

<https://helda.helsinki.fi>

Deforested and drained tropical peatland sites show poorer peat substrate quality and lower microbial biomass and activity than unmanaged swamp forest

Könönen, M.

2018-08

Könönen, M., Jauhiainen, J., Straková, P., Heinonsalo, J., Laiho, R., Kusin, K., Limin, S. & Vasander, H. 2018, 'Deforested and drained tropical peatland sites show poorer peat substrate quality and lower microbial biomass and activity than unmanaged swamp forest', *Soil Biology & Biochemistry*, vol. 123, pp. 229-241. <https://doi.org/10.1016/j.soilbio.2018.04.028>

<http://hdl.handle.net/10138/322438>

<https://doi.org/10.1016/j.soilbio.2018.04.028>

cc_by_nc_nd

acceptedVersion

Downloaded from Helda, University of Helsinki institutional repository.

This is an electronic reprint of the original article.

This reprint may differ from the original in pagination and typographic detail.

Please cite the original version.

1 **DEFORESTED AND DRAINED TROPICAL PEATLAND SITES SHOW POORER**
2 **PEAT SUBSTRATE QUALITY AND LOWER MICROBIAL BIOMASS AND**
3 **ACTIVITY THAN UNMANAGED SWAMP FOREST**

4 K n nen, M.¹, Jauhiainen, J.¹, Strakov, P.², Heinonsalo, J.³, Laiho, R.², Kusin K.⁴, Limin, S.⁴ † & Vasander, H.¹

5

6 ¹ University of Helsinki, Department of Forest Sciences, P.O. Box 27, 00014, Helsinki, Finland

7 ² Natural Resources Institute Finland (Luke), PO Box 2, FI 00791, Helsinki, Finland

8 ³University of Helsinki, Department of Food and Environmental Sciences, P.O. Box 56, 00014, Helsinki, Finland

9 ⁴CIMTROP, University of Palangka Raya, Palangka Raya 73112, Indonesia

10 **ABSTRACT**

11 Swamp forests on deep tropical peatlands have undergone extensive deforestation and draining for agriculture and
12 plantations, consequently becoming globally significant carbon (C) sources.

13 To study the effects of land-use change on peat as a biological environment, which directly affects decomposition
14 dynamics and greenhouse gas emissions, we sampled peat from four common land-use types representing different
15 management intensities in Central Kalimantan, Indonesia. The near-pristine *swamp forest* was used to describe
16 unmanaged conditions, and the three other sites in order of increasing management intensity were *reforested*;
17 *degraded*; and *agricultural*. We examined peat substrate quality (total C & nitrogen (N), dissolved organic C
18 (DOC) and N (DON), and organic matter quality characterized by infrared spectroscopy) and microbial biomass
19 and extracellular enzyme activity to describe both biotic and abiotic conditions in peat.

20 We found that the peat at altered sites was poorer in quality, i.e. decomposability, as demonstrated by the higher
21 intensity of aromatic and aliphatic compounds, and lower intensity of polysaccharides, and concentration of DOC,
22 total N and DON compared to the peat in the swamp forest. The observed differences in peat properties can be
23 linked to changes in litter input and decomposition conditions altered after deforestation and draining, as well as
24 increased leaching and fires. The quality of the peat substrate was directly related to its biotic properties, with
25 altered sites generally having lower microbial biomass and enzyme activity. However, irrespective of management

26 intensity or substrate quality, enzyme activity was limited primarily to the first 0–3 cm of the peat profile. Some
27 differences between wet and dry seasons were observed in enzyme activity especially in swamp forest, where the
28 most measured enzyme activities were higher in dry season.

29 Reforestation 6 years before our measurements had not yet restored enzyme activity in the peat to the level of the
30 swamp forest, although the topmost peat characteristics in the reforested site already resembled those in the swamp
31 forest. This is likely contributed to by the chemical weed control performed at the site, and the limited capacity of
32 the young plantation to produce litter to support peat formation and restore the quality and structure of the peat.
33 Therefore, we conclude that intensive land management, including deforestation and draining, leads to the surface
34 peat becoming poorer biological environment, and it may take long time to restore the peat properties.

35

36 **Keywords:** decomposability; land-use; microbial biomass; enzyme activity; peat properties; tropical peat

37

38 Corresponding author: Mari Könönen, maritkononen@gmail.com, Latokartanonkaari 7, P.O. Box 27, 00014,
39 Helsinki, Finland

40

41 1 INTRODUCTION

42 Peatlands in Southeast Asia form globally significant carbon (C) pools comprising 11–14% of C stored in peat
43 (Page et al. 2011). However, since 1990s land-use change, typically deforestation and draining to convert land for
44 agriculture and industry, has occurred extensively. In Peninsular Malaysia, Borneo and Sumatra, only 29% of the
45 original peat swamp forest area still remained forested in 2015, while only 6% of the forests lacked clear signs of
46 human impact (Miettinen et al. 2016). The largest remaining swamp forests in Southeast Asia (40% of the land
47 area) are located in Central Kalimantan, but the increasing demand for land, especially for cultivation, poses a
48 major threat to their existence (Miettinen et al. 2016). Due to land-use change towards drier and more sparsely
49 vegetated systems, altered tropical peatlands are globally significant C sources (IPCC, 2014). In 2015 in Peninsular
50 Malaysia, Borneo and Sumatra, approximately 78% of the total amount of C (146 Mt C yr⁻¹) released in
51 decomposition of peat was from small holder-farming and industrial plantations on peat soils (Miettinen et al.

52 2017). Altogether, drained tropical peatlands release approximately 200 Mt of C to the atmosphere per year
53 (Biancalani et al. 2014; Page & Hooijer 2016).

54 Carbon accumulation in soil takes place when the rate of litter deposition exceeds that of decomposition; if this
55 ratio reverses, C is released. Microbial decomposition of organic matter is mediated by extracellular enzymes, each
56 specified to degrade certain compounds. The primary abiotic factor affecting the decomposition rate in peatlands is
57 generally considered to be oxygen availability, which is limited by high water-table level (WT), but also substrate
58 quality, pH, nutrient availability, and especially in boreal latitudes also temperature may further constrain
59 decomposition (Clymo 1984; Laiho 2006). Additionally, microbial population and its composition, as a biotic
60 factor determines the potential decomposition activity and is usually limited by the one or more of the abiotic
61 factors. Microbial decomposition can accelerate due to WT drawdown improving oxygen availability (e.g.,
62 Freeman et al. 2001; Fenner & Freeman 2011; Ishikura et al. 2017), but it may also be suppressed by drought stress
63 (Kwon et al. 2013; Ishikura et al. 2017). The substrate quality can also be the primary factor limiting
64 decomposition, especially when the substrate is rich in polyphenols and other recalcitrant compounds (Berg 2000;
65 Strakova et al. 2011; Hoyos-Santillan et al. 2016). Therefore, land-use change that alters both vegetation and
66 draining, and thus influences both the substrate quality and WT, is likely to result in greatly modified environment
67 for decomposers. To promote more sustainable and C-neutral management of tropical peatlands, it is crucial to
68 increase our understanding of how peat decomposability and microbial decomposition activity (i.e. enzyme
69 activity) in peat varies between land management types.

70 Mature swamp forest with closed canopy, high biomass (Sulistiyanto 2004), relatively cool and unvarying
71 temperatures on the ground (Brady 1997; Jaya 2007) and usually moist peat up to the surface encloses abundant
72 organic substrate resources and good environment for decomposition. In the unmanaged swamp forests, high
73 amount of leaf litter is deposited and also mainly decomposed on the peat surface (Sulistiyanto 2004; Yule et al.
74 2009; Hoyos-Santillan et al. 2015). Woody litters; roots, trunks and branches, are more resistant to decomposition
75 and partly deposited below the WT, and therefore have a higher importance in peat formation in swamp forests
76 (Wüst et al. 2008; Hoyos-Santillan et al. 2015). The woody peat formed in swamp forests has a lignin concentration
77 of up to 72% of dry mass (Andriessse 1988; Könönen et al. 2015). Swamp forests are ombrotrophic ecosystems (i.e.,
78 rain fed), and cycling of the limited nutrient pools within the system is tight. The majority of the nutrients are
79 released from litter during decomposition at or close to the peat surface, and rapidly bound again to microbial and
80 plant biomass (Page et al. 1999). Therefore, nutrients are most concentrated in the surface peat (Page et al. 1999;
81 Andriessse 1988; Lampela et al. 2014). The slow water runoff in comparison to the amount of water received in

82 precipitation maintains a WT close to the peat surface. The forest floor is partly waterlogged in wet seasons
83 (Lampela et al. 2014). In dry seasons, aerobic decomposition processes are possible in the surface peat above the
84 WT, yet, the shading of the tree canopy reduces ground temperature and loss off moisture from peat. The high WT,
85 highest litter inputs at the peat surface especially of the easily decomposed leaf litter, and the highest availability of
86 nutrients in the surface peat all mean that the highest microbial biomass and activity is likely restricted to the very
87 surface layers in the unmanaged swamp forests. Yet, the fluctuation of WT between the wet and dry seasons can be
88 high, and the impacts of this fluctuation on the biomass and activity of the microbial community has not yet been
89 quantified for swamp forests.

90 Cultivation on peatlands requires removal of the original vegetation, i.e. deforestation, and draining, which ensures
91 sufficient aeration of the rooting zone necessary for the growth of most crop plants. Permanent draining also leads
92 to consolidation of the peat, improving its bearing capacity and thus enabling the use of cultivation machinery.
93 Deforestation reduces the litter deposition rate, as the volume of vegetation in all subsequent land use types (small-
94 holder farms, plantations, and abandoned areas that often develop to fern-covered shrub lands) is typically much
95 smaller than in swamp forest (Sulistiyanto 2004; Hoscilo et al. 2011; Blackham et al. 2014; Yule et al. 2016).
96 Although the stand biomass in mature oil palm and acacia plantations may be relatively high, harvesting and
97 intensive management practices (slash-and-burn, weeding, tilling) reduce litter deposition (Hertel et al. 2009; Smith
98 et al. 2012). The vastly reduced litter deposition may be one of the most important parameters influencing peat C-
99 dynamics as it reduces the potential for C-accumulation, irrespective of potential changes in decomposition rates. It
100 may also cause the intuitively paradoxal soil respiration patterns observed in some comparative studies, with the
101 highest respiration coming from the large mass of litter decomposing in unmanaged swamp forests, and clearly
102 lower rates observed for sites under intensive land use (Hirano et al. 2009; Jauhiainen et al. 2016b).

103 Draining enables aerobic decomposition deeper in the peat profile, consequently leading to C loss from a thicker
104 peat layer (Hirano et al. 2009; Jauhiainen et al. 2016a). Together the enhanced draining and replacing of the swamp
105 forest vegetation with no or patchy vegetation cover further lead to high peat surface temperatures due to the direct
106 solar radiation reaching to and being absorbed by the dark, bare soil surface (Jauhiainen et al. 2014). High surface
107 peat temperatures may cause drought stress thereby suppressing microbial decomposition activity (Kwon et al.
108 2013). Removal of the original vegetation cover and draining also upsets the tight nutrient cycle between litter and
109 live vegetation and may exacerbate the loss of nutrients, as it is known to increase the export of dissolved organic
110 carbon (DOC) (Page et al., 1999; Anshari et al. 2010; Moore et al. 2013; Evans et al. 2014). DOC and nutrient
111 leaching often take place in tandem (Nieminen et al. 2015). The impoverishment of already nutrient-poor peat

112 increases the need for fertilization for productive cultivation (Andriessse 1988). Poor soil quality for cultivation may
113 also lead to the abandonment of the deforested and drained degraded areas, where recurrent wildfires enhance the
114 impoverishment of peat by preventing regrowth of vegetation (Page & Hooijer 2016).

115 Altered WT and litter deposition rates, progressing peat decomposition, and fire occurrence all modify peat
116 properties in complex manners, and determine the quality of the peat as a biological environment for decomposers.
117 However, relatively little is still known about peat as a biological environment in the Tropics. Differences in
118 substrate quality, temperature climate and WT regime may lead to different patterns from the northern peatlands,
119 where the peat is primarily formed by *Sphagnum* mosses and *Carex* sedges. This study aims to provide insight into
120 the factors regulating decomposition in swamp forests subject to differing land management intensities. To do so,
121 we studied peat substrate quality (total C & nitrogen (N), DOC & DON, and organic matter quality characterized by
122 infrared spectroscopy) as well as microbial biomass and extracellular enzyme activity under the varying
123 environmental conditions of different land-use types in Central Kalimantan, Indonesia.

