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Control of soil pH on turnover of belowground organic matter in subalpine grassland

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Abstract: Grasslands store substantial amounts of carbon in the form of organic matter in soil and roots. At high latitudes and elevation, turnover of these materials is slow due to various interacting biotic and abiotic constraints. Reliable estimates on the future of belowground carbon storage in cold grassland soils thus require quantitative understanding of these factors. We studied carbon turnover of roots, labile coarse particulate organic matter (cPOM) and older non-cPOM along a natural pH gradient (3.9–5.9) in a subalpine grassland by utilizing soil fractionation and radiocarbon dating. Soil carbon stocks and root biomass, turnover, and decomposability did not scale with soil pH whereas mean residence times of both soil organic matter fractions significantly increased with declining pH. The effect was twice as strong for non-cPOM, which was also stronger enriched in ^{15}N at low pH. Considering roots as important precursors for cPOM, the weaker soil pH effect on cPOM turnover may have been driven by comparably high root pH values. At pH 5, long non-cPOM mean residence times were probably related to pH dependent changes in substrate availability. Differences in turnover along the pH gradient were not reflected in soil carbon stocks because aboveground productivity was lower under acidic conditions and, in turn, higher inputs from aboveground plant residues compensated for faster soil carbon turnover at less acidic pH. In summary, the study provides evidence for a strong and differential regulatory role of pH on the turnover of soil organic matter that needs consideration in studies aiming to quantify effects of changing environmental conditions on belowground carbon storage.

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1 **Control of soil pH on turnover of belowground organic matter in subalpine grassland**

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18

19 **Abstract**

20 Grasslands store substantial amounts of carbon in the form of organic matter in soil and roots.
21 At high latitudes and elevation, turnover of these materials is slow due to various interacting
22 biotic and abiotic constraints. Reliable estimates on the future of belowground carbon storage
23 in cold grassland soils thus require quantitative understanding of these factors. We studied
24 carbon turnover of roots, labile coarse particulate organic matter (cPOM) and older non-
25 cPOM along a natural pH gradient (3.9 – 5.9) in a subalpine grassland by utilizing soil frac-
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28 ter fractions significantly increased with declining pH. The effect was twice as strong for
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31 by comparably high root pH values. At $\text{pH} < 5$, long non-cPOM mean residence times were
32 probably related to pH dependent changes in substrate availability. Differences in turnover
33 along the pH gradient were not reflected in soil carbon stocks because aboveground produc-
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35 residues compensated for faster soil carbon turnover at less acidic pH. In summary, the study
36 provides evidence for a strong and differential regulatory role of pH on the turnover of soil
37 organic matter that needs consideration in studies aiming to quantify effects of changing en-
38 vironmental conditions on belowground carbon storage.

39

40

41 **Introduction**

42 The turnover of soil organic matter in grassland soils of cold climates is strongly limited by
43 temperature (Anderson 1991; Townsend et al. 1995). Climatic conditions also exert control
44 on vegetation communities, thereby altering amount and quality of the incoming plant resi-
45 dues (Hobbie et al. 2000). Both factors may induce accumulation of litter-like material, such
46 as particulate organic matter (POM) in soil (Leifeld et al. 2009), that is potentially sensitive
47 to rapid microbial oxidation once environmental conditions change. This has prompted con-
48 cern about the state and vulnerability of organic matter in cold climates such as, for example,
49 that in European mountain soils (Sjögersten et al. 2011).

50 However, there are various partially interacting mechanisms involved in controlling micro-
51 bial kinetics, and thus residue turnover in cold grassland soils is not influenced by tempera-
52 ture and litter quality alone. One often overlooked quantitative environmental factor, influ-
53 encing both vegetation and soil processes, is soil pH. Low pH values are common in areas
54 such as low-productive mountainous grasslands on silicate rocks, particularly when high rain-
55 fall and heather vegetation fosters podzolization (Bouma et al. 1969; Egli et al. 2003). Soil
56 pH is involved in many states and processes, such as enzyme activities (Sinsabaugh et al.
57 2008), dissolved organic carbon (DOC) and N availability (e.g. Kalbitz et al. 2000; Pietri and
58 Brookes 2008), and litter decomposition, as ascribed to the combinatory effect of enzyme
59 activities, decomposer community and Al^{3+} toxicity (Walse et al. 1998). Soil acidity modu-
60 lates alpine microbial (Eskelinen et al. 2009) and plant communities (Budge et al. 2011,
61 Körner 2003, Nilsson et al. 2002), and it affected compositions of both vegetation and soil
62 microbial biomass, even 70 years after cessation of a liming experiment in subalpine grass-
63 land (Spiegelberger et al. 2006).

64 The regulatory role of soil pH on decomposition rates is considered to be quite strong, with
65 rates varying by a factor of four in the pH range from 4.0 to 6.0 (Walse et al. 1998, Leifeld et
66 al. 2008). Much of the knowledge on the regulatory role of pH is based on in vitro activities
67 or controlled decomposition studies. Leifeld et al. (2008) used the soil radiocarbon signature
68 along a climate and pH gradient for the quantification of the pH effect on POM turnover after
69 correction for temperature. Hitherto, however, there is no study that explicitly addresses the
70 role of soil pH on the turnover time of belowground organic matter components, such as
71 POM or non-POM fractions in an otherwise homogeneous field situation under long-term
72 steady-state conditions. In addition, the role of pH on turnover of roots, the most important
73 source for soil carbon (Rasse et al. 2005), is not yet established. Given the long turnover
74 times of organic matter under cold and acid conditions of many decades, even for so-called
75 labile fractions (Leifeld et al. 2009), the issue evades being addressed by short-term con-
76 trolled experiments but can be approached by using natural pH gradients that existed for long
77 periods of time.

78 Here we study the role of soil pH as a modifier for soil and root carbon turnover in a steady-
79 state environment of a subalpine permanent grassland. We hypothesise that more acidic soils
80 should increase the mean residence times of belowground carbon and, subsequently, also
81 affect its distribution between labile and stable soil fractions.

82

83 **Material and Methods**

84 *Sites and sampling*

85 Alp Flix (46°30'60"N, 9°39'56"E) is located in the canton of Grisons, Switzerland, at an
86 elevation of ca. 2100 m asl. Mean annual temperature is +2.2°C, and mean annual rainfall
87 1050 mm. The site is located on a gentle slope and used as a permanent cattle pasture grazed

88 in the summer season from June to September. Soils are well-drained Leptosols and leptic
89 Cambisols developed on granite-diorite rock with loamy to clayey-loamy texture. The vegeta-
90 tion is typically an acidic Geo-Montani-Nardetum pasture, comprising patches with baso-
91 philic plant species typical for a calcareous Seslerio-Caricetum sempervirentis community.
92 Soil pH effect on vegetation community is visible, for example, at the species level, as in-
93 creasing abundance of basophilic *Plantago atrata* HOPPE, *Carex ornithopoda* WILD and *An-*
94 *thyllis vulneraria* L. with pH while abundance of pointers of acidity such as *Nardus stricta*
95 L., *Arnica montana* L. or *Gentiana acaulis* L. significantly decreases with pH (S. Bassin,
96 pers. communication).

97 The centre of the unfenced pasture extends over an area of ca. 1 ha. A soil survey in 2003
98 with 180 subplots across the field revealed a high variability in soil pH, ranging from 3.9 to
99 5.9 due to the irregular presence of limestone gravel originating from the adjacent calcareous
100 mountain tops. For further study we selected eight of these sites that span the whole pH range
101 but share similar soil texture.

