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DOI:

[10.1016/j.orggeochem.2016.01.010](https://doi.org/10.1016/j.orggeochem.2016.01.010)

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Document Version

Peer reviewed version

Citation for published version (Harvard):

Bendle, J & Wang, C 2016, 'Impacts of pH and temperature on soil bacterial 3-hydroxy fatty acids: development of novel terrestrial proxies', *Organic Geochemistry*, vol. 94, pp. 21-31.
<https://doi.org/10.1016/j.orggeochem.2016.01.010>

[Link to publication on Research at Birmingham portal](#)

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Checked for eligibility: 22/03/2016

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1 **Impacts of pH and temperature on soil bacterial 3-hydroxy fatty acids:**
2 **development of novel terrestrial proxies**

3

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19

20 **Abstract:** Gram-negative bacterial 3-hydroxy fatty acids (3-OH-FAs)
21 biomarkers are widespread in a variety of environments including both marine
22 and terrestrial sediments (including speleothems). In this study we analysed
23 the hydroxylated membrane lipids of 26 soil samples from an altitudinal
24 transect of Shennongjia Mountain (Mt.) in central China to study the
25 environmental factors controlling the relative distribution of 3-OH-FAs. Our
26 results show that both the ratio of the summed *iso* and *anteiso* to the total
27 amount of *normal* 3-OH-FAs (RIAN), and the ratio of summed *iso* and *anteiso*
28 to the total amount of all 3-OH-FAs (Branched Index) were primarily related to
29 the pH of soil ($R^2 = 0.70$ and 0.70 , respectively). Additionally, the *anteiso* to
30 *normal* 3-hydroxy fatty acids ratio of the C_{15} and C_{17} homologues (RAN₁₅ and
31 RAN₁₇) shows a significant negative correlation with mean annual air
32 temperature (MAAT) ($R^2=0.51$ and 0.48 , respectively). When comparing the 3-
33 OH-FA based indices with established glycerol dialkyl glycerol tetraether
34 (GDGT) based indices from the same soil samples, the RIAN and Branched
35 Index show strong linear correlations with the cyclisation ratio of branched
36 tetraethers (CBT) ($R^2 = 0.77$ and 0.74 , respectively), and the RAN₁₅ and RAN₁₇
37 show negative correlations with the MBT/CBT-MAAT (MBT, methylation index
38 of branched tetraethers) ($R^2 = 0.61$ and 0.36 , respectively). Our new field-based
39 correlations demonstrate the physiological response of Gram-negative bacterial
40 cell membranes to the external environment and suggest that 3-hydroxy fatty

41 acids can be applied in palaeoenvironmental studies to estimate past MAAT
42 and soil pH.

43

44 **Keywords:** proxy, 3-hydroxy fatty acid, soil, temperature, soil pH,
45 palaeoclimate

46

47 **1. Introduction**

48 A wide range of environmental information from both terrestrial and
49 marine realms is required from palaeoclimate archives to better understand
50 the climate system and to provide a palaeoclimatic context for predictions of
51 future rates of climate change, impact and Earth System sensitivity. To date,
52 various geochemical proxies based on inorganic and organic fossil remains have
53 been applied in order to reconstruct past environmental parameters. Organic
54 biomarkers have become widely deployed tools in the reconstruction of past
55 environmental conditions, due in part to: a) the sensitive physiological
56 responses of cell membranes and structural lipids to the external environment
57 and b) their relatively high preservation potential (Summons, 1993; Eglinton
58 and Eglinton, 2008). Since the 1960's a large array of lipid biomarkers with
59 applications in palaeoclimatology have been identified, including plant waxes,
60 hopanes, alkanes and glycerol dialkyl glycerol tetraethers (GDGTs). Two
61 proxies, U_{37}^K , (Brassell et al., 1986; Prahl and Wakeham, 1987; Sachs et al.,

62 2001; Haug et al., 2005) and TEX₈₆ (Schouten et al., 2002; Kim et al., 2008),
63 based on C₃₇ alkenones and GDGTs, respectively, have been widely employed
64 to calculate sea surface temperatures (SST) as far back as the Jurassic
65 (Jenkyns et al., 2012).

66 Numerous lipid biomarkers derived from terrestrial organic matter are
67 preserved in lacustrine (e.g. Castañeda and Schouten, 2011) and marine
68 (Pancost and Boot, 2004) archives. Commonly utilised biomarker groups
69 include higher plant derived *n*-alkyl compounds, terpenoids and lignins
70 (Pancost and Boot, 2004) and soil bacterial branched-GDGTs (Weijers et al.,
71 2007a). Such compounds can be used to reconstruct general changes in inputs
72 and provenance of terrestrial material (Pancost and Boot, 2004, Seki et al.,
73 2014). Compound specific isotopic analyses, particularly on higher plant waxes
74 have expanded the range of palaeoclimatic applications, for example, D/H
75 analysis is used to infer changes in past hydrological regimes (Sachse et al.,
76 2012 and reference therein) and the $\delta^{13}\text{C}$ analysis of higher plant biomarkers is
77 a powerful tool to constrain changes in C₃ vs C₄ vegetation (e.g. Hughen et al.,
78 2004). More recently, the bacterial GDGT based cyclization of branched
79 tetraether (CBT) proxy has been developed and applied for the reconstruction
80 of soil pH in terrestrial settings (Weijers et al., 2007b). In parallel, the
81 combination of CBT with the methylation of branched tetraethers (MBT) index
82 may be deployed to estimate past variations in mean annual air temperature
83 (MAAT) (Weijers et al., 2007b). However, overall, relatively less attention has