124 We hypothesized that: (i) deforested and drained sites have lower surface peat substrate quality, and thus lower
125 microbial biomass and enzyme activity, than unmanaged swamp forest. However, due to improved oxygen
126 conditions via WT drawdown, we also assumed that (ii) enzyme activity continues deeper in the peat profile at the
127 drained sites than in the swamp forest. Since decomposition in peatlands is also in great extent controlled by WT,
128 we assumed that (iii) the high WT during the wet season will suppress enzyme activity in the undrained swamp
129 forest, whereas the drought during the dry season will suppress enzyme activity at the drained sites. We expected
130 (iv) the reforestation of degraded peatland with a relatively high WT to restore peat quality and thus improve
131 conditions for decomposers, reflected as higher microbial biomass and enzyme activity.

132 **2 MATERIALS AND METHODS**

133 **2.1 Study area and sites**

134 Our study area was located on the upper parts of the Sabangau River catchment in Central Kalimantan, Indonesia.
135 (Fig. 1). During the period 2002–2010, the mean annual temperature in the area was 26.2 ± 0.3 °C and mean rainfall
136 2540 ± 596 mm yr⁻¹ (Sundari et al. 2012). The high precipitation received in the region is normally interrupted by a
137 dry season occurring from June to September. However, in recent decades, strong El Niño events have prolonged
138 the dry seasons and caused extreme droughts, which have led to the spread of wildfires. Our four study sites were

139 within c. 20 km distance in same catchment area on two ombrotrophic peat domes located on opposite sides with
140 similar distances from Sabangau River with peat depth exceeding 3 m at the sites.

141 At three of the sites, the vegetation and hydrology had been altered for at least 6 years and up to three decades prior
142 to the sampling. It was assumed that the conditions at these transformed sites had stabilized after land-use change;
143 as it has been considered to take five years before the peat physical properties and greenhouse gas dynamics
144 stabilize after deforestation and draining (Hooijer et al. 2012; IPCC 2014). The sites in order of increasing
145 management intensity were: near-pristine swamp forest (SF; 2°19'16.96" S, 113° 53'43.29" E), reforested site (RF,
146 2°18'54.94"S, 114° 03'31.76"E), degraded site (DO 2°19'25.11"S, 114°1'5.15" E), and agricultural site (AO; 02°
147 17' 25.21", E114° 0' 41.20") (Fig. 2). Degraded and agricultural site together are referred to as *open sites*.

148 **Figure 1.** A satellite photo of the study area with study sites indicated. The location of the study area in Southeast
149 Asia is shown on the inserted map. The dark green areas are swamp forests and lighter green areas deforested
150 areas with some ground vegetation. Copyrights of the photo: ©2016 Google.

151 **Figure 2.** Description of study sites with management intensity increasing from left to right.

152 In the swamp forest, the hydrology and original swamp forest vegetation were almost pristine. However, due to
153 selective logging prior to 1997 and remnants of small ditches, it cannot be considered fully intact. The topmost
154 layer of the tree canopy reached 35 m height, and ground vegetation consisted of a few, scattered *Pandanus spp.*
155 palm thickets (Page et al. 1999). The soil microtopography was characterized by open, low peat surfaces, which
156 formed pools during high WT, and higher hummock-like surfaces mainly around tree bases (Lampela et al. 2014).
157 Two of the altered sites, reforested and degraded, were clear-felled and drained in 1997 during the so-called Mega
158 Rice Project, a failed project attempting to convert 1 million hectares of peatlands into arable land (Page & Rieley
159 2005). The degraded site was abandoned following draining without further land use, while the other site was
160 reforested in 2008. The hydrology at both of these altered sites was affected by the same approximately 10-meter-
161 wide and 3–4-meter-deep canal dug in 1997. However, draining at the reforested site has been less efficient due to
162 its lower elevation, consequently leading to higher annual WT and frequent flooding events. Both of these sites had
163 burned after land-use change, but the reforested site not since 1998. The degraded site had burned recurrently with
164 the most intense fires occurring in 1997, 1999, 2002, 2006 and 2009. At the degraded site, at least 0.5 m of the
165 topmost peat had been lost due to fire (Hoscilo et al. 2011). At the time of sampling in the reforested site, 10-meter-
166 tall *Shorea balangeran* trees grew in rows at 3*3-meter intervals at a density of c. 500 trees ha⁻¹. Herbaceous
167 understorey vegetation was kept low by cutting and occasional application of herbicides containing mainly

168 glyphosate. The dominant vegetation at the degraded site was composed of ferns and scattered small trees with
169 crowns extending approximately 3 meters above the land surface. The most intensively managed site with the
170 longest history of management in this study was the agricultural site (AO), which was deforested and drained for
171 smallholder cultivation in the beginning of the 1980s. The main cultivated species were corn, tomatoes and cassava.
172 Controlled draining at the site maintained the WT below but rather close to the soil surface (Hirano et al. 2009).
173 The agricultural site has been repeatedly fertilized mainly with urea and by burning the surface peat and previous
174 crop residues. At all three altered sites, the microtopography was flat. Further information on peat properties at the
175 sites is presented in Table 1 and information on WT depth in Figure 3.

176 We chose managed sites that would be as comparable as possible in conditions where detailed soil and vegetation
177 analyses prior to land-use change were not available. Our view of the comparability was based on the distance to
178 the river, since the vegetation is known to gradually change with distance from the river due to the changes in
179 hydrology and nutrient conditions (Page et al. 1999; Sjögersten et al. 2011), and peat properties that were rather
180 uniform in the deepest layers studied at the sites, which best reflect the pre-draining properties as land-use impacts
181 are usually most evident in the surface layers (Könönen et al. 2015, 2016). All sites have been previously forested
182 judged by the high amount of wood and organic substrates characteristic to woody peat (e.g. Könönen et al. 2016).
183 However, since the site status before land-use change has not been recorded, we cannot rule out site-specific
184 variation in the properties studied here. This is a basic problem in nearly all space-for-time substitution studies on
185 land-use impacts (Oksanen 2001). This problem is even more emphasized in our study due to the limited number of
186 sites. However, since research in this region is logistically challenging we consider that our data are yet among the
187 best currently available for this area, and provide valuable information on peat as a biological environment in
188 managed former swamp forests. Recurrent burning of some sites may also be seen as a confounding factor in our
189 comparisons; however, since wild fires in this region are common and are known to target especially the degraded
190 drained sites, we consider that they should be accepted as a typical part of the land-use change impact in these sites.

191 **Figure 3.** Average monthly water-table level during the sampling year (2014) at the sites based on 1–2 monthly
192 monitoring events.

193 **Table 1.** Selected soil properties at the study sites corresponding to the sampling depths in this study. Data from this
194 and other studies (source mentioned under the table).

195 **2.2 Sampling**

196 We established five permanent square-shaped 1m² sampling plots at the swamp forest, degraded and reforested
197 sites. The permanent plots were located at approximately ten-meter intervals. In the swamp forest and degraded
198 site, four of the permanent plots were situated at the corners and one in the middle of a square-shaped sampling area
199 covering approximately 400 m². At the reforested site, the plots were arranged in a row along the midsection of the
200 reforested area covering approximately 300 m². At the agricultural site, permanent sample plots could not be
201 established because they would have disturbed farming, and samples were thus taken from three randomly chosen
202 spots.

203 Soil samples for determining enzyme activity were taken at the middle of the wet and at the end of the dry season,
204 mid-March and mid-September in 2014, respectively, except for the agricultural site where samples were only
205 taken at the end of the dry season. Sampling for total N and C, infrared spectroscopy, microbial biomass C and N,
206 and dissolved organic N and C (DON and DOC, respectively) analyses was carried out at the end of the dry season.
207 In the swamp forest and reforested site, samples were extracted from low, vegetation-free surfaces between the
208 trees, as they covered the majority of the land area. At the open sites (agricultural and degraded), both vegetated
209 and unvegetated surfaces were sampled (ferns at degraded site and maize at agricultural site).

210 The samples included the first 30 cm of the peat from the surface divided into four sections (0-3 cm, 3-10 cm, 10-
211 20 and 20-30 cm). The rationale for this was that we expected to see different patterns right at the surface than in
212 the deeper layers, since for instance the temperature variations are the most extreme in the topmost layer in the
213 managed sites. Non-volumetric samples from the topmost peat (0–3 cm) were collected by hand. Volumetric
214 samples (68.7 cm³ from 3–10 cm depth and 98.2 cm³ from 10–20 and 20–30 cm depth) were extracted from deeper
215 depths using a Russian peat auger 5 cm in diameter. The total number of undivided samples (n) per season was 5 at
216 the swamp forest, degraded and reforested sites, whereas at the agricultural site the total number of undivided
217 samples was 3.

218 Samples were sealed in plastic bags immediately after extraction. Since the time needed to transport samples from
219 the field to the laboratory located at the University of Helsinki, Finland was long, samples were stored in coolers or
220 in a refrigerator at 4 °C temperature for a maximum period of two weeks prior to analyses, if not mentioned
221 otherwise. Prior to analyses, the living roots were removed.

222 2.3 Laboratory analyses

223 2.3.1 Infrared spectroscopy

224 Subsamples were air-dried before milling to fine powder. Infrared spectra were obtained with a Bruker VERTEX
225 70 series FTIR (Fourier Transform InfraRed) spectrometer (Bruker Optics, Germany) equipped with a horizontal
226 attenuated total reflectance (ATR) sampling accessory. Dried and powdered samples were inserted directly on the
227 ATR crystal and a MIRacle high-pressure digital clamp was used to achieve even distribution and contact of the
228 sample and crystal. Each spectrum consisted of 65 averaged absorbance measurements between 4000 and 650 cm^{-1} ,
229 with a 4 cm^{-1} resolution. Summed absorbance values of the following bands were used as representative of the
230 different organic compounds in the ordination diagram of Figure 7: 2920 and 2850 cm^{-1} (wax, lipids); 1612, 1513,
231 1450 and 1265 cm^{-1} (lignin and other phenolics); 3340, 1150 and 1034 cm^{-1} (polysaccharides. Offsets in baseline
232 and slope between the different runs (samples) were removed by standard normal variate transformation and the
233 second derivative using the Unscrambler software (CAMO, Norway). The individual bands were assigned
234 according to Artz et al. (2008).

235 2.3.2 Total C and N and C/N-ratio

236 Total C and N concentrations (mg g^{-1} of dry mass) and, consequently, the C/N-ratio were determined from
237 approximately 0.3 g of homogenized, air-dried sample with LECO CHN-1000. The peat C and N pools for each
238 sampled layer were calculated by multiplying the C and N concentrations with the thickness and bulk density of the
239 layer, and transforming the values per m^2 . The peat bulk density used in these calculations is from Table 1.

240 2.3.3 Soil DOC and DON and microbial C and N

241 Two comparable samples, approximately 3 g of homogenized fresh peat, were prepared. One sample set was used
242 to measure the total DOC and DON concentrations of soil, and the other to measure soil microbial C and N after
243 chloroform (CHCl_3) fumigation (Vance et al. 1987). DOC and DON concentrations were measured from non-
244 fumigated samples after extraction with K_2SO_4 at the start of the fumigation process of the other sample set.
245 Fumigated samples were incubated at 25 °C for 24 hours with CHCl_3 prior to extracting with 0.05 M K_2SO_4 . A
246 TOC-analyser was used to analyse the C and N concentrations of the filtered (Whatman No. 42-filters) non-
247 fumigated and fumigated samples. To calculate the microbial C and N, the C and N concentrations of non-
248 fumigated samples were subtracted from the fumigated samples. To estimate the microbial biomass, a correction

249 factor of 2.64 was used for unrecovered microbial C (Vance et al. 1987) and a correction factor of 1.86 for
250 microbial N (Brookes et al. 1985).

