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Enhanced priming of old, not new soil carbon at elevated atmospheric CO₂

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23 Abstract

Rising atmospheric CO₂ concentrations accompanied by global warming and altered precipitation 24 patterns calls for assessment of long-term effects of these global changes on carbon (C) dynamics in 25 26 terrestrial ecosystems, as changes in net C exchange between soil and atmosphere will impact the atmospheric CO₂ concentration profoundly. In many ecosystems, including the heath/grassland 27 system studied here, increased plant production at elevated CO₂ increase fresh C input from litter 28 and root exudates to the soil and concurrently decrease soil N availability. Supply of labile C to the 29 soil may accelerate the decomposition of soil organic C (SOC), a phenomenon termed 'the priming 30 effect', and the priming effect is most pronounced at low soil N availability. Hence, we 31 hypothesized that priming of SOC decomposition in response to labile C addition would increase in 32 soil exposed to long-term elevated CO₂ exposure. Further, we hypothesized that long-term warming 33 would enhance SOC priming rates, whereas drought would decrease the priming response. 34 We incubated soil from a long-term, full-factorial climate change field experiment, with the factors 35 elevated atmospheric CO₂ concentration, warming and prolonged summer drought with either labile 36 C (sucrose) or water to assess the impact of labile C on SOC dynamics. We used sucrose with a 37 $^{13}C/^{12}C$ signature that is distinct from that of the native SOC, which allowed us to assess the 38 contribution of these two C sources to the CO₂ evolved. Sucrose induced priming of SOC, and the 39 priming response was higher in soil exposed to long-term elevated CO2 treatment. Drought tended 40 to decrease the priming response, whereas long-term warming did not affect the level of priming 41 significantly. 42

We were also able to assess whether SOC-derived primed C in elevated CO₂ soil was assimilated
before or after the initiation of the CO₂ treatment 8 years prior to sampling, because CO₂
concentrations were raised by fumigating the experimental plots with pure CO₂ that was ¹³Cdepleted compared to ambient CO₂. Surprisingly, we conclude that sucrose addition primed

decomposition of relatively old SOC fractions, i.e. SOC assimilated more than 8 years beforesampling.

49

50 **1. Introduction**

The global terrestrial soil organic carbon (SOC) pool is the largest terrestrial carbon (C) pool and constitutes a C stock that is more than twice the size of the atmospheric CO_2 -C pool (IPCC, 2013). Therefore, even relatively moderate fluctuations in net C exchange between soil and atmosphere will impact the CO_2 concentration in the atmosphere profoundly. Faced by rising atmospheric CO_2 levels and the anticipated climatic changes that will result from this rise, we need to better understand how such changes will influence SOC decomposition and CO_2 -release from terrestrial organic C pools.

At least two factors that can potentially alter SOC decomposition, namely nitrogen (N) availability 58 and input of fresh plant C, are expected to change with rising CO₂ levels. Supply of fresh plant 59 derived C into the soil matrix may accelerate the decomposition of SOC and decrease soil C stocks 60 (Fontaine et al., 2004); a phenomenon termed 'the priming effect'. Priming effects induced by root 61 exudation and rhizodeposition can cause an up to 350 % increase in SOC decomposition compared 62 63 to the root free soil (Cheng et al., 2014). Even so, most C-cycling models do not consider the influence of priming and living roots on SOC decomposition rates (Cheng et al., 2014), perhaps as a 64 result of limited knowledge about underlying mechanisms and factors influencing the magnitude of 65 priming. However, the few attempts that have been made to date to represent plant-induced priming 66 of SOC decomposition have resulted in improved model performance, also regarding global change 67 effects (Cheng et al., 2014, Perveen et al., 2014). It is therefore evident that models predicting 68 future climatic conditions and atmospheric CO₂ levels, as well as our means of mitigating the 69 effects of rising CO₂ levels, depend on more in-depth understanding of feedbacks between climate, 70 elevated atmospheric CO₂ levels, and SOC decomposition caused by priming. 71

73	Common plant physiological responses to elevated CO ₂ comprise enhanced photoassimilation of C,
74	increased root volume, and increased input of plant C to the soil in the form of root exudates and
75	rhizodeposits (Hungate et al., 1997; Adair et al., 2009; Albert et al., 2011; Arndal et al., 2013,
76	2014). Since the extent of priming seems to depend on the concentration of labile C inputs, with no
77	or low priming at low concentrations (Blagodatskaya & Kuzyakov, 2008; Gude et al., 2012) and
78	gradually increasing priming with increasing concentrations (Blagodatskaya & Kuzyakov, 2008;
79	Paterson & Sim, 2013) up to a point of saturation (Guenet et al., 2010; Xiao et al., 2015), it can be
80	expected that priming will increase at elevated atmospheric CO ₂ concentration due to higher inputs
81	of labile plant C to soils (Paterson et al., 2008). Accordingly, elevated atmospheric CO ₂ has been
82	shown to increase decomposition of SOC in grasslands (Xie et al., 2005; Niklaus & Falloon, 2006)
83	as well as forests (Phillips et al., 2012). However, there are also examples where input of labile
84	plant C resulted in negative priming (Sullivan & Hart, 2013; Cheng et al., 2014), i.e. inhibited the
85	decomposition of SOC. A possible reason to the contradictory findings is that the magnitude and
86	direction of priming is dependent on the nutrient status of the soil. In fact, it has been suggested that
87	the decomposition of SOC in response to input of labile C is driven by enhanced microbial demand
88	for nutrients retained in soil organic matter (SOM) (Paterson, 2009; Philips et al., 2012).
89	Accordingly, Bengtson et al. (2012) demonstrated a strong link between rhizodeposition, SOM
90	decomposition and gross N mineralization in a coniferous forest soil, while Fontaine et al. (2004,
91	2011) found that soil C losses caused by priming increased when soil microbes are nutrient limited.
92	In line with these findings, a recent meta-analysis concluded that input of labile C enhanced
93	decomposition of native SOM, but only in soils with low nitrogen (N) content (Zhang et al., 2013).
94	Since it has been observed that increased plant N demand at elevated atmospheric CO ₂ commonly
95	leads to decreased soil N availability (Luo et al., 2004; Larsen et al., 2011), this can also explain
96	why priming can be expected to increase under elevated atmospheric CO ₂ conditions.

