

# Peat decomposability in managed organic soils in relation to land-use, organic matter composition and temperature

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**Abstract.** Organic soils comprise a large yet fragile carbon (C) store in the global C cycle. Drainage, necessary for agriculture and forestry, triggers rapid decomposition of soil organic matter (SOM), typically increasing in the order forest < grassland < cropland. But there is also large variation in decomposition due to differences in hydrological conditions, climate, and specific management. Here we studied the role of SOM composition on peat decomposability in a variety of differently managed drained organic soils. We collected a total of 560 samples from 21 organic cropland, grassland and forest soils in Switzerland, monitored their CO<sub>2</sub> emission rates in lab incubation experiments over 6 months at two temperatures (10 and 20°C) and related them to various soil characteristics, including bulk density, pH, soil organic carbon (SOC) content, and elemental ratios (C/N, H/C and O/C). CO<sub>2</sub> release ranged from 6 to 195 mg CO<sub>2</sub>-C g<sup>-1</sup> SOC at 10°C and from 12 to 423 mg g<sup>-1</sup> at 20°C. This variation occurring under controlled conditions suggests, that besides soil water regime, weather and management, SOM composition may be an underestimated factor that determines CO<sub>2</sub> fluxes measured in field experiments. However, correlations between the investigated chemical SOM characteristics and CO<sub>2</sub> emissions were weak. The latter also did not show a dependence on land-use type, although peat under forest was decomposed the least. High CO<sub>2</sub> emissions in some topsoils were probably related to the accrual of labile crop residues. A comparison with published CO<sub>2</sub> rates from incubated mineral soils indicated no difference in SOM decomposability between these soil classes, suggesting that accumulation of recent, labile plant materials that presumably account for most of the evolved CO<sub>2</sub> is not systematically different between mineral and organic soils. In our data set, temperature sensitivity of decomposition (Q10 on average  $2.57 \pm 0.05$ ) was the same for all land-uses, but lowest below 60 cm in croplands and grasslands. This, in turn, indicates a relative accumulation of recalcitrant peat in topsoils.

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## 1. Introduction

Organic soils represent a major global sink for atmospheric carbon (C). Although they cover only 3 % of the earth's terrestrial surface (Tubiello et al., 2016), they store up to 30 % of the global soil organic carbon (SOC) pool (Parish et al., 2008). In Europe, more than 50 % of the former peatland area has been degraded by peat mining and conversion of land use, including drainage to improve their suitability for agriculture or forestry (Joosten, 2010). Drainage aerates the soil so that plants of interest for agriculture and forestry can grow and

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make these soils manageable. The change from anaerobic to aerobic conditions, however, triggers rapid decomposition of peat that had accumulated under the conditions of waterlogging. This transforms the former C-sink into a major source of atmospheric carbon dioxide (CO<sub>2</sub>) and makes peatlands an important contributor to global climate change (Freeman et al., 2004). Around 85% of the global annual CO<sub>2</sub> emission of 915 Mt CO<sub>2</sub>-C from drained peatlands are estimated to originate from organic soils now used as croplands (Tubiello et al., 2016). With rates of 6.5 - 9.4 t C ha<sup>-1</sup> a<sup>-1</sup> net CO<sub>2</sub> fluxes from organic soils now used as croplands were in average found to be higher than from organic soils under grassland, which were estimated to vary between 1.8 and 7.3 t C ha<sup>-1</sup> a<sup>-1</sup> (IPCC, 2014). However, recent studies reported emission rates of 7.6 ± 2.0 t C ha<sup>-1</sup> a<sup>-1</sup> on organic soils managed as grassland in Germany and thus much higher rates than previously found for this type of land-use (Tiemeyer et al., 2016). Drained organic soils under forest can act as both, net sinks or sources of atmospheric CO<sub>2</sub> (Cannell et al., 1993; Minkkinen and Laine, 1998; Minkkinen et al., 1999; Wüst-Galley et al., 2016), although they are in general considered to represent a source with average net CO<sub>2</sub> emissions of 2.0 – 3.3 t C ha<sup>-1</sup> a<sup>-1</sup> in the temperate zone (IPCC, 2014). Temperature and soil moisture regime, which depends among others on drainage depth, are main factors influencing peat decay in drained organic soils (Hogg et al., 1992; Berglund, 1995; Scanlon and Moore, 2000; Chimner and Cooper, 2003; Couwenberg et al., 2010; Leifeld et al., 2012). However, there are substantial differences in CO<sub>2</sub> emissions from organic soils with similar drainage and cultivation properties. Protection of organic matter (OM) against decomposition by mechanisms such as occlusion in aggregates and binding to mineral surfaces, which are important for the stabilization of OM in mineral soils (Six et al., 2002), are of minor importance in organic soils due to the lack or low abundance of minerals (Han et al., 2016). Therefore, the intrinsic decomposability of organic matter is considered another major factor influencing the rate of peat decomposition and a major reason of substantial variation in CO<sub>2</sub> emissions among different sites (Chimner and Cooper, 2003; Byrne and Farrell, 2005; Höper, 2007; Wickland and Neff, 2008; Reiche et al., 2010).