251 2.3.4 Extracellular enzyme activity

252 Extracellular activity of five enzymes was determined from peat with a fluorescence method modified from Pritsch
253 et al. (2011). The measured enzymes, their abbreviations and substrates used in analysis (in parentheses) were:

254 (i) b-Xylanase (Xyl, 4-methylumbelliferyl- β -D-xylopyranoside, 3.2.1.37),

255 (ii) b-Glucosidase (Gls, 4-Methylumbelliferyl- β -D-glucuronide, 3.2.1.21),

256 (iii) N-acetyl-b-glucosaminidase (Nag, 4-Methylumbelliferyl N-acetyl- β -D-glucosaminide, 3.2.1.14),

257 (iv) Phosphomonoesterase (Pho, 4-Methylumbelliferyl phosphate, 3.1.3.2),

258 (v) Arylsulfatase (Sulf, 4-Methylumbelliferyl sulfate potassium salt, 3.1.6.1)

259 The activity of these enzymes is used to describe the enzymatic decomposition activity related to degradation of C
260 (b-xylanase and b-glucosidase), N (N-acetyl-b-glucosaminidase), P (phosphomonoesterase) and S (arylsulfatase)
261 compounds.

262 One day (24 h) prior to the analyses, approximately 2 g of peat of each sample stored in a refrigerator was weighed
263 in small plastic bags ($10 \times 10 \text{ cm}^2$) and incubated in the sealed bags at room temperature (22°C) in darkness. On the
264 day of analyses, 7 mL of deionized water was added to the bags, mixed throughout for three minutes, and filtered
265 through a cellulose acetate filter (Whatman, $0.45 \mu\text{m}$). Then $50 \mu\text{L}$ of universal buffer prepared as in Pritsch et al.
266 (2011) (pH 4.5 for all substrates), $50 \mu\text{L}$ of filtered sample aliquot and $50 \mu\text{L}$ of enzyme substrate were added to a
267 microplate (Optiplate 96F HB, Perkin-Elmer) having separate microplates for each substrate. Plates were incubated
268 on a microplate shaker for 15 min (b-Glucosidase and N-acetyl-b-glucosaminidase) or 30 min (b-Xylanase,
269 Phosphomonoesterase and Arylsulfatase). After incubation, $100 \mu\text{L}$ of incubated aliquot was added on a microplate
270 on top of $100 \mu\text{L}$ TRIS-buffer (1M; pH 10-11) to stop the fluorescence assay. All used buffers were prepared as in
271 Pritsch et al. (2011). Fluorescence values were measured with a Wallac 1420 Victor2 (PerkinElmer, Inc., USA)
272 multilabel plate reader. The aliquot of substrate in deionized water was used as the background value in
273 calculations.

274 2.4 Data analyses

275 To test the first and second hypothesis, the differences in measured variables between sites at each sampling depth
276 were tested with one-way analysis of variance (ANOVA). When the effect of site was significant, ANOVA was
277 followed by Games-Howell post hoc test, which is suitable for testing differences between groups having unequal
278 sample sizes and heterogeneous variances, and when normality cannot be assumed. Due to values below detection
279 limits, enzymatic data had zero-values and were thus transformed as $\log(x+1)$ to improve the strength of statistical
280 tests. Differences were statistically significant if the p-value was less than 0.05.

281 Multidimensional ordination methods were applied to further analyze the data, where several variables were
282 presumably intercorrelated. Before analysis, the data were centered and standardized to make the different scales
283 commensurable. To test the first, second and fourth hypothesis, redundancy analysis (RDA) was used to test the
284 effect of land management type on substrate quality (the infrared spectroscopy-derived peat quality). In this RDA,
285 the interaction of site and depth was used as explanatory variable. The summed absorbance values of some main
286 compound groups, e.g. polysaccharides; waxes and lipids; lignin and other phenolics, were used as supplementary
287 variables to better visualize the correlation of substrate quality with the site and depth. This means that they were
288 not affecting the analysis as such, but their correlations with the ordination axes were projected in the graphs. To
289 test further the first, second, and third hypothesis, constrained redundancy analysis (RDA) with variation
290 partitioning was conducted to analyze how much of the total variation in enzyme activity was explained first by site
291 and depth, and then by the substrate quality (pH, and total C and N, DOC and DON, microbial C and N
292 concentration) and to analyze how much of the total variation (inertia) was explained by each environmental
293 variable. Last, we tested the effect of spectroscopy data-derived peat quality on enzyme activity from the dry season
294 (for which data from all sites were available) with RDA. In all analyses, a Monte Carlo permutation test with 999
295 permutations was applied to evaluate the significance of the canonical axes; permutations were constrained by the
296 covariates. All ordination analyses were made using Canoco 5 for Windows.

297 **3 RESULTS**

298 **3.1 Peat as an environment**

299 *3.1.1 Carbon and nitrogen*

300 The total C concentration of peat varied in the range 48.7–61.1 mg g⁻¹ (of dry peat) and the total N concentration in
301 the range 0.81–1.61 mg g⁻¹, depending on sampling depth and site (Fig. 4). The total C concentration was lowest in
302 the topmost peat (0–3 cm) and increased with depth at all sites (Fig. 4). In the degraded and reforested sites, the C

303 concentration was generally higher than at other sites throughout the sampling profile (Fig. 4). The N concentration
304 of swamp forest peat was higher and also rather constant at all sampled depths as compared to the altered sites,
305 where the N concentration showed a clear maximum in the topmost peat (Fig. 4). The C/N-ratio varied in the range
306 32–76, being at its highest deeper in the peat profile at all sites (Fig. 4). Generally, the ratio was lowest in the
307 swamp forest and highest in the agricultural site. The total C and N pools for the 30 cm peat layer sampled varied
308 between the sites from 16.8 to 22.8 kg C m⁻² and 0.30 to 0.51 kg N m⁻² (Fig. 4). The C pool was typically smaller
309 and the N pool was greatest in the swamp forest than at the other sites. The highest C pool was in agricultural site
310 and the lowest N pool was at reforested site.

311 The concentrations of DOC and DON extracted from the peat varied in the range 0.52–1.80 mg g⁻¹ and 0.15–0.50
312 mg g⁻¹ of peat dry mass, respectively (Fig. 4f & g). In the topmost peat layer (0–3 cm), the DOC concentration in
313 the swamp forest and reforested site was up to three times higher than at the open sites (Fig. 4f). The swamp forest,
314 however, showed the highest DOC concentration in the deepest layer studied (20–30 cm), while the reforested site
315 had its maximum in the topmost layer. The degraded site had the highest peat DOC concentration in the deepest
316 layer. The agricultural site showed a clear maximum in the 3–10 cm layer. The DON concentration was highest in
317 the topmost peat layer (0–3 cm) at all sites and decreased towards the deeper layers. DON concentration was
318 overall highest in the swamp forest (Fig. 4g). The DOC/DON –ratio was lowest in swamp forest throughout the
319 peat profile (Fig. 4h). The total N and DON concentration correlated positively (R=0.90, p=).

320 **Figure 4.** From top to bottom, peat total pool carbon and nitrogen pool; and total, dissolved and microbial
321 concentrations of carbon and nitrogen at various depths in peat during the dry season at sites. Bars indicate the
322 mean ± SE. Different letters next to bars indicate significant (p < 0.05) difference between sites at the same depth.
323 The number of samples for each depth was five (n=5) at swamp forest, reforested and degraded sites and three
324 (n=3) at agricultural sites.

325 3.1.2 Peat characterization with infrared spectroscopy

326 The strongest gradient in RDA, axis–1, explained 51.1% of the total variation in infrared spectroscopy data and
327 separated most of the layers at altered sites from the swamp forest (Fig 5). This indicates that generally, the peat in
328 all managed sites differed from the unmanaged swamp forest peat. The second strongest gradient, axis–2, explained
329 24.4% of the variation and separated the swamp forest and reforested site from the open sites. This indicates that
330 the peat of the reforested site was closer in some respects with the swamp forest peat than the peat of the open sites.
331 The topmost peat layer (0–3 cm) showed higher intensities of polysaccharide absorption bands at the swamp forest

332 and reforested site and higher intensities of absorption bands assigned to lignin and other phenolic compounds at
333 the open sites (Fig 6). Differences between forest-covered and open sites were most apparent in the surface peat (0–
334 10 cm), while deeper layers revealed less contrast. RDA also indicated that generally, the topmost peat layers
335 differed from deeper layers of the same site. The exception was the agricultural site, where peat chemical
336 composition varied little between the sampled depths.

337

338 **Figure 5.** Redundancy analysis summarizing the variation in peat quality characterized by infrared spectroscopy as
339 explained by site and depth (used as interaction factor). Axis 1 explained 51.1% and Axis-2 24.4% of the total
340 variation in peat quality characterized by infrared spectroscopy. Carbon compound groups were used as
341 supplementary variables, i.e., their correlation with the axes is shown but they did not affect the analysis. SF=
342 swamp forest, RF=reforested site, DO=degraded site, AO=agricultural site.

343 **Figure 6.** Peat carbon characteristics described by infrared spectra for each depth and site. Each carbon compound
344 group associated with a specific wavelength is indicated on the top of the respective peak (i.e., polysaccharides at
345 the wavenumber 3340).

346

347 **3.3 Microbial N and C**

348 At all sites, the microbial C and N concentrations were highest in the topmost peat (0–3 cm) and decreased with
349 depth (Fig. 4). Generally, the concentrations were highest in the swamp forest and decreased following the intensity
350 of land management, thus being lowest at the agricultural site. In the surface peat of the swamp forest,
351 concentrations of microbial C were 0.5–2.7 times higher and microbial N 0.82–2.46 times higher than at other sites
352 at the respective depths (Fig. 4).

353 **3.4 Enzyme activity**

354 Enzyme activity greatly depended on the site, sampling depth and substrate quality. The variation partitioning,
355 showed that of the total variation in enzymatic activity 23.4% was explained by the sampling depth and the
356 interaction of site and season together, and 19.0 % was explained by soil properties (pH, and total C and N, DOC
357 and DON, and microbial C and N). The enzyme activities positively correlated with the absorption intensities of the
358 bands assigned to polysaccharides, and the variation in spectroscopy data explained 22.2% of variation in enzyme
359 activities (data not shown).

360 If detected, extracellular enzyme activity was highest in the topmost peat (0–3 cm) and there was hardly any
361 activity at deeper depths. In the wet season, the measured extracellular enzyme activities in peat were significantly
362 higher in the swamp forest than at the degraded and reforested sites (Fig. 7, Supplementary table. 2). In the dry
363 season, the reforested site showed the lowest overall activity. Interestingly, in the dry season, the degraded and
364 agricultural sites had similar or higher C-related enzyme activity, slightly lower N- and much lower P related
365 activity than in the swamp forest.

366 In the swamp forest, where activities were detected in both seasons, some differences between the seasons
367 occurred. In the topmost peat, the enzyme activity related to N degradation (N-acetyl-b-glucosaminidase) was two
368 times higher during the wet season than the dry season. The activities of enzymes related to P (acid phosphatase)
369 and C (β -glucosidase) release, in turn, were 2.5 and 2 times lower, respectively, during the wet season than dry
370 season (Supplementary table 2).

371 The enzyme allocation clearly differed between land management types (Figure 8). In the swamp forest, allocation
372 to P acquisition was highest in both seasons, despite the contrasting differences in the activities of the enzymes
373 between seasons described above. In the degraded and reforested sites, the allocation patterns differed between
374 seasons, with higher allocation to C acquisition in the wet season and P acquisition in the dry season.

375 **Figure 7.** The extracellular enzyme activity in peat samples according to season (wet or dry). On the x-axis, activity
376 is expressed as $\mu\text{mol min}^{-1} \text{g}^{-1}$ (dry mass) and sampling depth (cm) is indicated on the y-axis. Enzyme activity at
377 the agricultural site was only measured during the dry season. SF= swamp forest, RF= reforested site, DO=
378 degraded site, AO= agricultural site. (mean \pm SE).

379 **Figure 8.** Enzyme allocation on the topmost (0-3 cm) peat during the wet and dry season at the sites.

380 **Supplementary table 2.** The mean extracellular enzymatic activity in peat samples according to season. (mean \pm
381 standard error). Different letters indicate significant ($p < 0.05$) difference.