However, in order to fully appreciate how priming will influence the net ecosystem exchange of C 98 in a high CO₂ world we also need to consider climatic parameters, such as temperature and 99 precipitation patterns that are also undergoing changes that are expected to continue for the decades 100 to come (IPCC, 2013). In Northern Europe we expect an annual mean temperature increase between 101 0.75 and 0.1 °C over the coming 20 years and more extreme precipitation patterns, for instance 102 prolonged summer droughts according to the IPCC RCP4.5 scenario (IPCC, 2013). Hence, 103 104 evaluation of elevated CO₂ impacts on priming of SOC decomposition must consider projected temperature and precipitation scenarios. 105 Low soil moisture generally reduces soil microbial activity (Moyano et al., 2013) and may also 106 107 reduce rhizosphere priming of SOC decomposition (Dijkstra & Cheng, 2007). It can, therefore, be expected that prolonged summer droughts will decrease priming. However, as plants have better 108 water use efficiency at elevated CO₂ (Field et al., 1995; Ainsworth & Rogers, 2007), drought 109 impacts on microbial activity are in some cases less severe when combined with elevated CO₂ 110 (Kassem et al., 2008). The combined effect of elevated CO₂ and drought on the magnitude of 111 112 priming is to our knowledge not known. Likewise, little is known about temperature effects on priming. The only study to date that has 113 systematically addressed the temperature dependency of priming found the process to be non-114 responsive to temperature variations (Ghee et al., 2013). In general, even moderate warming 115 enhances the activity of heterotrophic microbial SOM decomposers (Wang et al., 2014). Therefore, 116 in systems with low N availability temperature dependent stimulation of microbial activity could 117 enhance the need for microbial N acquisition through SOM decomposition and increase priming. 118 However, the priming response to warming may very well depend on soil moisture conditions, since 119 warming enhances evaporation. This could potentially exacerbate negative effects of drought on 120 soil microbial activity. 121

123	Previous studies demonstrated that elevated CO ₂ changed C turnover dynamics of different fractions
124	of SOM. Elevated CO ₂ increased the content of recently assimilated C in both coarse and fine
125	particulate fractions of SOM, but decreased the content of older C in more physically protected, fine
126	particulate organic matter and mineral-associated organic matter (Hofmockel et al., 2011). This
127	suggests that elevated CO ₂ elicits priming of older relatively stable rather than recent SOC pools. In
128	the current experiment we are able to test this hypothesis, as we raised the atmospheric CO_2
129	concentration by fumigating with CO ₂ that was ¹³ C-depleted compared to the naturally occurring
130	atmospheric CO ₂ . Therefore, C fixed in elevated CO ₂ treatments was ¹³ C-depleted compared to the
131	C assimilated before CO ₂ treatment started and compared to the C pools of ambient CO ₂ treatments
132	(Reinsch & Ambus, 2013). A comparison of the isotopic composition of primed SOC-derived CO ₂ -
133	C from elevated CO ₂ and ambient CO ₂ treatments can therefore reveal if primed C derives from C
134	fixed before or after the initiation of CO ₂ fumigation.

135

The aim of this study was to test the effects of long-term elevated CO₂ exposure, warming and 136 annual extended drought events on potential priming in a nutrient-poor temperate heath/grassland, 137 where long-term elevated CO₂ exposure has reduced the relative N content of organic inputs 138 139 (Larsen et al., 2011; Arndal et al., 2013, 2014; Vestergård et al., 2015). Moderate warming has also prolonged the plant growth season with two weeks in the spring at the site (Kongstad et al., 2012). 140 If an extended growth period also enhances plant N uptake over the season, this could potentially 141 142 intensify microbial N demand. We hypothesize, in accordance with other reports (van Groenigen et al., 2005; Xie et al., 2005; Niklaus & Falloon, 2006), that potential priming of soil C is enhanced in 143 soil exposed to elevated CO₂, where plant production and hence C input to the soil is enhanced and 144 the relative N availability has declined. We further hypothesize that warming enhances priming in 145 soil exposed to elevated CO₂, because we expect that an earlier onset of spring growth and 146

enhanced microbial activity under warming further reduced N availability. We hypothesize that
summer drought, which is expected to reduce microbial activity, will reduce priming in soil exposed
to elevated CO₂. Finally, we expect that warming augments this effect of drought. Further, we will
clarify if primed soil C is recently fixed or of older origin.