Although intrinsic decomposability of (SOM) cannot be addressed directly, useful indicators of the latter are the relative abundances of labile and recalcitrant C moieties, which shift towards progressively higher proportions of the recalcitrant C with decomposition (Beer et al., 2008; Tfaily et al., 2014) and result in selective enrichment and depletion of specific functionalities (Leifeld et al. 2017; McAnallen et al., 2017). It is important to recognize that during peat formation, most of the net primary production contained in the initial mass of plant residues are lost due to mineralization, and only 10-20% is transformed and accumulated as peat in the water saturated zone of a peat bog or fen (Clymo, 1984). Although decomposition acts slowly on accumulating peat of undisturbed, i.e. water saturated organic soils, it is believed that primarily the most labile OM moieties are lost. Due to fresh

peat layers accumulating on top of older ones, age and depletion in labile compounds increase with soil depth. Incubation studies of peat samples and carbon loss studies with undisturbed organic soils found smaller CO<sub>2</sub> emission rates from deeper peat layers, which was related to the absence of labile compounds i.e. a lower intrinsic decomposability of soil organic matter (SOM) (Hogg et al., 1992; Scanlon and Moore, 2000; Wang et al., 2010; Hardie et al., 2011; Leifeld et al., 2012). Using solid-state <sup>13</sup>C-NMR, DRIFT/FTIR spectroscopy, and pyrolysis-GC/MS, various studies of OM composition of undisturbed peat profiles have shown a gradual change with increasing depth towards a relative enrichment of compounds that are recalcitrant against decomposition under anoxic conditions, such as lignins and polyphenols (Freeman et al., 2004), while the contents of labile oxygen-rich compounds, such as polysaccharides, were found to decrease (Leifeld et al., 2012; Biester et al., 2014; Sjögersten et al., 2016).

Elemental ratios between oxygen (O), hydrogen (H), nitrogen (N) and carbon are widely used as indicators of the relative abundance of different groups of compounds such as phenols, lipids and polysaccharides, and proteins. Lignins and polyphenols have molar O/C ratios in the range of 0.2 – 0.6 and H/C ratios between 0.9 and 1.5, while the respective ratios of carbohydrates range from 0.8 to 0.9 for O/C and from 1.4 to 1.8 for H/C (Kim et al., 2003). In line with the molecular and spectroscopic analyses mentioned before, both ratios were found to decrease with increasing depth in peat (Klavins et al., 2008; Biester et al., 2014; Wüst-Galley et al., 2016). On the other hand, both, fresh plant residues and undisturbed peat usually have high C/N ratios (Loisel et al., 2012). When peat becomes exposed to oxic conditions, mineralization seems to lead to relative enrichment of N, explaining why decreased C/N ratios are found in organic topsoils compared to undrained peat layers or bottom layers of drained organic soils (Malmer and Holm, 1984; Kuhry and Vitt, 1996; Krueger et al., 2015). While undisturbed organic soils have a low bulk density, drainage leads to subsidence processes and increasing bulk densities in the topsoils (Rogiers et al., 2008; Leifeld et al., 2011a; b).

The temperature sensitivity of peat mineralization, as expressed by its Q<sub>10</sub> value, is a useful parameter to characterize the intrinsic decomposability of SOM (Hogg et al., 1992; Biasi et al., 2005; Davidson and Janssens, 2006; Conant et al., 2008; Boddy et al., 2008; Karhu et al., 2010; Hiltunen et al., 2013). In line with the biochemical and elemental evidence reviewed above, it was reported to increase with increasing resistance of peat soils against OM decomposition (Scanlon and Moore, 2000), soil depth and peat age (Hardie et al., 2011; Hiltunen et al., 2013).

Despite its likely important role in determining future C losses from drained peatland, the influence of SOM composition on peat decomposition in managed organic soils is not well studied. While decomposition rates seem to decline with increasing peat age i.e. profile depth, the oxic conditions, occurring after drainage onset,

lead to fast SOM decomposition. As for undisturbed organic soils, we expect that post drainage decomposition  
100 primarily acts on the most labile OM moieties. However, the much faster decomposition of labile SOM might  
alter the depth interaction found in undisturbed peat soils. Further, recent inputs from plant residues may supply  
the topsoils with labile OM. Around 20 % of carbon in organic soils under agriculture is derived from crop  
residues and thus more rapidly decomposing (Bader et al., 2017). The fractions of OM derived from peat and  
recent inputs and their decomposability in drained organic soils may, however, substantially vary with land-use,  
105 site conditions and time since land use conversion. (Schulze et al., 2009) reported that inputs of fresh organic  
matter residues were smaller in croplands than in grasslands or forests, suggesting that SOM might be on  
average more aged and thus less decomposable. In-situ measurements of CO<sub>2</sub> fluxes from managed organic soils  
reveal slower decomposition of peat under forest (IPCC, 2014). Together, smaller peat loss rates and higher  
residue input make us expect that SOM decomposition rates under controlled conditions are fastest in forest  
110 topsoils.

In this study, we analysed the relationship between SOM properties, specific decomposition rates (CO<sub>2</sub>-C mg<sup>-1</sup>  
SOC) and their temperature sensitivity of peat samples taken from depths between 0 and 200 cm of 21 drained  
organic soils in Switzerland managed as cropland, perennial grassland or forest. These sites embody three major  
after-uses for drained peatlands as occurring in Europe (Joosten 2010) and are also representative for the  
115 situation in Switzerland where most former peatlands are drained and managed (Wüst-Galley et al. 2015). We  
measured decomposition rates in incubation experiments under standardized lab conditions and interpreted the  
current decomposition status of peat using SOM properties such as i) carbon stocks, bulk densities and the  
elemental ratios O/C, H/C and C/N as well as ii) the temperature sensitivity towards decomposition, expecting  
that

- 120 1. Specific decomposition rates of SOM decline with depth
2. Specific decomposition rates of SOM in managed organic soils correlate with its composition and are  
inversely related to the temperature sensitivity of decomposition
3. Specific decomposition rates of topsoil SOM are largest in the forest and smallest in the croplands.

## 2. Methods

### 125 2.1 Sampling sites

The soil samples used for this study were taken from organic soils distributed across Switzerland that were  
identified using the map of Wüst et al. (2015). Apart from current land-use (grassland, cropland, forest), they  
differed in the type of drainage system (ditches in forest, pipes in crop- and grassland), time since drainage onset

and drainage intensity, altitude (MASL), and climate (Tab. 1). All sites were classified as fens, although we  
130 found bog-derived peat layers within the top 30 and 40 cm of the soil profiles at two sites (SK\_FL, K\_FL).  
Cropland management comprised rotations typical for Switzerland with maize, winter wheat, ley, and rape seed  
as major crops. Sites were conventionally tilled. Grasslands were used for cutting and haymaking, not grazing,  
and fertilized according to the Swiss Fertilization Recommendations (Flisch et al. 2009). Forest sites were  
managed and their vegetation was not peat-forming.