84 been paid to terrestrial environments, compared to the marine realm, due to
85 the historical paucity of ubiquitous biomarkers with quantitative
86 palaeoclimatic utility. Thus the discovery and development of new quantitative
87 terrestrial proxies is of major significance. Targets of particular value are
88 compounds preserved in both aquatic and terrestrial sediments, as this
89 facilitates the correlation and comparison of palaeoclimatic records between
90 marine and terrestrial environments (Pancost and Boot, 2004; Castañeda and
91 Schouten, 2011).

92 Lipopolysaccharide (LPS) is the main component of the outer membrane of
93 Gram-negative bacteria. Lipid A, a constituent part of LPS, consists of
94 glucosamine units and fatty acids, many of the latter are 3-hydroxy fatty acids
95 (3-OH-FAs), also known as ω -hydroxy fatty acids, with carbon numbers from
96 C₁₀ to C₁₈ (Fig. 1) (Wollenweber and Rietschel, 1990; Szponar et al., 2002;
97 Szponar et al., 2003). These are bound to the glucosamine unit either by ester
98 bonds or amide bonds (Wollenweber et al., 1982; Kumar et al., 2002). A
99 significant body of literature demonstrates that the dominant precursors for
100 C₁₀-C₁₈ 3-OH-FAs compounds in the environment are Gram-negative bacteria
101 (Wollenweber and Rietschel, 1990; Saraf et al., 1997; Szponar et al., 2002;
102 Keinänen et al., 2003; Szponar et al., 2003). Such that 3-OH-FAs in the C₁₀-C₁₈
103 range are accepted as diagnostic markers for the characterisation and
104 quantification of Gram-negative bacterial LPS (i.e. endotoxins) in clinical and
105 environmental studies (Sonesson et al., 1990; Mielniczuk et al., 1993; Saraf et

106 al., 1997; Szponar et al., 2002; Keinänen et al., 2003; Wakeham et al., 2003;
107 Lee et al., 2004; Ferrando et al., 2005; Kračnik et al., 2006; Lee et al., 2007).
108 However, one study suggests C₁₀-C₁₈ 3-OH-FAs are also produced by Gram-
109 positive *Lactobacillus plantarum* (Sjogren et al., 2003). Additionally, long chain
110 3-OH-FAs (C₂₆-C₃₀) are reportedly derived from microalgae of the class
111 Eustigmatophyceae (Volkman et al., 1998).

112 3-OH-FAs with carbon chain lengths from C₁₀ to C₁₈ have been used to
113 quantify and characterize the Gram-negative bacterial community in samples
114 from a diverse array of environments, including atmospheric aerosols (Lee et
115 al., 2004) and marine dissolved organic matter (DOM) (Wakeham et al., 2003).
116 However, thus far, the relationship between 3-OH-FAs and environmental
117 parameters has not been systematically investigated in soils or sediments with
118 the aim of exploring the possible utility of these ubiquitous fatty acids as
119 quantitative environmental proxies.

120 We explore the distribution of these microbial biomarkers on Mt.
121 Shennongjia, a national reserve located at the northwest of Hubei province,
122 central China (31°15'-31°57'N, 109°59'-110°58'E) (Fig. 2), to test whether 3-OH-
123 FAs record a signal of sensitive and differential physiological responses, by
124 Gram-negative bacteria, to ambient environmental conditions, and if novel
125 quantitative proxies could be independently established for
126 palaeoenvironmental reconstruction.

127

128 **2. Methods**

129 *2.1 Sampling site*

130 Mt. Shennongjia, with an altitude of 3105 m above sea level (m.a.s.l.), is
131 located in a climatic region dominated by the Asian monsoon. Five
132 meteorological stations established at different altitudes in this region provide
133 a precise altitudinal record of meteorological conditions. Moreover, a large
134 gradient of soil pH, MAAT and mean annual precipitation (MAP) prevails on
135 Mt. Shennongjia, making it a natural laboratory to test the relationship
136 between 3-OH-FAs and environmental parameters. Average climatic conditions
137 trend from warm and dry conditions at the base (315 m.a.s.l.) to cool and wet
138 conditions at the highest sampling site (2840 m.a.s.l.), with MAAT varying
139 from 1.9 °C to 14.7 °C; MAP from 1226mm to 3313mm and soil humidity from
140 11.6% to 55.6% (Supplementary data Table 1). Soil pH varies from 4.49 to 7.98,
141 however it has no causal relationship with altitude, MAAT, MAP or soil
142 humidity (Fig. 3), indicating the pH is an independent environmental factor,
143 likely controlled by changes in bedrock geology. Both MAAT ($R^2=0.995$) and
144 MAP ($R^2= 0.951$) are highly correlated to altitude (and thus co-vary), according
145 to the linear regressions between altitude and climatic factors reported by Li
146 and Manfred (2002) based on the climatic data from the local meteorological
147 station (Songpei, 930 m.a.s.l.) and the four subsidiary stations in the Mt.
148 Shennongjia area (Yangriwan, 460 m.a.s.l.; Dajiuhu, 1700 m.a.s.l.;
149 Changyanwu, 2300 m.a.s.l.; the mountain observation tower, 2930 m.a.s.l.).

