 underlying permafrost with associated hydrological shifts correlated to changes in
45 microbial composition, alpha, beta, and phylogenetic diversity associated with a
46 higher radiative forcing signature. These results support the complex role of
47 microbial interactions in mediating carbon budget changes and climate feedback in
48 response to climate forcing.

49

50 **Introduction**

51 Modern discontinuous permafrost is found in regions with a mean annual air
52 temperature between -5 °C and +2 °C and where the insulating properties of peat
53 enable persistence of permafrost above freezing temperatures (Shur and Jorgenson,
54 2007; Seppälä, 2011). As these regions experience climate change-induced
55 warming, they are approaching temperatures that are destabilizing permafrost
56 (Schuur et al., 2015). Permafrost degradation typically leads to significant loss of soil

57 carbon (C) through erosion, fire and microbial mineralisation (Osterkamp et al., 2009;
58 Mack et al., 2011). The area of permafrost at risk of thaw in the next century has
59 been estimated to be between 10^6 and 10^7 km², with the quantity of C potentially lost
60 ranging from $1-4 \times 10^{14}$ kg (McGuire et al., 2010; Schuur et al., 2015). Increasing
61 plant production in thawed systems may partially compensate for this loss but this is
62 poorly defined (Hicks Pries et al., 2013). Thus, the potential positive feedbacks to
63 climate change are not well constrained, and will vary depending on the emission
64 ratio of the greenhouse gases (GHGs) carbon dioxide (CO₂) and methane (CH₄), and
65 the time scale considered (Dorrepaal et al., 2009; Nazaries et al., 2013). Changes in
66 microbial community membership (e.g. methanogen to methanotroph ratio) will be a
67 significant factor controlling the CO₂ to CH₄ emission ratio (Hodgkins et al., 2014).
68 To examine the relationship between permafrost thaw, shifting GHG emissions, and
69 the microbial community, we investigated a natural *in situ* thaw gradient at Stordalen
70 Mire, northern Sweden, on the margin of the discontinuous permafrost zone.
71 Permafrost thaw has been causatively linked to changes in topography, vegetation,
72 and GHG emission at Stordalen (Christensen, 2004; Johansson et al., 2006;
73 Bäckstrand et al., 2008, 2010; Johansson and Åkerman, 2008), and elsewhere
74 (Turetsky et al., 2007; Olefeldt et al., 2013). Currently, the Mire is a partially
75 degraded complex of elevated, drained hummocks (palsas with intact permafrost)
76 and wet depressions (bogs with thinning permafrost and fens without any
77 permanently frozen ground), each representing different stages of thaw (Fig S1), and
78 each characterised by distinct vegetation (Bhiry and Robert, 2006; Johansson et al.,
79 2006). Changes in topography and vegetation (proxy for thaw) have been tracked
80 through the last 40 years and show a decrease in area covered by palsa, an
81 expansion of fens and a variable area covered by bogs (Christensen et al., 2004;
82 Malmer et al., 2005; Johansson et al., 2006). Part of the known palsa-cycle is the
83 external carving and internal collapse of palsas into the surrounding bog or fen,
84 though the time scale at which this happens varies depending on whether dome-

85 palsas or palsa-complexes/plateaus (as seen at Stordalen) are considered (Railton
86 and Sparling, 1973; Zoltai, 1993; Sollid and Sørbel, 1998; Gurney, 2001; Turetsky et
87 al., 2007; Seppälä, 2011; O'Donnell et al., 2012; Liebner et al., 2015). At Stordalen
88 complete collapse due to absence of permafrost results in a fen or lake, while partial
89 collapse due to permafrost thinning results in a bog (Johansson et al., 2006).
90 Photographs, topographical survey and GHG data comparing Stordalen in the 1970's
91 and 1980's to 2000's and 2010's show that for the particular area studied here the
92 palsa has degraded both externally and internally (Fig S1, S2). Bogs (sphagnum or
93 semi-wet) have expanded within the perimeter of the palsa-complex and around its
94 southern edge, while fens (eriophorum, wet, or tall-graminoid) have encroached from
95 the north, east, and west having converted the bog that once existed on the western
96 and eastern edges of the palsa, along with increases in GHG emissions (Rydén et
97 al., 1980; Malmer et al., 2005; Johansson et al., 2006; Bäckstrand et al., 2010). The
98 majority of bog samples were taken from the subsided section within the palsa-
99 complex, while the fen samples were taken from the western side of the complex,
100 which was recorded as a bog 40 years ago. As permafrost continues to disappear
101 from Stordalen over the coming decades, subsidence of the surface will likely
102 increase, driving the creation of more transient bog-type communities and
103 degradation into fens (Christensen, 2004; Parviainen and Luoto, 2007; Johansson
104 and Åkerman, 2008; Fronzek et al., 2010; Bosio et al., 2012; Jones et al., 2016). It is
105 also predicted that this region could be free of permafrost as early as 2050
106 (Parviainen and Luoto, 2007; Fronzek et al., 2010). The "natural experiment"
107 underway in the Mire presents a model ecosystem for investigating climate-driven
108 changes in lowland permafrost regions with high cryosequestered-C (Masing et al.,
109 2009).

110 As a model system, the Mire has been intensively studied over the last several
111 decades for permafrost thaw impacts on plant communities, hydrology, and
112 biogeochemistry – providing rich context for interpreting microbial communities. The

113 seasonally thawed peat layer (active layer) of Stordalen's palsas is drained, aerobic,
114 ombrotrophic (rain-fed) and isolated from nutrient-rich groundwater. The palsa sites'
115 low plant productivity and aerobic decomposition make them net emitters of
116 appreciable CO₂ and no CH₄, with a net C balance (NCB) of - 30 mgC/m²/day
117 (negative value indicates net C emissions, positive indicates net uptake; Bäckstrand
118 et al., 2010). In contrast, the bog sites (semi-wet in Johansson *et al* 2006) are
119 physically lower and collect rainwater, leading to partial inundation, and are
120 dominated by layered bryophytes (typically *Sphagnum* spp.). Although this results in
121 less lignin (a recalcitrant C compound not produced by sphagnum), C degradation is
122 still slow due to sphagnum's higher phenolic content (Freeman et al., 2004) and
123 extremely poor C:N ratios of up to 70:1 (Hodgkins et al., 2014). Partially anoxic
124 conditions permit microbial fermentation and CH₄ production (Nilsson and Bohlin,
125 1993; Hobbie et al., 2000). Mire bog sites have the lowest radiative forcing signature
126 (NCB in CO₂ eq. of - 8 mgC/m²/day; Bäckstrand et al., 2010), as fixation of C in the
127 bog peat is high compared to emission of C gases. Finally, the fully-thawed fen sites
128 (tall-graminoid in Johansson *et al* 2006) are the most subsided and are
129 minerotrophic (groundwater-fed). Vegetation succession results in dominance of
130 graminoids (sedges, rushes, reeds), with a subsequent shift in the litter preserved as
131 peat. Some graminoids enhance gas transport between inundated soil and the
132 atmosphere (Chanton et al., 1993) and due to high productivity, contribute
133 appreciable fresh labile organic litter and exudates (Wagner and Liebner, 2009). High
134 productivity results in the fens being the Mire's biggest gross C-sinks of the Mire,
135 however their high CH₄ emissions result in a net warming potential 7 and 26 times
136 greater than the palsa and bog respectively (NCB in CO₂ eq. of - 213 mgC/m²/day,
137 Bäckstrand et al., 2010; Christensen et al., 2012).

138 Here we explore the relationship between the biogeochemical differences among
139 palsa, bog, and fen and the active layer microbial community, via a temporal (over a
140 growing season) and spatial (across habitats, and with depth through the active

141 layer) community survey using SSU rRNA gene amplicon sequencing, pore-water
142 chemistry, peat chemistry, stable C isotope analyses, and CH₄ flux. Previous work
143 has demonstrated that permafrost thaw has an overall impact on Stordalen Mire's
144 microbiota (Mondav & Woodcroft et al, 2014; Hodgkins et al., 2014; McCalley et al.,
145 2014). Here, we deepen understanding of this impact by addressing the following
146 descriptors with respect to climate-induced thaw and correlated environmental
147 parameters: 1) dominant phyla, 2) beta diversity, 3) assemblage alpha diversity, 4)
148 phylogenetic distance to identify drivers of assembly processes, 5) community
149 network, and 6) C-cycling phylotype distribution.