## 135       **2.2 Soil sampling**

Between October 2013 and June 2015, we sampled in total 84 peat cores from all 21 sites (4 cores per site). All  
cores were taken to a maximum depth of 1 m. If the underlying mineral layer was reached before 1 m depth,  
coring was discontinued. We used a Belorussian peat corer (cuts a half-cylindrical undisturbed core of diameter  
4 cm) for soils with low bulk densities and a motorized Humax corer (cuts a cylindrical core of diameter 5 cm)  
140 for denser soils. The samples were stored at 4 °C for up to 2 months until analysis. We applied the method of  
(Rogiers et al., 2008) to account for soil compaction during sampling for any sample, and divided the cores into  
segments corresponding to 5 - 10 cm depth increments. This corresponded, depending on the type of soil corer  
used and length of the increment, to sample volumes of between 31 and 196 cm<sup>3</sup> per segment. In total this  
resulted in 1605 soil samples. Some cores had interlayers of mineral sediment, identified by a different color  
145 (grey), a high bulk density as well as their SOC content being lower than 150 g kg<sup>-1</sup>. These interlayers were  
excluded from the analysis. The soil of one site (BI\_FL) had no limnic layer and therefore was classified as a  
murshic histosol; all others were classified as murshic limnic histosols (WRB, 2014).

## **2.3 Soil analysis**

Soil pH was measured for two to three samples of each core (307 samples in total), using a flat surface electrode  
150 (pH 100, Extech Instruments, USA) calibrated at pH 7.00 and pH 4.01. Aliquots of fresh soil (10 g dry matter)  
were diluted in distilled water (2.5 parts water to 1 part material by mass), shaken, left for 20 hours and shaken  
again, before the pH measurements were carried out.

Prior to further chemical analysis, the samples were oven-dried at 105 °C and weighed to determine bulk density  
(g cm<sup>-3</sup>). The dried samples were ground for 2 min at 25 rotations s<sup>-1</sup> in a ball mill (Retsch MM400) and  
155 subsampled to determine total carbon (C<sub>tot</sub>), SOC, hydrogen (H), nitrogen (N) and oxygen (O) contents. C<sub>tot</sub>, H  
and N were analysed after dry combustion of ground subsamples in an elemental analyser (Hekatech, Germany).  
To determine SOC, we hydrolysed ground aliquots with 36 % HCl (acid fumigation) in a desiccator to remove  
any carbonates, before the samples were analysed in the elemental analyser. A third set of ground subsamples  
were used to determine the O contents by means of the same analyzer after pyrolysis at 1000°C. We corrected O

160 contents for inorganic O, assuming that all inorganic O was present in form of CaCO<sub>3</sub>. The O/C and H/C ratios given in this paper represent mole ratios, whereas the C/N ratios represent mass ratios. For analysis O/C ratios and H/C ratios of samples having a SOC content lower than 150 g kg<sup>-1</sup> were excluded from analysis. Soil carbon stocks (t C ha<sup>-1</sup>) refer to the organic horizons summed over each profile and thus do not include sediment layers that interspersed the profiles.

#### 165 **2.4 Incubation experiment**

We selected at least two soil segments of each soil core from depths between 0-30 cm, 30-60 cm and 60-100 cm for incubation to determine SOM decomposability. Each segment was divided in two subsegments whereas one subsegment was incubated at 10 °C and the other subsegment at 20 °C for between 6 and 13 months. From the one location (M\_CL) where we had taken cores of >100 cm length, we selected six additional samples from the  
170 depth below 100 cm for incubation, resulting in a total of 560 incubated samples. Prior to incubation, we thoroughly mixed every segment, removed visible roots and adjusted the water potential to -10 kPa, using a hanging water column. The sample weight was 53.9 ± 0.7 g (mean ± standard error) at -10 kPa. Following the method of (Chapman, 1971), we measured CO<sub>2</sub> emission rates by means of a Respicond VII analyser (Nordgren Inovations, Sweden) over three to four measurement cycles of several weeks between November 2013 and  
175 March 2016. The measurement principle is based on the change in electrical conductivity of the NaOH solution with increasing uptake of CO<sub>2</sub>. In each cycle, we vented the alkali CO<sub>2</sub> traps (NaOH 0.6 M) of the analyser regularly after 50-60 mg of CO<sub>2</sub> had been emitted to prevent O<sub>2</sub> deficiency. In addition, we exchanged the NaOH solution while the traps were vented. Between measurement cycles, we kept the soil samples at the same temperature and moisture level as during the cycles.

#### 180 **2.5 Data analysis**

We only used CO<sub>2</sub> data taken after the first 3 days of each measurement cycle for data analysis to avoid artefacts that might have resulted from moving the samples and adjusting their water content. Furthermore, we excluded all negative emission rate values (0.45 % out of 1700 CO<sub>2</sub> measurements taken on average per sample). Data gaps (83 % of the timeline) between measurement cycles were filled by means of interpolation using a robust  
185 linear regression on the log transformed data. The specific amount of SOC which was emitted from a sample as CO<sub>2</sub> during 10,000 hours of incubation at 10 or 20 °C [mg CO<sub>2</sub>-C g<sup>-1</sup> SOC], L, was calculated as

$$L = \frac{(CO_{2\ sample} - CO_{2\ blank}) \times \frac{12.01}{44.01}}{SOC_{sample} \times m_{sample}} \quad (1)$$

where CO<sub>2 sample</sub> is the amount of CO<sub>2</sub> emitted from the sample over 10,000 hours of incubation [mg CO<sub>2</sub>-C g<sup>-1</sup> SOC], CO<sub>2 blank</sub> is the median of ambient CO<sub>2</sub> accumulation collected in 6 blank vessels over more than 6

190 months and extrapolated to 10,000 hours (on average 27 mg),  $SOC_{\text{sample}}$  is the SOC content of the sample [ $g\ kg^{-1}$ ], and  $m_{\text{sample}}$  is the mass of the soil sample [kg].