150 The vertical vegetation distribution on Shennongjia Mountain is very distinct.
151 Based on the latest investigation by Zhao et al., (2005), the vegetation zones
152 along the elevation gradient were described as follows: evergreen broadleaved
153 forest zone at altitudes below 900 m.a.s.l.; mixed evergreen and deciduous
154 broadleaved forest between 900 and 1500 m.a.s.l.; deciduous broadleaved forest
155 zone between 1500 and 2000 m.a.s.l.; mixed conifer and deciduous broadleaved
156 forest between 2000 and 2400 m.a.s.l.; and sub-alpine conifer forest zone
157 (including sub-alpine shrubs and meadows) above altitudes of 2400 m.a.s.l.
158 (Zhao et al., 2005).

159 *2.2 Sample collection*

160 Twenty-six soil samples were collected along an altitude transect of Mt.
161 Shennongjia between 315 and 2840 m.a.s.l. at altitudinal intervals of ca. 200 m.
162 The topmost leaf-litter layer was removed before sampling. Samples from each
163 soil are derived from the depth intervals between 0 to 10 cm. The samples were
164 wrapped in pre-combusted aluminium foil and then stored with ice bags. Upon
165 arrival at the laboratory, the soils were stored at -20°C in a freezer before
166 freeze drying. The location of sampling sites was measured by a portable GPS
167 instrument (Supplementary data Table 1). Soil moisture was determined by
168 measuring the weight difference before and after freeze drying. Then the dry
169 samples were ground into powder with a pestle and mortar. A late Holocene
170 lake sediment sample was taken from a core collected from Tianchi Lake in

171 Gansu Province, China (Zhou et al., 2010) (Fig. 2). A stalagmite sub-sample
172 was obtained from the HS4 stalagmite which was collected from Heshang Cave,
173 Hubei province, China (Hu et al., 2008) (Fig. 2). A marine sediment sample was
174 collected from IODP Site M0060, in the Baltic Sea.

175 *2.3 Soil pH measurement*

176 Soil pH data either comes from or was measured following the method of
177 Yang et al. (2015). Soil samples were mixed with ultrapure water in a ratio of
178 1:2.5 (g/mL). After standing for 30 min, the supernatant pH was measured,
179 using a meter with a precision of ± 0.01 . The pH was measured three times and
180 the mean value was taken as the final pH.

181 *2.4 Extraction and clean-up methods*

182 The soil, stalagmite and marine sediment samples were subjected to acid
183 hydrolysis following an optimized acid digestion method (Wang et al., 2012).
184 10g of homogenized sample was mixed with 30 mL pre-cleaned HCl (3M), and
185 then refluxed under 130 °C for 3h. After cooling, the solution was extracted x3
186 with DCM, to yield the Total Lipid Extract (TLE). The lake sediment was
187 hydrolysed by 0.3M KOH methanolic solution containing 5% water, heating
188 under 70 °C for 2h in a closed test tube. The neutral fraction was extracted
189 with *n*-hexane:DCM (9:1, v/v) and then the acid fraction was extracted with
190 DCM after adjusting the pH of the residues below 2 with pre-cleaned HCl. The
191 TLE (soils, stalagmite and marine sediment) and acid fraction (lake sediment)

192 was methylated by $\text{BF}_3\text{-MeOH}$ solution at 70 °C for 1.5h. The resulting fatty
193 acid methyl esters (FAMEs) were separated into non-OH-FAMEs and OH-
194 FAMEs following the method described by Jenske and Vetter (2008). Non-OH-
195 FAMEs were eluted in the first fraction with a solvent mixture of n-hexane and
196 ethyl acetate (v/v =98:2), whereas OH-FAMEs were obtained by elution with
197 100% ethyl acetate. The OH-FAME fraction was further derivatised by BSTFA
198 (N, O-bis (trimethylsilyl) trifluoroacetamide) at 70 °C for 1.5h before further
199 analysis by gas chromatogram-mass spectrometer (GC-MS).

200 *2.5 Instrumentation*

201 The 3-OH-FAs from soils, stalagmite and marine sediment were analysed
202 by an Agilent 7890A gas chromatogram and 5975C mass spectrometer (GC-MS)
203 equipped with a ZB-5MS fused silica capillary column (60 m \times 0.25 mm \times 0.25
204 μm) at the China University of Geosciences (Wuhan). The GC oven
205 temperature was ramped from 70 °C to 200 °C at 10 °C/min, then to 310 °C at
206 3 °C/min, held at 310 °C for 47 min. The carrier gas was Helium (99.999%) and
207 the gas flow was 1.0 mL/min. The 3-OH-FAs from Tianchi Lake were analysed
208 by a 7890B gas chromatogram and 5977A mass spectrometer equipped with a
209 BP5MS fused silica capillary column (60 m \times 0.32 mm \times 0.25 μm) at the
210 University of Birmingham. The ionization energy of the mass spectrometer was
211 set at 70 eV. The 3-OH-FAs were identified based on their mass spectra and
212 relative retention times (Fig. 4). All the 3-OH-FAs TMSi esters show diagnostic

213 fragment ions, m/z 175 ($[\text{CH}_3]_3\text{SiO} = \text{CHCH}_2\text{CO}_2\text{CH}_3$), due to the cleavage
214 between C_3 and C_4 , and M-15 (base peak) results from a loss of a CH_3 group.
215 Other characteristic ions include m/z 103, 89, 133, 159, and M^+-31 (Eglinton et
216 al., 1968; Mielniczuk et al., 1993; Volkman et al., 1999). Samples were analysed
217 in duplicate or triplicate to obtain the analytical errors of the proxies. The
218 analytical errors are graphically illustrated in the relevant figures with error
219 bars.