102 In previous work at the same pasture, a 30 x 40 cm wide, 20 cm deep soil monolith was ex-
103 cavated in 2003 from each of the eight locations. Afterwards pits were filled with vegetated
104 soil monoliths from an adjacent site. Above-ground biomass yield of extracted monoliths was
105 measured in 2004 after one growing season for the purpose of a different study (Bassin et al.
106 2007). Our aboveground biomass data were taken from that study. For the present work, we
107 made use of the corresponding archived soil samples from each of these eight locations taken
108 in 2003 and took additional samples from the same sites in 2009. In 2003, soil samples (0-10
109 cm) had been taken from the four outer walls of the excavated soil monolith. These soil sam-
110 ples were used for measuring nutrients, texture, pH, for soil fractionation and for analyzing
111 ¹⁴C in soil fractions. In 2009, the site was re-visited to take 4 cores (diameter 6 cm, depth 10
112 cm) at distance of 0.5 m from each pit wall where monoliths have been excavated in 2003.

113 From these cores, soil bulk density and stone content were calculated and fine soil pH, C and
114 N concentrations were measured. In addition, pH and decomposability of biomass and ^{14}C
115 content of roots were measured. All concentrations are based on 105 °C oven-dried samples.

116

117 *Soil fractionation and chemical analysis*

118 Soils were oven-dried, sieved < 2 mm and analyzed for total C, N, texture (pipette method
119 after H_2O_2 treatment) and pH (2:1 in water). All carbon and nitrogen contents were measured
120 after combustion with an elemental analyzer (Hekatech Euro EA 3000, Wegberg, Germany).
121 Soils were free of carbonate. Extractable nutrients (P, K, Mg, Ca) were measured after treat-
122 ment with 1:10 NH_4 -acetate solution (FAL 1998).

123 The aim of soil fractionation was to retrieve two fractions that differ systematically in their
124 isotopic signature to allow meaningful calculation of SOC turnover times (see below). It is
125 based on previous studies with similar objectives where this approach proved reliable (Budge
126 et al. 2011; Conen et al. 2008; Leifeld et al. 2009). A light particulate organic matter fraction
127 > 63 μm was obtained after ultrasonic dispersion of the soil < 2 mm by applying an energy of
128 22 J ml^{-1} suspension, followed by density separation with Na-polytungstate ($d = 1.8 \text{ g cm}^{-3}$)
129 in a centrifuge. The light material was decanted and poured into a 63 μm nylon mesh bag.
130 After decantation the sediment in each test tube was stirred, centrifuged again and super-
131 natants were combined and rinsed with distilled water to remove the salt. We refer to this
132 material as coarse particulate organic matter (cPOM). The remaining material was not meas-
133 ured for its elemental contents and isotopic signature but these parameters were calculated
134 based on the respective values of the cPOM and the bulk soil considering mass conservation
135 (see eq. 1). This was done because in any fractionation some material is lost, for example in
136 solution, which may compromise the radiocarbon-derived turnover calculation. The remain-

137 ing material consists of some fine POM, mineral-associated and soluble organic matter. We
138 refer to the calculated differential fraction as non-cPOM.

139 Material < 63 μm was oxidized in a solution of 1 M NaOCl (Roth AG, Reinach), adjusted to
140 pH 8.0, at a soil-to-solution ratio of 1:50 (w/v) and agitation for six hours at 25°C. It corre-
141 sponded to 6.5% active Cl_2 (determined by iodometry). After, the samples were centrifuged
142 at 2000 g for 30 minutes and the supernatants removed. This procedure was repeated three
143 times. After the last treatment, the centrifugation pellets were intensively washed with deion-
144 ized H_2O to remove salts, and air-dried.

145 The specific surface area of material < 63 μm , before and after oxidation with NaOCl, was
146 determined by N_2 adsorption using a Quantachrome Nova 2200 surface analyzer and BET
147 isotherm.

148 Prior to X-ray diffraction analysis (XRD) and X-ray fluorescence spectrometry (XRF), the
149 samples were milled. Total elemental content of Fe, Mg, Al, Si, and P in the oxidized < 63
150 μm fraction was measured using wavelength dispersive XRF (ARL Optim'X, Thermo Elec-
151 tron Corp., Switzerland).

152 The mineralogical composition of the NaOCl oxidized fraction < 63 μm was determined us-
153 ing XRD on randomly oriented specimens. X-ray measurements were made using a Bragg-
154 Brentano diffractometer (BRUKER AXS D8, $\text{CuK}\alpha$ with theta compensating slits and graph-
155 ite monochromator) in the range of $2 \theta \approx 80^\circ$ with a step width of 0.02° and a counting
156 time of 2 s. A special frontloading preparation was carried out to hold the preferred orienta-
157 tion as low as possible in randomly oriented specimens (Kleeberg et al. 2008). In the range 2
158 $\theta \approx 15^\circ$ the measurements were carried with and without ethylene glycol solvation. The inter-
159 calation of ethylene glycol causes a typical shift of the (001) reflexes in the XRD pattern of
160 expandable clay minerals. The qualitative phase composition was determined on the basis of

161 the peak position and the relative intensities of the mineral phases were identified in compari-
162 son to the ICDD data base. The analysis was carried out with the software package DIF-
163 FRACplus (Bruker AXS). The quantitative mineralogical composition was estimated using
164 the peak heights of the XRD patterns of randomly oriented specimens.

165 Stable N isotope ratios were measured on cPOM and bulk soils by ion ratio mass spectrome-
166 try (Thermo Finnigan Delta plus XP coupled with an elemental analyzer Flash EA 1112 Se-
167 ries; accuracy 0.2 permil). The $\delta^{15}\text{N}$ of non-cPOM was calculated by mass balance as

$$168 \quad \delta^{15}\text{N}_{\text{non-cPOM}} = (\delta^{15}\text{N}_{\text{SOM}} - (\delta^{15}\text{N}_{\text{cPOM}} * s_{\text{cPOM}})) / (1 - s_{\text{cPOM}}) \quad [1]$$

169 where s_{cPOM} is the mass ratio of nitrogen bound in cPOM to nitrogen bound in SOM.

170 Bulk soil samples, cPOM, and roots that contained 0.5 to 1 mg of C were combusted and
171 graphitised for AMS measurements of radiocarbon content. These were measured at the AMS
172 facility of Laboratory of Ion Beam Physics of the Institute for Particle Physics of at the ETH
173 (the Swiss Federal Institute of Technology), Zurich (Synal 2007). The results were expressed
174 as percent Modern Carbon (pMC) calculated following the protocol of Stuiver and Polach
175 (1977). Radiocarbon content of non-cPOM was calculated by carbon mass balance in corre-
176 spondence to [1].

177

178 *¹⁴C-derived mean residence times of soil fractions, bulk soil, and roots*

179 A radiocarbon bomb model based on Harkness et al. (1986), but adjusted for time-lag effects,
180 was applied for calculating carbon mean residence times (MRTs) separately for roots, cPOM,
181 and non-cPOM. In the model, the ¹⁴C activity of the carbon can be expressed as

$$182 \quad A_t = A_{(t-1)} e^{-k} + (1 - e^{-k}) A_{(i-TL)} - A_{(t-1)} \lambda \quad [2]$$

183 where $A_{(t)}$ is the (measured) ^{14}C activity (pMC) of C in any fraction at time t , $A_{(t-1)}$ the ^{14}C
184 activity of the previous year; $A_{(i)}$ is the atmospheric ^{14}C activity corrected for the time-lag
185 (TL) between photosynthetic fixation and plant residue input into the soil pool, k the ex-
186 change rate constant of the respective C pool, and λ the ^{14}C decay constant ($1/8268 \text{ a}^{-1}$). Val-
187 ues for A_i were taken from the atmospheric ^{14}C record of Stuiver et al. (1998) for the period
188 from year 1511 to 1954 and from Levin and Kromer (2004) for the period 1959 to 2004. The
189 period between 1954 and 1959 was linearly interpolated. Data for 2004-2009 were provided
190 by I. Levin (pers. communication).

