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1 Management of grasslands by mowing versus grazing – impacts on soil organic matter

2 quality and microbial functioning

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14	Keyw	ords					
15	Grassl	and, grazing, mowing, non-cellulosic polysaccharides, lignin, microbial activity					
16							
17							
18	Highlights						
19	•	Grazing and mowing have contrasting effects on soil biogeochemical properties					
20	•	Grazing promoted more efficient microbial functioning					
21	•	Mowing leads to more degraded lignin than grazing					
22	•	Only microbial properties were sensitive to treatment effects in subsurface soil					
23							

Abstract

25 Although 30% of the European surface area is covered with grasslands, little is known 26 about the effect of their management on soil quality and biogeochemical cycling. Here, we 27 analysed soil from an experimental site in Western France, which had been under either 28 grazing or mowing regime for 13 years. We aimed to assess the effect of the two 29 management practices on the biogeochemical functioning of soil system. To this end we 30 compared soil organic matter (SOM) composition and microbial properties at two soil depths. 31 We analysed for elemental, lignin and non-cellulosic polysaccharide content and composition 32 and for microbial biomass, soil microbial respiration and enzyme activities. Our results 33 showed higher soil organic carbon (SOC) and nitrogen contents in the surface soil under 34 grazing as compared to mowing. Soil biogeochemical properties differed between grazing 35 and mowing treatments. In particular, soil under grazing showed lower lignin and higher 36 microbial biomass. Despite the similar non-cellulosic polysaccharide content under both 37 treatments, microbial community under mowing was characterised by higher enzyme 38 production per microbial biomass, leading to more degraded SOM in the mowing system as 39 compared to grazing. We conclude that grazing and mowing regimes impact differently the 40 biogeochemical soil functioning. Higher and more diverse carbon input under grazing 41 compared to mowing may lead to enhanced substrate availability and thus more efficient 42 microbial functioning, which could favour SOC sequestration through formation of microbial 43 products.

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45

1. Introduction

48 Dangerous climate change can only be avoided if we succeed to remove CO_2 from the atmosphere with negative emission technologies (IPCC, 2018). Soil organic carbon (SOC) 49 50 sequestration is a nature-based negative emission technology, which may be achievable at 51 scale through the introduction of sustainable management practices with permanent soil cover 52 (Rumpel et al., 2018). Permanent grasslands, which in Europe, occupy about 30% of the 53 agricultural area (Ec.europa.eu, 2018), are responsible for many ecosystem services including 54 forage for animal production and SOC storage (Havstad et al., 2007; Rumpel et al., 2015). 55 Biogeochemical cycling in grassland soils can be influenced by a variety of management 56 practices (Rumpel et al., 2015). The impact of these management practices on processes 57 impacting soil biogeochemical cycling via soil-plant interactions are poorly understood 58 (Dignac et al., 2017). These interactions result in contrasting effects of grassland 59 management on SOC storage potential (Post and Kwon, 2000; Rumpel et al., 2015; Smith et 60 al., 2008).

Grazing and mowing are the most frequently used grassland management practices. Both practices lead to defoliation (removal of plant aboveground tissue). Defoliation alters root exudation and C allocation in plants but the direction of these changes was found to be contrasting (Bazot et al., 2005; Gavrichkova et al., 2010; Medina-Roldán and Bardgett, 2011), related to different climatic and pedological conditions (Pineiro *et al.*, 2010).

Defoliation under grazing management is caused by herbivores during several days (Senapati et al., 2014). This process plays an important role in terms of carbon and nutrient return (Soussana et al., 2006), because about 50-70% of the ingested biomass is returned to soil in the form of excreta. In mowing systems, plant biomass is removed in a day with up to 20% of all cut biomass remaining as green litter in form of harvesting losses (Sanaullah et al., 2010). In order to compensate for nutrient exportation during defoliation events, mineral
fertilisers are applied in mowing systems.

73 Due to the different types of biomass returned in the two systems, the quality of 74 biomass input also varies. Mowing systems receive only plant residues while input in grazing 75 systems comprises additionally animal depositions. Dung and urine inputs are characterised 76 by lower C:N ratio, higher amount of easily available compounds (Dungait et al., 2009) and 77 relatively stable compounds, such as crude proteins and fats (Dungait et al., 2005; Ngo et al., 78 2011). Moreover, in grazing systems, there is a return of senescent brown litter, which 79 contains less N and less soluble compounds compared to the green litter returned as 80 harvesting losses in mowing systems (Sanaullah et al., 2010).

81 These differences may affect belowground processes (Wilson et al., 2018), SOC 82 formation and storage (Cotrufo et al., 2015; Rumpel et al., 2015) through their effect on the 83 soil microbial biomass and its activity (Liang et al., 2017). We therefore hypothesised that the 84 two management systems may lead to contrasting soil microbial functioning and affect 85 differently biogeochemical cycling. The effect of management has been analysed up to now 86 mainly in the first few centimetres of soil, although it has been shown that management can affect SOC stored down to 2 m depth (Tautges et al., 2019). We thus hypothesised that 87 88 grassland management affects SOC below the first centimetres.

We focused on an experimental site with grazing and mowing as two contrasting management practices under similar soil and climatic conditions. We aimed to evaluate the differences in biogeochemical cycling in soil under the two different management practices at two depths. To this end we analysed C and N contents, molecular signatures of polysaccharide and lignin monomers. These variables were compared to the functioning of the soil microbial communities, assessed by the analyses of soil microbial respiration, growth kinetic parameters and activity of 9 enzymes as well as microbial biomass C and N.

97 2. Materials and methods

98

2.1. Site description and soil sampling

99 The field experiment is located in Lusignan (southwest of France, 46°25'12,91"N; 100 0°07'29,35"E) at the national long-term experimental observatory SOERE ACBB 101 (Agroecosystems, Biogeochemical Cycles and Biodiversity). The mean annual temperature 102 and precipitation for the period 2006–2010 were 11.2°C and 773 mm, (Senapati et al., 2014). 103 The landscape is flat. The soil is classified as a Dystric Cambisol with loamy texture (Chabbi 104 et al., 2009).

105 The current study is focused on two permanent sown grasslands (each of about 3 ha in size), which were established in 2005 by sowing a mixture of three plant species (Lolium 106 107 perenne, Festuca arundinacea, Dactylis glomerata L.) in both treatments. In the grazing 108 system, legume Trifolium repens was included in the species mixture but covered only 5% of 109 grazed paddock in 2017. The mown grassland was cut four times per year with biomass 110 exported. To replace the exported nutrients, nitrogen (N) fertilizer was applied at rates between 170 and 380 kg N ha⁻¹ year⁻¹ (Puche et al., 2019). Grazing in the grazed paddock 111 112 took place from March to December with 50 days per year using 15 to 20 livestock units per 113 hectare. Grazed grasslands received less nitrogen fertilization (60-150 kg N ha⁻¹ year⁻¹, Puche 114 et al., 2019) because nitrogen losses were additionally returned by dung and urine and 115 through the presence of the leguminous species. In order to compare the treatments at similar 116 N status, fertilizer application rates were adjusted to maintain the Nitrogen Nutrition Index 117 between 0.9 and 1.0 for both treatments, close to non-limiting nitrogen nutrition to near 118 maximum plant production (Senapati et al., 2016). Moreover, both sites were limed regularly 119 in order to neutralize acid pH.