382

383 **4 DISCUSSION**

384 In order to improve the sustainability of tropical peatland management, decomposition dynamics and environmental
385 conditions resulting in C loss need to be understood. Therefore, we studied peat as a biological environment under
386 common land management types by focusing on both the major abiotic (substrate composition and moisture

387 conditions) and biotic (microbial biomass and enzyme activity) factors that influence organic matter decomposition
388 (Clymo 1984; Berg 2000; Laiho 2006; Hoyos-Santillan et al. 2016). The results of this study, which cover four land
389 management types but only few sampling events, reflect the current biotic and abiotic peat conditions modified up
390 to several decades by the present land management type. Therefore, the results cannot be used to estimate the
391 effects of land use change as a dynamic process, but instead they can be used to describe the differences between
392 land management types and to discuss of the potential effects of the changes in land management practices.

393 **4.1 Deforestation and draining lead to peat becoming poor biological environment**

394 As hypothesized, intensive land management including deforestation and draining have led to an overall lower
395 microbial biomass and enzyme activity. This was likely due to the peat quality showing advanced peat
396 decomposition degree at the altered sites, i.e., increased recalcitrance, being richer in aromatic and aliphatic
397 compounds and poorer in of polysaccharides, DOC and nutrients (total N, NO_3^- , NH_4^+ , DON) compared to the peat
398 in the undrained swamp forest. The observed changes in peat properties can be linked to changes in litter input,
399 conditions influencing decomposition in soil, and loss of the original surface peat in recurrent fires. In the swamp
400 forests, net formation of peat is supported by a high and continuous litter deposition rate into environment where
401 aerobic decomposition is seasonally reduced by high WT (Page et al. 1999; Wüst et al. 2008; Hoyos-Santillan et al.
402 2015), which was reflected in the high concentration of labile C compounds throughout the sampled peat profile in
403 the swamp forest. At open sites, decomposition progresses primarily in older peat substrates as the litter input rates
404 are low, hence the topmost peat is enriched with recalcitrant compounds as the labile compounds have already been
405 consumed during earlier period after the land-use change. Similar effects of land-use change on the carbon
406 compound characteristics of tropical peat have also been reported earlier (Könönen et al. 2016). The N and DON
407 concentrations were higher in the swamp forest than at the altered sites, which likely implies biomass removals,
408 fires and leaching leading to the losses of N from peat at altered sites.

409 In tropical peatlands, the availability of labile carbohydrates (Hoyos-Santillan et al. 2016; Jauhiainen et al. 2016b;
410 Sangok et al. 2017), along with N and P (Rejmánková 2001; Sjögersten et al. 2011) has been emphasized in the
411 regulation of organic matter decomposition, which is in line with our findings of generally lower enzyme activity in
412 the impoverished peat at the altered sites. The enzyme allocation patterns can be used to estimate the substrate
413 demand of microbes as well as availability of the nutrients (Sinsabaugh et al. 1994). Interesting differences were
414 observed between the sites in this respect. The high allocation to P-acquisition in the swamp forest indicates that P
415 is likely the limiting factor for microbial activity, despite this site having the highest peat P concentration (Könönen

416 et al. 2015). At the open sites, the acquisition of C was highest in the wet season and P in the dry season. This may
417 imply different limiting factors between the seasons, possibly caused by varying qualities or quantities of litter
418 input or released root exudates.

419 Draining of the swamp forest, where the peat contains a high concentration of labile carbohydrates (this study;
420 Könönen et al 2016; Sangok et al. 2017) and has relatively high pore space (Page et al. 1999; Lampela et al. 2014;
421 Könönen et al. 2015), has led to high C-loss especially during the first years after draining and deforestation, and to
422 subsequent decrease in the level of C-loss rate with time (Hooijer et al. 2012). Therefore, it appears that the WT is
423 the most important parameter limiting the decomposition processes when the labile carbohydrates are available, and
424 later especially in deforested areas with reduced input of new litter, with proceeding decomposition, the substrate
425 quality becomes more important parameter regulating decomposition of peat. The sites studied here appear to have
426 reached the latter stage. However, increased availability of labile substrates or decomposition-limiting nutrients, for
427 example through reforestation or fertilization, may again boost microbial decomposition and lead to increased
428 decomposition of the recalcitrant compounds through priming (Kuzyakov 2010; Dungait et al. 2012). The peat at
429 the agricultural site has been fertilized with urea and residues of plants burned as part of land management. This
430 may have boosted decomposition, consequently resulting in higher enzyme activity than at the other altered sites.
431 Both ex situ and in situ studies have demonstrated that the addition of N alone or together with a labile C source
432 (glutamate and glucose) to peat from intensively managed land-use types enhances decomposition rates in peat
433 (Jauhiainen et al. 2014, 2016b; Comeau et al. 2016). Additionally, in drainage-affected plantations with high-
434 biomass production, e.g. acacia and oil palm plantations, the fresh root litter input and N-binding ability of Acacia-
435 family may boost the decomposition processes deeper in peat; however, such information is still lacking.

436 The enzyme activity was limited primarily to the topmost 0–3 cm of the peat profile at all sites. In line with this
437 result, decrease in microbial enzyme activity and microbial population richness with increasing depth from the peat
438 surface has been reported previously from tropical peatlands (Jackson et al. 2009; Hoyos-Santillan et al. 2015). This
439 phenomenon has been linked to the lack of oxygen and reduction in substrate quality in deeper peat layers (Hoyos-
440 Santillan 2016; Jauhiainen et al. 2016b). Therefore, we had expected to find higher enzyme activity in deeper soil
441 layers at the altered sites due to the assumedly improved conditions for decomposition via increased soil aeration
442 following draining. The diffusion of oxygen to deeper layers may, however, be slow at the open drained sites,
443 where the surface peat has typically shown to have higher bulk density, higher proportion of small particles and
444 lower porosity than at swamp forest, and thus smaller proportional airspace in peat (Iiyama et al. 2010; Hooijer et
445 al. 2012; Könönen et al. 2015). Overall, it seems that draining and deforestation have made the surface peat a poor

446 biological environment, that does not inherently support high rates of peat decomposition in the studied sites. This
447 may appear beneficial from the point of view of conserving the peat C stores. However, when substantial inputs of
448 litter are lacking, even lower rates of decomposition will lead to loss in peat stores (Hooijer et al. 2010; Wijadasa et
449 al. 2017).

450 **4.2 Reforestation had not yet restored microbial activity**

451 In opposition to our hypothesis, reforestation had yet to restore microbial enzyme activity in peat, despite the fact
452 that the topmost peat characteristics were already rather similar in the swamp forest and the reforested site.
453 Potentially, glyphosate applied regularly to remove understorey vegetation has leached into the surface peat and
454 suppressed the activity of the microbial community (Zaller et al. 2014; Druille et al. 2016). Additionally, the trees
455 planted six years prior still likely require more time to significantly increase their biomass and produce coarse,
456 slowly decomposing root litter, which is the largest contributor to peat formation in swamp forest (Niiyama et al.
457 2010). Over time, if the vegetation cover will grow and increasingly promote the input of both above- and
458 belowground litters and associated labile carbohydrates, the accumulation of new surface peat with lower BD may
459 commence. This may eventually restore the quality and structure of the surface peat, yet, long-term monitoring will
460 be needed to evaluate the potential of reforestation to restore the C-sink function in degraded areas.

461 **4.3 Seasonality in microbial enzyme activity**

462 The samples to study the seasonal variation in microbial enzyme activity were collected in the end of the rainy and
463 dry seasons. Therefore, we assume that the measured enzyme activity reflect the acclimation of microbial
464 decomposition on longer lasting seasonal WT conditions. However, we acknowledge that the measurements reflect
465 a snapshot in both space and time, and may be affected by both spatial and temporal variation caused by other
466 factors as well. In any case, some seasonal differences were observed in the enzymatic decomposition activity. In
467 the swamp forest, enzyme activity was primarily lower during the wet season than the dry season. This observation
468 is in line with decreased CO₂ emissions released through decomposition when the WT lies close to the soil surface
469 (Fig. 2 in Jauhiainen et al. 2016 and references therein), and may in principle be caused by either low oxygen or
470 substrate availability. Interestingly, in the swamp forest the activity of N-acetyl-β-glucosaminidase, which is
471 involved in microbial breakdown of N-rich chitin, was higher during the wet season when peat was water saturated.
472 The N released in chitin breakdown, may partly contribute to the efficient denitrification and CH₄ consumption
473 noticed under the anaerobic conditions (Adji et al. 2014; Jauhiainen et al. 2016b), and thus the differences in
474 enzyme activities may also indicate that different parts of the microbial community with differing enzyme activity

475 profiles are active during the wet and dry seasons. Overall, however, the microbial communities and their activities
476 in tropical peatlands still warrants further and more thorough study.

477 Although we expected drought stress to suppress enzymatic activity at the degraded site, there the highest activity
478 actually occurred during the dry season. In the agricultural site, where only dry-season samples were taken, the
479 enzymatic activity was comparable to the degraded site. The absence of shading combined with the solar radiation-
480 absorbing capacity of dark soil at these sites elevates soil temperatures (Jauhiainen et al. 2014), which probably
481 influenced positively in activity in the recalcitrant, fire-affected peat. Notably, the surface peat properties at these
482 sites actually resembled those of biochar (Könönen et al. 2015, 2016), and accelerated decomposition of biochar
483 has been observed in temperatures over 40°C in substrate in aerobic conditions (Nguyen et al. 2010; Fang et al.
484 2015). Although the long-term average soil temperature during the dry season did not exceed 28°C in the degraded
485 site in this study (Table 1), it is very likely that the topmost peat temperatures can rise above 40°C momentarily
486 during the hottest hours of the day.

487 **4.5 Implications for C balance of tropical peatlands**

488 We conclude that the land management greatly modifies the biotic and abiotic conditions for decomposition in peat
489 as a result of altered litter input and WT level. Enzymatic activity is especially related to substrate quality
490 (compound composition and nutrient availability) and WT. The observed positive relation between microbial
491 decomposition activity and substrate quality may partly explain the relatively low CO₂ emissions from peat
492 reported for land uses characterized by low vegetation cover and extensive time since deforestation (Hirano et al.
493 2009; IPCC 2014). Yet, we suggest that deforestation and draining weaken both the *biotic and abiotic* conditions
494 supporting peat C stores, because new peat formation is negligible due to reduced litter input rates at deforested
495 sites, fire events (either controlled or uncontrolled) promote peat oxidation, and fertilizer applications on cultivated
496 lands may enhance decomposition of recalcitrant compounds.

497 Even though the relatively young reforested site still clearly differed from the swamp forest both functionally and in
498 regard to peat properties, over time and under seasonally high WT, the growth of forest vegetation may be expected
499 to increase litter input to peat, and eventually this ecosystem may become functionally capable of suppressing the
500 loss of C and increasingly supporting the formation of peat, however the time span for this to happen remains to be
501 evaluated.

502 The results of this study concern the biotic and abiotic conditions on the time of sampling at the annual WT extreme
503 on four sites each representing different land use types. To fully understand the annual dynamic in the
504 decomposition processes long-term monitoring is still needed.

505 **ACKNOWLEDGMENTS**

506 We acknowledge the Centre for International Cooperation in Sustainable Management of Tropical Peatland
507 (CIMTROP) for providing facilities and assistance during field sampling. We thank Marjut Wallner from the
508 University of Helsinki for her patience and expertise in the laboratory analyses. We acknowledge Meeri Pearson for
509 polishing the language of this paper.

510 **FUNDING**

511 This research was supported by the Academy of Finland –funded ‘Restoration Impact on Tropical Peat Carbon and
512 Nitrogen Dynamics’ -project (RETROPEAT; Decision number 253933), Jenny and Antti Wihuri Foundation,
513 Finnish Cultural Foundation, and University of Helsinki prize money for Peatlanders.

514 **5 REFERENCES**

515 Adjil, F., Hamada, Y., Darang, U., Limin, S., Hatano, R., 2010. Effect of plant-mediated oxygen supply and
516 drainage on greenhouse gas emission from a tropical peatland in Central Kalimantan, Indonesia. *Soil Science and*
517 *Plant Nutrition*, 60(2), 216-230.

518 Artz, R., Chapman, S., Robertson, A., Potts, J., Laggoun-Defarge, F., Gogo, S., Comont, L., Disnar, J-R., Francez,
519 A-J., 2008. FTIR spectroscopy can be used as a screening tool for organic matter quality in regenerating cutover
520 peatlands. *Soil Biology & Biochemistry* 40, 515-527.