191 Carbon mean residence times were calculated according to [2] by iteratively varying the
192 MRT until it matched the measured ^{14}C activity of the sample. This was done separately for
193 each of the three fractions (roots, cPOM, non-cPOM). Root MRT was directly derived from
194 the bomb model. Mean root MRTs over the eight sites and their respective confidence inter-
195 val (CI) were taken as time-lag of the carbon entering the soil when calculating MRT of
196 cPOM and non-cPOM. Another source of error considered in the estimate is the analytical
197 error of the AMS measurement given as one sigma of the pMC value. MRTs of carbon
198 cPOM and non-cPOM were calculated for any possible combination of means and CI (time-
199 lag) or σ (pMC) (i.e., $n=9$) to derive a more robust error estimate of carbon turnover, shown
200 as mean (\pm) one standard error.

201 Following Leifeld & Fuhrer (2009) and Torn et al. (2009), the flux F of carbon [$\text{t C ha}^{-1} \text{ a}^{-1}$]
202 through a fraction f (i.e., roots, cPOM, non-cPOM) under steady-state conditions equals the
203 input and was calculated as

$$204 \quad F_f = 1 / MRT_{fraction} \bullet poolsize_{fraction} \quad [3]$$

205 with $poolsize_{fraction}$ in [t C ha^{-1}], yielding the total flux F_t through the whole soil as the sum of
206 the single fluxes i

207 $F_t = \sum F_{fi}$ [4]

208 and the corresponding MRT [a] for SOC 0-10 cm

209 $MRT_{SOC} = SOC_{0-20} / F_t$ [5]

210 The flux through the root biomass was calculated according to [3, 4] but not added to the flux
211 through the whole soil. Mathematically, calculation of bulk soil MRTs based on bulk soil
212 radiocarbon content is possible with the same formula but implicitly assumes kinetics in ac-
213 cordance with a single-pool soil carbon turnover model. Such an assumption violates the evi-
214 dence of fractions being transformed at different rates and may overestimate soil carbon turn-
215 over times (e.g., Trumbore et al. 1997). Therefore, MRTs using two or more fractions of dif-
216 ferent MRT provide a better approximation of bulk carbon soil dynamics (Leifeld and Fuhrer
217 2009; Budge et al. 2011).

218

219 *Roots*

220 The four replicates from each of the eight sites sampled in 2009 were weighed, thoroughly
221 sieved field-moist over a 2 mm mesh and coarse roots were separated. Finer roots in material
222 passing the sieve was hand-picked using a pair of tweezers and combined with the first batch.
223 All roots were carefully washed, freeze-dried, and chopped. Replicates were analyzed sepa-
224 rately for C, N, pH (2:1 in water) as above but were bulked for ^{14}C analysis. For decomposi-
225 tion experiments, 0.8 g freeze-dried chopped roots (n = 32) were mixed with 19.2 g purified
226 quartz sand and inoculated with 4.2 ml solution inoculum (5.9 mg DOC l^{-1} ; extracted from
227 fresh soil from the same site) to reach 60% maximum water holding capacity. Six blanks
228 without roots were run in parallel. Samples were incubated in closed jars containing NaOH at
229 20 °C for 3 weeks and respiration was measured every 3 days by back-titration of the remain-
230 ing alkalinity.

231

232 *Statistics*

233 Errors of replicates are given as standard error of the mean (SE). Correlation between pa-
234 rameters was tested using Pearson's correlation coefficient and significant relationships are
235 marked with the respective error probability p. Quantitative effects of pH on carbon mean
236 residence times were studied by ordinary least squares regression and the coefficient of de-
237 termination. Regression statistics includes standard errors of regression coefficients and con-
238 fidence intervals of the regression line. A t-test was applied to test differences in carbon MRT
239 and inputs between groups of different soil acidity. All statistics was calculated using Statis-
240 tica 9.1, StatSoft Inc., USA.

241

242 **Results**

243 *Soil properties and soil pH relationship to organic matter and vegetation*

244 All samples were of similar texture (see Table 1) and mineralogical composition. The X-ray
245 diffractograms of the samples were almost identical (data not shown). XRD analysis revealed
246 that the $< 63 \mu\text{m}$ fraction was dominated by quartz, plagioclase, K-feldspar, mica, chlorite
247 and actinolite. Minor phases were epidote, rutile, titanite and mixed-layered clay minerals.
248 Major phyllosilicate phases were mica (biotite, muscovite, illite), chlorite, subordinate hydro-
249 biotite (regularly interstratified mica-vermiculite), and interstratified mica-smectite. Some
250 vermiculite was also present. HIV (hydroxy-interlayered vermiculite) and HIS (hydroxy-
251 interlayered smectite) could not be distinguished individually. All samples contained some
252 oxyhydroxides. Among them, lepidocrocite and traces of gibbsite could be identified.

253 The specific surface area of the fraction $< 0.63 \mu\text{m}$ averaged $9.8 (\pm 0.5) \text{ m}^2 \text{ g}^{-1}$ (NaOCl) and
254 was not related to pH either before or after oxidative treatment. From total element contents

255 (XRF) only total Mg significantly correlated with pH ($r = 0.86$, $p < 0.01$) (see supplementary
256 material). Extractable Ca and Mg was highly positively correlated with soil pH ($r = 0.97$ and
257 0.91 , $p < 0.001$ and $p < 0.01$, respectively) (see supplementary material).

258 Most organic matter characteristics of vegetation and soil were not related to soil pH (Table
259 1). Soil pH affected neither the total amount of SOC or roots, nor the composition, in terms of
260 distribution among soil fractions or C/N ratios (soil or roots). In addition, root degradability,
261 as measured in the incubation experiment, and root mean residence times did not scale with
262 pH. Root pH, however, significantly increased with increasing soil pH but was offset by al-
263 most two pH units. Aboveground biomass highly significantly increased with increasing pH
264 and vegetation composition also responded to pH with an increase in the fraction of forbs ($r =$
265 0.71 , $p < 0.05$), whereas the fraction of sedges revealed the opposite pattern ($r = -0.71$, $p <$
266 0.05). The $\delta^{15}\text{N}$ of non-cPOM significantly declined with pH.

267

268 *Mean residence time of soil fractions and roots*

269 Coarse particulate organic carbon (cPOC) turned over at decadal timescales whereas MRTs
270 of non-cPOC were 1.4 to 2.9 times longer. MRT of both fractions significantly increased with
271 soil acidity (Fig. 1). However, the slopes of the corresponding regression lines differed sig-
272 nificantly. A one pH-unit acidification caused MRT of cPOC to increase by 22% while the
273 same pH difference caused MRT of non-cPOC to increase by 86%. The result indicates that
274 MRT of non-cPOC responded more sensitively to soil acidity than MRT of cPOC (Fig. 2). In
275 other words, MRT of non-cPOC was on average $2.6 (\pm 0.15)$ times larger than MRT cPOC
276 below pH 4, whereas it was only $1.7 (\pm 0.17)$ times larger above pH 4 ($p < 0.01$, t-test). Fig. 2
277 also reveals that a different pH effect on the two soil fractions only occurred at pH below 6.1

278 (point of intersection). The mean age of root biomass was 8.7 (± 1.2) years. Soil pH has no
279 effect on the mean residence time of carbon in roots (Table 1).