120 Due to the large land requirements (3 ha for plots with cows), it was not possible to 121 establish and maintain a completely replicated field experiment including grazing treatment 122 for several decades. Limitations to generalization of the treatment effects due to the absence 123 of replication of the experiments were limited by choosing homogenous flat areas in close 124 proximity with similar land use history, climate, and soil type. Moreover, we carried out 125 baseline measurements, in form of geostatistical evaluation of the soils SOC and N contents and included initial SOC stocks as a co-variate. These data showed that both plots were 126 127 significantly different in initial SOC and N contents (n=28). The SOC contents on mowing plots varied between 9.9 and 13.7 mg g⁻¹ (average 12.0 ± 1.0 mg g⁻¹), while under grazing it 128 was between 11.9 and 19.1 mg g⁻¹ (average 14.8 ± 1.5 mg g⁻¹). N contents varied between 1.0 129 and 1.4 mg g⁻¹ (average 1.2 \pm 0.1 mg g⁻¹) under mowing, while under grazing the values 130 ranged between 1.2 and 1.9 mg g⁻¹ (average 1.5 ± 0.1 mg g⁻¹). These previous analyses 131 132 indicated on average non-significant differences in SOC stock changes between grazing and mowing after nine years of treatment (Crème et al., 2020). The study also showed 133 partitioning of the field into different zones with SOC gain and loss (Crème et al., 2020; Fig 134 135 S1, Supplementary materials).

136

137 Five replicated soil samples were taken from each of the two zones, giving a total of 10 replicated field samples per plot. Sampling took place in November 2017, 2 weeks and 5 138 139 months after the last grazing and mowing events, accordingly. The shortest distance between 140 samples was 25 m. Soil samples were collected with a mechanical auger (5cm Ø, 30cm) at 141 two depths: 0-10 cm (surface soil) and 20-30 cm (subsurface soil) giving in total 40 samples. All samples were sieved through a 2-mm mesh. Thereafter, half of the samples was air-dried 142 143 and ground for measurements of physicochemical analysis and the other half was stored at 4°C before microbial analyses. Because of dry field conditions prior to measurements of 144

microbiological analysis, soil samples were moistened by distilled water to adjust 50% of
WHC and pre-incubated at 22 °C for 7 days.

147 2.2. Soil general properties

Soil pH (H₂O) was measured in a soil:water suspension (1:2.5 weight/volume). Soil organic carbon (SOC), nitrogen (N) and stable isotopes (13 C and 15 N) contents were measured using a CHN auto-analyser (Flash EA, Thermo Electron Corporation, Bremen, Germany) coupled with an isotope ratio mass spectrometer. The isotopic ratios were calculated relative to the Pee Dee Belemnite Standard (PDB) for C and relative to atmospheric N₂ for nitrogen.

- 153
- 154 2.3. Soil chemical properties

155 Lignin was analysed by the alkaline cupric oxide (CuO) oxidation method (Hedges and 156 Ertel, 1982; Kögel and Bochter, 1985). Briefly, oxidation was carried out under alkaline 157 conditions (2M NaOH) at 172 °C for 4 hours using 500 mg of air-dried soil, 250 mg of CuO, 158 50 mg of ammonium ferrous hexahydrate and 50 mg of glucose. After cooling, samples were 159 acidified with 5 M HCl and left overnight for humic acid precipitation. Removal of humic 160 acids was conducted through centrifugation (10 min at 10000 rpm) and followed by 161 extraction of phenolic oxidation products with C18 reversed phase columns. The phenols 162 were derivatized with BSTFA and quantified as trimethylsilyl derivatives by gas 163 chromatography with a HP gas chromatograph (HP GC 6890) equipped with a flame 164 ionization detector and a SGE BPX-5 column (50 m length, 0.25 mm inner diameter, 0.32 µm 165 coating). Samples were injected in split mode (1:10). The GC oven temperature was programmed at 100 °C for 2 min, then increased from 100 to 172 °C at a heating rate of 8 °C 166 167 min⁻¹, from 172 to 184 °C at 4 °C min⁻¹, and from 184 to 300 °C at a rate of 10°C min⁻¹.

168 The internal standard ethylvanillin was added before the purification step to quantify 169 lignin recovery and the quantification standard phenylacetic acid was added before GC 170 analyses.

The total lignin content (mg g⁻¹ dry soil) of the sample was determined as the sum of phenolic oxidation products: vanillyl (V), syringyl (S) and p-coumaryl (C) in their acid (Ac), aldehyde (Al) and ketone forms. Lignin content was also expressed as lignin content per SOC (mg g⁻¹ SOC). Lignin decomposition was assessed by the ratios of S, C to V and (Ac/Al) ratios of V and S, which generally indicate decomposition state (Thevenot et al., 2010).

176 Non-cellulosic polysaccharides of plant and microbial origin (Kögel-Knabner, 2002) 177 were determined by gas chromatography after trifluoroacetic acid (TFA) hydrolysis and 178 reduction-acetylation using a method introduced by Rumpel and Dignac (2006) and modified 179 by Eder et al. (2010). The analysis was performed using 700 mg of soil samples. Briefly, 180 hydrolysis of non-cellulose polysaccharides was carried out at 105°C for 4 h with 10 ml of 4 181 M TFA. Thereafter, Myo-inositol was added as quantification standard to account for the 182 losses during the purification procedure. Removal of soil was performed by filtration through 183 glass fibre filters (Whatman GF/C 0.45 µm). Afterwards, TFA was evaporated using a 184 centrifugal Evaporator EZ-2 ENVI at 35°C for 4 hours and dry samples were left overnight in 185 the freezer. Thereafter, dry samples were dissolved in 0.5 ml of H₂O followed by the addition 186 of 0.9 EDTA in order to avoid co-precipitation of organic material with metal oxides and 187 hydroxides (Eder et al., 2010). One mL sodium borohydride (NaBH4) in dimethylsulfoxide 188 (20 g L-1) was added for reduction of polysaccharide monomers into alditol forms and kept 189 at 40°C for 1.5 hours. Then, acetylation was conducted by addition of 0.2 mL acetic acid, 2 190 mL of acetic anhydride and 0.2 mL Methylimidazole. Acetylated alditols were extracted by 1 191 ml of dichloromethane and quantified with a HP GC 6890 gas chromatograph equipped with a flame ionization detector. Separation was achieved with a 60 m fused silica capillary 192

193 column (SGE BPX 70, 0.32 mm internal diameter, 0.25 mm film thickness) under the 194 following temperature program: 170 to 250 °C at 8 °C.min⁻¹, followed by 12 min at 250 °C 195 (isothermal). Helium was used as the carrier gas at a flow rate of 1.0 mL min⁻¹. The injector was kept at 250 °C and the detector at 260 °C. The non-cellulosic polysaccharides content of 196 197 soil samples was determined as the sum of monosaccharides: C5 (pentoses: xylose, ribose 198 and arabinose), C6 (hexoses: glucose, galactose and mannose), and desoxyC6 199 (desoxyhexoses: fucose and rhamnose) (Kögel-Knabner, 2002). A higher C6/C5 ratio 200 generally indicates higher contribution of microbial sugars.