521 Andriess, J., 1988. Nature and management of tropical peat soils. *FAO Bulletin* 59, Food and Agriculture
522 Organization of the United Nations, Rome, 165 pp.

523 Anshari, G., Afifudin, M., Nuriman, M., Gusmayanti, E., Arianie, L., Susana, R., Nusantara, R., Sugardjito, J.,
524 Rafiastanto, A., 2010. Drainage and land use impacts on changes in selected peat properties and peat degradation
525 in West Kalimantan Province, Indonesia. *Biogeosciences* 7, 3403–3419.

- 526 Biancalani, R., Avagyan, A. (eds), 2014. Towards climate responsible peatland management practices: Part 1.
527 Rome, Italy: FAO.
- 528 Blackham, G., Webb, E., Corlett, R., 2014. Natural regeneration in a degraded tropical peatland, Central
529 Kalimantan, Indonesia: implications for forest restoration. *Forest Ecology and Management* 324, 8–15.
- 530 Berg, B., 2000. Litter decomposition and organic matter turnover in northern forest soils. *Forest Ecology and*
531 *Management* 133, 13–22.
- 532 Brookes, P., Landman, A., Pruden, G., Jenkinson, D., 1985. Chloroform fumigation and the release of soil-
533 nitrogen—a rapid direct extraction method to measure microbial biomass nitrogen in soil. *Soil Biology &*
534 *Biochemistry* 17, 837–842
- 535 Clymo, R., 1984. The limits to peat bog growth. *Philosophical Transactions of the Royal Society London B* 303,
536 605-654.
- 537 Comeau, L.-P., Hergoualc’h, K., Hartill, J., Smith, J., Verchot, L., Peak, D., Salim, A., 2016. How do the
538 heterotrophic and the total soil respiration of an oil palm plantation on peat respond to nitrogen fertilizer
539 application? *Geoderma* 268, 41–51
- 540 Druille. M., García-Parisi, P., Golluscio, R., Cavagnaro, F., Omacini, M., 2016. Repeated annual glyphosate
541 applications may impair beneficial soil microorganisms in temperate grassland. *Agriculture, Ecosystems &*
542 *Environment* 230, 184–190.
- 543 Dungait, J., Hopkins, D., Gregory, A., Whitmore, A., 2012. Soil organic matter turnover is governed by
544 accessibility not recalcitrance. *Global Change Biology* 18, 1781–1796.
- 545 Evans, C., Page, S.E., Jones, T., Moore, S., Gauci, V., Laiho, R., Hruška, J., Allott, T., Billett, M., Tipping, E.,
546 Freeman, C., Garnett, M., 2014. Contrasting vulnerability of drained tropical and high-latitude peatlands to fluvial
547 loss of stored carbon. *Global Biogeochemical Cycles* 28, 1215–1234.
- 548 Fang, Y., Singh, B., Singh, B., 2015. Effect of temperature on biochar priming effects and its stability in soils. *Soil*
549 *Biology & Biochemistry* 80, 136–145.
- 550 Fenner, N., Freeman, C., 2011. Drought-induced carbon loss in peatlands. *Nature Geoscience* 4, 895–900.

- 551 Freeman C, Ostle N, Kang H (2001) An enzymic 'latch' on a global carbon store. *Nature* 409:149
- 552 Hertel, D., Harteeveld, M., Leuschner, C., 2009. Conversion of a tropical forest into agroforest alters the fine root-
553 related carbon flux to the soil. *Soil Biology & Biochemistry* 41, 481–490.
- 554 Hirano, T., Page, SE., Kusin, K., Limin, S., Osaki, M., 2014. Carbon dioxide emissions through oxidative peat
555 decomposition on a burnt tropical peatland. *Global Change Biology* 20, 555–565.
- 556 Hooijer, A., Page, SE., Jauhiainen, J., Lee, W., Lu, X., Idris, A., Anshari, G., 2012. Subsidence and carbon loss in
557 drained tropical peatlands. *Biogeosciences*, 9, 1053–1071.
- 558 Hoscilo, A., Page, SE., Tansey, K., Rieley, J., 2011. Effect of repeated fires on land-cover change on peatland in
559 southern Central Kalimantan, Indonesia, from 1973 to 2005. *International Journal of Wildland Fire* 20, 578–588.
- 560 Hoyos-Santillan, J., Lomax, B., Large, D., Turner, B., Boom, A., Lopez, O., Sjögersten, S., 2016. Quality not
561 quantity: Organic matter composition controls of CO₂ and CH₄ fluxes in neotropical peat profiles. *Soil Biology &*
562 *Biochemistry* 103, 86–96.
- 563 Hoyos-Santillan, J., Lomax, B., Large, D., Turner, B., Boom, A., Lopez, O., Sjögersten, S., 2015. Getting to the
564 root of the problem: litter decomposition and peat formation in lowland Neotropical peatlands. *Biogeochemistry*
565 126, 115–129.
- 566 Ishikura, K., Yamada, H., Toma, Y., Takakai, F., Morishita, F., Darung, U., Limin, A., Limin, S., Hatano, R., 2017.
567 Effect of groundwater level fluctuation on soil respiration rate of tropical peatland in Central Kalimantan,
568 Indonesia. *Soil Science and Plant Nutrition*, 63(1), 1–13.
- 569 Iiyama, I., Osawa, K., 2010. Surface O₂ influx related to soil O₂ profiles in a drained tropical peatland. *Soil science*
570 *and plant nutrition* 56, 517–520.
- 571 IPCC (2014), 2013 Supplement to the 2006 IPCC guidelines for national greenhouse gas inventories: wetlands. In:
572 Hiraishi, T., Krug, T., Tanabe, K., Srivastava, N., Baasansuren, J., Fukuda, M., Troxler, TG. (eds). IPCC, Geneva,
573 353 pp.
- 574 Jackson, C., Liew, K., Yule, C., 2009. Structural and Functional Changes with Depth in Microbial Communities in
575 a Tropical Malaysian Peat Swamp Forest. *Microbial Ecology* 57, 402–412.

576 Jauhiainen, J., Page SE., Vasander, H., 2016a. GHG dynamics in degraded and restored tropical peatlands. *Mires*
577 *and Peat* 17(6), 1–12.

578 Jauhiainen, J., Silvennoinen, H., Könönen, M., Limin, S., Vasander, H., 2016b. Management driven changes in
579 carbon mineralization dynamics of tropical peat. *Biogeochemistry* 129(1), 115–132.

580 Jauhiainen, J., Kerojoki, O., Silvennoinen, H., Limin, S., Vasander, H., 2014. Heterotrophic respiration in drained
581 tropical peat is greatly affected by temperature—a passive ecosystem cooling experiment. *Environmental Research*
582 *Letters* 9 (105013), 18pp.

583 Jauhiainen, J., Takahashi, H., Heikkinen, J., Martikainen, P., Vasander, H., 2005. Carbon fluxes from a tropical peat
584 swamp forest floor. *Global Change Biology* 11, 1788–1797.

585 Knicker, H., 2007. How does fire affect the nature and stability of soil organic nitrogen and carbon? A review.
586 *Biogeochemistry* 85, 91–118.

587 Kuzyakov, Y., 2010. Priming effects: Interactions between living and dead organic matter. *Soil Biology &*
588 *Biochemistry* 42, 1363–1371.

589 Kwon, M., Haraguchi, A., Kang, H., 2013. Long-term water regime differentiates changes in decomposition and
590 microbial properties in tropical peat soils exposed to the short-term drought. *Soil Biology & Biochemistry* 60, 33–
591 44.

592 Könönen, M., Jauhiainen, J., Laiho, R., Spetz, P., Kusin, K., Limin, S., Vasander, H., 2016. Land use increases the
593 recalcitrance of tropical peat. *Wetlands Ecology and Management* 24(6), 717–731.

594 Könönen, M., Jauhiainen, J., Laiho, R., Kusin, K., Vasander, H., 2015. Physical and chemical properties of tropical
595 peat under stabilised land uses. *Mires and Peat* 16(8), 1–13.

596 Laiho, R., 2006. Decomposition in peatlands: reconciling seemingly contrasting results on the impacts of lowered
597 water levels. *Soil Biology & Biochemistry* 38, 2011–2024.

598 Lampela, M., Jauhiainen, J., Vasander, H., 2014. Surface peat structure and chemistry in a tropical peat swamp
599 forest. *Plant and Soil* 382, 329–347.

600 Miettinen, J., Hooijer, A., Vernummen, R., Liew, S., Page, SE., 2017. From carbon sink to carbon source: extensive
601 peat oxidation in insular Southeast Asia since 1990. *Environmental Research Letters* 12, 1–10.

602 Miettinen, J., Shi, C., Lies, S., 2016. Land cover distribution in the peatlands of Peninsular Malaysia, Sumatra and
603 Borneo in 2015 with changes since 1990. *Global Ecology and Conservation* 6, 67–78.

604 Moore, S., Evans, C., Page, SE., Garnett, M., Jones, T., Freeman, C., Hooijer, A., Wiltshire, A., Limin, S., Gauci,
605 V., 2013. Deep instability of deforested tropical peatlands revealed by fluvial organic carbon fluxes. *Nature* 493,
606 660–663.

607 Nguyen, B., Lehmann, J., Hockaday, W., Joseph, S., Masiello, C., 2010. Temperature sensitivity of black carbon
608 decomposition and oxidation. *Environmental science and technology* 44(9), 3324–3331

609 Nieminen, M., Koskinen, M., Sarkkola, S., Lauren A., Kaila, A., Kiikkila, O., Nieminen, T., Ukonmaanaho, L.,
610 2015. Dissolved organic carbon export from harvested peatland forests with differing site characteristics. *Water,*
611 *Air and Soil Pollution*. 226: 181.

612 Niiyama, K., Kajimoto, T., Matsuura, Y., Yamashita, T., Matsuo, N., Yashiro, Y., Ripin, A., Kassim, R., Noor, S.
613 (2010). Estimation of root biomass based on excavation of individual root systems in a primary dipterocarp forest in
614 Pasoh Forest Reserve, Peninsular Malaysia. *Journal of Tropical Ecology* 26, 271–284

615 Oksanen, L. 2001. Logic of experiments in ecology: is pseudoreplication a pseudoissue? *Oikos* 94: 27–38.

616 Page, SE., Hooijer, A., 2016. In the line of fire: the peatlands of Southeast Asia. *Philosophical Transactions of the*
617 *Royal Society B: Biological Sciences* 371, 20150176.

618 Page, SE., Rieley, J., Banks, C., 2011. Global and regional importance of the tropical peatland carbon pool. *Global*
619 *Change Biology* 17, 798–818.

620 Page, SE., Rieley, J., (ed) 2005. *The Mega Rice Project In Central Kalimantan, Indonesia 1995-1999*. In: *Wise Use*
621 *of Tropical Peatlands: Focus on Southeast Asia*. Published by: ALTEERRA - Wageningen University and Research
622 Centre and the EU INCO - STRAPEAT and RESTORPEAT Partnerships. ISBN: 903-270-347-1

623 Page, SE., Rieley, J., Shoty, Ø., Weiss D., 1999. Interdependence of peat and vegetation in a tropical peat swamp
624 forest. *Philosophical Transactions of the Royal Society B* 354, 1885–1897.

625 Pritsch, K., Courty, P., Churin, J-L., Cloutier-Hurteau, B., Ali, M., Damon, C., Duchemin, M., Egli, S., Ernst, J.,
626 Fraissinet-Tachet, L., Kuhar, F., Legname, E., Marmeisse, R., Müller, A., Nikolova, P., Peter, M., Plassard, C.,
627 Richard, F., Schloter, M., Selosse, M-A., Franc, A., Garbaye, J., 2011. Optimized assay and storage conditions for
628 enzyme activity profiling of ectomycorrhizae. *Mycorrhiza* 21(7), 589–600.

629 Rejmánková, E., 2001. Effect of experimental phosphorus enrichment on oligotrophic tropical marshes in Belize,
630 Central America. *Plant and Soil* 236, 33–53.

631 Sinsabaugh, R., 1994. Enzymic analysis of microbial pattern and process, *Biol. Fertil. Soils*, 17, 69–74.

632 Sjögersten, S., Cheesman, A., Lopez, O., Turner, B., 2011. Biogeochemical processes along a nutrient gradient in a
633 tropical ombrotrophic peatland. *Biogeochemistry* 104, 147–163.