220 **3. Results and discussion:**

221 *3.1 Distribution of 3-OH-FAs*

222 A total of 26 soil samples from Mt. Shennongjia were analysed. The carbon
223 number of the 3-OH-FAs ranges from C_{10} to C_{18} , including *iso*- C_{11} , C_{13} , C_{15} , C_{16} ,
224 C_{17} and *anteiso*- C_{13} , C_{15} , C_{17} 3-OH-FAs. *n*- C_{14} is the dominant homologue (Fig.
225 5). The distribution of the Mt. Shennongjia 3-OH-FAs is akin to that derived
226 from the LPS component of the outer bacterial membrane of Gram-negative
227 bacteria (Klok et al., 1988). Thus we assume that the 3-OH-FAs measured in
228 the Mt. Shennongjia soils originate from the soil dwelling consortia of Gram-
229 negative bacteria. Furthermore, the suite of 3-OH-FAs compounds detected is
230 similar to that reported from stalagmites (Blyth et al., 2006; Huang et al., 2008;
231 Wang et al., 2012), marine DOM (Wakeham et al., 2003) and lake sediments
232 (Matsuda and Koyama, 1977; Zhang et al., 2014), although the dominant
233 homologue varies between C_{12} , C_{14} to C_{16} in these different sample types, and

234 the relative abundance of each individual compound fluctuates from sample to
235 sample.

236 *3.2 pH impact on 3-OH-FAs and potential proxies*

237 Organic geochemical method development work on acid digestion of
238 speleothem and cave samples from Heshang cave, located ca. 120 km from Mt.
239 Shennongjia in central China (Wang et al., 2012; Huang et al., 2008), revealed
240 that a suite of 3-OH-FAs were readily extractable and relatively abundant
241 compared to established palaeoclimate biomarkers (e.g. plant waxes). This
242 prompted an investigation of the distributions of these compounds along the Mt.
243 Shennongjia altitudinal gradient and the current study of their empirical
244 relationship to environmental parameters. Below we discuss in more detail the
245 most promising 3-OH-FA indices we have identified. In Table 3 in the
246 Supplementary data we include a list of all the 3-OH-FA based indices we
247 tested, including those which showed low or insignificant correlations with
248 environmental parameters (MAAT, soil pH, MAP, soil moisture and altitude).

249 The first group of indices we discuss are those which show relatively high
250 correlations with soil pH. Recent work has demonstrated that pH is a key
251 environmental parameter in controlling soil bacterial community structure and
252 diversity (Bååth and Anderson, 2003; Lauber et al., 2009; Griffiths et al., 2011;
253 Shen et al., 2013; Zhang et al., 2015). In particular, Giotis et al. (2007) found
254 that a strain of Gram-negative bacterium increased/decreased the proportion of
255 branched-chain fatty acids in higher pH/lower pH conditions. Our results from

256 the Mt. Shennongjia transect show that the ratio of the total sum of *iso* and
257 *anteiso* 3-OH-FAs to the total amount of *normal* 3-OH-FAs i.e., the Branching
258 Ratio (equation 1), has a positive correlation with the pH value of soils (Fig. 6a).
259 The Branching Ratio is defined as follows:

$$260 \text{ Branching Ratio} = (I + A)/N \quad (1)$$

261 Where I represents the sum of all the *iso* 3-OH-FAs, A represents the sum
262 of all the *anteiso* 3-OH-FAs, and N represents the sum of all the *normal* 3-OH-
263 FAs.

264 When plotting the Branching Ratio against the pH value of the soils, there
265 is an exponential relationship between the two ($R^2= 0.76$), with the Branching
266 Ratio increasing significantly from 0.31 at pH 4.49 to 0.61 at pH 7.98 (Fig. 6a).
267 Notably, the Branching Ratio shows no obvious correlation with MAAT, MAP
268 or soil humidity (Fig. 7a-c, Supplementary data Table 3).

269 The fact that pH on Mt. Shennongjia does not correlate with other
270 measured parameters (MAAT, MAP, soil humidity) precludes problems of co-
271 variance and gives us confidence that the Branching Ratio does primarily
272 record a signal of environmental pH.

273 Equation (1) and Figure 6a clearly indicate proportionally less branched 3-
274 OH-FAs, including *iso* and *anteiso* isomers, when pH decreases, and thus a
275 lower pH yields a lower Branching Ratio value. This is consistent with the
276 general observation that bacteria can alter the branching and cyclicity of their
277 fatty acid membrane lipids in response to ambient environmental factors

278 (Denich et al., 2003). Branching in fatty acids increases the fluidity (Russell
279 and Fukunaga, 1990) and permeability (McElhaney et al., 1973) of the
280 cytoplasmic membrane.