150

151 **Results and discussion**

152 *Dominant phyla of each thaw stage*

153 Dominant phyla can inform on geochemical correlations and community functionality,
154 reflecting overall habitat conditions. *Acidobacteria* and *Proteobacteria* were
155 ubiquitous phyla across the Mire (Fig 1). Dominant palsa phyla also included
156 *Actinobacteria* and Candidate bacterial phylum "WD272" (WPS-2), the abundances
157 of which decreased across the thaw gradient (palsa>bog>fen). Surface bog samples
158 had similar community composition to palsa samples being dominated by
159 *Acidobacteria* and *Proteobacteria* suggesting that site classification, which was
160 identified by vegetation, is not the only environmental correlate important to microbial
161 community assembly in the Mire. Bog samples at or below the waterline (midpoint
162 and deepest) retained similar proportions of *Acidobacteria* and *Actinobacteria* as
163 palsa samples. *Proteobacteria* however, were relatively less abundant, likely due to
164 lower C lability (Goberna et al., 2014; Hodgkins et al., 2014), and were replaced by
165 *Euryarchaeota* in deeper anoxic samples. The shift in phylum ratios from palsa-like
166 to fen-like supports the transitional nature of this thawing site. The most abundant
167 phyla in the fen were *Euryarchaeota*, *Proteobacteria*, *Bacteroidetes*, and *Chloroflexi*

168 followed by *Acidobacteria*. *Woesearchaeota* (DHVEG-6) were only detected in fen
169 samples.

170 **<Fig 1, 110 mm wide>**

171 *Beta diversity and environmental correlates of site assemblages*

172 Microbial assemblages were significantly different (anosim $r^2_{adj} = 0.90$, $p < 0.001$) at
173 the sample OTU level between sites. Whole community analysis by nonmetric
174 multidimensional scaling (NMDS) (Fig 2a, stress = 0.082, $r^2 = 0.99$) clustered
175 samples by site. Samples separated along the primary NMDS axis according to
176 hydrological states with ombrotrophic (palsa and bog) samples clustered together left
177 of the origin and the minerotrophic fen samples to the far right. The secondary axis
178 separated samples according to depth from surface with the two ombrotrophic site
179 samples diverging from each other with depth. The palsa and surface bog
180 assemblages overlapped at both OTU (Fig 2) and phyla level (Fig 1). Sharing of
181 species across the palsa and surface bog (aerobic ombrotrophic) is likely due to
182 local dispersal and seen in other methanogenic soils (Kim and Liesack, 2015). Local
183 dispersal mechanisms include transport by burrowing lemmings, palsa dome runoff
184 washing microbes into lower altitudes, local aerial dispersal (Bowers et al., 2011).
185 Ubiquitous deposition across the mire via precipitation may also contribute (Christner
186 et al., 2008). These more ubiquitous microbes likely persist through environmental
187 filtering including oxygenation, acidity, ombrotrophy, and bryophyte presence (Brettar
188 et al., 2011; King et al., 2012).

189 C-fixing autotrophs were associated with all samples except the deepest bog and
190 fen. Bacterial methanotrophs, while distributed across the Mire, were not associated
191 with deep fen or deep bog samples, supporting their known preference for aerobic
192 and microaerobic habitats. Both autotrophs and methanotrophs were unique to
193 individual sites with only a few OTUs shared across the surface bog and palsa
194 samples. The majority of methanogenic phylotypes were clustered exclusively with

195 fen samples, though a secondary cluster of methanogens while more tightly
196 associated to deep bog samples were detected across both bog and fen.
197 Relative increases in δD_{H_2O} and δD_{CH_4} were correlated with palsa sample
198 assemblages (Fig 2a&b). This is indicative of methanotrophy as supported both by
199 the detection of methanotrophic phylotypes but also by the negative CH_4 flux (uptake
200 of CH_4 from atmosphere) recorded by the palsa auto-chambers. Increased peat C:N
201 ratio, pore-water C:N, and porewater DOC were correlated with the bog
202 assemblages supporting previous studies finding higher DOC in run-off from
203 ombrotrophic regions of the Mire (Nilsson, 2006; Kokfelt et al., 2010) Pore-water N
204 content decreased in relation to deeper bog samples. Increased pH, CH_4 flux from
205 the auto-chambers, flux $\delta^{13}C_{CH_4}$ from the autochambers, pore-water $\delta^{13}C_{CH_4}$, and
206 water table depth (WTD) were positively correlated with fen samples (Fig 2a). Pore-
207 water CH_4 concentration, pore-water CO_2 concentration, CO_2 gas from peat
208 samples, and pore-water total C increased with deeper (below watertable) bog and
209 fen samples. Increased CH_4 in porewater and flux measurements were correlated to
210 presence of detected methanogens (Fig 2b) supporting established linkage between
211 detected abundance and metabolic activity of these microbes (Mondav et al., 2014).

212 **<Fig 2, 110 mm wide>**

213 *Microbial assemblage alpha diversity*

214 The number of unique OTUs per normalised (N=2000) sample ranged from 309 in
215 the merged anoxic bog samples from august 2011 up to 1226 in the combined
216 surface fen samples from june 2011 (Fig 3), with a cross-Mire mean OTU richness of
217 721 (s.d. 204 OTUs). The total richness was 9700 across the Mire for the 42 merged
218 and normalised samples. The percentage of OTUs that could not be taxonomically
219 classified either to or below order level was typical, at 40%, supporting that
220 environmental microbes are still appreciably under-characterised. Total, archaeal,
221 and bacterial assemblage alpha diversity as measured by richness (observed
222 OTUs), Fisher alpha (Fisher et al., 1943), Shannon entropy (Shannon and Weaver,

223 1949), and Heip's evenness (Heip et al., 1974) varied between thaw stages, with
224 there being a significant difference ($p < 0.05$) between sites as measured by Kruskal-
225 Wallace test (K-W) (Fig 3). For total assemblage alpha diversity the bog had lowest
226 (richness and fisher alpha), and was significantly lower than the palsa (shannons
227 entropy and heips evenness) K-W post-hoc test for significance (K-Wmc, $p < 0.001$).
228 The fen had highest archaeal alpha diversity (richness, fisher, and Shannon) while
229 the palsa had most even archaeal assemblage (K-Wmc, $p < 0.001$). Apart from the
230 exception of archaeal evenness the bog had lower or lowest alpha diversity of the
231 three sites. Archaeal evenness in the bog site covers a wide range from
232 assemblages with high evenness similar to that found in palsa samples but also
233 includes assemblages that were more highly dominated than those found in the fen
234 (Fig 3). Evenness is an important property of methane producing communities where
235 higher evenness of fen assemblies, compared to bog samples, may constitute a
236 feedback mechanism by which higher CH_4 production is enabled (Galand et al.,
237 2003; Godin et al., 2012). Depth of sample was related to decreases in all alpha
238 diversity metrics (total and bacterial, range: $-0.46 < \rho < -0.36$, $p < 0.01$). Bacterial
239 ($r^2_{\text{adj}} = 0.92$, $p < 0.001$) and total richness ($r^2_{\text{adj}} = 0.92$, $p < 0.001$) decreased with depth
240 in the bog while Archaeal richness increased ($r^2_{\text{adj}} = 0.69$, $p < 0.001$) (Fig 3, Equations
241 S1 a-c). Higher DOC correlated to lower richness (total $\rho = -0.76$, bacterial $\rho = -0.70$,
242 and archaeal $\rho = -0.74$: $p < 0.001$) (Fig S3). Bacterial richness was correlated to
243 decreased porewater CO_2 ($\rho = -0.60$, $p < 0.001$, Fig S3). Archaeal richness was
244 positively correlated to distance below water-table ($\rho = 0.83$, $p < 0.001$, Fig S3). The
245 number of singletons observed in each site directly correlated with richness and
246 varied between sites ($r^2_{\text{adj}} = 0.97$, $p < 0.001$, Fig S4, Equations S1 d-g).