151

Addition of labile carbohydrates is a common method to assess and compare potential priming activity between different soils and treatments (Wu et al., 1993; Zyakun & Dilly, 2005; Garcia-Pausas & Paterson, 2011; Paterson & Sim, 2013; Reinsch et al., 2013). If the ${}^{13}C/{}^{12}C$ signature of the added carbohydrate is distinct from that of native SOC it is possible to assess the contribution of these two C sources to the respiratory CO₂ evolved. In the present study we estimated potential priming by incubating soils with labile sucrose, a common constituent of root exudates (Grayston et al., 1998), with a ${}^{13}C/{}^{12}C$ ratio that is distinct from the isotopic ratio of the native SOC.

159

160 2. Materials and Methods

161

162 *2.1 Field site*

Our field experiment was carried out in an unmanaged temperate heath/grassland in North Zealand 163 c. 50 km northwest of Copenhagen, Denmark (55°53'N, 11°58' E). The soil is a Cambic Arenosol 164 (FAO classification) developed on a nutrient-poor sandy deposit. The organic layer is 5-10 cm thick 165 and the pH is around 5. From 1975-2005 the average annual precipitation was 610 mm and the 166 mean annual temperature was 8 °C (Danish Meteorological Institute). From 2005 to 2013 the mean 167 annual precipitation was 742 mm with a range between 648 and 894 mm and the mean annual 168 temperature was 9.7 °C with a range between 7 and 10 °C. The prevailing species are the grass 169 Deschampsia flexuosa (c. 70 % coverage) and the dwarf shrub Calluna vulgaris (c. 30 % coverage) 170 intermixed with other grasses, herbs, mosses and lichens (Kongstad et al., 2012). 171

173 2.2 Climate manipulation treatments

Climate and CO₂ manipulations, aimed to simulate climatic conditions and atmospheric CO₂ levels 174 that are predicted for Denmark in 2075, were initiated in October 2005. The global change factors 175 drought (D), warming (T) and CO₂ concentration (CO₂) were manipulated individually and in all 176 possible combinations. The CO₂ concentration was increased with 120 ppm based on the average of 177 predicted concentrations in 2075 in five atmospheric CO₂ stabilization scenarios (SP450, SP550, 178 SP650, SP750 and SP1000) (IPCC 2007). The temperature increase chosen was the average of 179 predicted temperature responses for Northern Europe (IPCC 2007), and drought manipulations were 180 also based on the predictions in the IPCC 2007 report. The CO₂ concentration was increased in-situ 181 182 via the free-air carbon dioxide enrichment (FACE) technique in octagon shaped plots (octagons) 183 during daytime hours. Extended spring/summer droughts were imposed using moveable curtains to exclude precipitation for a period of ~1 month during spring/early summer each year. Drought 184 curtains reduced precipitation by 7.6 ± 2.1 % (mean \pm SD) annually. In 2013, the drought period 185 was conducted between 29th of April and 27th of May. From 27th of May to sampling, June 5th-6th, 186 the site only received a few mm precipitation, effectively extending the drought period until 187 sampling. Passive night time warming was achieved via moveable curtains that covered the 188 experimental plots during night time hours and prevented heat loss to the atmosphere. The warming 189 effect at 20 cm above ground surface ranged between 0.5 °C and 1.5 °C over the year (Scherber et 190 al., 2013). Warming curtains were withdrawn during rainfall. 191

The experiment is a full-factorial split plot design organized in 6 blocks. One block contains two octagons each of 6.8 m diameter, one exposed to ambient CO_2 concentration and one exposed to elevated CO_2 concentration, respectively. Each octagon is divided into four plots, which amounts to a total of 48 plots. Within each octagon, one plot is subjected to the drought treatment, one is subjected to warming, and a third plot is subjected to the combined drought and warming treatment. 197 The treatment, which is not subjected to any of the global change treatments, represents ambient 198 conditions (A). For further details regarding online measurements, treatments and experimental 199 setup, see Mikkelsen et al. (2008).

Characteristics of the soil in the treatments at sampling in June 2013 are shown in Table 1. SOC content in 0-10 cm depth was higher at elevated CO_2 than at ambient CO_2 , and at elevated CO_2 drought further increased the SOC content.

203

204 *2.3 Soil sampling and incubation*

Within each of the 48 plots, an undisturbed area of 0.5 m × 0.5 m was selected for this experiment. Areas were chosen to contain an approximately equal amount of *C. vulgaris* and grasses (mainly *D. flexuosa*) at initiation of the experimental setup in 2003. Unfortunately, an error in the experimental procedures forced us to discard samples from one of the blocks, and we therefore present data from five blocks (n = 5).

Soil cores were taken on the 5th-6th of June 2013 with an 8.7 cm diameter cylinder auger to 10 cm 210 depth. The soil was sieved (2 mm) in the field to separate roots from the soil and kept under cool 211 conditions (5 °C) until use. Two weeks after sampling, two 117 mL serum flasks per soil sample 212 were each added 4 g (fw) soil. We added 10 mL water or 10 mL of sucrose solution (4 g L^{-1}) to 213 each of the paired flasks, respectively. Assuming an average bulk density of 1.24 g cm⁻³ in the 10 214 top cm of the soil the sucrose added corresponds to an input of 572 g C m⁻². This is well above the 215 total annual C input to the soil, given that the annual net primary production roughly corresponds to 216 350 g C m⁻² (Chapin III et al., 2002). Hence, during the incubation, the soil was fully water 217 saturated, and microorganisms were at no risk of experiencing C limitation. We used sugar cane-218 derived sucrose with a ${}^{13}C/{}^{12}C$ ratio of $\delta^{13}C = -12$ ‰, as it is distinct from the $\delta^{13}C$ value (-28.5 ‰) 219 of the C3 plants in the area (Reinsch & Ambus, 2013). This enables us to distinguish between CO₂ 220 derived from the added sucrose and from SOC. To eliminate the initial CO₂-content of the flasks, 221