201 2.4. Soil microbial properties

Microbial biomass C (MBC) and nitrogen (MBN) were determined by the chloroform 202 203 fumigation-extraction method (Vance et al., 1987). Dissolved organic C and N in fumigated 204 and non-fumigated soil samples were extracted in 0.05 M K₂SO₄ and were measured using a 205 multi C/N analyzer (multi C/N analyser 2100S, Analytic Jena). MBC and MBN were 206 calculated with a conversion factor of 0.45 (Jenkinson et al., 2004). For measuring soil 207 microbial respiration (SMR) a half gram of soil sample was placed in 2 ml Eppendorf tubes. 208 The CO₂ efflux was trapped in 3 ml of 0.1 M NaOH and determined by conductometry. The 209 metabolic quotient (qCO₂), reflecting decomposition activity (Anderson, 2003; Anderson and 210 Domsch, 1993), was calculated as soil microbial respiration expressed per gram of microbial biomass carbon: $qCO_2 = SMR/MBC$ (µg CO₂-C g⁻¹ MBC h⁻¹). 211

We used microbial growth kinetics technique as an approach to estimate microbial biomass activity state (Blagodatskaya and Kuzyakov, 2013). This approach is based on soil respiratory response to unlimited nutrient amendments (Panikov and Sizova, 1996). For this purpose, soil samples were treated with a solution (0.1 ml per g of dw soil) containing per g soil: 10 mg glucose, 1.9 mg (NH₄)₂SO₄, 3.8 mg MgSO₄*7H₂O, 0.11 mg K₂HPO₄ and 1.68 mg KH₂PO₄ for surface soil samples and 10 mg glucose, 1.9 mg (NH₄)₂SO₄, 3.8 mg MgSO₄*7H₂O, 0.53 mg K₂HPO₄ and 1.35 mg KH₂PO₄ for subsurface soil samples. The amount of mineral salts was preliminary selected in order to avoid soil pH change of more than 0.1 units after addition. For active microbial biomass (AMB) and specific growth rate calculation, the results of substrate induced respiration rate were fitted with a model proposed by Panikov and Sizova (Panikov and Sizova, 1996; Wutzler et al., 2012):

 $CO_2(t) = A + B * \exp(\mu * t) \tag{1}$

224 In order to estimate catabolic (decomposition) activity in regards to specific substrates 225 in soil, we measured extracellular enzyme activity using the fluorometric technique (Koch et 226 al., 2007; Marx et al., 2005; Razavi et al., 2015). Nine types of fluorogenic substrates based 227 on 4-methylumbelliferone (MUF) and 7-amino-4-methylcoumarin (AMC) were used: (1) MUF- α -D-glucopyranoside for α -glucosidase, (2) MUF- β -D-glucopyranoside for β -228 229 MUF- β -D-galactopyranoside for β -galactosidase, (4) MUF- β -Dglucosidase, (3) 230 xylopyranoside for β -xylosidase, (5) MUF- β -D-cellobioside for β -cellobiohydrolase, (6) 231 MUF-N-acetyl-β-D-glucosamide for chitinase, (7) Leucine-AMC for leucine aminopeptidase, (8) MUF-heptanoate for lipase and (9) MUF-phosphate for phosphatase. Saturation 232 233 concentrations of fluorogenic substrates were determined in preliminary experiments and comprised 20 μ mol g⁻¹ soil for all enzymes except lipase with 60 μ mol g⁻¹ soil. Briefly, a 234 water extract of soil (1:10) was homogenised by low-energy sonication (40 J s⁻¹ output 235 236 energy) for 60 s. Thereafter 50 ml of the soil suspension were added to 150 ml of each 237 substrate solution in a 96-well microplate. Fluorescence was measured at an excitation 238 wavelength of 355 nm and an emission wavelength of 460 nm (Victor3 1420-050 Multilabel 239 Counter, PerkinElmer, USA).

241 2.5. Statistical analysis

242 All results are presented as arithmetic means with standard error. The statistical 243 analyses were conducted by using R (Studio Version 1.1.447). We used analyses of 244 covariance (ANCOVA) to test treatment effect, depth effect and their interactions using 245 chemical and microbial variables with initial SOC stock as a covariate. The initial SOC 246 stocks data was obtained from exactly the same sampling points based on the geostatistical 247 evaluation before the beginning of the experiment. This procedure allowed us to account for 248 the lack of field replication by taking into account the original difference between the grazed 249 and mowed plots. In order to obtain better understanding of treatment and depth effects, non-250 transformed data (except C and N contents) were subjected to Principal Component Analysis 251 (PCA) and the results were also tested by ANCOVA with initial SOC stock as a covariate. 252 The equations were fitted by non-linear regression, using Model Maker-3 software (SB 253 technology Ltd.).

254

255 **3.** Results

256 *3.1.* Soil properties

257 Soil physicochemical properties are presented in Table 1. The pH was not controlled by 258 initial SOC stock (P=0.70). Lower pH was found for both treatments in surface soil compared 259 to subsurface soil, although, the lowest pH value was recorded in surface soils under mowing 260 treatment. SOC and N contents were nearly twice as high in the surface soil compared to the 261 subsurface soil under both treatments. Even if SOC and N contents were dependant on initial 262 SOC stock (P=0.03 and 0.02, respectively), there were still significant effects of depth 263 (P<0.001) and treatment (P<0.001) after correction by using it as covariate. C:N ratio differed 264 only between soil depths (P<0.001) showing slightly higher C:N ratios in surface soils as compared to subsurface soils. δ^{13} C followed the same pattern as SOC content and the highest enrichment was recorded for the surface soil of the grazing treatment (depth effect P<0.001 and treatment effect P=0.002). The δ^{15} N did not differ between the treatments and was enriched in surface soils compared to subsurface soils.

269

270 *3.2. Specific SOM compounds*

271 Non-cellulosic polysaccharide (NCP) content was not affected by initial SOC stock 272 (P=0.52) and there was treatment × depth interaction (Table 2, P<0.001). Grazing resulted in 273 higher NCP content in both depths compared to mowing. The NCP content per SOC (mg g⁻¹ 274 soil C) was affected only by depth (P=0.002). Concerning the NCP monomers ratio, C6/C5 275 and Man/Xyl ratios were controlled by initial SOC stock (P=0.03 and 0.04, respectively), 276 consequently, after ANCOVA application the treatment effect was varnished while depth 277 effect remained significant (Table 2, P<0.001). All NCP monomers ratios were higher in 278 subsurface soil compared to surface soil under both treatments.

279 Lignin content was not affected by initial SOC stock correction (P=0.82), so the effects 280 of depth (P<0.001), treatment (P<0.001) and their interactions (P=0.04) remained significant 281 (Table 3). Lignin content was higher in surface soils than in subsurface soils and was higher 282 under grazing compared to mowing as well. Correcting for initial SOC stock caused the 283 elimination of all effects on lignin content per SOC content. The C/V ratio was affected only by depth (P=0.006) showing higher values in surface soils than in subsurface soils. The S/V 284 285 ratio was greater under grazing treatment than under mowing treatment at both depths even 286 after correction by initial SOC stock (Table 3, P=0.01). Based on the presence of treatment \times 287 depth interaction (Ac/Al)_V and (Ac/Al)_S ratios were lower in the surface soil of grazing 288 treatment as compared to mowing treatment (P<0.001). In contrast to surface soils, treatments 289 did not show any effects on these lignin ratios in subsurface soils.

291

3.3. Soil microbial properties

The soil microbial respiration (SMR) ranged between 0.2 and 0.7 µg CO₂ –C g⁻¹ h⁻¹ 292 293 with highest values in the surface soil under grazing treatment (Fig. 1A). After correcting for 294 initial SOC stock, treatment × depth interaction effect on SMR was significant (Table S1, 295 Supplementary materials, P<0.001). Soil microbial respiration per SOC was around 33% 296 higher in the surface soil under grazing as compared to mowing (Fig. 1B). In contrast, it was 297 greater in the subsurface soil under mowing than under grazing treatment. Including initial 298 SOC stock as covariate resulted only in significant effect of treatment × depth interaction on 299 soil microbial respiration per SOC (Table S1, Supplementary materials, P=0.004).

MBC per SOC was highest in the surface soil under grazing (20 μ g C mg⁻¹ SOC, Fig. 1C). Mowing treatment resulted in two times lower MBC per SOC in the surface soil compared to grazing treatment. After correction for initial SOC stock, treatment (P<0.001) and their interaction (P<0.001) showed significant effects on qCO₂. Mowing treatment resulted in higher qCO₂ at both depths as compared to grazing treatment (Fig. 1D, P=0.02).