247 <Fig 3, 80 mm wide>

248 *Site assembly dynamics*

249 Links have been drawn between a community's diversity, functional and phylogenetic
250 redundancy (robustness), and the community's ability to maintain function during

251 change (resistance), and to recover original state if the disturbance is removed
252 (resilience) (Shade et al., 2012; Venail and Vives, 2013). While the degree to which
253 phylogenetic and/or functional diversity influence resistance is not fully elucidated,
254 recent studies support (Werner et al., 2011; Singh et al., 2014), these are important
255 properties to consider with climate change and permafrost thaw, two interacting
256 press disturbances at Stordalen (Shade et al., 2012; Hayes et al., 2014). A press
257 disturbance is a change in the environment that persists for a long period of time, in
258 comparison to a pulse disturbance which is a change that decreases suddenly after
259 a short period of time. Phylogenetic robustness can be measured by various
260 diversity relationships including phylogenetic distance between OTUs (PD); nearest
261 taxon index (NTI), which examines phylogenetic clustering of closely related
262 phylotypes; and mean relatedness index (NRI) which examines variance of
263 phylotype distance within an assemblage. Further, these indices can indicate the
264 relative degree to which stochastic or deterministic processes contribute to
265 community assembly (Wang et al., 2013). The Mire as a whole and each site
266 individually had positive correlation ($r^2_{adj} = 0.73$ to 0.94 , $p < 0.001$) between overall
267 phylogenetic diversity (PD) and richness (Eqs S2a-d, Fig S5). Because PD and
268 richness were auto-correlated, subsequent analyses examined PD/OTU so as to
269 examine only the difference due to diversity and not an artefact of abundance
270 counts. Fen PD/OTU was significantly ($p < 0.001$) higher than both bog and palsa
271 assemblages (Fig 4). Measuring assemblage net relatedness (NRI, equivalent to -
272 sesMPD) by OTU phylogenetic distance from sample mean as generated by the null
273 model examines clustering over a whole tree. Greater emphasis is placed on
274 changes towards the tree root compared to other measures such as NTI, which
275 examines diversity at the tips of the phylogenetic tree. Negative values of NRI
276 indicate expansion of the tree via increased branching at higher-level tree nodes i.e.
277 even-dispersal, while positive values indicate filling in of internal phylogenetic tree
278 nodes i.e. clustering. Palsa assemblages had uniformly high NRI, ($0.1 < NRI < 0.4$,

279 Fig 4) indicative of phylogenetic clustering. A deterministic factor associated with
280 clustering specific to the palsa (and surface bog) is dry-ombrotrophy which increases
281 habitat isolation in heterogeneous soil environments (Kraft et al., 2007; Jones et al.,
282 2009; Stegen et al., 2013; Quiroga et al., 2015). Fen assemblage NRIs were
283 significantly lower than the other sites (KW, $p < 0.001$, Fig 4) and were neutral to
284 negative. Bog assemblage NRIs varied ($0 < \text{NRI} < 0.4$) from neutral in deeper
285 samples to higher than some Palsa in surface samples (Fig 4, Fig S5, Eq S4). The
286 lower NRI in the fen indicates assemblages have broader representation across the
287 bacterial and archaeal domains with less clustering than predicted by the null model
288 i.e. phylogenetic even dispersal. Even dispersal can indicate an assemblage less
289 affected by deterministic processes such as habitat filtering or isolation (Webb et al.,
290 2002; Horner-Devine and Bohannon, 2006), while being more affected by stochastic
291 processes such as dispersal and drift or controversially, by competition (Mayfield and
292 Levine, 2010). All assemblage NTI values were above zero, i.e. more clustered than
293 predicted with the null model, a result seen in early stage successional forests and
294 freshwater mesocosms, (Horner-Devine and Bohannon, 2006; Whitfeld et al., 2012).
295 The bog and fen had greater phylogenetic tip clustering than the palsa (KW $p < 0.05$,
296 Fig 4) and was, in the bog, related to depth (Eq S3). Greater tip-clustering as seen in
297 the bog and fen indicates greater genomic diversification within 'species'-
298 populations. This could be achieved through horizontal gene transfer (HGT) or
299 endogenous mutation enabling more closely related organisms to coexist (Goberna
300 et al., 2014), though there is not yet data to address effective population sizes, or the
301 relative frequencies of HGT and endogenous genome mutation in these habitats.
302 The ratio of NRI to NTI indicates the level (tree:tip) at which phylogenetic diversity is
303 affected by assembly processes. Fen assemblages had significantly greater
304 clustering at the whole tree level than tree tip when compared to the other sites
305 (NRI/NTI, Fig 4, $p < 0.01$). Conversely, the palsa assemblages displayed greater
306 clustering towards the tree's tips. The bog NRI/NTI was in-between the palsa and the

307 fen. This shift in where the assemblage diversity lies supports clustering through
308 local species divergence being a property of the ombrotrophic mire sites while even-
309 dispersal is more prevalent in the minerotrophic fen. The clustering shift from tip to
310 whole tree diversity is also seen at a smaller scale within the bog site (Fig S5, Eq S5)
311 where surface samples have higher tip clustering and deeper samples have more
312 evenly distributed diversity. That the mid depth bog samples were taken at the
313 waterline supports that a main factor regulating this shift is inundation by water.
314 Phylogenetic diversity of Mire assemblages as measured through PD, NRI, and NTI
315 showed the bog grouping alternately with the palsa or the fen supporting that the bog
316 may be an intermediate site undergoing transition from a palsa-like assemblage to
317 fen-like assemblage due to a shift from ombrotrophy to minerotrophy. These four
318 phylogenetic distance analyses support that each site has a unique overall
319 phylogenetic diversity profile, thus giving support to there being differences in
320 assembly and evolution of the microbial community across sites and therefore thaw
321 stages (Stegen et al., 2012).

322 Examining correlations between environmental parameters and phylogenetic
323 diversity can potentially inform on the relative contribution, directly or indirectly, of
324 environmental to assembly processes. Phylogenetic clustering related to
325 environmental filtering is predicted to be evident in environments with poor nutrient
326 availability or parameters considered to increase selection such as high acidity.
327 Phylogenetic evenness, conversely, is expected to be evident in environments with
328 high resource availability where competition becomes more dominant in assembly
329 processes. Increasing soil pH, ratio of methanogens to methylotrophs, CH₄ flux, and
330 distance below watertable were significantly correlated to increasing PD/OTU and
331 decreasing NRI and NRI/NTI ($\rho \geq \pm 0.60$, $p < 0.001$, Table S2, Fig S6) i.e. associated
332 with phylogenetic even dispersal. Greater depletion of ¹³C_{CH₄}, higher porewater
333 DOC, and higher porewater C:N ratios were correlated to decreasing PD/OTU and
334 increasing NRI, and NRI/NTI i.e. phylogenetic clustering ($\rho \geq \pm 0.60$, $p < 0.001$, Table

335 S2). Only bog and fen porewater DOC and C:N were analysed as there was
336 insufficient moisture in the palsa samples. Conditions associated with environmental
337 filtering (acidity) and isolation (ombrotrophy) may here be linked to phylogenetic
338 clustering (Kraft et al., 2007). Stochastic mechanisms specific to the fen include
339 inflow of runoff from the raised palsa and bog, minerotrophy, and local water mixing
340 which reduces isolation (Putkinen et al., 2012). Conditions associated with
341 phylogenetic even-dispersal may therefore be linked to warming potential (increased
342 CH₄ flux) and increases in acetotrophic methanogens (less depletion of ¹³C in CH₄
343 emissions) in methanogenic soils. Clustering is an emerging characteristic of soil
344 communities (Lozupone and Knight, 2007; Auguet et al., 2010) that is currently
345 connected to habitat filtering (Kraft et al., 2007; Shade and Handelsman, 2012)
346 though some recent evidence also supports a role for biotic filtering (Goberna et al.,
347 2014). The correlations between higher phylogenetic diversity, including even-
348 dispersal, and CH₄ flux corroborate findings from reactors and environmental
349 systems (Werner et al., 2011; Yavitt et al., 2011) that the structure of microbial
350 communities may be significant to global CH₄ budgets.