the flasks were sealed, evacuated (< 10% air remaining) and refilled to atmospheric pressure with 222 CO₂-free atmospheric air (Alphagaz Luft 1, Air Liquide, Denmark). Evacuation and refilling was 223 repeated twice and finally 20 mL extra CO₂-free air was added. We incubated the flasks on a shaker 224 (4 h, 20 °C). Following incubation, we sampled 19 mL headspace gas with a syringe and flushed an 225 evacuated 5.9 ml Exetainer vial (Labco Scientific, High Wycombe, UK) with the gas sample 226 leaving the vial at atmospheric pressure for subsequent analyses of CO₂ concentration and isotopic 227 ¹³C/¹²C ratio. Likewise, we sampled gas from four background control flasks, which only contained 228 10 mL water or 10 mL sucrose solution. 229

230

231 2.4 Soil and sucrose analysis

We dried 10 g soil at 103 °C to determine gravimetric soil water content. The total C content and 232 ¹³C/¹²C isotopic ratio of soil and sucrose were measured in dried samples by Dumas combustion 233 (1020 °C) on an elemental analyser (CE 1110, Thermo Electron, Milan, Italy) coupled in continuous 234 flow mode to a Finnigan MAT Delta PLUS isotope ratio mass spectrometer (Thermo Scientific, 235 Bremen, Germany). Homogenized portions of 2 mg (sucrose) or 15 mg (soil) were weighed out into 236 tin combustion cups for elemental analysis. Acetanilide (Merck, Darmstadt, Germany) and soil 237 standards (Elemental Microanalysis, Okehampton, UK) were used for elemental analyser mass 238 calibration. As working standard for isotope ratio analysis we used pure CO₂ gas calibrated against 239 certified reference ¹³C-sucrose (IAEA, Vienna, Austria). Performance of analysis (Oa/Oc) was 240 assessed by the inclusion of reference samples of biological origin (Peach leaves (NIST 1547), 241 National Institute of Standards and Technology, Gaithersburg, MD, USA). 242

243

244 2.5 CO_2 concentration and ${}^{13}C/{}^{12}C$ isotopic ratio

245 CO_2 concentrations and isotopic ${}^{13}C/{}^{12}C$ ratio were analysed on a DeltaV Advantage Isotope Ratio 246 Mass Spectrometer (Thermo Scientific, Bremen, Germany) coupled in continuous flow mode to a GasBench II. Two calibration gas mixtures of CO₂ in synthetic air were included in the analytical runs, *viz.* 362 ppm CO₂ at $\delta^{13}C = -2.7 \%$ vs. Vienna Pee Dee Belemnite (VPDB) and 356 ppm CO₂ at $\delta^{13}C = -29.3 \%$ (Messer Denmark, Padborg, Denmark). CO₂ concentrations and $\delta^{13}C$ were corrected according to the values measured in the background control treatments.

251

252 2.6 Data analysis

Because we eliminated the initial CO_2 content of the flasks before the soil incubation, we assessed

respiration rates based on the CO_2 concentration measured after the incubation period. We

calculated the sucrose-induced respiration as the difference between CO₂-C evolved in sucrose-

amended and non-amended flasks.

257

- We calculated the proportion of SOC-derived respiratory C (P_{SOC}) and sucrose-derived respiratory C (P_{suc}) in sucrose-amended flasks using a two end-member-mixing-model:
- 260

261 [Equation 1]
$$P_{!"\#} = \frac{!!"!_{!"\#\$}!!"!_{!"\#\$}!!"!_{!"\#}}{!!"!_{!"\#\$}!!"!_{!"\#}},$$

262

264

where $\delta^{13}C_{samp}$ denotes the isotopic value of the CO₂ evolved in flasks amended with sucrose,

266 $\delta^{13}C_{SOC}$ denotes the isotopic value of soil C, and $\delta^{13}C_{suc}$ is the isotopic value of the added sucrose (-267 12 ‰).

269	By calculating the total SOC-derived CO ₂ evolved in sucrose-amended flasks and subtracting the
270	basal respiratory CO ₂ evolved, i.e. the CO ₂ produced in non-amended flasks, we can calculate the
271	amount of SOC primed in response to sucrose addition as follows:
272	
273	[Equation 3] C primed = $P_{!"\#} \times [CO_{!"\#\$}] - [CO_{!"!\#}]$,
274	
275	where $[CO_{2suc}]$ and $[CO_{2H2O}]$ denote the CO_2 evolved in flasks with and without sucrose,
276	respectively. We present the priming effect as the increase in SOC-derived CO ₂ with sucrose-
277	amendment in relation to the SOC content of individual samples.
278	
279	To assess whether primed C for elevated CO ₂ octagons derived from C assimilated before or after
280	the initiation of the FACE in 2003, we calculated the $\delta^{13}C$ of the primed SOC-derived CO ₂ ,
281	$\delta^{13}C_{primed}$. First, we calculated the $\delta^{13}C$ of the sucrose-induced respiratory CO ₂ , i.e. the additional
282	CO ₂ produced from sucrose and from SOC priming in response to sucrose addition, as the
283	difference in ¹³ C respired with and without sucrose divided by the difference in CO ₂ produced with
284	and without sucrose:
285	
286	$[Equation 4] \delta^{!"}C \text{ of sucrose-induced CO}_{!}\text{-production} = \frac{[!!!_{!"\#\$}] \times !^{!"}!_{!"\#\$}] - [!!!_{!"\#\$}] \times !^{!"}!_{!"\#\$}}{[!!_{!"\#\$}] - [!!!_{!"!\#!}]},$
287	
288	where subscripts "suc" and "H2O" denote CO ₂ evolved in flasks with and without sucrose addition,
289	respectively.
290	At the same time, $\delta^{13}C$ of sucrose-induced CO ₂ -production can also be expressed in relation to the
291	proportional contribution of sucrose and SOC to the sucrose-induced CO ₂ -production:
292	