634 Smith, D., Townsend, T., Choy, A., Hardy, I., Sjögersten, S., 2012. Short-term soil carbon sink potential of oil palm
635 plantations. *Global Change Biology Bioenergy* 4, 588–596.

636 Straková, P., Niemi, R., Freeman, C., Peltoniemi, K., Toberman, H., Heiskanen, I., Fritze, H., Laiho, R., 2011.
637 Litter type affects the activity of aerobic decomposers in a boreal peatland more than site nutrient and water table
638 regimes. *Biogeosciences* 8, 2741–2755.

639 Sulistiyanto, Y., 2004. Nutrient dynamics in different sub-types of peat swamp forest in central Kalimantan,
640 Indonesia. PhD thesis, University of Nottingham, UK. 388 pp.

641 Sundari, S., Hirano, T., Yamada, H., Kusin, K., Limin, S., 2012. Effect of groundwater level on soil respiration in
642 tropical peat swamp forests. *Journal of Agricultural and Forest Meteorology* 68, 121–134.

643 Turner, B., Romero, T., 2010. Stability of hydrolytic enzyme activity and microbial phosphorus during storage of
644 tropical rain forest soils. *Soil Biology & Biochemistry* 42, 459–465.

645 Yang, J., Tian, H., Tao, B., Ren, W., Lu, C., Pan, S., Wang, Y., Liu, Y., 2015. Century-scale patterns and trends of
646 global pyrogenic carbon emissions and fire influences on terrestrial carbon balance, *Global Biogeochemical Cycles*,
647 29, 1549–1566.

648 Yule, C., Gomez, L., 2009. Leaf litter decomposition in a tropical peat swamp forest in Peninsular Malaysia.
649 *Wetland Ecology and Management* 17, 231–241.

- 650 Vance, E., Brookes, P., Jenkinson, D., 1987. An extraction method for measuring soil microbial biomass-C. Soil
651 Biology and Biochemistry 19, 703–707.
- 652 Wüst, R., Jacobsen, G., van der Gaast, H., Smith, A., 2008. Comparison of radiocarbon ages from different organic
653 fractions in tropical peat cores: insights from Kalimantan, Indonesia. Radiocarbon 50(3), 359–372.
- 654 Yule, C., Gomez, L., 2009. Leaf litter decomposition in a tropical peat swamp forest in Peninsular Malaysia.
655 Wetlands Ecology and Management 17, 231–241.
- 656 Zaller, J., Heigl, F., Ruess, L., Grabmaier, A., 2014. Glyphosate herbicide affects belowground interactions
657 between earthworms and symbiotic mycorrhizal fungi in a model ecosystem. Scientific Reports 4(5634), 1–8.

Table 1. Selected soil properties at the study sites corresponding to the sampling depths in this study. Data from this and other studies (source mentioned under the table).

	Swamp forest				Reforested site				Degraded site				Agricultural site			
	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30
Sampling depth, cm	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30	0-5	5-15	15-25	25-30
BD, g cm ⁻³ ⁽¹⁾	0.11	0.10	0.11	0.11	0.13	0.11	0.10	0.09	0.17	0.12	0.13	0.12	0.13	0.12	0.10	0.9
pH ⁽²⁾	3.5	3.49	3.49	3.46	3.4	3.41	3.43	3.43	3.55	3.47	3.52	3.48	3.43	3.44	3.46	3.44
Moisture %	76.6	86.2	87.3	87.3	70.7	74.6	80.8	82.4	42.7	64.1	75.4	80.7	-	-	-	-
Moisture %	81.4	82.7	85.6	88.6	63.3	77.0	79.8	83.1	19.3	66.5	65.2	79.3	34.1	78.9	81.9	83.8
Moisture, %	88.3	88.2	87.4	87.0	76.8	81.7	83.6	84.3	76.4	81.9	84.4	84.9	83.7	85.0	86.9	88.1
Soil temp, °C	25.4	25.6	25.4	25.6	27.5	27.5	27.3	27.1	27.9	27.4	27.2	27.3	-	-	-	-
	24.4	24.6	24.7	24.5	26.3	26.6	26.6	26.5	27.2	26.9	27.2	27.2	-	-	-	-
NO ₃ -N, mg g ⁻¹	0.016	-	0.013	-	0.007	-	0.008	-	0.008	-	0.007	-	-	-	-	-
	0.050	-	0.047	0.056	0.007	-	0.016	0.010	0.008	-	0.008	0.008	-	-	-	-
NH ₄ ⁺ -N, mg g ⁻¹	0.092	-	0.068	-	0.041	-	0.033	-	0.020	-	0.024	-	-	-	-	-
	0.154	-	0.160	0.122	0.038	-	0.037	0.028	0.019	-	0.029	0.030	-	-	-	-

¹⁾ Peat dry bulk density, soil temperature and moisture are from data collected monthly for 1+ year (Jyrki Jauhainen, unpublished). Soil temperature for wet season was measured in March and dry season in August by loggers set to 1-hour recording intervals. The soil temperatures at the agricultural and degraded sites can be assumed to be similar due to their close location and similar shading effect.

²⁾ pH was determined from peat: deionized water solution (1:2) in field conditions. Measured from the samples collected for this study.

³⁾ Moisture concentration was determined by drying samples at 105 °C for 24 h. Wet and dry season moisture was measured from the samples collected for this study

⁴⁾ Total NO₃-N and NH₄⁺-N were extracted from approximately 2 g of fresh peat (dry mass concentration approximately 24 ± 14 %) with 2M KCl and analysed with Lachat Instrument (Method 12-107-06-2-A for NH₄ and 12-107-04-1-E for NO₃). Wet season samples were collected in March and dry season samples in September in 2014.

Supplementary Table 1. The mean extracellular enzymatic activity in peat samples according to season. (mean \pm standard error). Different letters indicate significant ($p < 0.05$) difference between sites at the respective depths.

ENZYME	Season	Depth, cm	Swamp forest	Reforested site	Degraded site	Agricultural site
b-Xylanase	wet	0-3	10.3 \pm 2.19 ^a	2.90 \pm 0.50 ^b	0.21 \pm 0.70 ^c	–
		3-10	2.83 \pm 0.65 ^a	1.81 \pm 0.85 ^a	0 \pm 0 ^b	–
		10-20	1.71 \pm 0.65 ^a	1.52 \pm 1.22 ^{ab}	0.04 \pm 0.29 ^b	–
		20-30	1.51 \pm 0.68 ^a	1.52 \pm 0.75 ^{ab}	1.32 \pm 0.66 ^b	–
	dry	0-3	10.9 \pm 2.55 ^a	0.86 \pm 0.85 ^b	20.6 \pm 4.30 ^{ab}	19.10 \pm 5.41 ^{ab}
		3-10	3.10 \pm 0.50 ^a	1.11 \pm 0.76 ^{ab}	0.01 \pm 0.21 ^b	1.66 \pm 1.12 ^{ab}
		10-20	3.22 \pm 0.54 ^a	0.99 \pm 0.58 ^{ab}	0.02 \pm 0.18 ^b	0.96 \pm 0.80 ^{ab}
		20-30	3.68 \pm 0.32 ^a	0.93 \pm 0.49 ^{ab}	0.03 \pm 0.18 ^b	1.96 \pm 1.54
b-Glucosidase	wet	0-3	16.56 \pm 3.97 ^a	2.28 \pm 0.29 ^{ab}	0.01 \pm 0.16 ^b	–
		3-10	4.50 \pm 0.53 ^a	3.77 \pm 0.69 ^b	0.59 \pm 0.92 ^c	–
		10-20	4.18 \pm 0.39 ^a	3.77 \pm 0.37 ^a	0.75 \pm 0.70 ^b	–
		20-30	3.59 \pm 0.70 ^a	3.56 \pm 0.69 ^a	0.38 \pm 0.61 ^b	–
	dry	0-3	43.16 \pm 7.02 ^a	7.76 \pm 1.41 ^b	43.05 \pm 5.90 ^a	39.39 \pm 7.62 ^a
		3-10	2.33 \pm 0.77 ^a	3.03 \pm 0.51 ^a	0 \pm 0 ^b	0.52 \pm 0.92 ^{ab}
		10-20	1.35 \pm 1.63 ^b	2.54 \pm 0.56 ^a	0 \pm 0 ^c	0.43 \pm 0.57 ^{bc}
		20-30	2.95 \pm 3.49 ^{ab}	3.59 \pm 0.69 ^a	0 \pm 0 ^b	0.09 \pm 0.42 ^{bc}
N-acetyl-b-glucosaminidase	wet	0-3	72.92 \pm 7.98 ^a	3.27 \pm 1.09 ^b	0.09 \pm 0.43 ^c	–
		3-10	3.94 \pm 1.48 ^a	3.02 \pm 1.34 ^b	0.48 \pm 0.93 ^c	–
		10-20	3.73 \pm 0.81 ^a	1.16 \pm 0.86 ^b	0.90 \pm 0.96 ^b	–
		20-30	1.68 \pm 1.18	0.87 \pm 0.70	1.94 \pm 0.65	–
	dry	0-3	36.10 \pm 2.90 ^a	6.87 \pm 2.14 ^b	12.32 \pm 3.42 ^b	22.38 \pm 5.98 ^{ab}
		3-10	3.39 \pm 2.42 ^{ab}	0.45 \pm 1.32 ^b	0.23 \pm 0.55 ^b	5.12 \pm 1.06 ^a
		10-20	0.78 \pm 1.26 ^a	0.01 \pm 0.24 ^a	0.40 \pm 0.79 ^a	4.22 \pm 0.35 ^b
		20-30	0.49 \pm 1.50	0.04 \pm 0.29	0.63 \pm 1.36	3.81 \pm 1.84
Phospho-monoesterase	wet	0-3	861.4 \pm 1.83 ^a	0 \pm 0 ^b	0 \pm 0 ^b	–
		3-10	68.97 \pm 11.89 ^a	9.54 \pm 6.91 ^b	7.31 \pm 5.95 ^b	–
		10-20	58.66 \pm 11.37 ^a	0 \pm 0 ^b	0 \pm 0 ^b	–
		20-30	4.08 \pm 4.04 ^a	0 \pm 0 ^b	0 \pm 0 ^b	–
	dry	0-3	1696.3 \pm 39.01 ^a	40.81 \pm 4.63 ^b	80.24 \pm 1.52 ^b	75.13 \pm 10.72 ^b
		3-10	103.07 \pm 4.44 ^a	0.23 \pm 1.08 ^b	6.80 \pm 1.08 ^c	2.15 \pm 2.54 ^d
		10-20	77.10 \pm 7.18 ^a	0.35 \pm 1.32 ^b	9.69 \pm 2.10 ^c	7.05 \pm 2.39 ^d
		20-30	52.61 \pm 7.91 ^a	0 \pm 0 ^b	5.23 \pm 3.36 ^c	14.33 \pm 3.78 ^c
Arylsulfatase	wet	0-3	6.09 \pm 2.37	1.67 \pm 1.27	0.26 \pm 1.01	–
		3-10	5.82 \pm 0.57 ^a	3.05 \pm 0.62 ^b	0.00 \pm 0.09 ^c	–
		10-20	5.92 \pm 0.55 ^a	3.07 \pm 0.57 ^b	0.12 \pm 0.35 ^c	–
		20-30	5.61 \pm 0.39 ^a	2.53 \pm 0.29 ^b	0.24 \pm 0.48 ^c	–
	dry	0-3	4.94 \pm 3.32 ^a	1.60 \pm 0.25 ^a	0.82 \pm 0.51 ^b	3.15 \pm 0.01 ^{ab}
		3-10	1.67 \pm 0.48 ^a	1.38 \pm 0.49 ^a	0.22 \pm 0.51 ^b	2.20 \pm 0.74 ^a
		10-20	1.33 \pm 0.62 ^{ab}	1.24 \pm 0.31 ^b	0.14 \pm 0.81 ^c	1.92 \pm 0.40 ^a
		20-30	1.81 \pm 0.47	1.34 \pm 0.25	0.25 \pm 0.39	2.18 \pm 0.05