305 Microbial C:N ratio ranged between 4.9 and 6.4. It was affected by treatments in all 306 depths showing higher values under mowing (Fig. 2A). After taking into account initial SOC 307 stock, the treatment effect was still significant (Table S1, Supplementary materials, P<0.001). 308 Active microbial biomass was also higher under mowing at both depths compared to grazing 309 treatment (Fig. 2B, P=0.02). The highest specific microbial growth rate (Fig. 2C) was 310 recorded in subsurface soils without difference between treatments. But in surface soils, the 311 specific microbial growth rate was higher under grazing than under mowing (Fig. 2C). 312 However, ANCOVA with initial SOC stock as covariate decreased the significance treatment effects (P=0.05) on specific microbial growth rate but increased the depth effect Table S1, 313 314 Supplementary materials, P<0.001).

315 Treatment effect on absolute enzyme activities is presented only for leucine 316 aminopeptidase in surface soil and chitinase and phosphatase in subsurface soil (Fig S2, 317 Supplementary materials). When the initial SOC stock was used as covariate, treatment 318 differences between enzyme activities per MBC were observed for all enzymes (except 319 leucine aminopeptidase) in surface soil. Soil under mowing treatment showed 2-2.5 times 320 higher enzyme activity per MBC under mowing compared to soil under grazing (Fig. 3). The 321 differences between treatments were more pronounced in surface soil for activities of 322 chitinase, β -galactosidase, β -glucosidase and phosphatase (Table S1, Supplementary 323 materials).

324

325

3.4. Principal component analysis

Principal component analysis based on SOC normalised data of all soil properties 326 327 showed that the first two factors explained 54.4% of the variation (Fig. 4). The first 328 component (Dim1) was related to microbial functioning, as it was strongly associated with 329 the soil microbial properties MBC and MBN per SOC in negative direction. The positive 330 direction was related to the lipase activity per MBC. The second component (Dim2) was 331 explained by variables related to polysaccharides. It was positively correlated with enzymes 332 participating in polysaccharide degradation and negatively with polysaccharide ratios. The 333 clustering of samples allowed to separate surface soil and subsurface soil samples along both 334 axes, while surface soil samples were additionally separated by treatments along the first axes 335 (Fig. 4). Subsurface soil samples were differentiated from surface soil by high neutral 336 polysaccharide monomer ratio, low enzymes activities per MBC, MBC and MBN per SOC. 337 Treatments in surface soil were separated by C- and N-cycle enzyme activity and MBC and 338 MBN per SOC. We also applied ANCOVA with initial SOC stock as a covariate on new 339 PCA coordinates which resulted in significant effects of treatment, depth and their

interaction. Treatment effect was more pronounced on Dim1, while Dim2 was more affectedby depth.

342

343 **4. Discussion**

344 *4.1. Effect of grazing and mowing on chemical properties of surface soil*

Since the primary factor of SOM formation is organic matter input (Fujisaki et al., 345 346 2018; Kögel-Knabner, 2002), higher SOC and N contents in the surface soil under grazing 347 system might be explained by greater C input compared to mowing systems. This was shown 348 through ecosystem flux measurements at these plots (Senapati et al., 2014). Moreover, dung 349 return comprising about 50-80% of plant biomass could also favour higher SOC and N 350 content under grazing (Soussana et al., 2006). Even if mowing leads to some biomass input in 351 the form of plant material lost during grass removal (Sanaullah et al., 2010), the amount is 352 not enough to reach a similar input level than under grazing. Additionally, the lower pH 353 under mowing could contribute to indirect losses of SOC via changing C cycle and microbial 354 functioning (Kemmitt et al., 2006). Consequently, our results suggest that temperate loamy 355 soil under grazing is more prone to higher SOC contents when compared to mowing.

356 With regards to the biogeochemical composition of SOC, we did not find any 357 differences in non-cellulosic polysaccharide concentrations. These results are in agreement 358 with other studies showing that the soils' polysaccharide content is more or less stable and 359 even plant removal does not have a strong effect on the total polysaccharide concentrations 360 (Marchus et al., 2018). Soil lignin content, in contrast, was lower under grazing than mowing. 361 As lignin is a biomarker for plant-derived organic matter and more difficult to decompose, 362 because it requires a specific enzyme system (Buswell et al., 1987; Thevenot et al., 2010), 363 lower exportation of plant biomass and lignin input via dung deposition in soil under grazing 364 would suggest the opposite trend. However, dung contains only small amounts of lignin 365 (Dungait et al., 2005), which is relatively instable being degraded during one year (Dungait et 366 al., 2008). All lignin parameters (except the C/V ratio) suggested that lignin was less 367 degraded in the grazing than the mowing system. More acid pH in fertilised mowing systems 368 could have favoured the activity of lignin-degrading fungi (Couto et al., 2006). In mowing 369 systems microbial activity is fuelled exclusively by plant litter, whereas in grazing systems 370 organic matter input is supplied also by animal depositions. We hypothesise that this could 371 lead to contrasting quantitative lignin inputs, but could also impact its decomposition. Our 372 data show that lignin degradation in the mowing system is slower and less complete than in 373 the grazing system, leading to accumulation of partially degraded lignin molecules (Filley et 374 al., 2006). Therefore, lignin in the mowing system was characterised by a higher state of 375 degradation and at the same time its contribution to SOC was higher as compared to the 376 grazing system.

377

4.2. Effect of grazing treatment on biological properties of surface soil

378 Higher maturity and sustainability of the grazing system was shown by higher MBC per SOC together with a lower qCO₂ (Anderson and Domsch, 2010). Higher qCO₂ in the mowing 379 380 system indicates that the microbial communities were less efficient and respired more C to 381 maintain metabolic activity as compared to those under grazing (Anderson, 2003). 382 Microorganisms are the main SOM decomposers leading to release of greenhouse gases and 383 nutrients in natural as well as in managed soils (Bardgett et al., 2008; Gougoulias et al., 384 2014). This is particularly relevant for grazed pastures. Higher soil microbial respiration and 385 microbial CO₂ –C per unit SOC (soil microbial respiration per SOC) in the grazing system 386 was probably related to dung input with a huge amount of easily available compounds (Chu 387 et al., 2007; Marinari et al., 2000).

388 Contrary to our expectations, absolute enzyme activity did not differ among the 389 treatments, even after normalisation by SOC. A treatment effect was only observed after 390 normalisation by MBC, which expresses microbial activity in terms of enzyme production. 391 The enzymatic activities per MBC were higher in the mowing system as compared to the 392 grazing one, indicating that microorganisms in mowed soil produced enzymes more actively 393 than those under grazing. Microbial communities in the mowing system stayed active and 394 were investing in enzyme production probably to adapt to less decomposable organic 395 materials with higher lignin contents (see above). This maintenance of active state requires a 396 lot of energy, consequently, it could change C-cycling rates and decomposition of SOM 397 (Schimel and Schaeffer, 2012; Wang et al., 2014).

398 Microbial communities in the mowed soil are probably characterised by a higher 399 contribution of fungi than those of the grazed soil because we recorded a higher C:N ratio of 400 the microbial biomass (Joergensen and Emmerling, 2006) and more acid pH. Lower specific 401 growth rates in the mowing system may indicate relative domination of K-strategists in the 402 microbial community, which are more adapted to nutrient poor conditions (Strickland and 403 Rousk, 2010; Xu et al., 2017) and the decomposition of specific substances, such as plant 404 material containing high amounts of biopolymers (Fontaine et al., 2003). As illustrated by 405 lower enzyme activity per MBC, microorganisms in the grazing system invested less energy 406 for the degradation of complex compounds than those of the mowing system, most probably 407 because of higher availability of easily decomposable substrates. These conditions favour r-408 strategists (Fierer et al., 2007; Xu et al., 2017) and thus stimulate microbial activity, as shown 409 by higher MBC per SOC and higher soil microbial respiration under the grazing as compared 410 to mowing system. As a consequence, the biogeochemical soil functioning under the two 411 management practices is quite different. This may affect significantly SOM formation, which 412 is favoured in systems with intensive microbial processing of C input (Kallenbach et al., 413 2016; Liang et al., 2017) thus corroborating the high SOC contents observed under grazing.