[Equation 5] $\delta^{!"}C$ of sucrose-induced CO₁-production = $P^*_{!"\#} \times \delta^{!"}C_{!"\#} + P^*_{!"\#} \times \delta^{!"}C_{!"\#\$\%\&}$,

where P_{suc}^* and P_{SOC}^* denote the proportions of sucrose-derived and SOC-derived CO₂ in the sucrose-induced respiration, respectively. We know all the variables of equation 4, and by combining equations 4 and 5, we can express $\delta^{13}C_{primed}$ as follows:

We know that $\delta^{13}C_{suc}$ is -12 ‰, so we must calculate P^*_{suc} . We calculate P^*_{suc} based on the sucrose-

derived CO₂, CO_{2sucrose-derived}, and the total sucrose-induced respiration, i.e. the difference between CO₂ evolved in sucrose-amended and non-amended flasks. This is possible, because we can calculate CO_{2sucrose-derived} as the product of the proportion of the sucrose-derived respiratory CO₂ (P_{suc}) in sucrose-amended flasks and the CO₂ evolved in sucrose-amended flasks:

308

309 [Equation 7]
$$P_{1'''}^* = \frac{\left[!''_{1'''} * \$ \% \$''' \cdot derived \right]}{\left[! !_{1'''} * \$ \right] ! \left[!''_{1'''} * \right]} = P_{1'''} \times \frac{\left[!''_{1'''} * \$ \right]}{\left[!''_{1'''} * \$ \right] ! \left[!''_{1'''} * \$ \right]}$$

310

311 We know P_{suc} from equation 1; hence:

312

313 [Equation 8]
$$P_{1''\#}^* = \frac{I_{1''}I_{1''\#}I_{1''}I_{1''}I_{1''}}{I_{1''}I_{1''}I_{1''}I_{1''}} \times \frac{[I_{1''}I_{1''}I_{1''}]}{[I_{1''}I_{1''}I_{1''}]}$$

and by inserting equation 8 in equation 6, we find that $\delta^{13}C$ of the primed SOC, $\delta^{13}C_{\text{primed}}$ can be calculated accordingly:

317

319

For one of the soil samples from the ambient treatment the δ^{13} C value of CO₂ evolved in the flasks was unexpectedly high, which suggests that the CO₂ partly originated from carbonate C. We therefore omitted this sample from data analyses. We tested the effects of elevated CO₂, drought and warming on all response variables with full factorial three-way ANOVAs. Homogeneity of variance was assessed with the Brown-Forsythe test. Data for basal respiration rate were log transformed prior to analysis to obtain homogeneity of variance. All statistical analyses were executed in Sigma Plot version 13.0.

327

328 **3. Results**

329 The soil-weight-specific basal respiration and sucrose-induced respiration, i.e. the extra CO₂ produced in sucrose-amended flasks compared to non-amended flasks, was on average c. 50 % 330 331 higher in soil from elevated CO₂ plots, and this increase was more pronounced when drought and elevated CO₂ were combined (Table 2 and 3). This is in line with the higher SOC content at 332 elevated CO₂, which was also highest when elevated CO₂ and drought were combined (Table 1). In 333 contrast, warming did not affect basal or sucrose-induced respiration. The basal decomposition of 334 SOC, expressed as the respiration activity per g SOC, was independent of treatments (Table 2 and 335 3). Likewise, the SOC-specific sucrose-induced respiration activity did not differ between 336 337 treatments (Table 2 and 3).

Sucrose addition enhanced decomposition of native SOC (Fig. 1), hence priming occurred. At ambient CO₂, sucrose enhanced the SOC decomposition rate with 35-49 μ g C g SOC⁻¹ h⁻¹. The priming effect was significantly higher at elevated CO₂ (Table 3), where sucrose addition enhanced SOC decomposition rate with 43-59 μ g C g SOC⁻¹ h⁻¹. There was a tendency towards reduced priming in soils exposed to drought (*P*=0.11), but warming did not affect the level of priming (Fig. 1, Table 3).

345

As a consequence of the eight years of FACE with pure ¹³C-depleted CO₂, the δ^{13} C of the total CO₂ efflux from elevated CO₂ soil (Fig. 2a) was significantly 3.0-5.4 ‰ lower than from ambient CO₂ soil in flasks without sucrose (Table 3). Addition of sucrose with the higher δ^{13} C of -12 ‰ raised the δ^{13} C of the CO₂ evolved during the four-hour incubation. Nevertheless, CO₂ from elevated CO₂ octagons in the sucrose-amended flasks was still significantly 1.2-2.0 ‰ lower than CO₂ evolved from ambient CO₂ soil (Fig. 2b, Table 3).