351 **<Fig 4, 80 mm wide>**

352 *Network topology and community interactions*

353 Microbial networks can be described mathematically by topological indices. Common
354 indices include degree, modularity, betweenness, and closeness. Degree describes
355 the level of connectedness between phylotypes by counting the number of
356 phylotypes that co-occur. Modularity identifies if sub-networks of co-occurrence exist
357 within a larger community network and is thought to be an indicator of resilience.
358 Betweenness Centrality provides information on how critical a phylotype is to the
359 connectedness of a network. Closeness Centrality describes how closely a phylotype
360 is connected to all others in the same module. Redundancy (e.g similar metabolic
361 strategies) is also a useful descriptor of co-existing organisms. Degree, closeness
362 and redundancy in microbial networks provide information on the community's

363 robustness and, potentially, ability to resist change. OTU Table B₂₀₀₀ had 9 700
364 unique phylotypes, 93% sparsity, average inverse Simpsons (n_{eff}) of 122 per sample
365 which was too sparse to obtain meaningful network information from. Restricting the
366 dataset to OTUs that were present in at least 15 samples (one third of total) left 257
367 unique OTUs, a table with 49% sparsity, and an average n_{eff} of 26, statistics that
368 provide assurance that network interactions could be correctly identified while
369 minimising type I errors (Friedman and Alm, 2012; Berry and Widder, 2014; Weiss et
370 al., 2016). Retaining only OTU pairs that were significantly correlated in at least two
371 of the network analyses from MENA, fastLSA, CoNet (Pearsons, Spearman, Bray,
372 KBL), and SparCC reduced the dataset to 123 OTUs with 265 significant pairwise
373 correlations (Table S3). The network had low degree (average 4.3 per node), a
374 maximum path of 14, and low checkerboard (C-score= 0.387). The C-score was
375 compared to the null model and found to be different (null model C-score = 0.338)
376 with a 97.5% CI and $p < 0.001$ supporting non-random OTU co-occurrence patterns
377 and the presence of a microbial network. However the work by Berry and Widder
378 (2014) examining the effect of filtering on sensitivity indicates that type II errors may
379 be as high as 0.5 based on 16-20% of the network potentially being habitat
380 specialists (Table S4). Analysis and visualisation of this network (Fig 5) revealed a
381 community consisting of 8 modules.

382 The detection of phylotypes in different environments, and their network topological
383 description, enables exploration of community metabolic roles of microbial lineages
384 (Foster et al., 2008). Highly connected phylotypes sometimes called hubs or
385 keystones (high degree, high closeness, low betweenness) are predicted to perform
386 key metabolic steps within microbial communities. In the identified network at
387 Stordalen only hubs with high degree, high closeness but with high betweenness
388 were identified (Fig 5, Table S3 & S4). Due to their high betweenness and close
389 phylogenetic relatedness to adjacent OTUs these hubs exhibit qualities associated
390 with redundancy or 'niche overlap' and therefore may have little effect if removed i.e.

391 they are unlikely keystone species. Keystone species, while sometimes described
392 statistically as hubs (Faust and Raes, 2012; Berry and Widder, 2014), are
393 ecologically those that would cause (disproportionate) disruption to a network if lost.
394 In the larger modules of the network, there were a few phylotypes (putative
395 keystones marked on Fig 5, Table S3, S4) that if removed would fragment the
396 network and/or were the only phylotypes identified as associated with a critical
397 metabolic process. The loss of any of the identified keystone phylotypes from any of
398 the three sites, past or future, could affect significant changes in C, N, S, or Fe
399 cycles at Stordalen. Identification of keystone species is problematic if there is a high
400 degree of type II errors or, as is expected with environmental phylogenetic-amplicon
401 surveys, there is limited information available on their phenotypes. Statistically, the
402 predicted keystones in this network cover the full range of betweenness and
403 closeness scores but none had high degree, supporting that high degree is a poor
404 predictor of 'keystoneness' in soil microbial communities.

405 Module 'A' consisted of 25 phylotypes (19 Acidobacteria, 4 Actinobacteria, 2
406 Euryarchaeota) that were dominant in the bog (Fig 5, Table S4). Two of the
407 *Acidobacteriaceae* (subgroup I Acidobacteria) phylotypes were identified as hubs.
408 The potential keystones (based on topology) were another *Acidobacteraceae* and
409 the less abundant of the two *Ca. Methanoflorens* (RCII) phylotypes. *Ca.*
410 *Methanoflorens* is a hydrogeno/formatotrophic methanogen of the *Methanocellales*
411 order that prefer low H² concentration, and are oxygen and acid tolerant (Sakai et al.,
412 2010; Lü and Lu, 2012; Mondav et al., 2014; Lyu and Lu, 2015). One of the three
413 *Acidobacteria* identified at genus level was a phylotype of *Ca. Solibacter*. *Solibacter*
414 are capable of degrading complex-C molecules such as cellulose, hemicellulose,
415 pectin, chitin, and starch (Ward et al., 2009; Pankratov et al., 2012), a useful
416 phenotype in the sphagnum-peat of the bog. Another was an *Acidobacterium* that
417 may be able to reduce ferric iron (Coupland and Johnson, 2008). The last of the
418 genus level identified was an *Acidcapsa* which likely preferentially utilise bi-products

419 of sphagnum degradation such as xylose or cellobiose, but if necessary could
420 directly degrade starch and pectin (Kulichevskaya et al., 2012; Matsuo et al., 2016).
421 The four *Acidimicrobiales* identified might contribute to Fe-cycling and likely capable
422 of degrading complex polymers (Kämpfer, 2010; Stackebrandt, 2014). The remainder
423 of module A (unknown *Acidobacteraceae*) are likely chemoheterotrophs that either
424 degrade sphagnum derived polymers or their hydrolysed bi-products (Campbell,
425 2014). Eighty percent of module A phylotypes had significant positive correlations (ρ
426 $\geq \pm 0.60$, $p_{ad} < 0.001$, Table S5) with porewater N, DOC concentration.

427 Module B^{total} was the most phylogenetically and phenotypically mixed module, it was
428 further divided into two sub-modules: B and B'. Thirty-four of module B phylotypes
429 were only detected in the fen, eight in both bog and fen samples, and three in both
430 fen and palsa samples. Module B hubs included both of the *Bacteroidetes*
431 phylotypes, a *Woesearchaeota* (DHVEG-6) and a *Methanosaeta*, and a
432 *Methanoregula* and a *Bathyarchaeota* (Msc. Crenarchaeota Grp) (Fig 5, Table S4).
433 The *Bacteroidetes* phylotypes are likely anaerobic, organotrophs with a preference
434 for sugar molecules (Krieg et al., 2010). *Methanosaeta* are obligate acetotrophic
435 methanogens and their higher substrate (acetate) affinity may be one mechanism
436 enabling them to compete in the fen (Westermann et al., 1989; Ferry, 2010; Liu et al.,
437 2010) as would the higher pH and reduction in inhibitory phenolics compared to
438 conditions in the bog. A *Methanoregula* phylotype was also identified as a hub, the
439 only potentially hydrogenotrophic (&/or formatotrophic) methanogen in module B. Its
440 requirement for and greater tolerance of acetate, might assist *Methanosaeta* by
441 using up some of the available acetate (Smith and Ingram-Smith, 2007; Bräuer et al.,
442 2011; Oren, 2014b). No phenotypic data is yet available for *Woesearchaeota*. and
443 the *Bathyarchaeota* so far described were either methanogenic, methanotrophic, or
444 organoheterotrophic (Butterfield et al., 2016; Lazar et al., 2016). So while no
445 predictions can be made as to whether these phylotypes are e.g. methanogens, it is
446 evident that they are important methanogenic-community members. One of the