To determine Q10 values we applied the method used by others (e.g., Hogg et al., 1992; Scanlon and Moore, 2000; Wang et al., 2010; Wetterstedt et al., 2010; Hardie et al., 2011), dividing the 10,000 h length of the incubation period at 10 °C by the time span over which samples incubated at 20 °C emitted the same amount of  
195  $CO_2\text{-C}$  per mg SOC as those incubated at 10 °C emitted during 10,000 h. Given that the same amount of SOC is lost at both temperatures, changes in OM composition during incubation are also assumed to be the same and thus differences in the rates are assumed to reflect only the influence of temperature and not that of differences in composition. Q10 values are known to depend on incubation temperatures. In order to compare our results with those of other studies we calculated the activation energy ( $E_a$  in  $kJ\ mol^{-1}$ ) required for decomposition of SOC,  
200 using Q10 values:

While  $R$  is the gas constant ( $8.314\ J\ K^{-1}\ mol^{-1}$ ) and  $T$  is the temperature used for incubation (K).

$$EA = \frac{R \times \frac{\ln(Q_{10})}{\left(\frac{1}{T_1} - \frac{1}{T_2}\right)}}{1000} \quad (1)$$

Mixed linear models were used to analyse the effects of the various soil parameters on SOM mineralization and their interactions with land-use. The function `lmer` from the package `lme4` (Bates et al., 2015) was implemented  
205 using the software R (R core Team, 2015) to run mixed linear models. Heteroscedasticity or departure from normality was assessed graphically. In order to avoid heteroscedasticity, we log-transformed topsoil C-stocks and bulk density data. We tested whether the factor “land-use” had a significant influence on the variation of each of the analysed variables ( $\alpha = 0.05$ ). To do this, the following two mixed models, [2] and [3], were run for each dependent variable and compared using an ANOVA.

$$210 \quad \text{variable} \sim \text{land.use} + \text{random effects} \quad (2)$$

$$\text{variable} \sim \text{random effects} \quad (3)$$

Sampling depth, sampling location and site repetition were included as random effects to account for the dependence among segments of the same core and among cores from the same sampling location, respectively. In addition, we included bulk density, SOC, nitrogen, hydrogen and oxygen contents, as well as the emitted  $CO_2$   
215 as further random effects, given that there was no collinearity with the tested variable and that the Aikaikes criterion (AIC) of the models revealed smaller scores with additional random effects. The additional random effects used for each model are given in table 2 and 3. Further, we determined the significance of land-use specific differences (CL vs FL, CL vs GL, FL vs GL), using a least square means test for linear models (`lsmeans` package).

220 We used the same approach to test the influence of the factor ‘soil depth’ on the target variables with interactions between the three sampling depths (0-30 cm, 30-60 cm, >60cm), using the model

$$\text{variable} \sim \text{depth.interval} + \text{random effects} \quad (4)$$

in addition to Equation 3. To determine the significance of depth-specific differences, we used least square means test as mentioned before.



225 **3. Results**

**3.1 SOM characteristics**

Soil pH, SOC content, C/N ratio and bulk density showed significant land-use effects (Fig. 1; Tab. 2; Tab. S1):  
The lowest soil pH values were found in the forest topsoil samples, whereas SOC content and C/N ratio were the  
highest in these samples. Bulk density was highest in the cropland topsoils. Below 30 cm depth, soil pH, SOC  
230 content, C/N ratio and bulk density showed no land-use effect.

In the forest soil profiles, soil pH overall increased with depth, whereas it decreased in the grassland and  
cropland soils (Tab. 3). Also bulk density decreased with depth in the grassland and cropland soils, while SOC  
content and C/N ratio increased. In the forest soils, SOC content, bulk density and C/N ratio did not differ  
between topsoil (0-30 cm depth) and subsoil samples (30-60 cm depth); however, below 60 cm depth SOC was  
235 slightly lower than above, while bulk density and C/N ratio were higher than above 60 cm depth (Fig. 1, Tab. 3).  
The cumulated topsoil C stocks showed no land-use effects, but tended to be larger in cropland and forest than in  
grassland soils over the entire profile (Fig. 1; Tab. 2).

The molar H/C and O/C ratios of the organic matter fell between the typical values of the ratios for  
carbohydrates and lignin, which is displayed in a Van Krevelen plot (Fig. 2). The lowest values of both ratios  
240 were found in the forest soils, the highest in the grassland and cropland topsoils. Both ratios were lower in the  
topsoils than in the subsoils of the cropland and grassland sites, while there was no difference between the two  
depths in the forest soils (Tab. 3). At depths below 30 cm, the O/C ratio was lower in the forest soils than in the  
other soils, but without land-use effect in the H/C ratio.

245 **3.2 CO<sub>2</sub> emissions and Q<sub>10</sub>**

The samples incubated at 10 °C emitted  $32.56 \pm 1.39$  mg CO<sub>2</sub>-C g<sup>-1</sup> SOC, while samples incubated at 20 °C  
emitted  $74.06 \pm 2.98$  mg CO<sub>2</sub>-C g<sup>-1</sup> SOC (Fig. 1). At 10°C we did not observe a land-use effect on CO<sub>2</sub> emission  
(Fig. 1, Tab. 2), but at 20 °C the topsoil samples from croplands emitted less CO<sub>2</sub> than those from forests. This  
effect occurred due to extra ordinarily high emissions of the samples from two grassland and two forest sites  
250 (VW\_GL, VW\_F, SK\_GL, SK\_F) (Fig. 3). Those four sites experienced the least intensive drainage.  
Furthermore, these sampling sites were situated at high altitude in a pre-alpine environment with lower mean  
annual temperatures and higher precipitation than at the other sites (Tab. 1). In pairwise comparisons between  
adjacent sites of different land-use (i.e., VW\_GL vs VW\_FL, SK\_GL vs SK\_FL, C\_CL vs C\_GL and G\_CL vs  
G\_GL and G\_FL (Tab. 1)), land-use effects were only found for the last site (Fig. 3).