280 Both the difference in MRT and in $\delta^{15}\text{N}$ between non-cPOM and cPOM (Table 1) were nega-
281 tively related to soil pH ($r = -0.70$, $p = 0.053$ and $r = -0.79$, $p < 0.05$, respectively). A greater
282 age of non-cPOM relative to cPOM thus coincided with a stronger enrichment of ^{15}N in that
283 fraction (Fig. 3). The relationship in Fig. 3 was significantly positive ($r = 0.83$, $p < 0.05$).

284

285 *Carbon flux through soil components*

286 Determination of carbon mass in roots, non-cPOM, cPOM, and bulk soil together with their
287 respective mean residence times allowed calculation of annual carbon fluxes through the
288 various belowground fractions (Fig. 4). At pH below 5.0, the carbon flux through cPOC and
289 non-cPOC fractions was similar whereas above pH 5.0, significantly more carbon annually
290 passed through the non-cPOC fraction. In contrast, carbon input delivered to the soil by root
291 turnover was independent of pH.

292

293 **Discussion**

294 *Carbon mean residence times and accrual of cPOM*

295 We found decadal to centennial carbon mean residence times in soil of our subalpine grass-
296 land pH gradient. Such long mean residence times are in line with previous studies on subal-
297 pine and alpine grasslands (Budge et al. 2011; Leifeld et al. 2009; Neff et al. 2002; Wang et
298 al. 2005). As a matter of principle this attribute may be related to factors controlling soil or-
299 ganic matter turnover, such as low temperatures, typical for sites at the treeline. The pH de-
300 pendency of many soil exoenzymes (Sinsabaugh et al. 2008) may be a major mechanism be-
301 hind the observed relationship between SOC turnover and pH. When low temperature coin-

302 cides with acidic soil, a frequent combination in mountain regions of humid climates, these
303 two factors act in concert. In addition to temperature and pH, smaller availabilities of Ca and
304 Mg at low pH may limit the overall microbial activity in our soil.

305 High proportions of cPOM of on average 25 percent were indicated in this study and seem
306 typical for subalpine and alpine environments (Budge et al. 2011; Leifeld et al. 2009; Neff et
307 al. 2002; Wang et al. 2008). Primarily this pattern might be related to a higher contribution of
308 roots to belowground SOM, as compared to temperate soils (Leifeld et al. 2009), i.e., it might
309 reflect pathways of carbon input. A high proportion of cPOM may also be indicative for fac-
310 tors that affect turnover rates of cPOM in a different way than those affecting turnover rates
311 of non-cPOM because otherwise total carbon stocks, but not the distribution of carbon among
312 fractions, would differ. In previous work (Leifeld et al. 2009), temperature was not found to
313 act differently on cPOM relative to non-cPOM turnover along a grassland elevation gradient.
314 We argue that the observed accrual of cPOM in cold grassland soil is also not caused by di-
315 rect effects of soil acidity on its decomposition as cPOM content did not scale with pH and
316 low pH was more limiting for the turnover of non-cPOM. The proportion of cPOM to SOM
317 would thus be expected to be maximum at high pH because of the relatively stronger stimula-
318 tion of non-cPOM decomposition. Hence, high cPOM content in cold grasslands may be
319 mainly driven by other mechanisms such as i) a higher contribution of roots to belowground
320 inputs, ii) vegetation-induced poor litter qualities as compared to warmer and fertilized, less
321 acidic sites, and ii) subsequent preferential feeding by macro-decomposers and shifts in mi-
322 crobrial decomposer communities (Eskelinen et al. 2009 and Seeber et al. 2009).

323

324 *Differential response of belowground carbon fractions dynamics to pH*

325 The most prominent observation was the differential effect of pH on turnover of the various
326 belowground carbon fractions under otherwise similar environmental conditions. The differ-

327 ential pH effect on cPOC vs. non-cPOC turnover may be explained by two factors. First, the
328 higher pH maintained by roots may attenuate pH limitation on enzyme activities. Root pH
329 varied by only 0.7 units whereas soil pH varied by 2 units. Considering that roots are the
330 main source for cPOM, cPOM turnover rate may benefit less from higher soil pH than non-
331 cPOM because of a higher pH of its feedstock. Therefore, soil pH seems to be an unreliable
332 predictor for pH controls on belowground plant residue decomposition. The stability of root
333 pH across the soil pH gradient may also be one reason for the small variability in root turn-
334 over. Additionally, root quality seems largely unaffected by soil pH or vegetation community
335 as both root C/N and root degradability, in the incubation experiment, revealed no trend
336 across sites despite marked differences in vegetation composition and productivity. Second, a
337 relatively strong reduction of non-cPOM turnover below pH 5 may be related to the contribu-
338 tion of mineral associated OM as a potentially stable component of our non-cPOM fraction
339 and the availability and nature of soluble OM as a potentially labile component of our non-
340 cPOM fraction. Because OM solubility, *inter alia*, depends on its surface charge density, it
341 typically correlates positively with soil solution pH (Kalbitz et al. 2000), supporting a larger
342 microbial availability at higher pH. In addition, at pH < 5 various organic compounds can
343 intercalate into interlayer spaces of 2:1 phyllosilicates, an effective SOM stabilization
344 mechanism, because their degree of dissociation is small (von Lützow et al 2006). Further-
345 more, complexation of SOM by reactive inorganic hydroxyls via ligand exchange, another
346 powerful stabilization mechanism, usually increases with decreasing pH as it is limited to
347 protonated hydroxyl groups (Kaiser and Guggenberger 2007; von Lützow et al. 2006).
348 Mechanisms related to the nature, and thus inherent degradability, of the substrate may exert
349 additional control on mean residence times. Adsorptive mineral association is selective to the
350 nature of the organic molecule (Kalbitz et al. 2000). The molecular composition is partially
351 driven by the vegetation community which was strongly graded along pH in our case (see site

352 description). Co-precipitation of dissolved OM (DOM) by aluminium, another proposed sta-
353 bilization mechanism in acid soil, tended to be selective and preferential for compounds high
354 in aromaticity but low in N in samples from a forest soil (Scheel et al. 2007). In the latter
355 study, co-precipitation was shown to be greater at pH 3.8 vs. pH 4.5 and DOM mineraliza-
356 tion, and thus turnover, was higher at pH 4.5 which is in line with our results. Together, these
357 stabilizing mechanisms may act specifically on the turnover of non-cPOM which includes
358 mineral-associated OM, reducing the exchange rate and thus the microbial availability of OM
359 at low pH. This is in line with the much longer MRT of non-cPOM in soil of greater acidity.
360 Concentrations and thus availability of DOM, however, may be higher at low pH in contrast
361 to its genuine solubility due to a decline in the degree of metal-organic complexation with
362 increasing acidity (proton competition; Guggenberger et al. 1994). Our data indicate that the
363 latter mechanism may be of minor importance but that a high pH supports DOM availability
364 and thus turnover.

365 Differences in soil pH often go along with differences in soil mineralogy and the latter exerts
366 control on the stabilization of mineral associated organic matter (Denef et al. 2004; Mikutta
367 et al. 2009). However, there is no indication for differences in soil texture, mineralogy, bulk
368 elemental composition, or the surface area of the fine soil fraction across our pH gradient. We
369 therefore consider possible effects induced by differences in soil mineralogy on altering or-
370 ganic matter (OM) turnover rates to be negligible at these sites.