414 *4.3.* Less pronounced treatment effects in subsurface soil

415 Treatment effects on soil properties were less pronounced in subsurface soil compared 416 to surface soil. Enhanced leaching and activity of soil fauna (Bohlen et al., 2004; Rumpel and 417 Kögel-Knabner, 2011) promote nutrient transport to subsurface soil under grazing which 418 resulted in higher SOC and N contents in subsurface soil under grazing than the one under 419 mowing. Treatment effects in the subsurface soil were neither observed for non-cellulosic 420 polysaccharide content and origin nor for lignin content or its degradation status. Since 421 lignins are typical indicators of plant input (Kögel-Knabner, 2002), this could indicate that 422 grazing and mowing have only small effects on plant rooting behaviour at lower depths.

423 On the other hand, the treatment effects on MBC and MBN was also observable in 424 subsurface soil. Soil microbial respiration did not differ between the treatments but microbial 425 CO₂ –C per SOC and qCO₂ were higher in the subsurface soil under mowing indicating that 426 the microbial communities used C inefficiently, similarly to surface soil. Higher 427 galactosidase activity in the subsurface soil of the mowing treatment is related to higher 428 contribution of galactose monomers in grass roots compared to grass leaves (Schädel et al., 429 2010). As lipase is hydrolysing triglycerides, higher lipase activity in the subsurface soil 430 indicates accumulation of lipid compounds at depth, which probably serve as C source for 431 microorganisms under C-limiting conditions (Heitkötter et al., 2017).

The absence of treatment separation for the subsurface soils on the PCA plot might indicate that in deeper soil probably more time is required to make treatment effects observable. It was interesting to note that chemical properties related to SOM composition were not sensitive to treatment effects in the subsurface soil, whereas microbial properties were. This is in agreement with other studies, which showed that microbial properties are most sensitive to changes introduced by management activities (Allison and Martiny, 2008; Bending et al., 2004).

440 **5.** Conclusions

441 In this study we investigated the effect of grazing and mowing treatments on soil 442 biogeochemical and microbial properties. Our data indicated significant differences in the soil 443 organic matter composition as well as microbial functioning of the two treatments. Both plots 444 were also characterized by contrasting SOC contents and pH values. The grazing system was 445 characterized by (1) more efficient microbial community and (2) less decomposed organic 446 matter as compared to the mowing system. We conclude that the harvesting regime by 447 grazing or mowing affects the biogeochemical functioning of grassland soils. Even though 448 both systems are favorable to SOC storage, grazing might be preferable to mowing because it 449 leads to better substrate quality and more efficient microbial functioning. Although SOM 450 changes were only evident in surface soil, microbial properties suggest that these processes 451 are also occurring in subsurface soil.

452

453 **6.** Conflicts of interests

454 We state that there is no conflict of interests.

455

456 **7.** Acknowledgements

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467 **8. References**

- Abdalla, M., Hastings, A., Chadwick, D.R., Jones, D.L., Evans, C.D., Jones, M.B., Rees,
 R.M., Smith, P., 2018. Critical review of the impacts of grazing intensity on soil organic
 carbon storage and other soil quality indicators in extensively managed grasslands.
 Agric. Ecosyst. Environ. 253, 62–81. https://doi.org/10.1016/j.agee.2017.10.023
- Allison, S.D., Martiny, J.B.H., 2008. In the Light of Evolution, In the Light of Evolution.
 National Academies Press, Washington, D.C. https://doi.org/10.17226/12501
- Anderson, T.-H., 2003. Microbial eco-physiological indicators to asses soil quality. Agric.
 Ecosyst. Environ. 98, 285–293. https://doi.org/10.1016/S0167-8809(03)00088-4
- 476 Anderson, T.-H., Domsch, K.H., 2010. Soil microbial biomass: The eco-physiological
 477 approach. Soil Biol. Biochem. 42, 2039–2043.
 478 https://doi.org/10.1016/j.soilbio.2010.06.026
- Anderson, T.-H., Domsch, K.H., 1993. The metabolic quotient for CO2 (qCO2) as a specific
 activity parameter to assess the effects of environmental conditions, such as ph, on the
 microbial biomass of forest soils, Soil Biology and Biochemistry. Pergamon.
 https://doi.org/10.1016/0038-0717(93)90140-7
- Bardgett, R.D., Freeman, C., Ostle, N.J., 2008. Microbial contributions to climate change
 through carbon cycle feedbacks. ISME J. 2, 805–814.
 https://doi.org/10.1038/ismej.2008.58
- Bazot, S., Mikola, J., Nguyen, C., Robin, C., 2005. Defoliation-induced changes in carbon
 allocation and root soluble carbon concentration in field-grown Lolium perenne plants:
 Do they affect carbon availability, microbes and animal trophic groups in soil? Funct.
- 489 Ecol. 19, 886–896. https://doi.org/10.1111/j.1365-2435.2005.01037.x
- Bending, G.D., Turner, M.K., Rayns, F., Marx, M.-C., Wood, M., 2004. Microbial and
 biochemical soil quality indicators and their potential for differentiating areas under
 contrasting agricultural management regimes. Soil Biol. Biochem. 36, 1785–1792.
 https://doi.org/10.1016/J.SOILBIO.2004.04.035
- 494Blagodatskaya, E., Kuzyakov, Y., 2013. Active microorganisms in soil: Critical review of495estimationcriteriaandapproaches.SoilBiol.Biochem.

- 496 https://doi.org/10.1016/j.soilbio.2013.08.024
- Bohlen, P.J., Pelletier, D.M., Groffman, P.M., Fahey, T.J., Fisk, M.C., 2004. Influence of
 Earthworm Invasion on Redistribution and Retention of Soil Carbon and Nitrogen in
 Northern Temperate Forests. Ecosystems 7, 13–27. https://doi.org/10.1007/s10021-0030127-y
- 501 Buswell, J.A., Odier, E., Kirk, T.K., 1987. Lignin Biodegradation. Crit. Rev. Biotechnol. 6,
 502 1–60. https://doi.org/10.3109/07388558709086984
- 503 Chabbi, A., Kögel-Knabner, I., Rumpel, C., 2009. Stabilised carbon in subsoil horizons is
 504 located in spatially distinct parts of the soil profile. Soil Biol. Biochem. 41, 256–261.
 505 https://doi.org/10.1016/j.soilbio.2008.10.033
- 506 Chu, H., Lin, X., Fujii, T., Morimoto, S., Yagi, K., Hu, J., Zhang, J., 2007. Soil microbial
 507 biomass, dehydrogenase activity, bacterial community structure in response to long-term
 508 fertilizer management. Soil Biol. Biochem. 39, 2971–2976.
 509 https://doi.org/10.1016/J.SOILBIO.2007.05.031
- Cotrufo, M.F., Soong, J.L., Horton, A.J., Campbell, E.E., Haddix, M.L., Wall, D.H., Parton,
 W.J., 2015. Formation of soil organic matter via biochemical and physical pathways of
 litter mass loss. Nat. Geosci. 8, 776–779. https://doi.org/10.1038/ngeo2520
- Couto, S.R., Moldes, D., Sanromán, M.A., 2006. Optimum stability conditions of pH and
 temperature for ligninase and manganese-dependent peroxidase from Phanerochaete
 chrysosporium. Application to in vitro decolorization of Poly R-478 by MnP. World J.
 Microbiol. Biotechnol. 22, 607–612. https://doi.org/10.1007/s11274-005-9078-0
- 517 Dignac, M.-F., Derrien, D., Barré, P., Barot, S., Cécillon, L., Chenu, C., Chevallier, T.,
 518 Freschet, G.T., Garnier, P., Guenet, B., Hedde, M., Klumpp, K., Lashermes, G., Maron,
 519 P.-A., Nunan, N., Roumet, C., Basile-Doelsch, I., 2017. Increasing soil carbon storage:
 520 mechanisms, effects of agricultural practices and proxies. A review. Agron. Sustain.
 521 Dev. 37, 14. https://doi.org/10.1007/s13593-017-0421-2
- Dungait, J.A.J., Bol, R., Bull, I.D., Evershed, R.P., 2009. Tracking the fate of dung-derived
 carbohydrates in a temperate grassland soil using compound-specific stable isotope
 analysis. Org. Geochem. 40, 1210–1218.
 https://doi.org/10.1016/j.orggeochem.2009.08.001
- Dungait, J.A.J., Bol, R., Evershed, R.P., 2005. Quantification of dung carbon incorporation in
 a temperate grassland soil following spring application using bulk stable carbon isotope
 determinations. Isotopes Environ. Health Stud. 41, 3–11.
 https://doi.org/10.1080/10256010500053516