447 methanogenic phylotypes connected to the Woesearchaeota and Bathyarchaeota
448 was a methylotrophic-methanogen of the *Methanomassiliicoccus* genus (Borrel et
449 al., 2013). Module B and B' were connected by the co-occurrence of one of these
450 *Bathyarchaeota* and a *Methanobacterium* respectively, both possible keystones. The
451 other putative keystones were an actinobacterial phylotype of the *Gaiellales* order of
452 unknown but likely heterotrophic metabolism, and the chemolithoautotrophic
453 *Nitrospiraceae* (Daims, 2014). It is probable that the *Nitrospiraceae* phylotype, which
454 was not detected in the bog, contributes to the module by C-fixation (Daims, 2014).
455 Two other module B phylotypes not in the bog were a chemoorganotrophic
456 (putative complex-C degrader) *Myxococcales* genus *Haliangium* (Kim and
457 Liesack, 2015), and an uncultured *Chloroflexi* KD4-96. The thirteen OTUs of
458 module B' were either not detected or detected at very low abundance in the palsa
459 (Fig 5, Table S4). All B' phylotypes were methanogens from either the
460 hydrogeno/formatotrophic *Methanobacterium* genus or the metabolically flexible
461 *Methanosarcina* genus (Oren, 2014a, 2014c). Module B', and to a lesser extent B,
462 displays the functional redundancy and phylogenetic clustering characteristic of soil
463 communities and in particular methanogenic soils (Embree et al., 2014). Module B
464 phylotypes (87 %) were strongly correlated ($\rho \geq \pm 0.60$, $p_{\text{adj}} < 0.001$, Table S5) to
465 phylogenetic even dispersal (NRI), pH, CH₄ flux, and decreasing porewater N.
466 Module C consisted mainly of *Xanthomonadales* an order of obligate aerobic
467 *Gammaproteobacteria* capable of degrading complex organic molecules and
468 participating in methyl / H syntrophic partnerships with methylotrophs (Kim and
469 Liesack, 2015). The co-presence of the keystone methylotrophic proteobacterial
470 *Methylotenera* (Doronina et al., 2014) phylotype supports the possibility that such
471 partnerships occur at Stordalen and are common in the aerobic partition of
472 methanogenic soils. A second keystone phylotype was the C-fixing verrucomicrobial
473 methanotroph *Methylacidiphilum* (Hedlund, 2010; Sharp et al., 2013). The final
474 keystone was the putative ferric iron reducing, H₂/CO₂ producing *Acidocella*. which

475 may also have a syntrophic partner within module C (Coupland and Johnson, 2008;
476 Johnson and Hallberg, 2008). Most phylotypes (12 of 16) were correlated to
477 increased clustering (NRI) and acidity (Table S5).

478 Module D consisted of 14 OTUs, nine *Acidobacteria*, four *Gammaproteobacteria*
479 (*Xanthomonadales*), and one *Verrucomicrobia* all of whom were dominant in the
480 palsa. Three keystones were identified, a subdivision 2 *Acidobacteria*, a *Bryobacter*
481 (subdivision 3 *Acidobacteria*), and a putatively ferric iron reducing *Acidobacterium*
482 (subdivision 1 *Acidobacteria*) phylotype. *Bryobacter* are chemoheterotrophs with a
483 preference for sugars (Dedysh et al., 2016). This module appears to have members
484 that together degrade complex-C molecules, polysaccharides and simple sugars
485 (Hedlund, 2010; Dedysh et al., 2016; Yang et al., 2016). Twelve of the phylotypes
486 had significant correlation ($\rho \geq \pm 0.60$, $p_{\text{adj}} < 0.001$, Table S5) with distance above
487 frozen ground/watertable. Module E had five *Syntrophobacterales* phylotypes,
488 putative producers of substrates for methanogenesis (acetate, H₂, and formate) and
489 capable of N fixation (Embree et al., 2014; Lin et al., 2014), with four from the
490 *Smithella* genus and one from *Syntrophus*. None of these five were detected in palsa
491 samples. The final OTU of the module was detected across the mire and identified
492 as belonging to the *Verrucomicrobia* OPB35 soil group which is thought to degrade
493 polysaccharides (Hedlund, 2010; Yang et al., 2016). All of module E phylotypes were
494 significantly correlated with increasing CH₄ flux and decreasing NRI ($\rho \geq \pm 0.60$,
495 $p_{\text{adj}} < 0.001$, Table S5).

496 Overall, the clustering of closely related phylotypes in modules is consistent with the
497 phylogenetic diversity results. The network associations of methanogen phylotypes
498 were complex and modular, with most falling into sub-network B' (present in bog and
499 fen and low abundance in palsa), followed by B (fen only, except for the more
500 cosmopolitan *Methanomasilliococcus* phylotype). The two outlying *Ca.*
501 *Methanoflorens* methanogens in module A (fen and bog) were associated with
502 *Acidobacteria* and *Actinobacteria* which are typical of peat environments and The

503 presence of permafrost fits the known distribution of this genus (Mondav et al., 2014)
504 and its predicted phenotype. The putative C-fixing autotrophs were scattered through
505 the network and only moderately connected, supporting their phylogenetically-based
506 assignments to a primary trophic role.

507 **<Fig 5, 169 mm wide>**

508 *Known C-cycling phylotypes*

509 Distinct shifts in relative abundances of putative methanogens, and methylotrophs
510 were evident across the Mire. Relative abundance of methanogens increased across
511 the thaw gradient (palsa<bog<fen, Table S6) (K-Wmc, $p < 0.001$). Methanogens were
512 strongly associated with the deepest bog and fen samples (Table S6, Fig S7). All but
513 one of the obligate acetotrophic methanogens (*Methanosaeta*) were detected
514 exclusively in the fen (Fig S7), the other one was found in a single palsa sample.
515 Apart from the anomalous *Methanosaeta* detected in the palsa, this is consistent with
516 reported acetotrophic sensitivity to low pH due to reduction of the ΔG (Gibbs free
517 energy) of the acetotrophic methanogenesis pathway (Kotsyurbenko et al., 2007).
518 The possibility of a divergent metabolism may explain the presence of the
519 *Methanosaeta* in the palsa. Other methanogenic phylotypes detected in apparently
520 aerobic samples (above the water line) may have been enabled by micro-anaerobic-
521 habitats, oxidative resistance as seen in some *Methanocellales* (Angel et al., 2011),
522 or association with an anaerobic host gut (Paul et al., 2012). Putative
523 methano/methylotrophic phylotypes were distributed across all samples (Fig S7) and
524 were highest in the bog samples (K-Wmc, $p < 0.01$, Table S6) likely accounting for the
525 lower CH₄ flux despite the abundance of methanogens. Some methylotrophs were
526 detected below the waterline in bog and fen samples (Fig S7) and likely exist in
527 micro-aerobic spaces enabled by plant root gas transport (Colmer, 2003). The
528 relative ratio of methanogen to methanotroph phylotypes differed significantly
529 between sites (K-Wmc, $p < 0.01$), increasing across the thaw gradient
530 (palsa<bog<fen, Table S6). Due to the polyphyletic distribution of autotrophic and

531 methanotrophic metabolisms the assignment of function requires identification to
532 family or genera level, while most methanogens can be identified at class or order
533 level. It is therefore likely that abundances and richness of autotrophic and
534 methanotrophic microbes described here are underestimated more than
535 methanogens. The shifting C-cycling phylotype patterns described here, especially
536 the methanogen to methanotroph ratio provide detail of biogenic methane production
537 and consumption that support reported site C-budgets (Bäckstrand et al., 2010), CH₄
538 emissions and CH₄ isotope ratios (McCalley et al., 2014) from Stordalen.

539 The presence of permafrost maintains the ombrotrophic, acidic environment (Natali
540 et al., 2011; Tveit et al., 2013) which correlated with a diverse, rich, phylogenetically-
541 clustered, and autotroph-abundant palsa community. The bog sites (thawing
542 permafrost) had significantly lower richness, diversity, evenness and estimated
543 population size than the palsa, the fen, and also other peat bog sites (Lin et al.,
544 2012; Serkebaeva et al., 2013; Tveit et al., 2013). Collectively, this suggests a
545 structural response to ecosystem disturbance (Degens et al., 2001) caused by site
546 inundation as a consequence of subsidence caused by permafrost thaw (Rydén et
547 al., 1980; Johansson and Åkerman, 2008). Correlations between diversity estimates
548 (alpha, beta, and phylogenetic) and distance of sample above or below the water
549 table support that site inundation (and therefore thaw) is a mechanistic driver of
550 community structure and function and that deterministic processes were the main
551 drivers of community composition and assembly in this and other bogs (Quiroga et
552 al., 2015). Complete loss of permafrost in the fen was correlated to assemblages
553 with highest richness, alpha diversity, beta diversity, and phylogenetic even-
554 dispersion.