281 We suggest that the observation of a decreasing Branching Ratio at lower
282 pH reflects chemiosmotic coupling, i.e. the production of fewer branched
283 homologues, producing a less fluid / more impermeable membrane to
284 counteract steeper proton gradients. The existence and maintenance of a
285 proton gradient over bacterial cell membranes is vital for the energy supply of
286 a cell (Mitchell, 1966) and involves the trapping of proton conducting water
287 molecules in the lipid core of the membranes (Nagle and Morowitz, 1978;
288 Wikström et al., 2015). The high significance of the exponential regression
289 supports this hypothesis. The proton gradient over the bacterial cell
290 membranes will be largely determined by ambient proton concentrations and
291 pH is a nonlinear function, being the negative logarithm of ambient proton
292 concentrations. Given the exponential relationship between pH and the
293 Branching Ratio (Fig. 6a) and the definition of pH as the negative logarithm of
294 the proton concentration, it is possible to obtain a linear relationship between
295 the two by defining an alternative index:

$$296 \text{RIAN} = -\log(\text{Branching Ratio}) \quad (2)$$

297 When plotting the ratio of the total sum of *iso* and *anteiso* 3-OH-FAs to the
298 total amount of *normal* 3-OH-FAs (RIAN) against the pH of the soils resulted
299 in the following linear correlation (Fig. 6b):

300 $RIAN = 1.11 - 0.10 \times pH$ ($R^2 = 0.70$, $p < 0.001$) (3)

301 Thus we propose the following novel pH proxy for application to terrestrial
302 palaeoclimatic archives:

303 $pH = 11.10 - 10.00 \times RIAN$ ($R^2 = 0.70$, $p < 0.001$, $RMSE = 0.54$) (4)

304 In addition to Branching Ratio and RIAN, we find that the ratio of
305 summed branched homologues to the sum of all 3-OH-FA homologues
306 (Branched Index) and the ratio of summed *iso* to summed *normal* 3-OH-FA
307 homologues (RIN) also show strong correlations with soil pH ($R^2 = 0.70$ and
308 $R^2 = 0.67$, respectively) (Fig. 6c, d, Supplementary data Table 3). The equations
309 for the Branched Index and RIN are:

310 $Branched\ Index = (I + A) / (I + A + N)$ (5)

311 $RIN = I / N$ (6)

312 Where I represents the sum of all the *iso* 3-OH-FAs, A represents the sum
313 of all the *anteiso* 3-OH-FAs, and N represents the sum of all the *normal* 3-OH-
314 FAs. The possible advantages of these alternative indices are that the
315 Branched Index is bounded at values between 0 and 1 (the Branching ratio and
316 the RIAN are unbounded), whereas RIN only utilises the *normal* and *iso*
317 homologues and does not require measurement of the *anteiso* homologues. RIN
318 may prove to have a practical advantage as the *anteiso* homologues occur in the
319 lowest abundance in our samples (see Figure 5) and may be hard to accurately
320 integrate in some environmental samples where the overall abundance or
321 preservation of 3-OH-FAs is lower.

322 All the ratios and indices presented show positive or negative correlations
323 ($R^2= 0.67$ to 0.76 , $p<0.001$) with pH (Fig. 6) but show no obvious correlation
324 with MAAT, MAP or soil humidity (Fig. 7 and Supplementary data Table 3).
325 All the ratios and indices appear to be independent measures of the
326 decreased/increased degree of branching of 3-OH-FAs with lower/higher pH.
327 As discussed above, for the Branching Ratio, this suggests a causal relationship
328 with soil pH which we argue reflects chemiosmotic coupling, i.e. the production
329 of fewer or more branched homologues to control membrane
330 fluidity/permeability in response to proton gradients across bacterial cell
331 membranes. This is comparable with the suggestion of Weijers et al. (2007b)
332 that a lower/higher degree of methylation of branched GDGTs in lower/higher
333 pH conditions reflects chemiosmotic coupling and is consistent with the finding
334 of Bardy et al. (2009) that the contribution of branched C₁₅ and C₁₇ alkanolic
335 acids relative to their linear homologues decreased with pH in a podzolic
336 sequence in the Amazon basin.

337 Based on the linear correlations showed in Fig. 6c, d, we obtain the
338 following equations with pH for the Branched Index and RIN:

339 Branched Index = $-0.03 + 0.05 \times \text{pH}$ (7)

340 RIN = $-0.21 + 0.08 \times \text{pH}$ (8)

341 Thus we propose the additional novel pH proxies for application to
342 terrestrial palaeoclimatic archives:

343 $\text{pH} = 0.60 + 20.00 \times \text{Branched Index} (R^2= 0.70, p<0.001, \text{RMSE}= 0.54) \quad (9)$

344 $\text{pH} = 2.63 + 12.50 \times \text{RIN} (R^2= 0.67, p<0.001, \text{RMSE}= 0.56) \quad (10)$

345 At this early stage of development of 3-OH-FA based proxies for
346 palaeoenvironmental applications, we recommend that the RAN, Branched
347 Index and RIN should all be measured in samples, as all of them clearly have
348 potential as pH proxies and only further work can constrain which may be
349 most reliable or practicable.