371

372 *¹⁵N enrichment as a function of soil pH*

373 A longer MRT of non-cPOM coincided with a stronger ¹⁵N enrichment in non-cPOM relative
374 to cPOM. This enrichment is most probably a result of isotope discrimination processes along
375 microbial transformation pathways and corresponds to previous studies showing that non-
376 cPOM and mineral-associated OM is microbially more transformed than POM (e.g., Conen et

377 al. 2008; Kramer et al. 2003; Tiessen et al. 1984). Interestingly, the ^{15}N signature of cPOM
378 did not change with pH whereas that of non-cPOM increased with declining pH, i.e., the de-
379 gree of microbial transformation of stabilized OM was larger in acidic soil. At the same time,
380 we calculate significantly smaller carbon inputs into the non-cPOM fraction at pH below 5.
381 The difference in input was about $14 \text{ g C m}^{-2} \text{ a}^{-1}$ between sites below and above pH 5, and
382 corresponded well to the difference in aboveground productivity of about $35 \text{ g dry matter m}^{-2}$
383 a^{-1} . Hence the higher delivery rate by the vegetation caused by the larger aboveground pro-
384 ductivity, at higher pH, may be one reason behind the finding that organic matter recovered
385 in the non-cPOM fraction at higher pH had a $\delta^{15}\text{N}$ signature more closely to that of plants.
386 The nature of the substrate may also play a role in OM stabilization, resulting in a pH-
387 dependency of $\delta^{15}\text{N}$ in the non-cPOM fraction. The various soil organic N pools differ sub-
388 stantially in their isotopic signature (Yano et al. 2010) and a preferential adsorption of any of
389 these compound classes at low pH might result in a systematic shift in $\delta^{15}\text{N}$ of non-cPOM.
390 With our data set we cannot unravel the mechanisms behind the isotopic systematics but find-
391 ings point toward a differentiation in the type of molecules involved, as well as in rates of
392 carbon delivery and mechanisms and strengths of mineral association considering that non-
393 cPOM also includes mineral-associated OM.

394

395 **Conclusions**

396 A comparison of pH response factors from this study with previous work confirms a strong
397 pH-dependency of soil carbon turnover rate (Fig. 5). Eskelinen et al. (2009) argued soil pH to
398 be the ultimate factor driving vegetation and microbial community patterns in tundra soil. We
399 add that pH is a key driver for the turnover of organic matter in cold grassland soil because
400 the previously stated strong dependency of turnover rates on pH has now been quantified and
401 confirmed under long-term steady-state field conditions. We argue that soil pH should be an

402 integrative part of global carbon and nitrogen turnover modelling. Soil acidity exerts stronger
403 control on turnover of older non-cPOM than on residue decomposition, albeit the effect is
404 significant in both cases. This differential effect is related to the pH of the corresponding
405 feedstock, or the solution in its vicinity, and to pH-dependent stabilization of mineral associ-
406 ated OM.

407

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414

415 **References**

416

417 Anderson JM (1991) The effects of climate change on decomposition processes in grassland
418 and coniferous forests. *Ecological Applications* 1: 326-347

419 Bassin S, Volk M, Suter M, Buchmann N, Fuhrer J (2007) Nitrogen but not ozone affects
420 productivity and species composition of subalpine grassland after 3 year of treatment. *New*
421 *Phytologist* 175: 523–534

422 Bouma J, Hoeks J, van der Plas L and van Scherrenburg B (1969) Genesis and morphology
423 of some alpine podzol profiles. *J. Soil Sci.* 20: 384-398

424 Budge K, Leifeld J, Hiltbrunner E, Fuhrer J (2011) Alpine grassland soils contain large pro-
425 portion of labile carbon but indicate long turnover times. *Biogeosciences* 8: 1911-1923.

426 Conen F, Zimmermann M, Leifeld J, Seth B, Alewell C (2008) Relative stability of soil car-
427 bon revealed by shifts in delta N-15 and C : N Ratio. *Biogeosciences* 5: 123-128

428 Deneff K, Six J, Merckx R, Paustian K (2004) Carbon sequestration in microaggregates of no-
429 tillage soils with different clay mineralogy. *Soil Sci. Soc. Am. J.* 68: 1935-1944

430 Egli M, Mirabella A, Sartori G, Fitze P (2003) Weathering rates as a function of climate: re-
431 sults from a climosequence of the Val Genova (Trentino, Italian Alps). *Geoderma* 111: 99-
432 121

433 Eskelinen A, Stark S, Mannisto M (2009) Links between plant community composition, soil
434 organic matter quality and microbial communities in contrasting tundra habitats. *Oecologia*
435 161: 113-123

436 FAL (1998) *Methodenbuch für Boden-, Pflanzen- und Lysimeterwasseruntersuchungen*. Eid-
437 genössische Forschungsanstalt für Agrarökologie und Landbau. Schriftenreihe der FAL, 27,
438 Zurich, Switzerland

439 Guggenberger G, Glaser B, Zech W (1994) Heavy-metal binding by hydrophobic and
440 hydrophilic dissolved organic carbon fractions in a Spodosol-A and Spodosol-B horizon. *Water Air and Soil Pollution* 72: 111-127

442 Harkness DD, Harrison AF, Bacon PJ (1986) The temporal distribution of bomb C-14 in a
443 forest soil. *Radiocarbon* 28: 328-337

444 Hobbie SE, Schimel JP, Trumbore SE, Randerson JR (2000) Controls over carbon storage
445 and turnover in high-latitude soils. *Glob. Change Biol.* 6: 196-210

446 Kalbitz K, Solinger S, Park JH, Michalzik B, Matzner E (2000) Controls on the dynamics of
447 dissolved organic matter in soils: A review. *Soil Science* 165: 277-304

448 Kaiser K, Guggenberger G (2007) Sorptive stabilization of organic matter by microporous
449 Goethite: sorption into small pores vs. surface complexation. *European Journal of Soil*
450 *Science* 58: 45-59

451 Kleeberg R, Monecke T, Hillier S. (2008) Preferred orientation of mineral grains in sample
452 mounts for quantitative XRD measurements: How random are powder samples? *Clays and*
453 *Clay Minerals* 56: 404-415

454 Körner C. (2003) *Alpine plant life*. Springer, Heidelberg

455 Kramer MG, Sollins P, Sletten RS, Swart PK (2003) N isotope fractionation and measures of
456 organic matter alteration during decomposition. *Ecology* 84: 2021-2025

457 Leifeld J, Zimmermann M, Fuhrer J (2008) Simulating decomposition of labile soil organic
458 carbon: Effects of pH. *Soil Biology & Biochemistry* 40: 2948-2951

459 Leifeld J and Fuhrer J (2009) Long-term management effects on soil organic matter in two
460 cold, high-elevation grasslands: clues from fractionation and radiocarbon dating. *Eur. J. Soil*
461 *Sci.* 60: 230-239

462 Leifeld J, Zimmermann M, Fuhrer J, Conen F (2009) Storage and turnover of carbon in grass-
463 land soils along an elevation gradient in the Swiss Alps. *Global Change Biology* 15: 668-679

464 Levin I, Kromer B (2004) The tropospheric (CO₂)-C-14 level in mid-latitudes of the North-
465 ern Hemisphere (1959-2003). *Radiocarbon* 46: 1261-1272

466 Mikutta R, Schaumann GE, Gildemeister D, Bonneville S, Kramer MG, Chorover J, Chad-
467 wick OA, Guggenberger G (2009) Biogeochemistry of mineral-organic associations across a
468 long-term mineralogical soil gradient (0.3-4100 kyr), Hawaiian Islands. *Geochimica et Cos-
469 mochimica Acta* 73: 2034-2060