555 As the permafrost thaws, causing subsidence, Palsas and transitory bogs at
556 Stordalen mire are expected to give way to fens. At Stordalen the transition from bog
557 to fen is accompanied by community diversification and proliferation of methanogens
558 and a decrease in the relative ratio of methanotrophs. It appears at Stordalen, as

559 found elsewhere under similar conditions (Liebner et al., 2015), that at the point of
560 inundation a regime shift is initiated and that beyond this point, the community does
561 not recover but instead shifts towards a new stable state as found in the fen. The fen
562 assemblage has qualities indicative of greater stability (high redundancy, evenness,
563 richness, and diversity) coincident with an altered C-budget of dramatically higher
564 warming potential. The combination of changes predicted by climate models, the
565 trajectory suggested in biogeochemical and vegetation surveys of the Mire, and the
566 details of the microbial community C-cycling shifts detailed here suggest that the
567 mires along the Torneträsk valley will increasingly add to radiative climate forcing via
568 increased CH₄ flux over the coming decades as more land is converted to fen.
569 Longer-term outcomes of climate change in this region are projected to eventually
570 include some drying (e.g. terrestrialisation, the conversion of wetlands to drier areas
571 (Payette et al., 2004)) and expansion of the dwarf forests (Rundqvist et al., 2011). If
572 these fundamental habitat shifts occur, they are extremely likely to drive further
573 changes in the microbial communities and C budgets of the region.

574

575 **Experimental Procedures**

576 *Field site and sampling*

577 Samples were taken from the active layer of an individual palsa thaw sequence in
578 Stordalen Mire, subarctic Sweden (68.35N, 19.04E), with three stages of permafrost
579 degradation evidenced by topographical and vegetative characteristics (intact,
580 thawing, and thawed; Fig S1). The intact permafrost was represented by a raised
581 section of the palsa (palsa site, Fig S2); the thawing transition site was an
582 elevationally depressed region within the palsa (bog site); and the thawed permafrost
583 was a thermokarst feature with no detectable permafrost and thus no apparent
584 seasonal active layer (fen site) (Rydén et al., 1980; Malmer et al., 2005). The palsa
585 site was an ombrotrophic, drained, raised peat (altitude 351 m.a.s.l (Jackowicz-
586 Korczyński et al., 2010)) with tundra vegetation including *Betula nana* and *Empetrum*

587 *hermaphroditum*, interspersed with *Eriophorum vaginatum*, *Rubus chamaemorus*,
588 lichens, and mosses. The bog site was a wet ombrotrophic depression sunken ~1 m
589 below the palsa. Vegetation was predominately *Sphagnum spp.* with *E. vaginatum*.
590 Water table varied seasonally from 5 cm above to 30 cm below the vegetation
591 surface and was perched above the local groundwater. The average pH of the
592 ombrotrophic sites was 4.2 +/- 0.3 sd. The fen site was a minerotrophic, waterlogged
593 fen ~2 m below the palsa, vegetation was dominated by *Eriophorum angustifolium*
594 with some *Sphagnum spp.* and *Equisetum spp.*, and open water was present. The
595 water table was always within 5 cm of the peat surface, sometimes being above it,
596 and average pH of 5.7 +/- 0.1 sd.

597 Soil cores were taken in August/September 2010 and June, July, August, and
598 October 2011 (Table S1, Table S7). On each of the five sampling dates, between two
599 and four cores were taken from each of the three sites (Fig S1). In 2010 four cores
600 were taken from palsa and bog sites, in October 2011 two cores were taken from the
601 fen site, and all remaining sampling dates and locations had three cores sampled.
602 Samples cut from cores taken from the same site, at either the same depth in cm or
603 the same ecologically significant depth (e.g. depth relative to water table) were
604 designated technical replicates. Samples taken at different depths were analysed as
605 treatments (Samples taken at different depths were selected based on ecologically
606 pre-determined indicators such as at the water table for the bog, Fig S2). Porewater,
607 peat, flux and isotope measurements taken simultaneously to the microbial samples
608 were described in Mondav et al. (2014), Hodgkins et al. (2014) and McCalley et al
609 (2014) but are detailed again in the Supplementary Methods.

610 *SSU rRNA gene amplicon sequencing and analysis*

611 Microbial community was surveyed by small subunit (SSU) rRNA gene amplicon
612 sequencing (Mondav et al., 2014). Briefly, DNA was amplified with tagged primers for
613 V6-V8 region of the SSU rRNA gene with the 926F (AACTYAAAKGAATTGRCGG)
614 and 1392wR (ACGGGCGGTGWGTRC) primers, in duplicate reactions, pooled, and

615 sequenced with the 454 Ti GS (LifeSciences, Carlsbad). Sequences are available
616 from SRA under accession SRA096214 (McCalley et al., 2014; Mondav et al., 2014),
617 SRR numbers and primer details are listed in detail in Table S1. Sequences were
618 cleaned (pre-processed) with MacQIIME v1.9.1 then analysed at an operational
619 taxonomic unit (OTU) of 97% identity. A detailed description of pre-processing
620 methods can be found in the Supplementary Methods. Cleaned sequences were
621 assigned taxonomy using the open picking method and the SILVA database and a
622 phylogenetic tree made with Fasttree2 and manually rooted between the archaeal
623 and bacterial domains (Caporaso, Bittinger, et al., 2010; Caporaso, Kuczynski, et al.,
624 2010; Edgar, 2010; Price et al., 2010; Huson and Scornavacca, 2012; McDonald et
625 al., 2012; Quast et al., 2013). Phylotype lineage obtained by assignment of reads to
626 taxon identity was utilised to assign putative C-metabolism. Phylotypes that were
627 assigned to lineages with known methanogen, methanotroph/methylotroph, and C-
628 fixing members were manually checked before C-metabolism was assigned.

629 *Microbial assemblages*

630 Phylum level analysis was obtained by collapsing the normalised OTU table in
631 Qiime, and phyla detected in more than one sample and also present at over 1% in
632 at least one sample were graphed in MS Excel. To investigate dissimilarity of sample
633 and site assemblages a non-parametric ordination (NMDS) was done in R v3.3.1 (R-
634 Core-team, 2011) using the RStudio v0.99.903 (RStudio, 2012) IDE with the vegan
635 package v2.4-1 (Oksanen et al., 2013) and plotted using gplots v3.0.1 and with
636 scales v0.2.3 (Warnes et al., 2011). Environmental variables and parameters were
637 fitted to the NMDS and factors with significant ($p < 0.05$) correlation plotted. Alpha
638 diversity metrics were generated in Qiime (richness, singletons, Shannon diversity,
639 Fisher alpha, Heip's evenness, Simpson's dominance and Chao1 estimates) and
640 analysed in R using the non-parametric Kruskal-Wallis (K-W) followed by post-hoc
641 testing with Kruskal-Wallis multiple comparison (K-Wmc) testing using R package
642 pgirmess v1.6.4 (Giraudoux, 2012). Correlation analyses were done using non-

643 parametric Spearman and linear regression and differences between sites were
644 checked for significance with the largest p value obtained reported. Images were
645 processed for publication in Inkscape 0.91. Analysis of the phylogenetic diversity and
646 distance (PD, NRI, NTI) were calculated using the distance tree output from QIIME,
647 and correlation and equations calculated in R with picante v1.6-0 (Gotelli, 2000;
648 Faith, 2006; Kembel et al., 2010). OTUs present in less than 15 samples were
649 removed and the resultant OTU table analysed for pairwise interactions in MENA
650 (Deng et al., 2012), fastLSA (Durno et al., 2013), CoNet v1.1.1 (Faust et al., 2012)
651 and SparCC (Friedman and Alm, 2012). All networks were loaded into R and OTU
652 pairs that were identified as present in at least two of the four networks were retained
653 and the network analysed in CytoScape v3.4.0. See Supplementary Methods for
654 detailed description of all methods.

655

656 Supplementary Information is available as a separate download and includes
657 Supplementary Methods, Figs S1-S7, Equations S1-S5, and Tables S1-S7.

658

659 **Acknowledgements**

660 Many thanks to Andrew C Barnes, Brian Lanoil, and James Prosser for critical
661 comments on a previous version of the manuscript. Sincere thanks to the two
662 anonymous reviewers who helped me greatly improve this manuscript. RM was
663 supported by an Australian Postgraduate Award Scholarship and a Swedish
664 Vetenskapsrådet grant. JPC, PMC, SF, SH, CKM, SRS, and VIR were supported by
665 the US Department of Energy, Office of Biological and Environmental Research
666 under the Genomic Science program (Award DE-SC0004632).

667

668 **Author contributions**

669 JPC, PMC, SF, SRS, VIR designed the project. RM designed and performed all
670 bioinformatics analyses and visualizations. RM wrote the paper in consultation with
671 all authors.