350

351 *3.3 Temperature impact on 3-OH-FAs and potential proxies*

352 In addition to the novel pH proxies described above, we found two indices
353 that have potential as novel temperature proxies, the ratio of *anteiso* to *normal*
354 C_{15} 3-OH-FA (RAN_{15}) and the ratio of *anteiso* to *normal* C_{17} 3-OH-FA (RAN_{17}).
355 RAN_{15} and RAN_{17} are defined as follows:

356 $\text{RAN}_{15} = \alpha\text{-}C_{15} / n\text{-}C_{15} \text{ 3-OH-FA} \quad (11)$

357 $\text{RAN}_{17} = \alpha\text{-}C_{17} / n\text{-}C_{17} \text{ 3-OH-FA} \quad (12)$

358 RAN_{15} shows a linear relationship with MAAT and MAP ($R^2 = 0.51$ and
359 0.50 , respectively) (Fig. 8a, b). A similar result was also found in RAN_{17} ($R^2 =$
360 0.48 and 0.48 , respectively) (Fig. 8c, d). It is not surprising that both MAAT
361 and MAP show a linear relationship with RAN_{15} and RAN_{17} , because both
362 parameters strongly co-vary with elevation on Mt. Shennongjia. It has been
363 suggested that precipitation could be an important environmental control on
364 soil bacterial lipids in semi-arid to arid regions. Although initially proposed as

365 being a function of MAAT and pH, recent work has highlighted that the GDGT
366 based MBT/CBT-MAAT index is significantly influenced by precipitation/ soil
367 moisture in the semi-arid western USA, where MAP is below 700-800 mm yr⁻¹
368 (Dirghangi et al., 2013), in the semi-arid Iberian Peninsula (Menges et al., 2014)
369 and in China (Yang et al., 2014). Yang et al. (2014) found complexities in the
370 relationship of the MBT and CBT indices to MAAT in alkaline and arid soils in
371 China, in contrast to their positive correlation in more acidic soils in the
372 complete Chinese, or global, datasets. Our research area is characterised by
373 relatively acidic to neutral soils (pH 4.5 - 8.0), and a moist-humid climate,
374 where MAP is above 1000 mm yr⁻¹, even on the drier, lower slopes of the
375 mountain. Therefore, we suggest precipitation/soil moisture is unlikely to be an
376 ecologically limiting factor that significantly affects the distribution of the
377 membrane lipids. In support of this assumption we found that both RAN₁₅ and
378 RAN₁₇ showed very weak correlations with soil humidity measurements (R^2 =
379 0.19 and 0.16, respectively, see Supplementary data Table 3), although we note
380 that such measurements only represent the conditions at the time of sampling
381 and not necessarily the average, mean annual conditions. Furthermore, RAN₁₅
382 and RAN₁₇ show significant correlations with the GDGT-based MBT/CBT-
383 MAAT proxy published by Yang et al. (2015) on the same soil samples (R^2 =
384 0.61 and 0.36, respectively) (Fig. 9a, b). Thus we assume that MAAT is the
385 dominant parameter that affects these ratios even though the impact of MAP
386 could not be entirely excluded. The ratios of both RAN₁₅ and RAN₁₇ increase

387 with decreasing environmental temperature (Fig. 8a, c). It has been observed
 388 that *anteiso* fatty acids have a lower melting point than *normal* fatty acids
 389 (Kaneda, 1991; Suutari and Laakso, 1994). Thus in order to maintain
 390 membrane fluidity, bacteria may increase the proportion of *anteiso* 3-OH-FAs
 391 (increasing the RAN indices) with decreasing temperature. This hypothesis is
 392 supported by the fact that we found a significant relationship between ratio of
 393 *anteiso* to *normal* C₁₅ 3-OH-FA and temperature, but a much less significant
 394 relationship between *iso* to *normal* C₁₅ 3-OH-FA (see Supplementary data
 395 Table 3). *Anteiso*-branched fatty acids have greater fluidizing properties and
 396 disturb packing order to a greater extent than *iso*-branched fatty acids (Russell,
 397 1995). This is conferred by the *anteiso*-methyl branch being located on the third
 398 carbon from the methyl terminus while the *iso*-methyl branch is positioned on
 399 the second carbon from the end of the chain (Russell, 1984).

400 Based on the linear correlation showed in Fig. 8, we obtain the following
 401 equations:

$$402 \text{ RAN}_{15} = 7.60 - 0.33 \times \text{MAAT} \quad (13)$$

$$403 \text{ MAAT} = 23.03 - 3.03 \times \text{RAN}_{15} \text{ (R}^2 = 0.51, \text{ p} < 0.001, \text{ RMSE} = 2.6 \text{ }^\circ\text{C)} \quad (14)$$

$$404 \text{ RAN}_{17} = 2.90 - 0.11 \times \text{MAAT} \quad (15)$$

$$405 \text{ MAAT} = 26.36 - 9.09 \times \text{RAN}_{17} \text{ (R}^2 = 0.48, \text{ p} < 0.001, \text{ RMSE} = 2.7 \text{ }^\circ\text{C)} \quad (16)$$

406 The relationships of both RAN₁₅ and RAN₁₇ (equations 13 and equation 15)
 407 to MAAT are similar (see Fig. 8), although RAN₁₇ has somewhat more scatter.