Figure 1

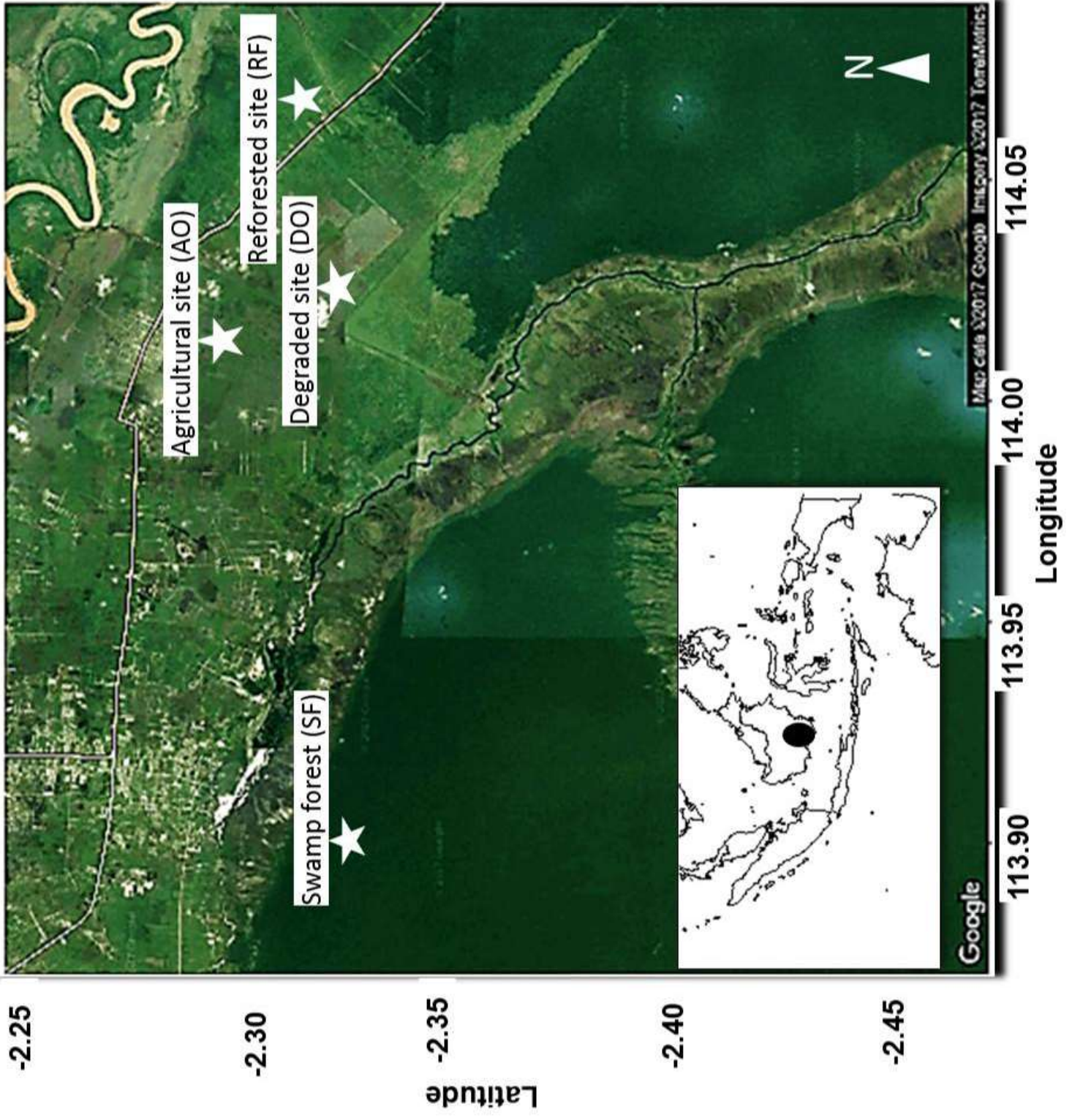






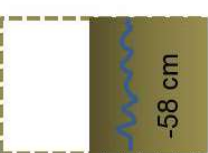



Figure 2

Intensity of land management increases

Site	Swamp forest	Reforested site	Degraded site	Agricultural site
Management history	2°19'16.96"S, 113°53'54.29"E Selective logging practices prior to 1997, when small ditches were also dug	2°20'43.24"S, 114°3'31.76"E Drained and clear-felled in 1997, fires only prior to 1998, reforested in 2008, addition of herbicides	2°19'25.11"S, 114°1'5.15"E Drained and clear-felled in 1997. Burned repeatedly since then (1997, 1999, 2006, 2009)	2°17'25.21"S, 114°0'41.20"E Deforested and clear-felled and under smallholder cultivation since 1980s
Current vegetation	Near-pristine peat swamp forest	<i>Shorea balangeran</i> trees 2000 trees ha ⁻¹	Ferns and scattered trees	During sampling maize, cassava, tomatoes
Photos from field				
Original peat surface indicated with dotted line and current surface with solid box. Annual median WTL represented by wavy line. (1)				
Causes of peat deposit depletion at altered sites	No depletion caused by management	Peat subsidence	Peat loss ~0.5m due to fires	Peat loss ~0.5m due to fires and tilling
Reported CO ₂ emissions from related sites	Considered as C-neutral (IPCC, 2014)	Shallow drained plantation 1.5 tons CO ₂ ha ⁻¹ yr ⁻¹ (IPCC, 2014)	Degraded peatlands 5.0–5.3 tons CO ₂ ha ⁻¹ yr ⁻¹ (Hirano et al. 2012; IPCC, 2014)	Croplands 14 tons CO ₂ ha ⁻¹ yr ⁻¹ (IPCC, 2014)

(1)WT measured twice monthly in swamp forest, reforested site and degraded site, and between March and August in agricultural site in 2014.