672

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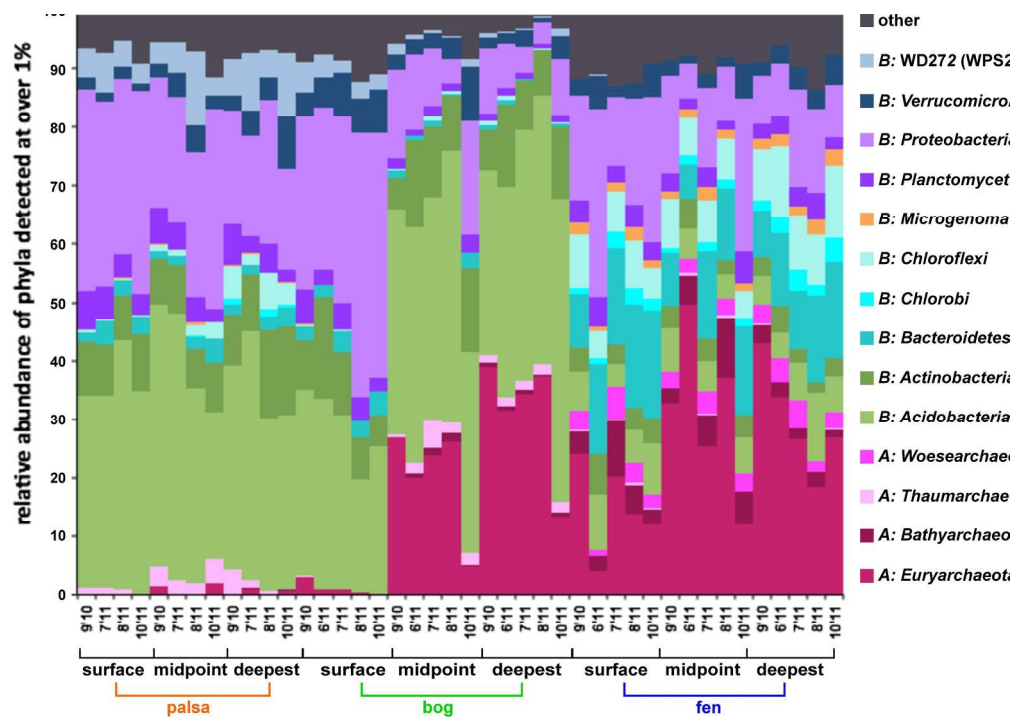


Fig 1 Mean relative abundance of phyla present at over 1 %. Samples grouped by site, then depth, then date. Date is marked along the horizontal axis as m'yy.

210x148mm (300 x 300 DPI)

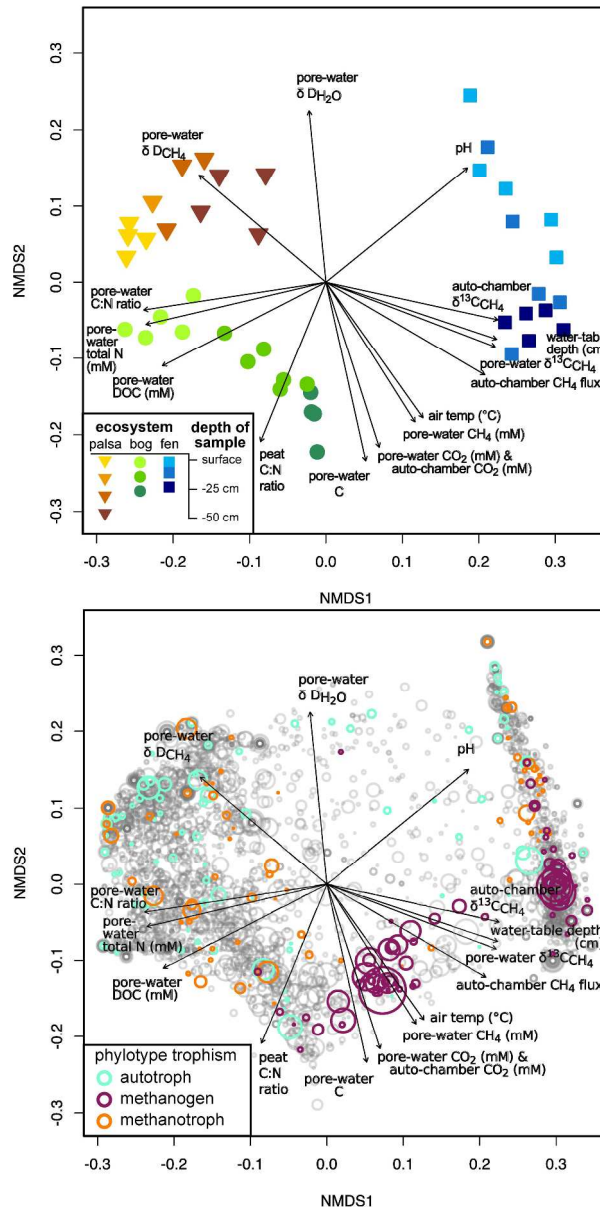


Fig 2 NMDS analysis of sample dissimilarity in community phylotype space, and environmental correlates (Stress = 0.0818, $r^2 = 0.993$). a) The clustering of sites based on their community composition; sites are distinguished by shape and colored to show depth of sample; b) The relative positional contribution of phylotypes to this clustering, with phylotypes plotted as circles with diameter scaled to the log of the mean abundance, and colored to show putative C-cycle metabolism. Plotted vectors on a) & b) are measured environmental variables that had significant correlation to differences in assemblage composition ($p < 0.01$), with the terminal arrow indicating the direction of strongest change without reference to sign (+ or -).

285x572mm (300 x 300 DPI)

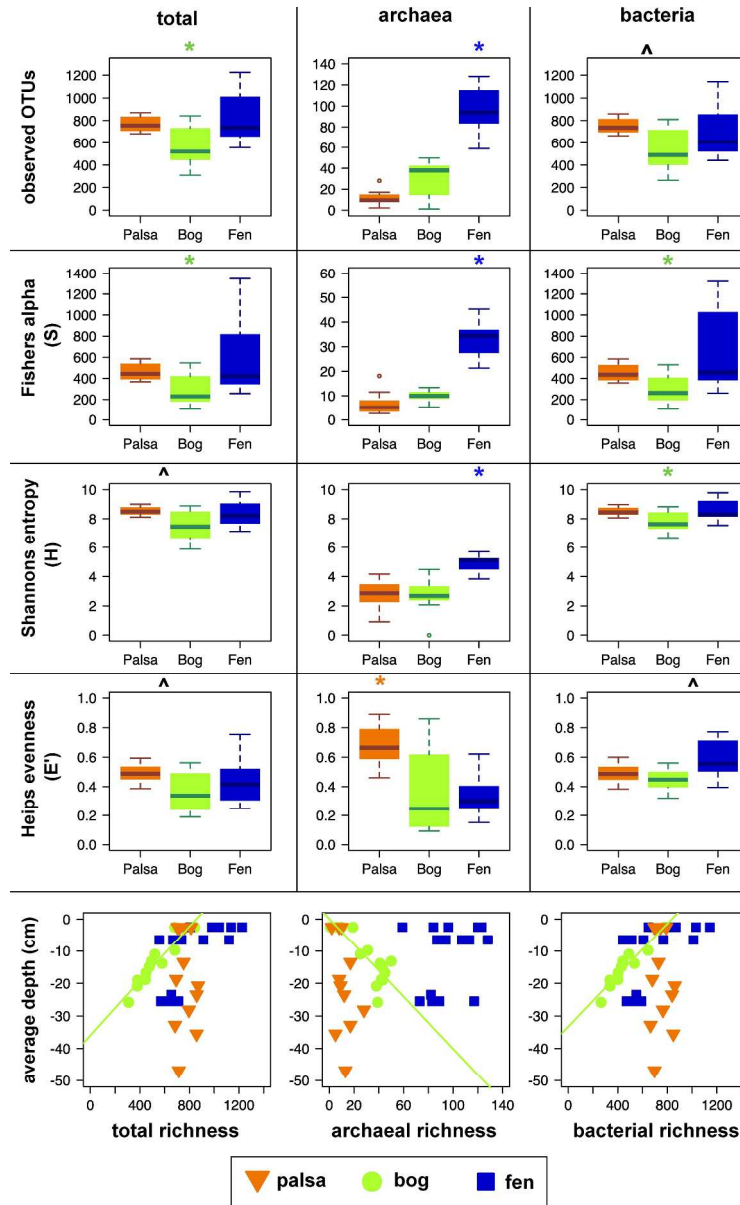


Fig 3 By-site comparison of alpha diversity metrics. a) The distribution of observed richness (top), Fisher's alpha-S (2nd from top), Shannon's entropy-H (2nd from bottom), Heip's evenness (bottom), on all 97 % OTUs (left), archaeal OTUs (middle) and bacterial OTUs (right) in combined normalized (N=2000) samples. Significant differences were measured by Kruskal-Wallis post-hoc test (* $p < 0.001$, where * designates a site significantly different from the other two sites and ^ designates a significant difference between the two adjacent sites only)). b) The distribution of OTU richness with sample depth for all OTUs (left), archaeal OTUs (centre), bacterial OTUs (right). Green trend lines show the correlations between richness and depth of sample in the bog site.