255 At 10 °C, CO<sub>2</sub> emissions of the topsoil samples from all sites together were higher than from samples taken at 30  
to 60 cm depth, independent of land-use (Tab. 3). Analysing the influence of depth separately by land-use type,  
this effect was only manifested in grassland and forest, but not in cropland soils. We found no overall depth  
effect at 20 °C, but CO<sub>2</sub> emissions of topsoil samples from forests were higher than those of samples taken at  
lower depths, whereas we found the opposite for the cropland soils. Despite the just mentioned depth effects, the  
260 general relationship between emissions and soil depth was weak and not consistent in its sign (Tab. 4).  
Over the course of the incubation, CO<sub>2</sub> emissions increased for 40 % of the samples, as revealed in Tab. S1 by  
positive slopes of the regression lines. These increases were independent of land-use. In total, the CO<sub>2</sub> emissions  
from these samples were almost 50 % higher than those from the other samples that instead showed a trend of  
decreasing emissions.

265 Mean Q10 values were  $2.57 \pm 0.05$ . The Q10 did not differ between the three land use types. It was lower below  
60 cm depth in the cropland and grassland, but not in the forest soils (Fig. 1; Tab. 3). Activation energies (E<sub>a</sub>)  
calculated from Q10 values ranged around 48.1 and 123.5 kJ mol<sup>-1</sup> and like Q10 values decreased with depth.  
There were significant relationships between CO<sub>2</sub> emission and SOC content, bulk density and C/N ratio in  
general but they were weak (Tab. 4). The Q10 values showed similar relationships to these soil variables as CO<sub>2</sub>  
270 emission.

## 4. Discussion

### 4.1 SOM characteristics

The SOC contents, bulk densities and C/N ratios found in the deeper parts of soil profiles presented here were  
275 close to values that are typical for undisturbed peat (Grover and Baldock, 2012; Loisel et al., 2014). They also  
indicate that soils of our study sites were characteristic for European fens and resembled typical properties of  
managed organic soils (Berglund, 1995; Kechavarzi et al., 2010; Eickenscheidt et al., 2015; Krueger et al., 2015;  
Wüst-Galley et al., 2016; Brouns et al., 2016). Several studies assume that deeper layer peat of managed organic  
soils is less decomposed (Ewing and Vepraskas, 2006; Rogiers et al., 2008; Leifeld et al., 2011b; b; Krueger et  
280 al., 2015; Wüst-Galley et al., 2016). We therefore interpret the different SOM characteristics found in the  
topsoils of our samples as indicators of an advanced decomposition triggered by drainage.

The land-use specific differences manifested in different topsoil SOC contents and C/N ratios (highest under  
forest), and topsoil bulk densities (lowest under forest). The higher forest C/N ratios might be explained by  
absence of the use of N fertilizers and lower bulk densities by lower traffic with field machinery. In addition,

285 differences in C/N between land-use types may also suggest that peat decomposition was less advanced in forests compared to croplands and grasslands. Further, depth effects are lowest in forest soils, indicating a lower impact of soil management that could also result in a lower decomposition of forest topsoils. The relatively high carbon stocks found in cropland top soils are most likely the result of subsidence after drainage and compaction from field traffic, leading to increased soil bulk density in the uppermost layers. This effect, with respect to C  
290 stocks, overrides the overall much smaller C concentration in agriculturally managed organic soils.

The H/C and O/C ratios in the deeper layers of the studied soils were similar to those found in undisturbed bogs and drained bogs used for forestry in Switzerland (Zaccone et al., 2007; Wüst-Galley et al., 2016). They indicate an enrichment of polyphenols and aromatic carbon with depth, which is in line with the current understanding of peat development in peatlands (Cocoza et al., 2003; Zaccone et al., 2007; Klavins et al., 2008; Delarue et al.,  
295 2011; Leifeld et al., 2012, 2017). The increased H/C and O/C ratios in the grass- and cropland topsoils can be attributed to inputs of fresh plant litter to the topsoil via above- and belowground residues, as such residues are rich in carbohydrates (Koegel-Knabner, 2002). In a previous study, in which we used stable and radiocarbon isotopes to label the SOC of two of the studied soils (C\_CL and C\_GL in Table 1), at least 20 % of topsoil organic matter was no peat, but derived from recent plant litter (Bader et al., 2017). The results further indicated  
300 that the OM derived from these fresh plant residues were a source of labile C that contributed more to decomposition than the old, peat-derived SOM.

The H/C and O/C ratios reflect the mixing ratio of these two SOM sources. The H/C and O/C ratios in forest topsoils were lower than of those under cropland and grassland and did not change with depth. Interpreting these lower H/C and O/C ratios in the forest top-soils as indicators of more advanced peat decomposition (Klavins et al., 2008; Leifeld et al., 2012; Biester et al., 2014; Wüst-Galley et al., 2016) would be in contradiction to our  
305 conjecture that land management effects on peat decomposition, revealed by SOC, bulk density and C/N ratio, are less pronounced for forests. We rather argue that reason for the low H/C and O/C ratio in the forest soils is a higher abundance of lignin rich (wood derived) plant residues. A second mechanism for comparably higher O/C and H/C ratios in cropland and grassland soils, could be that peat loss in the uppermost layers was higher under agriculture than under forest, resulting in a relatively higher share of SOM from recent inputs. Considering all  
310 the available evidence SOM characteristics, we conclude that peat decomposition is less advanced in forest soils than in agricultural soils, in line also with field flux measurements on managed organic soils that typically show faster decomposition in croplands and grasslands than in forests (IPCC, 2014).