408 GDGT data have been previously published from 19 of our 26 soil samples
409 (Yang et al., 2015). Thus, we can directly compare our 3-OH-FA based proxies
410 with established GDGT based proxies (CBT and MBT/CBT). Our RIAN and
411 Branched Index proxies for pH show high linear correlation with the GDGT-
412 based CBT (Fig. 9c, d) suggesting all three proxies have the same dominant
413 control, namely pH. Furthermore RAN₁₅ and RAN₁₇ based on 3-OH-FA show a
414 linear correlation with the GDGT-based MBT/CBT-MAAT proxy (Fig. 9a, b)
415 although this is significantly higher for RAN₁₅. It is important to note that,
416 unlike the current MBT/CBT-MAAT proxy, our proposed 3-OH-FA derived
417 temperature proxies are independent from pH.

418 In addition to the ratios, indices and proposed novel proxies presented
419 above we explored a full range of 3-OH-FA distributions (e.g. Average Chain
420 Length of 3-OH-FAs) versus environmental parameters in the samples
421 obtained from Mt. Shennongjia. Above we present only the most significant
422 correlations and findings, but include all results in the Supplementary Data,
423 Table 3.

424 **4. Wide occurrence of 3-OH-FAs in other settings**

425 We undertook an initial investigation to confirm the preservation of 3-OH-
426 FAs on Quaternary time scales in several palaeoclimatic archives: a lake
427 sediment sample dated to 1984±30 yr B.P. from Tianchi Lake, Gansu province,
428 China, a speleothem sample dated to 8645±78 yr B.P. from Heshang Cave,
429 China and a last glacial marine sediment sample from the 81 mbsf from IODP

430 Site M0060, Baltic Sea. The distribution of 3-OH-FAs varied between samples,
431 but the suite of C₁₀ to C₁₈ *normal*, plus certain *iso*- and *anteiso*- 3-OH-FAs
432 homologues, were all present in measurable concentrations (Fig. 10). Notably,
433 monounsaturated 3-OH-FAs with even carbon numbers (C₁₂, C₁₄, C₁₆, C₁₈) were
434 uniquely found in the Tianchi Lake sediment, suggesting either: a) a unique
435 source of 3-OH-FAs in that lake environment or; b) greater preservation of the
436 more labile unsaturated homologues (Fig. 10, Supplementary data Table 4).

437 The variations in the 3-OH-FA signatures between the different settings
438 are likely due to controls by environmental and climatic parameters on
439 membrane lipid production by bacteria (as suggested for the altitudinal
440 transect of modern soils in this paper). Moreover, the origin and preservational
441 pathways of 3-OH-FAs in some settings could be complex. For example, 3-OH-
442 FAs in lake sediments may be produced *in situ* and/or may be derived from the
443 surrounding soils, this may complicate the application of 3-OH-FAs as
444 temperature/pH proxies in lakes. In general, we can not discount the influence
445 on the 3-OH-FA signatures of unknown, site-specific, factors related to the
446 differences in depositional setting or variations in populations of the Gram-
447 negative bacterial producer. Thus specific calibrations are likely required for
448 applications to a diverse range of palaeoclimatic archives. However, the
449 preservation of the same suite of 3-OH-FAs in such different depositional
450 environments, hints at a potentially wide applicability of these microbial
451 proxies in a variety of environmental settings.

452 **5. Conclusion**

453 In summary, 3-OH-FAs in surface soils collected from an altitudinal
454 transect on Mt. Shennongjia were examined to explore their relationships with
455 environmental parameters. The RIAN, Branched index and RIN indices are
456 highly correlated with soil pH. Furthermore, the RAN₁₅ and RAN₁₇ ratios
457 exhibit significant correlations with MAAT and MAP. As precipitation is not
458 likely to be an ecologically limiting factor in the moist-humid environment of
459 Mt. Shennongjia we assume that MAAT is the dominant control. Notably, the
460 3-OH-FA based temperature proxies RAN₁₅ and RAN₁₇, are not pH dependent,
461 which should be an advantage in environments where pH is highly variable
462 and could be a confounding variable. Our discovery of new independent proxies
463 for pH and MAAT from an altitudinal transect of surface soils from Mt.
464 Shennongjia has potentially wide implications for palaeoclimatic and
465 environmental studies. 3-OH-FA proxies could be used in a variety of
466 environmental settings (See Fig. 10). Multi-proxy terrestrial reconstructions of
467 pH and temperature could be established by comparing 3-OH-FAs with GDGT
468 based proxies. Gram-negative bacteria have a wide distribution in natural
469 environment (Gupta, 1998), and 3-OH-FAs have been identified in diverse
470 environments, including marine and terrestrial settings and even in
471 atmospheric aerosols (Wakeham et al., 2003; Lee et al., 2004; Huang et al.,
472 2008). In particular, these compounds are easy to identify and precisely
473 quantify using GC-MS and GC-FID systems. This makes it possible to utilize a

474 small amount of sample weight and to gain high-resolution palaeo-records, for
475 example even from stalagmite archives (Blyth et al., 2006; Huang et al., 2008;
476 Wang et al., 2012). Additionally, measurement of 3-OH-FAs requires only
477 standard GC-MS and GC-FID systems and can be readily adopted by most
478 organic geochemistry laboratories (without the need for investment in
479 additional, expensive equipment). It is clear that 3-OH-FAs have hitherto
480 unrealized potential as palaeoclimate proxies. We hope this paper opens up
481 new avenues of research on 3-OH-FAs, including culture studies, empirical
482 calibrations (both global and regional) and application to an array of
483 palaeoclimatic archives (e.g. lakes, speleothems, marine records).