Figure3

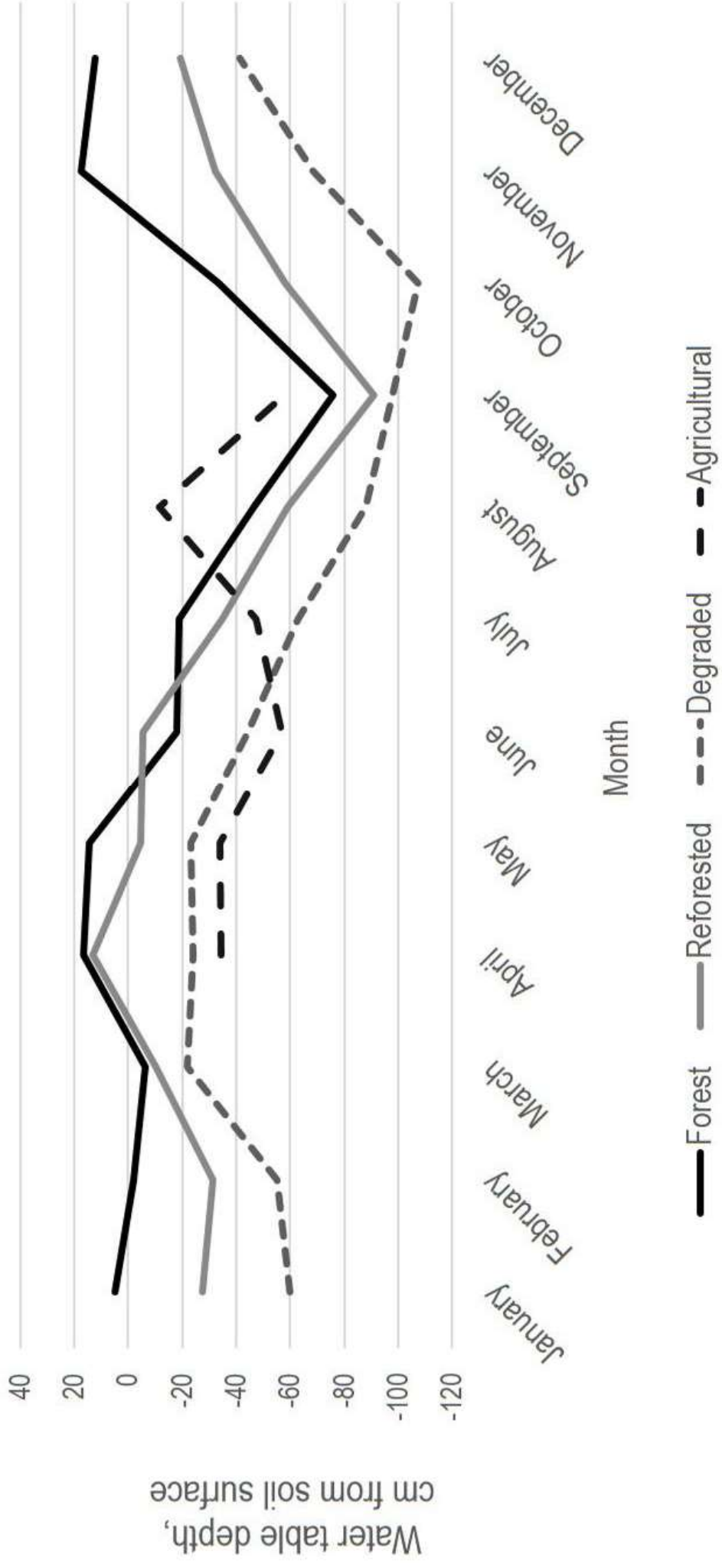


Figure 4

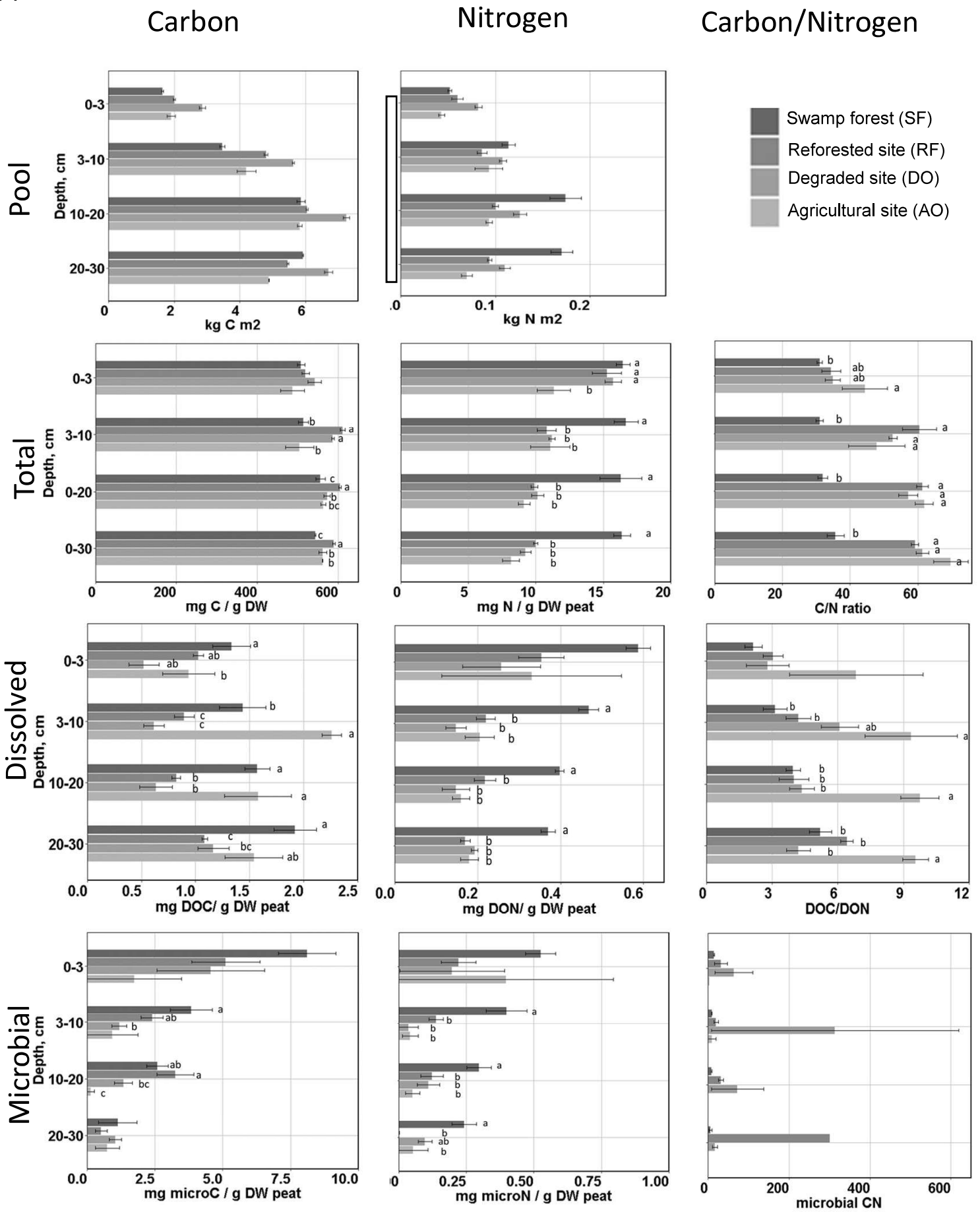


Figure 5_earlier numb 6

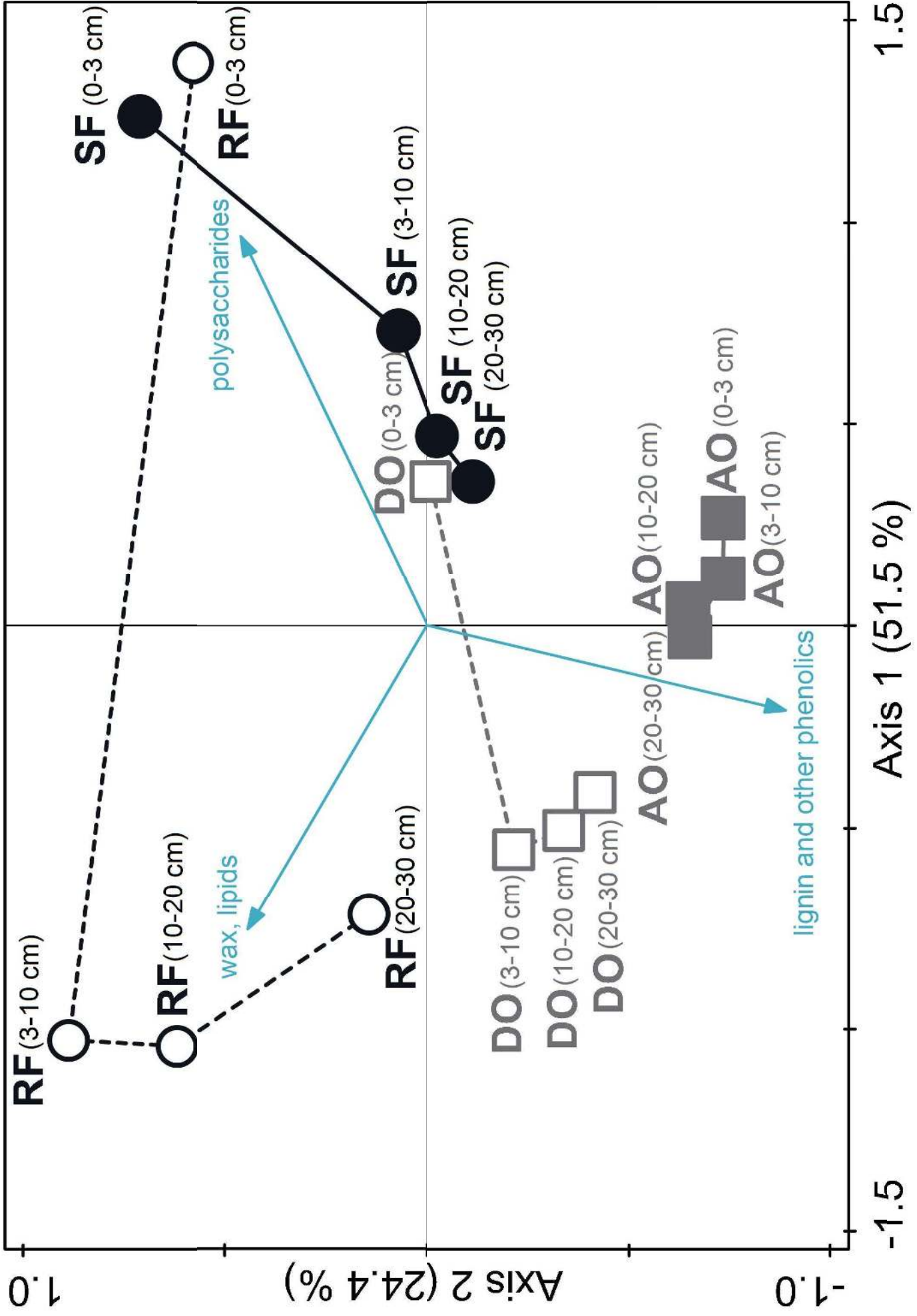


Figure 6_previous numb 7

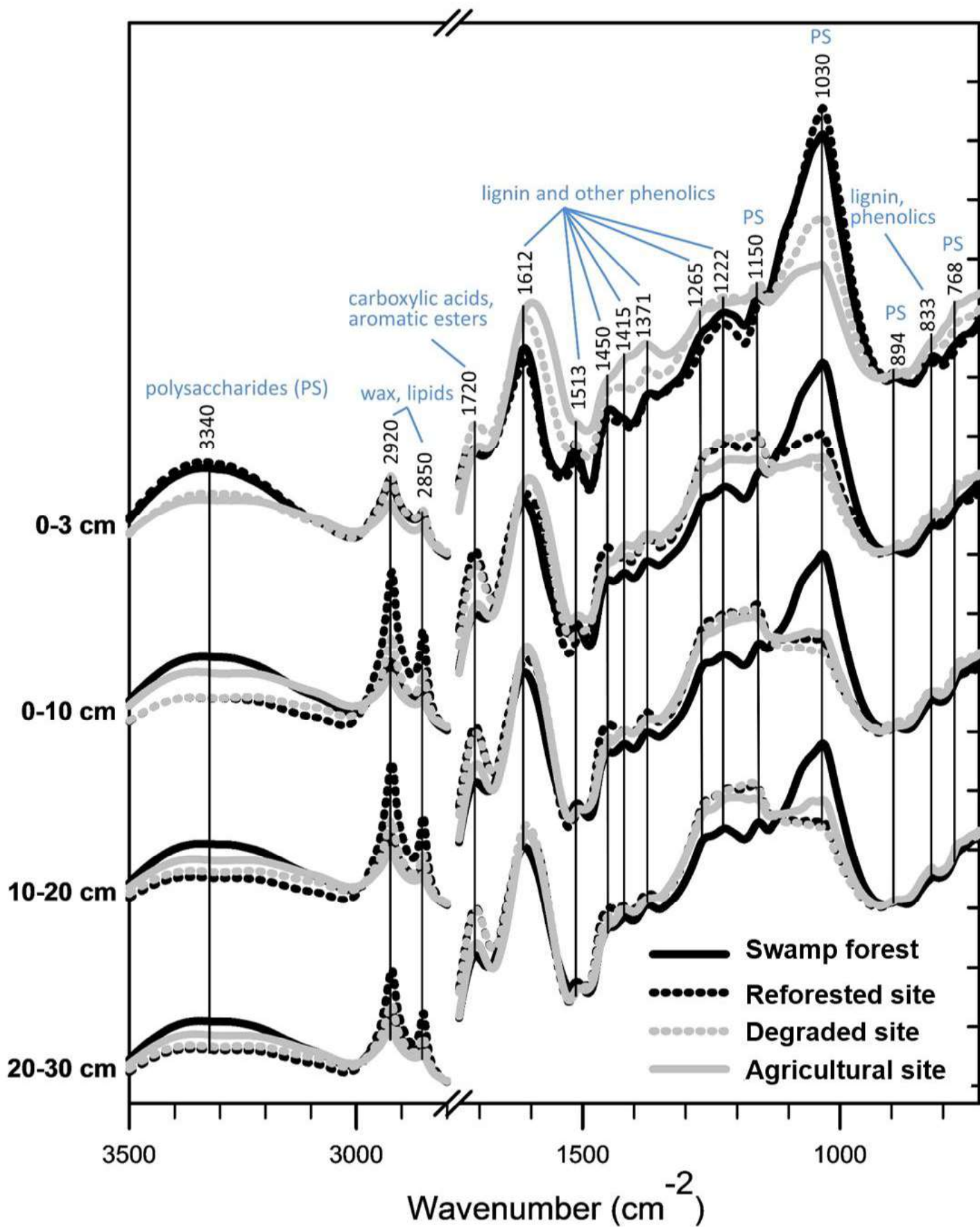


Figure 7_previous numb 8

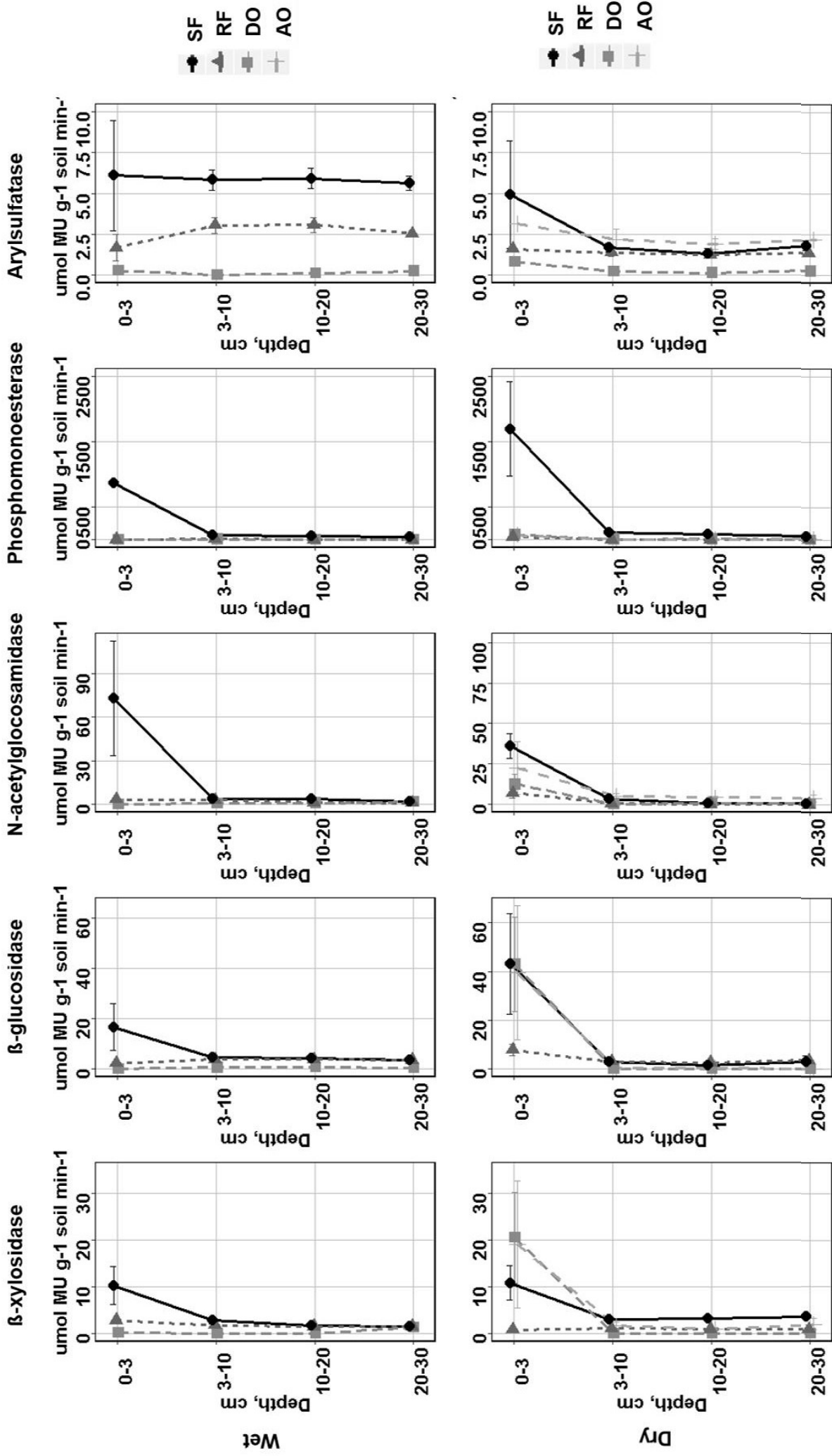


Figure 8

Figure 8