In contrast to the total CO₂ efflux (Fig. 2), the δ^{13} C of primed SOC-derived CO₂-C from elevated

353 CO₂ soil was not lower than the δ^{13} C of primed SOC-derived CO₂-C from ambient CO₂ soil

(Fig. 3). At ambient CO₂, the δ^{13} C of primed CO₂-C released from soils that were not exposed to

warming was significantly lower than the δ^{13} C of primed C in all other treatments (Fig. 3, Table 3).

356

357 **4. Discussion**

The soil C content in the upper 0-10 cm of the soil profile had increased with 12-22 % after eight years of elevated CO_2 exposure in treatments without experimental drought exposure, and drought further increased soil C content at elevated CO_2 (Table 1), which is consistent with the increased root production at elevated CO_2 recorded in 2009-2010 at the same field site (Arndal et al., 2013). This build-up of organic C resulted in larger basal and sucrose-induced respiration activities expressed per soil weight, whereas the SOC-specific respiration did not respond to any of the treatments (Table 2). Stimulating effects of elevated CO₂ on soil respiration rates (Fig. 1a) have
been reported at the current site of this investigation (Selsted et al., 2012), and are also well
described from other studies (Zak et al., 2000, van Groenigen et al., 2014).

367

As we hypothesized, the priming effect was more pronounced in soils exposed to elevated CO₂ 368 (Fig. 1). With the increased soil C content and C:N ratios of aboveground (Vestergård et al., 2015) 369 and belowground (Arndal et al., 2013, 2014) organic inputs at elevated CO₂, it is likely that the 370 microbial N demand increased. Consequently, the enhanced priming and mineralization of SOC 371 372 may be a result of increased microbial N mining (Dijkstra et al. 2013; Chen et al. 2014). In line with this, soils from elevated CO₂ plots at our field site exhibited higher activity of enzymes involved in 373 374 SOC degradation (Partavian et al., 2015). In a previous laboratory set-up with soil from the current 375 field site, Reinsch et al. (2013) assessed the temporal development of glucose-induced priming over two weeks at 8°C and also found positive priming induced by labile C (glucose), with stronger 376 effects at elevated CO₂. The similar outcomes of the two studies demonstrate that the short-term (4 377 h) immediate priming response to labile C input, i.e. the priming capacity of the inherent microbial 378 379 community prior to microbial growth on the added labile substrate, is a relevant indicator also of long-term priming effects. While Reinsch et al. (2013) only examined the occurrence of priming in 380 soils from a subset of the field treatments, i.e. the ambient (A), elevated CO₂ (CO₂) and the full 381 combination of all treatment factors (TDCO₂), we assessed the effects of all possible combinations 382 of the global change factors, i.e. elevated CO₂, warming and drought, on priming responses. In the 383 longer-term experiment, the priming effect diminished in soils exposed to elevated CO₂, drought 384 and warming in combination (Reinsch et al., 2013). In the present study we did not find a 385 comparable significant interaction between the three global change factors and potential priming of 386 SOC, although we note that the priming response in soils exposed to drought tended to be lower 387 than in soils that were not subjected to experimental drought. 388

389 In our investigation, we incubated soil samples from the different field treatments under standardized conditions with respect to moisture, temperature and CO₂. Hence, we address whether 390 the long-term field manipulation of climate accommodated changes in the microbial decomposition 391 392 of SOC, which could be caused by altered availability and quality of SOC and N and/or altered microbial community activity or composition. We sampled the soil immediately after the annual 393 394 drought treatment, where the water content in drought plots was still significantly reduced (Table 1), and we hypothesized that reduced microbial activity after the drought would impair 395 priming. However, contrary to our expectation, both basal respiration and sucrose-induced 396 respiration per soil weight were enhanced by drought in combination with elevated CO₂, and 397 drought tended to increase the SOC-specific respiration (Table 2). This probably reflects a high 398 399 turnover of drought-decimated microorganisms upon re-wetting in the incubation experiment 400 (Groffman & Tiedje, 1988). On the other hand, drought tended to reduce the sucrose-induced priming of SOC (Fig. 1). The stimulation of microbial activity upon re-wetting of soils after a 401 402 severe drought event thus appears uncoupled from the microbial priming of SOC in response to labile C input. 403

Contrary to our hypothesis, long-term warming did not affect basal respiration, sucrose induced 404 respiration (Table 2), or potential priming (Fig. 1). In the field, warming enhanced microbial 405 abundance (Larsen et al., 2011, Haugwitz et al., 2014) and initiated earlier plant growth in the 406 spring (Kongstad et al., 2012). We expected this to result in decreased N availability, which would 407 be reflected in increased priming (Fontaine et al., 2004, 2011; Zhang et al., 2013), but we found no 408 evidence for this hypothesis. A possible reason is that eight years of warming and earlier onset of 409 spring growth of plants did not decrease soil N availability sufficiently to influence priming. 410 Further, at the field site the warming treatment only raised mean soil temperatures at 5 cm depth by 411 0.1-0.2 °C over the 3 months preceding the soil sampling (Vestergård et al., 2015). This is hardly a 412 temperature increase that would stimulate microbial activity considerably. 413