470 Neff JC, Townsend AR, Gleixner G, Lehman SJ, Turnbull J, Bowman WD (2002) Variable
471 effects of nitrogen additions on the stability and turnover of soil carbon. *Nature* 419: 915-917

472 Nilsson MC, Wardle DA, Zackrisson O, Jaderlund A (2002) Effects of alleviation of ecologi-
473 cal stresses on an alpine tundra community over an eight-year period. *Oikos* 97: 3-17

474 Pietri JCA, Brookes PC (2008) Nitrogen mineralisation along a pH gradient of a silty loam
475 UK soil. *Soil Biology & Biochemistry* 40: 797-802

476 Rasse DP, Rumpel C, Dignac MF (2005) Is soil carbon mostly root carbon? Mechanisms for
477 a specific stabilisation. *Plant and Soil* 269: 341-356

478 Scheel T, Dorfler C, Kalbitz K (2007) Precipitation of dissolved organic matter by aluminum
479 stabilizes carbon in acidic forest soils. *Soil Sci. Soc. Am. J.* 71: 64-74

480 Seeber J, Langel R, Meyer E, Traugott M (2009) Dwarf shrub litter as a food source for ma-
481 cro-decomposers in alpine pastureland. *Applied Soil Ecology* 41: 178-184

482 Sinsabaugh RL, Lauber CL, Weintraub MN, Ahmed B, Allison SD, Crenshaw C, Contosta
483 AR, Cusack D, Frey S, Gallo ME, Gartner TB, Hobbie SE, Holland K, Keeler BL, Powers

484 JS, Stursova M, Takacs-Vesbach C, Waldrop MP, Wallenstein MD, Zak DR, Zeglin LH
485 (2008) Stoichiometry of soil enzyme activity at global scale. *Ecology Letters* 11: 1252-1264

486 Sjögersten S, Alewell C, Cécillon L, Hagedorn F, Jandl R, Leifeld J, Martinsen V,
487 Schindlbacher A, Sebastià M-T, Van Miegroet H (2011) Mountain soils in a changing cli-
488 mate – vulnerability of C stocks and ecosystem feedbacks. In Jandl R et al. (eds): *Soil carbon*
489 *balance in sensitive ecosystems in Europe: Relevance to policy issues*. Wiley-Blackwell, in
490 press

491 Spiegelberger T, Hegg O, Matthies D, Hedlund K, Schaffner U (2006) Long-term effects of
492 short-term perturbation in a subalpine grassland. *Ecology* 87: 1939-1944

493 Stuiver M, Polach HA (1977) Reporting of C-14 data - discussion. *Radiocarbon* 19: 355-363

494 Stuiver M, Reimer PJ, Braziunas TF (1998) High-precision radiocarbon age calibration for
495 terrestrial and marine samples. *Radiocarbon* 40: 1127-1151.

496 Synal HA, Stocker M, Suter M (2007) MICADAS: A new compact radiocarbon AMS sys-
497 tem. *Nuclear Instruments & Methods in Physics Research Section B-Beam Interactions with*
498 *Materials and Atoms* 259: 7-13

499 Tiessen H, Karamanos RE, Stewart JWB, Selles F (1984) Natural N-15 abundance as an in-
500 dicator of soil organic-matter transformations in native and cultivated Soils. *Soil Sci. Soc.*
501 *Am. J.* 48: 312-315

502 Torn MS, Swanston CW, Castanha C, Trumbore SE (2009) Storage and turnover of organic
503 matter in soil. In: Senesi N, Xing B & Huang PM (eds) *Biophysico-chemical processes in-*
504 *volving natural nonliving organic matter in environmental systems*. John Wiley & Sons, Inc.
505 p 219-272

506 Townsend AR, Vitousek PM, Trumbore SE (1995) Soil organic-matter dynamics along gra-
507 dients in temperature and land-use on the Island of Hawaii. *Ecology* 76: 721-733

508 Trumbore S E (1997) Potential responses of soil organic carbon to global environmental
509 change. *Proc. Natl. Acad. Sci. USA*, 94, 8284–8291.

510 Von Lützow M, Kögel-Knabner I, Ekschmitt K, Matzner E, Guggenberger G, Marschner B,
511 Flessa H (2006) Stabilization of organic matter in temperate soils: mechanisms and their
512 relevance under different soil conditions - a review. *European Journal of Soil Science* 57:
513 426-445

514 Walse C, Berg, B, Sverdrup, H (1998) Review and synthesis of experimental data on organic
515 matter decomposition with respect to the effect of temperature, moisture, and acidity. *Envi-
516 ronmental Reviews* 6: 25-40

517 Wang L, Ouyang H, Zhou CP, Zhang F, Song MH, Tian YQ (2005) Soil organic matter dy-
518 namics along a vertical vegetation gradient in the Gongga Mountain on the Tibetan Plateau.
519 *Journal of Integrative Plant Biology* 47: 411-420

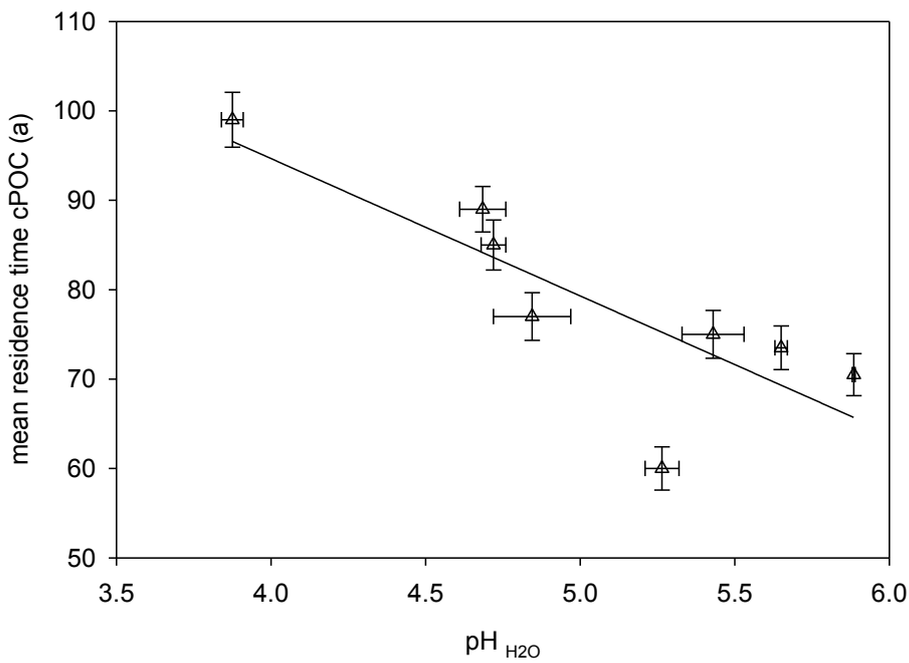
520 Wang G, Li Y, Wang Y, Wu Q (2008) Effects of permafrost thawing on vegetation and soil
521 carbon pool losses on the Qinghai-Tibet Plateau, China. *Geoderma* 143: 143-152

522 Yano Y, Shaver GR, Giblin AE, Rastetter EB (2010) Depleted ^{15}N in hydrolysable-N of arc-
523 tic soils and its implication for mycorrhizal fungi-plant interaction. *Biogeochemistry* 97: 183-
524 194