315 **4.2 CO<sub>2</sub> emissions and temperature sensitivity of decomposition**

The studied soils emitted, on average, ca. 5-10 % of their SOC (20 °C) as CO<sub>2</sub>, calculated for an incubation duration of one year. The advanced decomposition state of many of the samples might give reason to expect that CO<sub>2</sub> rates are below that of more intact peat or mineral topsoils that contain a larger fraction of recent plant residues. To understand whether SOM in the studied organic soils is particularly stable, we compared its average  
320 daily carbon loss with data from studies that used undisturbed to extensively managed organic soils, or mineral soils (Tab. 5; Fig. 3). Indeed, our values are on average below those from other organic soil studies. However, their range overlaps with the uncertainty of the regression line that is plotted through results from studies from other, mostly intact or little degraded organic soils. Hence, the pronounced oxidative decomposition after long periods of drainage might result in a relatively smaller labile SOC pool, but the large variability between  
325 experimental set-ups, incubation lengths, and water contents among incubation studies prevents a stronger line of interpretation. Interestingly, the regression lines modelled for organic and mineral soils did not deviate significantly from each other. Therefore, the pools size of labile carbon, indicated by the decomposition rates, seem not to differ between these soil classes. This comparison suggests that accumulation of recent, labile plant materials that presumably account for most of the evolved CO<sub>2</sub> is not systematically different between mineral  
330 and organic soils.

Samples showing an increase in CO<sub>2</sub> emission rate over time were predominantly of subsoil origin, where SOC contents and C/N ratios indicate a lower decomposition than in the topsoil. Furthermore, based on the information we have on land-use and drainage depths, it appears that most of these samples were taken from soil layers that were protected from intensive decomposition by water saturation. The long incubation period in our  
335 study might have given aerobic decomposer communities time to develop and grow, whereas time might not have been sufficient in shorter studies.

Like other studies on extensively managed or undisturbed organic soils that investigated depth interaction of decomposition rates in the top 30 to 50 cm (Hogg et al., 1992; Scanlon and Moore, 2000; Wang et al., 2010; Hardie et al., 2011), we found a decrease of specific CO<sub>2</sub> release with depth. However, the relationship between  
340 CO<sub>2</sub> emissions and depth was rather weak in our case and not consistent for both incubation temperatures and the different land-uses. Compared to the studies on unmanaged organic soils, reporting declines of a factor 2 to 30 (Hogg et al., 1992; Scanlon and Moore, 2000; Wang et al., 2010; Hardie et al., 2011), our differences were substantially smaller. Drainage and decadal agricultural use of the studied soils led to more intense decomposition processes in the topsoil, resulting in little depth interaction or, for croplands, sometimes maybe  
345 even a reversal of decomposability. Further, the abundance and decomposability of crop residues has to be

considered as a substantial CO<sub>2</sub> source. For two topsoils (C\_GL and C\_CL), Bader et al. (2017) showed that at least 20% of the SOM is crop residue derived and responsible for 40% of the emitted CO<sub>2</sub>. Assuming that the abundance of crop and plant residues is highest in topsoils, it might be possible that decomposability of peat derived SOM either does not depend on depth or topsoil peat decomposes at smaller rates. Therefore, we cannot confirm our first hypothesis of decreasing decomposition rates with depth.

As Tab. 5 shows, the Q10 values found in our study ( $2.74 \pm 0.06$ ) were higher than Q10 values found elsewhere for similar sampling depths but in unmanaged organic soils (Chapman and Thurlow, 1998; Hamdi et al., 2013; Hardie et al., 2011; Hiltunen et al., 2013; Hogg et al., 1992; Scanlon and Moore, 2000; Yavitt et al., 2000). Also the temperature independent Ea was higher in the studied samples ( $69.4 \pm 3 \text{ kJ mol}^{-1}$ ) than in most other studies on undisturbed organic soils ( $47.4 \pm 7.2 \text{ kJ mol}^{-1}$ ) (Tab. 5). However, three studies (Chapman, 1971; Hardie et al., 2011; Hogg et al., 1992) found similar or higher Ea values in northern organic soils. In the case of Chapman and Thurlow (1998) they were also managed as grassland or forest, whereas the other studies used peat from undisturbed organic soils. Nevertheless, the high Ea of the studied samples might reflect the change in chemical peat composition with decomposition after drainage towards higher recalcitrance. In contrast to other studies on unmanaged organic soils reporting no trend or increasing Q10 values with depth (Scanlon and Moore, 2000; Wang et al., 2010; Hardie et al., 2011; Hiltunen et al., 2013), the cropland and grassland profiles in our study had a lower Q10 below the 60 cm depth. Various studies on SOM decomposition used Q10 values as an indicator of SOM recalcitrance (Hogg et al., 1992; Biasi et al., 2005; Davidson and Janssens, 2006; Conant et al., 2008; Hartley and Ineson, 2008; Conant et al., 2011; Hiltunen et al., 2013). Considering that the presence of labile crop residues would decrease Q10 in the topsoil rather than in the subsoil, the higher topsoil Q10 may be explained by an extended accumulation of recalcitrant moieties. This proposed high abundance of recalcitrant moieties in topsoils of degrading organic soils is in line with the reported differences in SOM composition in different layers as well with the pattern of CO<sub>2</sub> emissions. The latter show no substantial difference with depth and indicate that a higher fraction of recent and labile plant residues in topsoils is counterbalanced by a high recalcitrance of the highly degraded peat. Comparing radiocarbon concentrations in SOC and emitted CO<sub>2</sub> of two sites also used for this study (C\_CL, C\_GL), Bader et al. (2017) estimated that SOC from plant residue inputs is more labile than peat. The measured radiocarbon contents for SOC were 75 to 80 pMC and therefore indicated that peat of the topsoil must have experienced a substantial decomposition.