- Dungait, J.A.J., Stear, N.A., van Dongen, B.E., Bol, R., Evershed, R.P., 2008. Off-line
 pyrolysis and compound-specific stable carbon isotope analysis of lignin moieties: a
 new method for determining the fate of lignin residues in soil. Rapid Commun. Mass
 Spectrom. 22, 1631–1639. https://doi.org/10.1002/rcm.3454
- Ec.europa.eu, 2018. Share of main land types in utilised agricultural area (UAA) by NUTS 2
 regions [WWW Document]. URL https://ec.europa.eu/eurostat/data/database (accessed
 1.23.19).
- Eder, E., Spielvogel, S., Kölbl, A., Albert, G., Kögel-Knabner, I., 2010. Analysis of
 hydrolysable neutral sugars in mineral soils: Improvement of alditol acetylation for gas
 chromatographic separation and measurement. Org. Geochem. 41, 580–585.
 https://doi.org/10.1016/J.ORGGEOCHEM.2010.02.009
- 541 Eivazi, F., Tabatabai, M.A., 1990. Factors affecting glucosidase and galactosidase activities
 542 in soils. Soil Biol. Biochem. 22, 891–897. https://doi.org/10.1016/0038-0717(90)90126543 K
- 544 Fierer, N., Bradford, M.A., Jackson, R.B., 2007. Toward an ecological classification of soil
 545 bacteria. Ecology 88, 1354–1364. https://doi.org/10.1890/05-1839
- Filley, T.R., Nierop, K.G.J., Wang, Y., 2006. The contribution of polyhydroxyl aromatic
 compounds to tetramethylammonium hydroxide lignin-based proxies. Org. Geochem.
 37, 711–727. https://doi.org/10.1016/J.ORGGEOCHEM.2006.01.005
- Fujisaki, K., Chevallier, T., Chapuis-Lardy, L., Albrecht, A., Razafimbelo, T., Masse, D.,
 Ndour, Y.B., Chotte, J.-L., 2018. Soil carbon stock changes in tropical croplands are
 mainly driven by carbon inputs: A synthesis. Agric. Ecosyst. Environ. 259, 147–158.
 https://doi.org/10.1016/j.agee.2017.12.008
- Gavrichkova, O., Moscatelli, M.C., Kuzyakov, Y., Grego, S., Valentini, R., 2010. Influence
 of defoliation on CO2efflux from soil and microbial activity in a Mediterranean
 grassland. Agric. Ecosyst. Environ. 136, 87–96.
 https://doi.org/10.1016/j.agee.2009.11.015
- Gougoulias, C., Clark, J.M., Shaw, L.J., 2014. The role of soil microbes in the global carbon
 cycle: tracking the below-ground microbial processing of plant-derived carbon for
 manipulating carbon dynamics in agricultural systems. J. Sci. Food Agric. 94, 2362–
 2371. https://doi.org/10.1002/jsfa.6577
- Havstad, K.M., Peters, D.P.C., Skaggs, R., Brown, J., Bestelmeyer, B., Fredrickson, E.,
 Herrick, J., Wright, J., 2007. Ecological services to and from rangelands of the United
 States. Ecol. Econ. 64, 261–268. https://doi.org/10.1016/J.ECOLECON.2007.08.005

- Hedges, J.I., Ertel, J.R., 1982. Characterization of lignin by gas capillary chromatography of
 cupric oxide oxidation products. Anal. Chem. 54, 174–178.
 https://doi.org/10.1021/ac00239a007
- Heitkötter, J., Heinze, S., Marschner, B., 2017. Relevance of substrate quality and nutrients
 for microbial C-turnover in top- and subsoil of a Dystric Cambisol. Geoderma 302, 89–
 99. https://doi.org/10.1016/j.geoderma.2017.04.029
- 570 IPCC, 2018. Special Report. Global Warming of 1.5 °C. https://www.ipcc.ch/sr15/.
- Jenkinson, D.S., Brookes, P.C., Powlson, D.S., 2004. Measuring soil microbial biomass. Soil
 Biol. Biochem. 36, 5–7. https://doi.org/10.1016/J.SOILBIO.2003.10.002
- Joergensen, R.G., Emmerling, C., 2006. Methods for evaluating human impact on soil
 microorganisms based on their activity, biomass, and diversity in agricultural soils. J.
 Plant Nutr. Soil Sci. 169, 295–309. https://doi.org/10.1002/jpln.200521941
- Kallenbach, C.M., Frey, S.D., Grandy, A.S., 2016. Direct evidence for microbial-derived soil
 organic matter formation and its ecophysiological controls. Nat. Commun. 7, 1–10.
 https://doi.org/10.1038/ncomms13630
- Kemmitt, S.J., Wright, D., Goulding, K.W.T., Jones, D.L., 2006. pH regulation of carbon and
 nitrogen dynamics in two agricultural soils. Soil Biol. Biochem. 38, 898–911.
 https://doi.org/10.1016/j.soilbio.2005.08.006
- Koch, O., Tscherko, D., Kandeler, E., 2007. Temperature sensitivity of microbial respiration,
 nitrogen mineralization, and potential soil enzyme activities in organic alpine soils.
 Global Biogeochem. Cycles 21. https://doi.org/10.1029/2007GB002983
- 585 Kögel-Knabner, I., 2002. The macromolecular organic composition of plant and microbial
 586 residues as inputs to soil organic matter. Soil Biol. Biochem. 34, 139–162.
 587 https://doi.org/10.1016/S0038-0717(01)00158-4
- Kögel, I., Bochter, R., 1985. Characterization of lignin in forest humus layers by highperformance liquid chromatography of cupric oxide oxidation products. Soil Biol.
 Biochem. 17, 637–640. https://doi.org/10.1016/0038-0717(85)90040-9
- Liang, C., Schimel, J.P., Jastrow, J.D., 2017. The importance of anabolism in microbial
 control over soil carbon storage. Nat. Microbiol. 2, 17105.
 https://doi.org/10.1038/nmicrobiol.2017.105
- Marchus, K.A., Blankinship, J.C., Schimel, J.P., 2018. Environmental controls on
 extracellular polysaccharide accumulation in a California grassland soil. Soil Biol.
 Biochem. https://doi.org/10.1016/j.soilbio.2018.07.009
- 597 Marinari, S., Masciandaro, G., Ceccanti, B., Grego, S., 2000. Influence of organic and