484 **Acknowledgements**

485 This work was supported by the Natural Science Foundation of China (grant
486 No. 41330103, 41130207, 41201203), and the 111 project (grant No. B08030).
487 We thank the China Scholarship Council (CSC) (File NO. 201306410031) for
488 supporting Canfa Wang's study at the University of Birmingham. We thank
489 the University of Birmingham Dynamic Investment Fund (DIF) for supporting
490 the establishment of the new Birmingham Molecular Climatology laboratory.
491 Two anonymous reviewers and the associate editor Dr. Klaas G.J. Nierop are
492 thanked for their constructive comment which helped to improve the
493 manuscript.

494

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729

730 **Figure captions**

731

732 **Fig. 1** General structure of lipopolysaccharide (LPS) from Gram-negative
733 bacteria (Alexander and Rietschel, 2001). LPS is characterized by three main
734 units: the O-polysaccharides chains, the core oligosaccharide and lipid A. The
735 repeating subunits of the O-polysaccharides are composed of between one and
736 eight glycosyl residues and differ between strains by virtue of differing sugars,
737 sequence, chemical linkage, substitution and the ring forms utilised. The outer
738 core is inclined to contain common sugars such as hexoses or hexosamines etc.
739 The inner core contains the unusual sugars 3-deoxy-D-manno-octulosonic acid
740 (Kdo) and D-glycero-D-manno-heptose (Hep) (Erridge et al., 2002). Lipid A, the
741 innermost part of LPS, consists of two glucosamine (GlcN) moieties, with
742 attached acyl chains ("fatty acids") by either amide bonds or ester bonds, and
743 normally contains one phosphate group on each GlcN (Raetz et al., 2009).

744

745 **Fig. 2** Regional map, illustrating the location of Shennongjia Mountain,
746 Heshang Cave and Tianchi Lake.

747

748 **Fig. 3** Cross plots showing the relationship of soil pH in samples from Mt.
749 Shennongjia with soil humidity, Mean Annual Air Temperature (MAAT), Mean
750 Annual Precipitation (MAP) and altitude.

751

752 **Fig. 4** Mass spectrum of the C₁₆ 3-OH-FA TMSi ester. The m/z 175 fragment is
753 due to the cleavage between C₃ and C₄, and the [M-15] base peak results from a
754 loss of a CH₃ group.

755

756 **Fig. 5** Extracted ion chromatograph (m/z 175) showing the composition and
757 distribution of 3-OH-FAs in the Mt. Shenongjia soil sample collected at 832
758 m.a.s.l. (see sample SNJ 11-4 in the Supplementary data Table 1 for more
759 detailed information). Red circles represent the *normal* 3-OH-FAs, yellow
760 squares represent the *iso* 3-OH-FAs, grey triangles represent the *anteiso* 3-OH-
761 FAs. The carbon numbers range from C₁₀ to C₁₈, including *iso* C₁₁, C₁₃, C₁₄, C₁₅,
762 C₁₆ and *anteiso* C₁₁, C₁₃ C₁₅ C₁₇.

763

764 **Fig. 6** The relationship between 3-OH-FAs indices and pH. (a) Exponential
765 correlation between the Branching Ratio and pH (R²= 0.76, p<0.001). (b) Linear
766 correlation between RIAN and soil pH (R²=0.70, p<0.001). (c) Linear correlation
767 between Branched Index and pH (R²= 0.70, p<0.001). (d) Linear correlation
768 between RIN and pH (R²= 0.67, p<0.001).

769

770 **Fig. 7** Cross plots showing the relationship between Branching Ratio and
771 Branched Index to environmental parameters (MAT, MAP, and soil humidity).

772

773 **Fig. 8** The relationship between 3-OH-FA ratios and environmental factors. (a)
774 The RAN₁₅ shows negative linear relationship with MAAT ($R^2= 0.51$, $p<0.001$)
775 and (b) positive linear relationship with MAP ($R^2= 0.50$, $p<0.001$). (c) The
776 RAN₁₇ shows negative linear relationship with MAAT ($R^2= 0.48$, $p<0.001$) and
777 (d) positive linear relationship with MAP ($R^2= 0.48$, $p<0.001$).

778

779 **Fig. 9** Cross plots showing the correlation between certain 3-OH-FA based and
780 GDGT based proxies.

781

782 **Fig. 10** Extracted ion chromatogram (m/z 175) showing the distribution of 3-
783 OH-FAs in contrasting geological samples. Red circles represent the *normal* 3-
784 OH-FAs, yellow squares represent the *iso* 3-OH-FAs, grey triangles represent
785 the *anteiso* 3-OH-FAs and white circles represent the monounsaturated 3-OH-
786 FAs. (a) The composition and distribution of 3-OH-FAs in a sediment sample
787 from Tianchi Lake. (b) The distribution of 3-OH-FAs in a Heshang Cave
788 stalagmite sample. (c) The distribution of 3-OH-FAs in Baltic Sea sediment
789 sample.

790