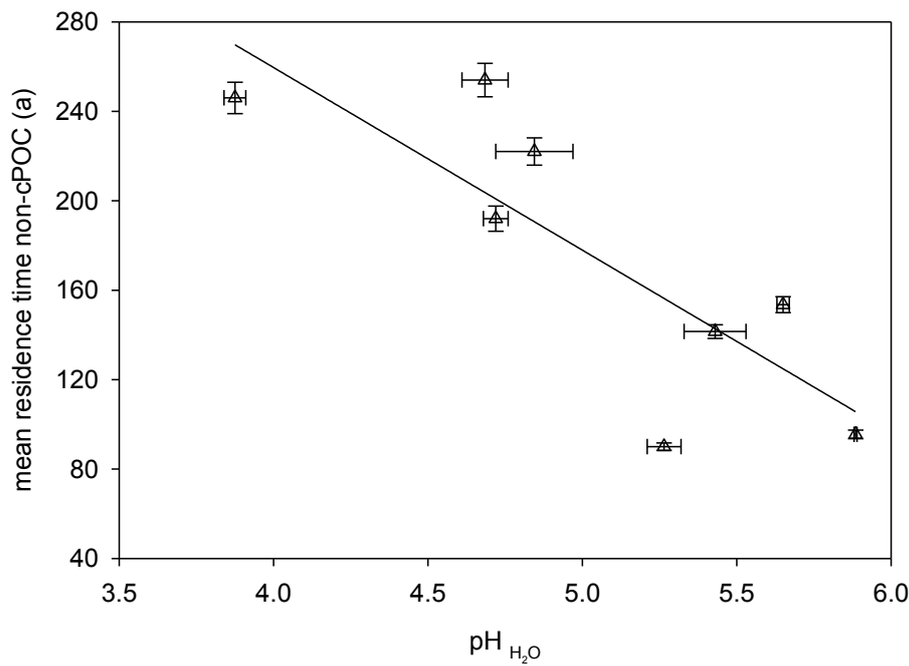
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526 Table 1. Soil and biomass characteristics along the pH gradient. Errors in parenthesis are one
 527 SE. Last column shows correlation coefficient to variable 'soil pH'. n.s. non significant; n.d.
 528 not determined. pMC percent modern carbon.

Soil pH		3.9 (<0.1)	4.7 (0.1)	4.7 (<0.1)	4.9 (0.1)	5.3 (0.1)	5.5 (0.1)	5.7 (<0.1)	5.9 (<0.1)	r
Clay	mg g ⁻¹	290	290	330	260	310	310	340	310	n.s.
Sand	mg g ⁻¹	350	360	270	370	310	300	290	320	n.s.
SOC	%	12.8 (0.9)	12.8 (1.4)	13.5 (1.0)	10.6 (1.3)	8.5 (0.3)	9.7 (0.5)	11.6 (0.8)	10.1 (0.2)	n.s.
SOC	kg m ⁻²	5.42 (0.46)	6.09 (0.48)	4.86 (0.29)	5.25 (0.61)	5.00 (0.11)	5.03 (0.37)	4.54 (0.69)	4.75 (0.33)	n.s.
C/N soil		13.0 (0.34)	15.3 (0.78)	13.2 (0.11)	14.2 (0.72)	11.7 (0.15)	12.0 (0.15)	11.7 (0.20)	12.1 (0.13)	n.s.
cPOC / SOC		0.26	0.31	0.31	0.23	0.23	0.24	0.18	0.22	n.s.
C/N cPOM		17.3	19.9	14.9	19.6	20.5	18.0	17.1	20.8	n.s.
Root-C	kg m ⁻²	0.45 (0.10)	0.58 (0.07)	0.33 (0.10)	0.73 (0.32)	0.27 (0.07)	0.3 (0.04)	0.29 (0.05)	0.43 (0.13)	n.s.
Root biom.	kg m ⁻²	1.02 (0.24)	1.24 (0.17)	0.71 (0.20)	1.55 (0.65)	0.64 (0.20)	0.66 (0.10)	0.65 (0.13)	1.01 (0.34)	n.s.
Root pH		5.22 (0.03)	4.94 (0.04)	5.14 (0.04)	5.18 (0.01)	5.34 (0.02)	5.42 (0.02)	5.68 (<0.01)	5.70 (0.01)	0.77*
C/N roots		32.2 (1.9)	47.6 (3.1)	48.0 (6.5)	46.2 (4.9)	34.3 (2.2)	34.5 (3.2)	42.8 (6.0)	52.4 (5.1)	n.s.
root decay	d ⁻¹ * 1000	7.1 (0.4)	6.9 (0.1)	7.3 (0.1)	6.7 (0.5)	7.0 (0.3)	6.9 (0.3)	7.7 (0.2)	6.1 (0.1)	n.s.
root MRT	years	7.7	9.2	8.6	17.0	6.1	7.4	7.1	6.8	n.s.
abovegrd. biomass	g m ⁻²	74.2	109.5	105.0	110.5	126.1	140.9	147.3	124.3	0.90**
δ ¹⁵ N cPOM	‰	2.1	0.3	1.2	-0.6	0.2	1.4	1.5	0.8	n.s.
δ ¹⁵ N non-cPOM	‰	4.7	4.1	3.8	2.2	2.0	2.8	2.7	2.4	-0.77*
pMC SOC		103.9	104.3	105.8	104.7	110.4	107.3	106.5	109.6	n.d.
pMC cPOC		108.8	109.7	110.0	110.9	113.0	111.1	111.3	111.6	n.d.
pMC non-cPOC		102.1	101.9	103.8	102.9	109.6	106.0	105.4	109.1	n.d.



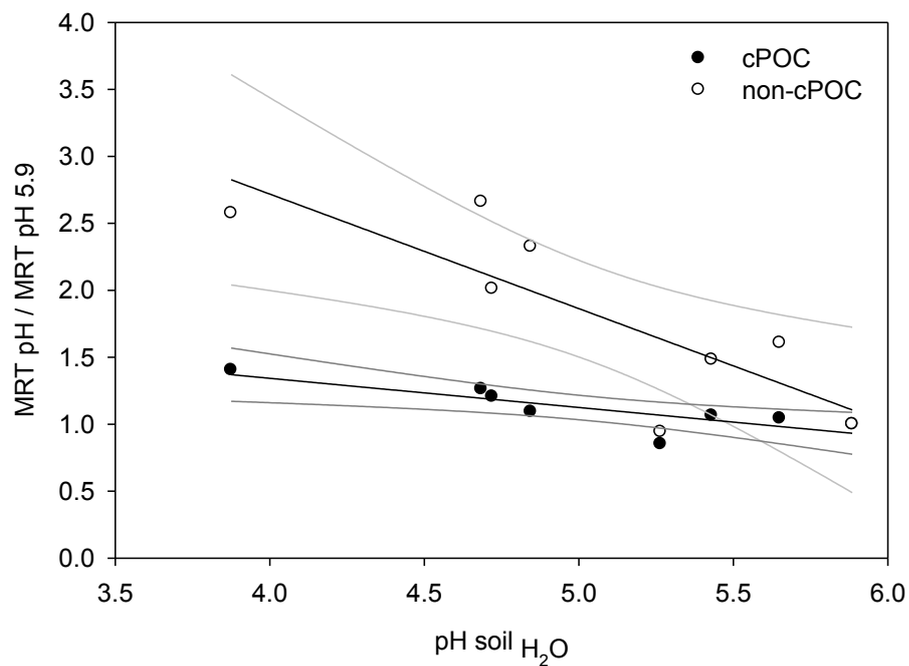
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530