- 598 mineral fertilisers on soil biological and physical properties. Bioresour. Technol. 72, 9–
 599 17. https://doi.org/10.1016/S0960-8524(99)00094-2
- Marx, M.-C., Kandeler, E., Wood, M., Wermbter, N., Jarvis, S.C., 2005. Exploring the
 enzymatic landscape: distribution and kinetics of hydrolytic enzymes in soil particle-size
 fractions. Soil Biol. Biochem. 37, 35–48.
 https://doi.org/10.1016/J.SOILBIO.2004.05.024
- McSherry, M.E., Ritchie, M.E., 2013. Effects of grazing on grassland soil carbon: a global
 review. Glob. Chang. Biol. 19, 1347–1357. https://doi.org/10.1111/gcb.12144
- Medina-Roldán, E., Bardgett, R.D., 2011. Plant and soil responses to defoliation: A
 comparative study of grass species with contrasting life history strategies. Plant Soil
 344, 377–388. https://doi.org/10.1007/s11104-011-0756-4
- Ngo, P.T., Rumpel, C., Dignac, M.-F., Billou, D., Duc, T.T., Jouquet, P., 2011.
 Transformation of buffalo manure by composting or vermicomposting to rehabilitate
 degraded tropical soils. Ecol. Eng. 37, 269–276.
 https://doi.org/10.1016/j.ecoleng.2010.11.011
- Panikov, N.S., Sizova, M. V., 1996. A kinetic method for estimating the biomass of microbial
 functional groups in soil. J. Microbiol. Methods 24, 219–230.
 https://doi.org/10.1016/0167-7012(95)00074-7
- Piñeiro, G., Paruelo, J.M., Oesterheld, M., Jobbágy, E.G., 2010. Pathways of Grazing Effects
 on Soil Organic Carbon and Nitrogen. Rangel. Ecol. Manag. 63, 109–119.
 https://doi.org/10.2111/08-255.1
- Post, W.M., Kwon, K.C., 2000. Soil carbon sequestration and land-use change: processes and
 potential. Glob. Chang. Biol. 6, 317–327. https://doi.org/10.1046/j.13652486.2000.00308.x
- Puche, N.J.B., Senapati, N., Klumpp, K., Fléchard, C.R., Kirschbaum, M.U.F., Chabbi, A.,
 2019. Modelling carbon and water fluxes of managed grasslands: comparing flux
 variability and net carbon budgets between grazed and mowed systems. Agronomy.
- Razavi, B.S., Blagodatskaya, E., Kuzyakov, Y., 2015. Nonlinear temperature sensitivity of
 enzyme kinetics explains canceling effect—a case study on loamy haplic Luvisol. Front.
 Microbiol. 6. https://doi.org/10.3389/fmicb.2015.01126
- Rumpel, C., Crème, A., Ngo, P., Velásquez, G., Mora, M., Chabbi, A., 2015. The impact of
 grassland management on biogeochemical cycles involving carbon, nitrogen and
 phosphorus. J. soil Sci. plant Nutr. 15(2), 353–371. https://doi.org/10.4067/S071895162015005000034

- Rumpel, C., Dignac, M.-F., 2006. Gas chromatographic analysis of monosaccharides in a
 forest soil profile: Analysis by gas chromatography after trifluoroacetic acid hydrolysis
 and reduction–acetylation. Soil Biol. Biochem. 38, 1478–1481.
 https://doi.org/10.1016/J.SOILBIO.2005.09.017
- Rumpel, C., Kögel-Knabner, I., 2011. Deep soil organic matter---a key but poorly understood
 component of terrestrial C cycle. Plant Soil 338, 143–158.
 https://doi.org/10.1007/s11104-010-0391-5
- Rumpel, C., Lehmann, J., Chabbi, A., 2018. '4 per 1,000' initiative will boost soil carbon for
 climate and food security. Nat. 2020 5537686.
- Sanaullah, M., Chabbi, A., Lemaire, G., Charrier, X., Rumpel, C., 2010. How does plant leaf
 senescence of grassland species influence decomposition kinetics and litter compounds
 dynamics? Nutr. Cycl. Agroecosystems 88, 159–171. https://doi.org/10.1007/s10705009-9323-2
- Schädel, C., Blöchl, A., Richter, A., Hoch, G., 2010. Quantification and monosaccharide
 composition of hemicelluloses from different plant functional types. Plant Physiol.
 Biochem. 48, 1–8. https://doi.org/10.1016/J.PLAPHY.2009.09.008
- 648 Schimel, J.P., Schaeffer, S.M., 2012. Microbial control over carbon cycling in soil. Front.
 649 Microbiol. https://doi.org/10.3389/fmicb.2012.00348
- Senapati, N., Chabbi, A., Gastal, F., Smith, P., Mascher, N., Loubet, B., Cellier, P., Naisse,
 C., 2014. Net carbon storage measured in a mowed and grazed temperate sown
 grassland shows potential for carbon sequestration under grazed system. Carbon Manag.

653 5, 131–144. https://doi.org/10.1080/17583004.2014.912863

- Senapati, N., Chabbi, A., Giostri, A.F., Yeluripati, J.B., Smith, P., 2016. Modelling nitrous
 oxide emissions from mown-grass and grain-cropping systems: Testing and sensitivity
 analysis of DailyDayCent using high frequency measurements. Sci. Total Environ. 572,
 955–977. https://doi.org/10.1016/j.scitotenv.2016.07.226
- Smith, P., Martino, D., Cai, Z., Gwary, D., Janzen, H., Kumar, P., McCarl, B., Ogle, S.,
 O'Mara, F., Rice, C., Scholes, B., Sirotenko, O., Howden, M., McAllister, T., Pan, G.,
 Romanenkov, V., Schneider, U., Towprayoon, S., Wattenbach, M., Smith, J., 2008.
 Greenhouse gas mitigation in agriculture. Philos. Trans. R. Soc. B Biol. Sci. 363, 789–
 813. https://doi.org/10.1098/rstb.2007.2184
- Soussana, J.-F., Loiseau, P., Vuichard, N., Ceschia, E., Balesdent, J., Chevallier, T.,
 Arrouays, D., 2006. Carbon cycling and sequestration opportunities in temperate
 grasslands. Soil Use Manag. 20, 219–230. https://doi.org/10.1111/j.1475-

666 2743.2004.tb00362.x

- 667 Strickland, M.S., Rousk, J., 2010. Considering fungal: Bacterial dominance in soils 668 Methods, controls, and ecosystem implications. Soil Biol. Biochem. 42, 1385–1395.
 669 https://doi.org/10.1016/j.soilbio.2010.05.007
- Tautges, N.E., Chiartas, J.L., Gaudin, A.C.M., O'Geen, A.T., Herrera, I., Scow, K.M., 2019.
 Deep soil inventories reveal that impacts of cover crops and compost on soil carbon
 sequestration differ in surface and subsurface soils. Glob. Chang. Biol. 25, 3753–3766.
 https://doi.org/10.1111/gcb.14762
- Thevenot, M., Dignac, M.-F.F., Rumpel, C., 2010. Fate of lignins in soils: A review. Soil
 Biol. Biochem. 42, 1200–1211. https://doi.org/10.1016/J.SOILBIO.2010.03.017
- Vance, E.D., Brookes, P.C., Jenkinson, D.S., 1987. An extraction method for measuring soil
 microbial biomass C. Soil Biol. Biochem. 19, 703–707. https://doi.org/10.1016/00380717(87)90052-6
- Wang, G., Mayes, M.A., Gu, L., Schadt, C.W., 2014. Representation of dormant and active
 microbial dynamics for ecosystem modeling. PLoS One 9.
 https://doi.org/10.1371/journal.pone.0089252
- Wilson, C.H., Strickland, M.S., Hutchings, J.A., Bianchi, T.S., Flory, S.L., 2018. Grazing
 enhances belowground carbon allocation, microbial biomass, and soil carbon in a
 subtropical grassland. Glob. Chang. Biol. 24, 2997–3009.
 https://doi.org/10.1111/gcb.14070
- Wutzler, T., Blagodatsky, S.A., Blagodatskaya, E., Kuzyakov, Y., 2012. Soil microbial
 biomass and its activity estimated by kinetic respiration analysis Statistical guidelines.
 Soil Biol. Biochem. 45, 102–112. https://doi.org/10.1016/j.soilbio.2011.10.004
- Ku, S., Silveira, M.L., Inglett, K.S., Sollenberger, L.E., Gerber, S., 2017. Soil microbial
 community responses to long-term land use intensification in subtropical grazing lands.
- 691 Geoderma 293, 73–81. https://doi.org/10.1016/j.geoderma.2017.01.019