It is remarkable that despite the controlled conditions in our incubation experiment the variation in cumulative loss of initial SOC of between 0.6 and 42.3 % (Fig. 3), was similar to or even larger than that observed in field flux measurements (IPCC, 2014). This large variability suggests that the composition of SOM is of similar

importance as drainage, climate and other site factors in controlling CO<sub>2</sub> emissions from drained organic soils. Nevertheless, the relationships between the measured SOM parameters to assess the biochemical decomposability, CO<sub>2</sub> emissions and Q10 values were rather weak and thus do not support our second hypothesis. This stands in contrast to other studies which concluded that chemical composition is a major factor of SOM decomposability in organic soils (Scanlon and Moore, 2000; Koch et al., 2007; Reiche et al., 2010; Hardie et al., 2011; Leifeld et al., 2012). However, these studies focused mainly on single profiles of undisturbed or extensively used organic soils. A recent study investigated relationships between SOM parameters and decomposition rates of German organic soils under controlled conditions (Säurich et al., 2017). These authors mostly studied strongly disturbed fens with similar properties to the soils in our study. Besides SOC contents, soil pH and C/N ratios, Säurich et al. (2017) focused on other soil nutrients, stable isotopes and microbial biomass. In line with our results, they could not identify strong proxies for SOC decomposition by means of simple chemical attributes.

In order to explain the weak relationships between SOM composition and CO<sub>2</sub> release it should be considered that, in our case, the emitted CO<sub>2</sub> comprised only 3.2-7.4 % of the total SOC, while the analysed SOM parameters in this and other studies represent bulk SOM. Our methods allowed to gain a broad overview on the chemical composition of SOM, while decomposition might more tightly be bound to the abundance of specific OM moieties.

Although land-use affected SOM characteristics, such as elemental contents and their ratios, the amount of CO<sub>2</sub> emitted from the soils did not differ among the three types of land-use. We therefore have to reject also our third hypothesis of a higher SOM decomposition rate in forest topsoils. We assume that long-lasting drainage and management might have resulted in an equivalent decomposition of most of the labile OM, along with its intrinsic decomposability.

## 400        **5. Conclusions**

Chemical characteristics of SOM indicated an advanced peat decomposition in the uppermost layers of drained organic soils used as cropland or grasslands. Under controlled moisture and temperature conditions, CO<sub>2</sub> emissions from peat samples had a similar variability, as found for in situ CO<sub>2</sub> flux experiments on drained organic soils. Therefore, carbon loss from drained organic soils cannot be explained entirely by climate or drainage depth. However, simple chemical characteristics of SOM, as used in this study, were not specific enough to explain the variability in CO<sub>2</sub> emissions or the temperature sensitivity of decomposition under

controlled conditions. Despite that CO<sub>2</sub> emissions were occasionally higher in topsoils, probably derived from accrual of labile plant residues, the remarkable decrease of Q10 values with depth suggested that the relative content of recalcitrant peat derived SOM was high in topsoils of managed organic soils, indicating an advanced degradation in these uppermost layers. It is therefore necessary to quantify the fraction of peat derived SOM throughout a drained soil profile, in order to verify this assumption. Yet we understand from the similar magnitude of CO<sub>2</sub> emission rates found above and below 30 cm depth that future peat loss will occur at similar or even faster rates, assuming an increasing mean annual temperature.

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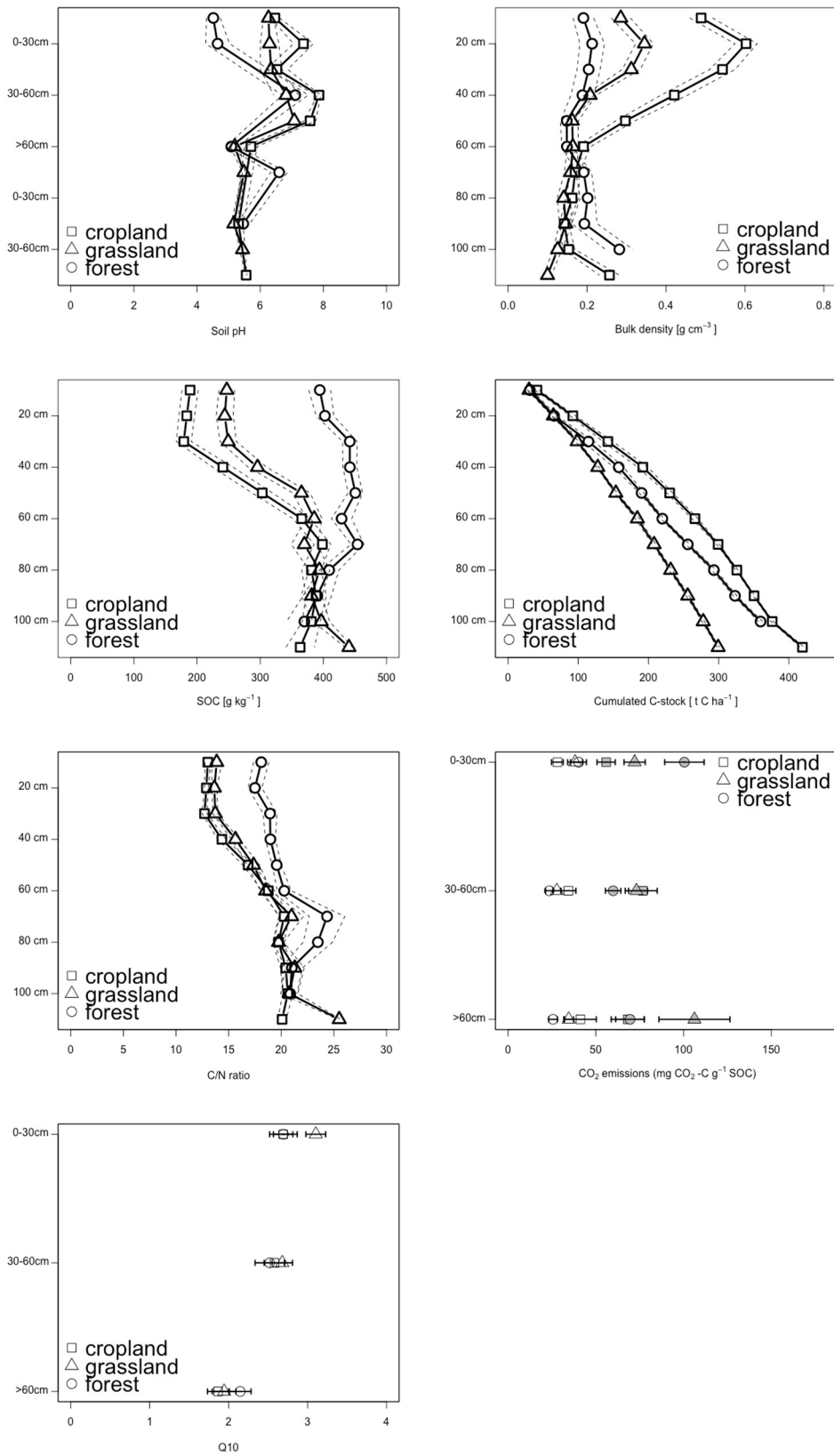
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**Figure 1**