415	It is remarkable that the δ^{13} C of respiratory CO ₂ derived from SOC priming in soils exposed to eight
416	years of elevated CO_2 with reduced ¹³ C was not lower than in ambient CO_2 soils (Fig. 3). In contrast,
417	the isotopic composition of total respired CO_2 from soils exposed to elevated CO_2 was ¹³ C-
418	depleted compared to CO ₂ evolved from ambient CO ₂ soils (Fig. 2). This shows that C assimilated
419	in the elevated CO ₂ treatments was indeed decomposed in the soil basal respiration, whereas this
420	pool of newly assimilated C was not subject to primed decomposition; hence its decomposition was
421	apparently not energy limited. This implies that the primed C was assimilated more than eight years
422	before sampling. We can therefore add evidence to support previous statements that elevated CO ₂
423	induces decomposition of older soil C (van Groenigen et al., 2005; Xie et al., 2005; Niklaus &
424	Falloon, 2006). Likewise, elevated CO ₂ enhanced the formation of coarse particulate SOM (fresh
425	SOM) and decreased the fraction of physically protected SOM (old SOM) in forest soil (Hofmockel
426	et al., 2011) and in prairie soil (Procter et al., 2015). Given that old SOM pools contain significant,
427	yet (to a large extend) physically and chemically protected N stocks, this lends support to the
428	hypothesis that priming in response to labile C supply is a mechanism by which (some)
429	microorganisms gain access to a reservoir of N to meet their enhanced N demand under conditions
430	of ample C supply (Dijkstra et al., 2013; Chen et al., 2014). If enhanced priming at elevated CO ₂ is
431	caused by increased microbial N demand, because more SOM with lower relative N content enters
432	the system at elevated CO ₂ , it is reasonable that priming should be directed towards SOM pools
433	with a higher N content, i.e. SOM pools incorporated into the system before the elevated CO_2
434	treatment was initiated.
435	Bulk SOC encompasses different pools of SOC of varying age and particle size, and the $\delta^{13}C$ of
436	these different pools vary considerably (Gerzabek et al., 2001). As expected, the SOC δ^{13} C was
437	decreased from -27.8 ‰ in ambient CO ₂ soil to -29.3 ‰ in elevated CO ₂ soil, whereas drought and

438 warming did not affect the isotopic composition of SOC. The lower δ^{13} C of the C primed in the

ambient plots and plots subjected to drought as a single factor compared to the other treatments
(Fig. 3), therefore suggests that sucrose-amendment primed the decomposition of different SOC
pools in the different treatments.

442 It has been argued that short-term incubations as employed in the current study reflects 'apparent' rather than 'real' priming effects. Theoretically, apparent priming is a state, where the initial 443 444 enhanced respiratory pulse induced by labile C addition, derives from turn-over of microbial biomass C rather than decomposition of SOC, i.e. part of the inherent microbial biomass C pool is 445 substituted by the added labile C. 'Real priming', on the other hand, describes the enhanced 446 447 decomposition of SOC after prolonged incubation with labile C (Blagodatskaya & Kuzyakov, 2008; Blagodatsky et al., 2010). We argue, though, that the finding that the primed C was at least eight 448 449 years old is strong indication that even in our short-term incubation study, the addition of labile C 450 resulted in real priming; i.e. the enhanced decomposition of SOC. If the sucrose-induced priming did indeed represent apparent priming, it would imply that the pool-substituted microbial biomass C 451 was more than eight years old. Microbial biomass turnover is on average much faster than eight 452 453 years, and it is quite unlikely that microorganisms grow preferentially on older C pools. Therefore, we find it most plausible that the enhanced soil-derived CO₂-C flux represents real priming of SOC 454 rather than pool substitution of microbial biomass C. 455

456 It has been suggested that increased primary production at elevated CO₂ will enhance C

457 sequestration in terrestrial ecosystems and thus counteract the rise in atmospheric CO_2

458 concentration (Oren et al., 2001; Jastrow et al., 2005; Houghton, 2007). However, this and other

459 studies (Carney et al., 2007; van Groenigen et al., 2014) demonstrate that ecosystems exposed to

460 elevated CO₂ concentrations will be more prone to SOC decomposition triggered by labile C input.

461 This will thus reduce the anticipated increase in C sequestration. In our heath/grassland system

462 elevated CO₂ did enhance the C input to the system and hence the SOC pool (Table 1). However,

463 we demonstrate that labile C inputs accelerate the turnover of older SOC pools and alter C

464	dynamics of the system under elevated CO ₂ . Therefore, in the longer term, the net C balance of this
465	and other systems in a high CO_2 world will depend on the extent to which the build up of new
466	organic C will compensate for the increased loss of older organic C pools.
467	
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731 Figure legends

732 Figure 1

733 Mean priming effect assessed as the increase in SOC decomposition rate in soils amended with

sucrose compared to soils incubated with water. Soils were collected from field plots in a Danish

grass/heathland exposed to 8 years of ambient conditions (A), annually repeated spring/early

summer drought (D), warming (T), elevated atmospheric CO₂ (CO2) and all possible combinations

of single factors. Error bars depict SE. n=4-5.

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Figure 2

Mean δ^{13} C values of CO₂-C respired during incubation of soils collected from field plots in a

741 Danish grass/heathland exposed to 8 years of ambient conditions (A), annually repeated

spring/early summer drought (D), warming (T), elevated atmospheric CO₂ (CO2) and all possible

combinations of single factors. Soils were incubated with water (a) or with a sucrose solution (b).

Error bars depict SE. n=4-5.