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Tables 695

696 Table 1. General soil properties under two grassland management practices (grazing and

mowing) at in surface soil (0-10 cm) and subsurface soil (20-30 cm). 697

	Treatment	pН	SOC content	Ν	δ ¹³ C	$\delta^{15}N$	C:N ratio
			mg g ⁻¹	mg g ⁻¹	%0	%0	
G . A	Grazing	5.95±0.09	21.4±0.81	2.2±0.09	-27.4±0.06	4.9±0.13	9.6±0.05
Surface soil	Mowing	5.51±0.08	14.6±0.51	1.5±0.05	-27.0±0.05	5.0±0.09	9.6±0.07
	Grazing	5.99±0.12	11.8±0.62	1.3±0.06	-26.7±0.06	6.2±0.08	9.2±0.06
Subsurface soil	Mowing	6.01±0.13	8.6±0.44	0.9±0.05	-26.3±0.10	6.4±0.10	9.1±0.07
ANCOVA, F value (P v SOC stocks Treatment Depth Treatment×l	y alues) in 2005 Depth	0.15 (0.70) 3.37 (0.08) 5.89 (0.02) 4.27 (0.04)	5.31 (0.03) 30.3 (<0.001) 181.8 (<0.001) 0.68 (0.41)	6.26 (0.02) 28.7 (<0.001) 153.8 (<0.001) 0.86 (0.36)	6.86 (0.01) 17.2 (0.002) 132.5 (<0.001) 0.28 (0.06)	1.90 (0.18) 0.35 (0.56) 157.9 (<0.001) 0.25 (0.62)	1.33 (0.26) 1.19 (0.28) 52.5 (<0.001) 0.37 (0.55)

698 699 Values are shown as the average of ten replicates and ±SE. Significant differences between the treatments are

indicated by capital case letters. Lower case letters show significant differences with depth (P < 0.05).

Table 2. Non-cellulosic polysaccharides (NCP) signature in soil under two grassland

	Treatment	NCP content	NCP content per	NCP monomers ratios			
			SUC	C6/C5 ¹	DesoxyC6/C5 ²	Man/Xyl ³	
		mg g ⁻¹	mg g ⁻¹ SOC				
Surface soil	Grazing	6.61±0.23	308.98±6.3	0.80±0.02	0.35±0.01	0.54±0.02	
5011	Mowing	4.45±0.18	306.63±11.5	0.84±0.02	0.34±0.01	0.61±0.02	
Subsurface	Grazing	3.09±0.15	263.39±6.4	1.03±0.02	0.43±0.01	0.87±0.03	
soil	Mowing	2.50±0.11	292.41±10.5	1.01±0.02	0.46±0.01	0.91±0.03	
ANCOVA, F value (P va	alues)	0.43 (0.52)					
SOC stocks in 2005		36.6 (<0.001)	2.50 (0.12)	5.42 (0.03)	3.81 (0.06)	4.74 (0.04)	
Treatment		241.1	0.11 (0.74)	0.87 (0.36)	0.01 (0.91)	0.64 (0.43)	
Depth		(<0.001)	11.5 (0.002)	122.2 (<0.001)	102.8 (<0.001)	166.3 (<0.001)	
Treatment×Depth		19.7 (<0.001)	3.17 (0.08)	3.14 (0.09)	3.18 (0.08)	0.52 (0.48)	

management practices (grazing and mowing) at two depths (0-10 cm and 20-30 cm).

704 705 $^{1}C6/C5$ – the ratio of C6- to C5- sugar monomers, $^{2}DesoxyC6/C5$ – the ratio of desoxy C6- to desoxy C5- sugar monomers, $^{3}Man/Xyl$ - the ratio of mannose to xylose. These ratios indicate the origin of non-cellulosic polysaccharides (microbial or plant).

709 Table 3. Lignin signature in soil under two grassland management practices (grazing and

	Treatment	Lignin content	Lignin content	Lignin monomers ratios			
			per SOC	C/V	S/V	(Ac/Al)v	(Ac/Al)s
		mg g ⁻¹	mg g ⁻¹ SOC				
Surface soil	Grazing	0.35±0.01	16.31±0.64	0.45±0.03	1.34±0.02	0.53±0.02	0.46±0.01
C-1	Mowing	0.26±0.01	17.86±0.43	0.45±0.03	1.24±0.02	0.65±0.02	0.57±0.02
soil	Grazing	0.19±0.02	16.22±0.57	0.37±0.02	1.33±0.03	0.66±0.01	0.54±0.02
	Mowing	0.16±0.01	18.86±0.89	0.37±0.03	1.30±0.02	0.63±0.03	0.56±0.02
ANCOVA, F value (P va	alues)						
SOC stocks i	n 2005	0.05 (0.82)	10.0 (0.002)	1 50 (0 22)	1 00 (0 17)	0.09 (0.78)	18.6 (<0.001)
Treatment		15.3(<0.02)	10.9(0.002) 1 14 (0 29)	0.64(0.22)	7.99 (0.17)	2.08(0.78) 2.83(0.10)	2.02 (0.10
Depth Treatment×Depth		96.3 (<0.001) 4.83 (0.04)	0.62(0.44) 0.88(0.36)	8.58 (0.006) 0.015 (0.90)	0.95 (0.34) 2.19 (0.15)	9.13 (0.005) 16.3 (<0.001)	13.1 (<0.001)

710 mowing) in surface soil (0-10 cm) and subsurface soil (20-30 cm).

711 712 C/V - the ratio of cinnamyl phenols to syringyl phenols; S/V - the ratio of syringyl phenols to vanillyl

phenols; (Ac/Al)V - acid to aldehyde ratio of vanillyl phenols; (Ac/Al)s - acid to aldehyde ratio of syringyl 713 phenols. These ratios are indicators of lignin degradation state in soil.

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Figure 1. (A) Soil microbial respiration (SMR), (B) soil microbial respiration (SMR) per soil organic carbon (SOC), (C) microbial biomass carbon (MBC) per soil organic carbon (SOC) and (D) metabolic quotient (qCO₂) in soil under two grassland management practices (grazing and mowing) in surface soil (0-10 cm) and subsurface soil (20-30 cm). Significant differences between the treatments are indicated by *, ** and ***, representing probability at the 5, 1, and 0.1% levels, respectively.





Figure 2. (A) Microbial C:N ratio, (B) the percentage of active microbial biomass (AMB) and (C) specific microbial growth rate (μ) in soil under two grassland management practices (grazing and mowing) in surface soil (0-10 cm) and subsurface soil (20-30 cm). Significant differences between the treatments are indicated by *, ** and ***, representing probability at the 5, 1, and 0.1% levels, respectively.

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Figure 3. Boxplot of enzyme activity per unit of microbial biomass C (MBC) for nine enzymes under two grassland management practices (grazing and mowing) in surface soil (0-10 cm) and subsurface soil (20-30 cm). Significant differences between the treatments are indicated by *, ** and ***, representing probability at the 5, 1, and 0.1% levels, respectively.



Figure 4. Principal component analysis (PCA) for soil under grazing and mowing in surface soil (0-10 cm) and in subsurface soil (20-30 cm). Only variables with quality of representation (cos²) higher than 0.75 was shown on PCA plot.