531 Fig. 1. Mean residence time of cPOC (top) and non-cPOC (bottom) along the pH gradient
 532 and corresponding regression line. Regression equations (± 1 SE): MRT cPOC = $156 (\pm 22) -$
 533 $15.4 (\pm 4.4) * \text{pH}$ [$R^2 = 0.68, p = 0.012$]; MRT non-cPOC = $586 (\pm 118) - 82 (\pm 24) * \text{pH}$ [R^2
 534 $= 0.67, p = 0.013$]

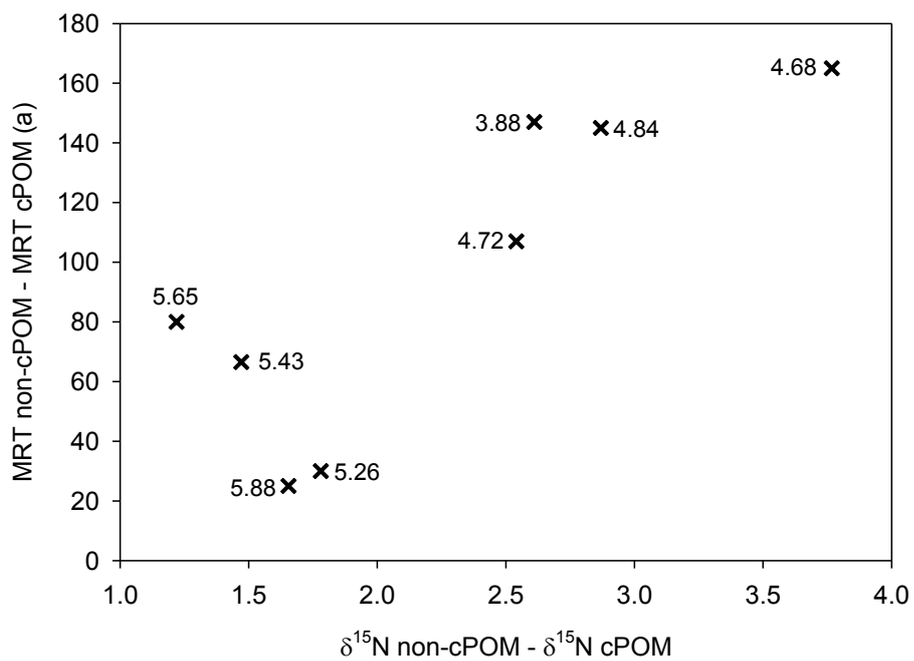
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536

537 Fig. 2. Mean residence times (MRT) of cPOC and non-cPOC at different pH values relative
 538 to MRT at pH 5.9. Light grey lines are 95% confidence intervals of the regression line. Point
 539 of intersection is at pH 6.1. The regression for cPOC is $2.21 (\pm 0.31) - 0.22 (\pm 0.06) \text{ pH} (\pm 1$
 540 $\text{SE})$ and for non-cPOC $6.14 (\pm 1.24) - 0.86 (\pm 0.24) \text{ pH}$. Slopes are significantly different.

541

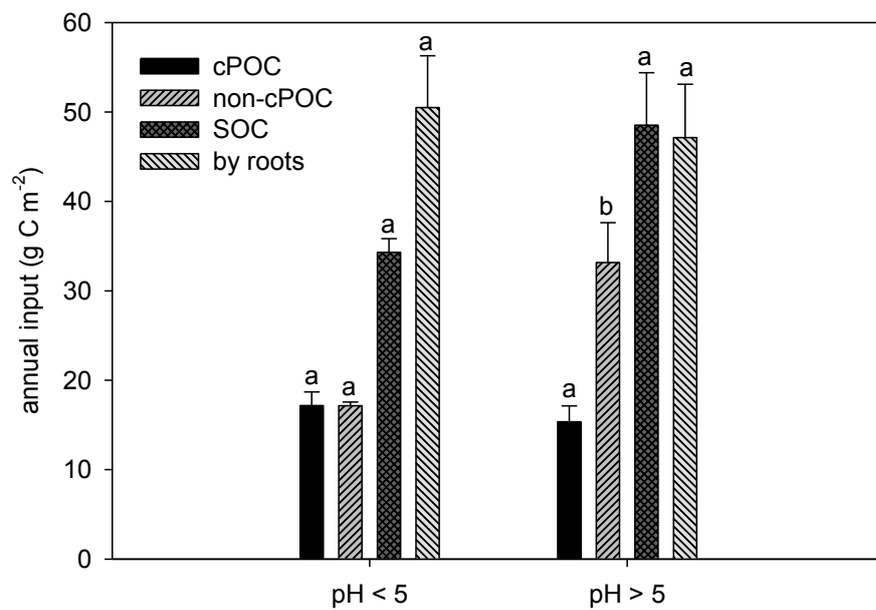


542

543 Fig. 3. Difference in $\delta^{15}\text{N}$ (non-cPOM-cPOM) relative to difference in carbon mean residence

544 time (non-cPOM-cPOM). Numbers next to symbols show pH value.

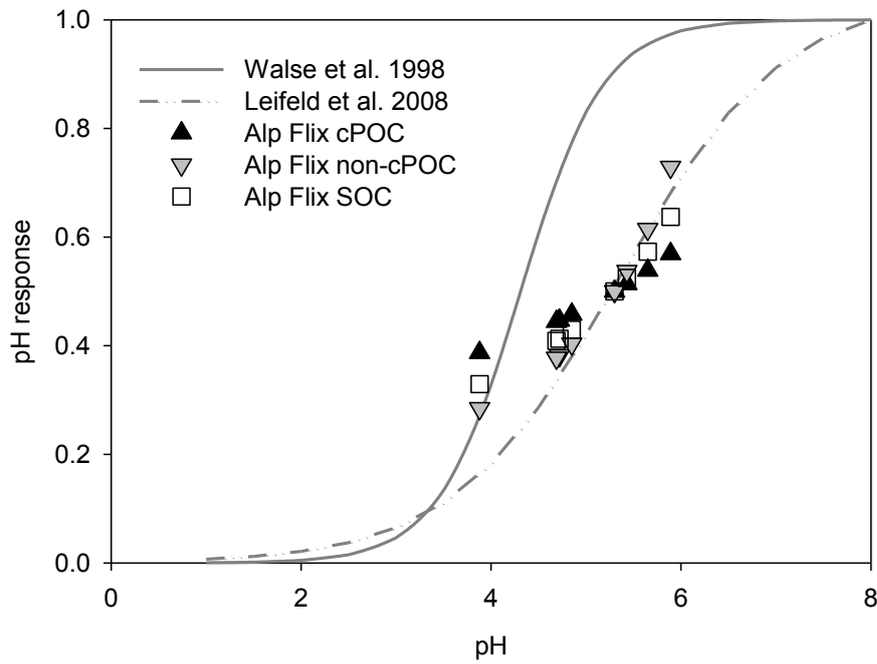
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546

547 Fig. 4. Annual carbon input to cPOC, non-cPOC, SOC and carbon delivered by root turnover
 548 below and above a soil pH of 5.0. Different letters indicate significant difference between pH
 549 class ($p < 0.05$, t-test). Error bars show 1 SE.

550



552

553 Fig. 5. Comparison of pH response functions for litter decomposition (Walse et al. 1998),
 554 cPOC turnover (Leifeld et al. 2008), and for bulk soil carbon, cPOC and non-cPOC from this
 555 study. The midpoint of the sigmoid (i.e., pH response = 0.5) was assumed to be the same as
 556 in the function of Leifeld et al. (2008).