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Figure 3

Mean δ^{13} C values of SOC-derived CO₂-C primed by sucrose addition during incubation of soils collected from field plots in a Danish grass/heathland exposed to 8 years of ambient conditions (A), annually repeated spring/early summer drought (D), warming (T), elevated atmospheric CO₂ (CO2) and all possible combinations of single factors. Treatments A and D, marked with asterisk, are significantly different from the other treatments (Tukey *P*<0.05). Error bars depict SE. *n*=4-5.

Table 1. Mean water content and C content in soils collected from field plots in a Danish grass/heathland exposed to 8 years of ambient conditions (A), annually repeated spring/early summer drought (D), warming (T), elevated CO_2 (CO_2) and all possible combinations of single factors. SE depicted in parentheses. n=4-5.

	Soil wate	er content (%)	Soil C c	Soil C content (%)				
A	10.30	(0.42)	3.15	(0.29)				
T	8.04 10.69	(1.00)	3.08	(0.32)				
TD	6.71	(0.53)	2.89	(0.23)				
CO ₂	11.86	(0.50)	3.52	(0.33)				
DCO ₂	9.27	(0.69)	4.80	(0.41)				
TCO ₂	11.68	(1.16)	3.65	(0.60)				
TDCO ₂	11.24	(2.50)	5.75	(1.28)				
Treatment effects	P _{co2} =0.02 P _D =0.012	24	P _{CO2} =0. P _{CO2xD} =	002 0.038				

Table 2. Mean basal and sucrose-induced respiration in relation to soil dry weight and soil organic C (SOC) content during incubation of soils collected from field plots in a Danish grass/heathland exposed to 8 years of ambient conditions (A), annually repeated spring/early summer drought (D), warming (T), elevated atmospheric CO_2 (CO_2) and all possible combinations of single factors. Sucrose-induced respiration is the difference between respiration activity in sucrose-amended and control samples. SE depicted in parentheses. n=4-5.

			Sucrose-in	duced	SOC-specifi	c basal	SOC-specific sucrose-			
	Basal respiration		respiration	า	respiration		induced respiration			
	(µg CO ₂ -	C g soil ⁻¹ h ⁻¹)	(µg CO ₂ -C	g soil ⁻¹ h ⁻¹)	(μg CO ₂ -C g	$sOC^{-1} h^{-1}$)	(µg CO ₂ -C g SOC ⁻¹ h ⁻¹)			
А	2.05	(0.38)	5.07	(0.51)	63.44	(6.45)	160.51	(3.57)		
D	2.00	(0.31)	4.69	(0.78)	65.63	(10.02)	158.85	(30.94)		
Т	1.97	(0.24)	5.00	(0.53)	65.34	(4.99)	166.12	(9.85)		
TD	1.84	(0.21)	3.98	(0.37)	64.05	(4.95)	138.67	(8.36)		
CO2	2.17	(0.16)	5.76	(0.73)	62.76	(4.45)	162.63	(9.50)		
DCO ₂	3.51	(0.47)	8.23	(1.04)	72.45	(5.20)	169.88	(11.04)		
TCO ₂	2.27	(0.49)	5.59	(1.03)	61.22	(5.32)	152.83	(13.68)		
TDCO ₂	4.39	(0.81)	8.49 (1.10)		80.18	80.18 (6.41)		(17.79)		

Table 3. ANOVA table of effects of elevated CO₂ (CO₂), annually repeated spring/early summer drought (D), warming (T) and their interactions on respiration activity, sucrose-induced respiration activity, priming of soil organic C, isotopic composition of respiratory CO₂ evolved during 4 h incubation without or with sucrose and isotopic composition of SOC-derived primed C.

		Soil-weight specific basal respiration		reight Soil-weight specific fic basal sucrose-induced ration respiration		SOC-specific basal respiration		SOC-specific sucrose-induced respiration		Priming		δ^{13} C‰ of CO ₂ without sucrose		δ^{13} C‰ of CO ₂ with sucrose		δ^{13} C‰ of primed SOC-derived CO ₂	
Source	df	F	Р	F	Р	F	Р	F	Ρ	F	Ρ	F	Р	F	Р	F	Р
CO ₂	1	13.70	<0.001	16.15	<0.001	1.23	0.275	0.42	0.520	8.61	0.006	65.22	<0.001	47.14	<0.001	5.97	0.020
D	1	5.88	0.021	2.93	0.097	2.61	0.116	0.369	0.548	2.65	0.114	4.13	0.051	0.02	0.888	3.58	0.068
Т	1	0.04	0.835	0.09	0.771	0.16	0.691	0.46	0.504	0.77	0.386	1.02	0.320	0.07	0.793	1.27	0.269
DxCO ₂	1	7.75	0.009	8.47	0.007	2.80	0.105	1.65	0.209	0.72	0.402	0.15	0.698	0.29	0.594	0.51	0.481
TxCO ₂	1	0.29	0.592	0.14	0.709	0.02	0.900	0.10	0.758	2.32	0.138	3.03	0.091	1.44	0.239	4.40	0.044
TxD	1	0.13	0.718	0.01	0.926	0.14	0.711	0.13	0.721	1.71	0.200	0.25	0.621	1.02	0.320	0.12	0.730
TxDxCO ₂	1	0.32	0.576	0.21	0.650	0.41	0.528	0.18	0.678	0.12	0.735	0.18	0.675	0.32	0.578	0.53	0.472
Error	31																

Figure 1







Figure 3