301x488mm (300 x 300 DPI)

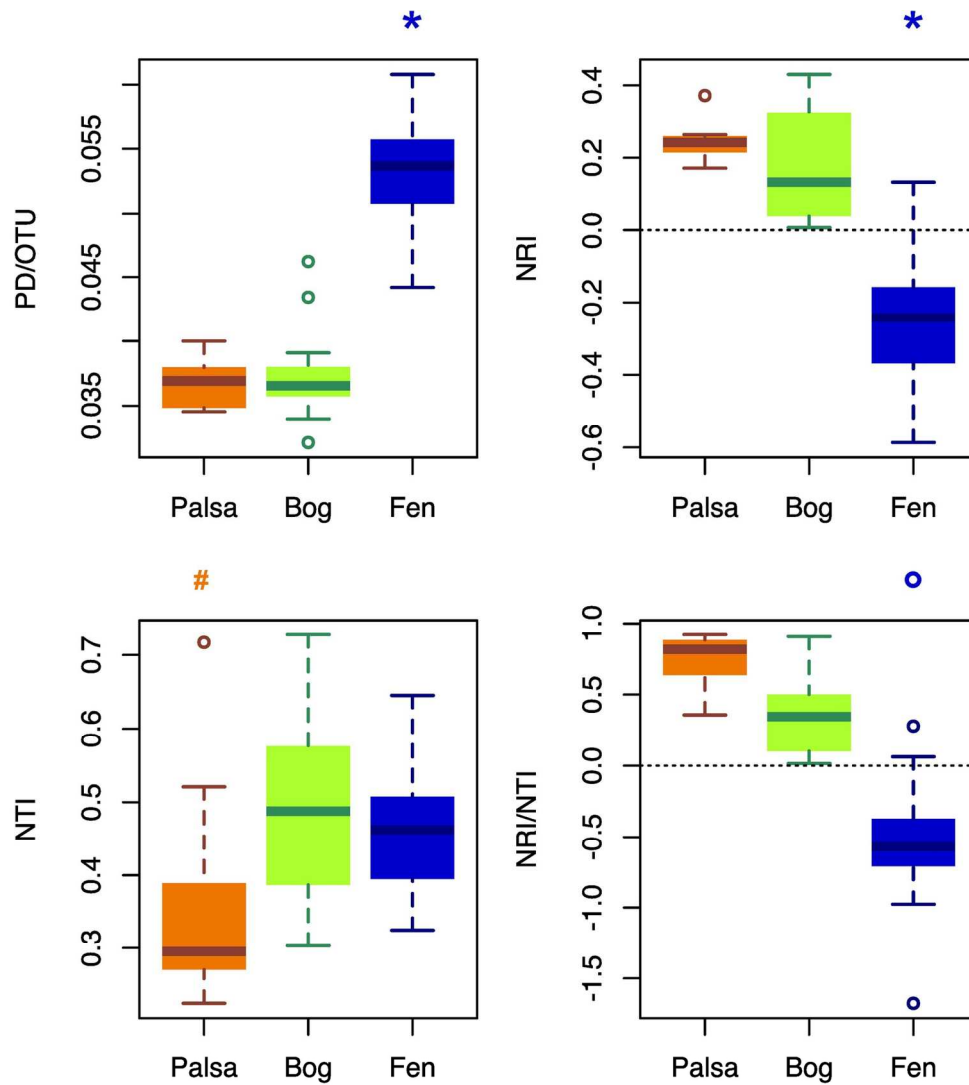


Fig 4 By-site comparison of phylogenetic diversity. topleft: faiths phylogenetic distance per OTU (PD/OTU); bottom left: Nearest taxon index (NTI); top right: Net relatedness Index (NRI), bottom right: NRI/NTI ratio.

147x167mm (300 x 300 DPI)

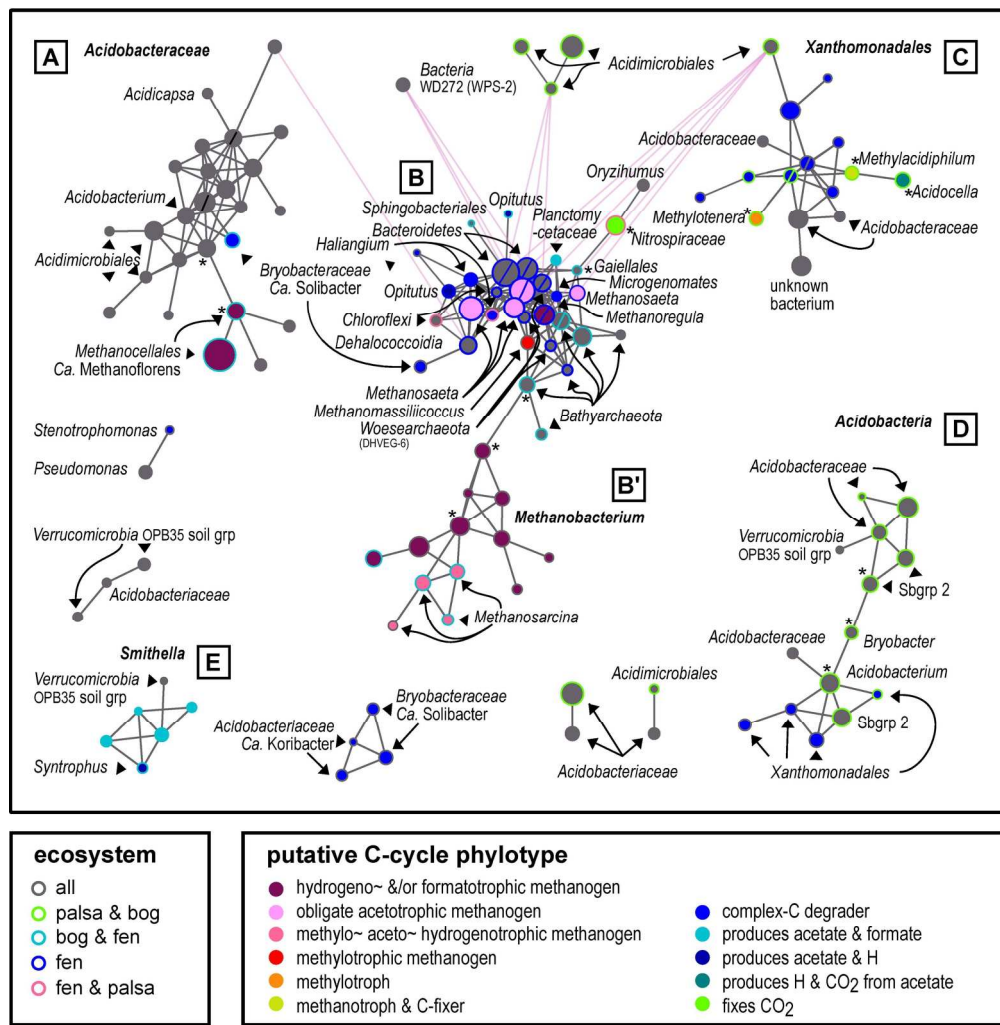


Fig 5 Network of phylotypes present in at least 15 samples with significant pairwise correlation. Circles indicate individual OTUs; circle diameter is a function of the log of the mean abundance. Circle colour denotes phylogenetically-inferred metabolism, and the colour of the outline denotes ecosystem presence. Circles with * are putative keystones and those with a / across the middle are putative hubs. Grey edges indicate co-occurrence, while pink edges indicate mutual-exclusion.

181x184mm (300 x 300 DPI)