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Fig. 1. Soil pH, bulk density, SOC content, cumulated C-stocks, C/N ratios,  $\text{CO}_2$  emissions and temperature sensitivity (Q10) displayed for the three land-use types (cropland, grassland and forest) in relation to the profile depth (cm, y-axis).  $\text{CO}_2$

emissions are displayed at 10 (open symbols) and 20°C (black symbols), while the area between dashed lines and error bars represents the standard errors of the mean.

**Figure 2**

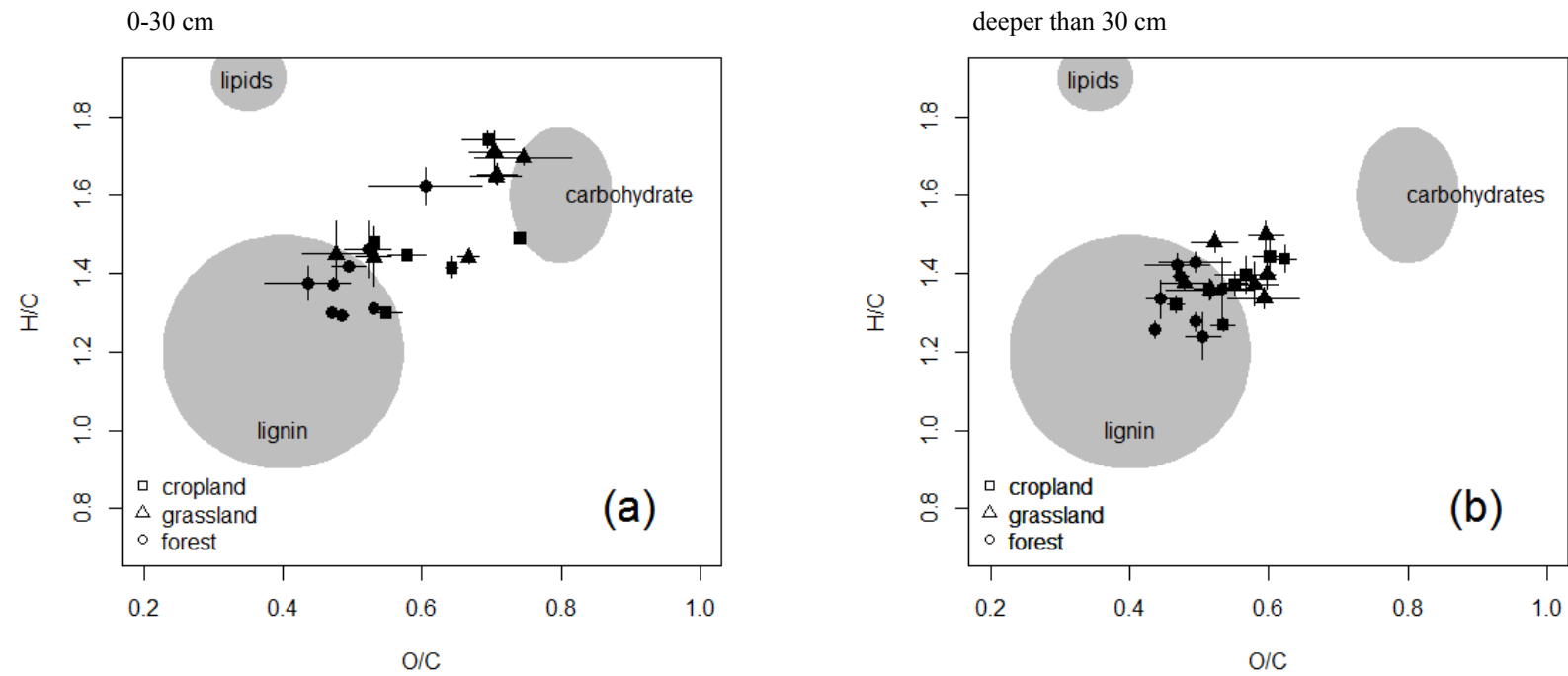


Fig. 2. Van Krevelen plots of samples from (a) the upper 30 cm and (b) depths below 30 cm. Symbols represent averages for relevant core segments from each site; black bars represent the standard error of the mean; grey surfaces represent the range of O/C and H/C for lignin, carbohydrates, and lipids, adapted from Preston and Schmidt (2006).

Figure 3

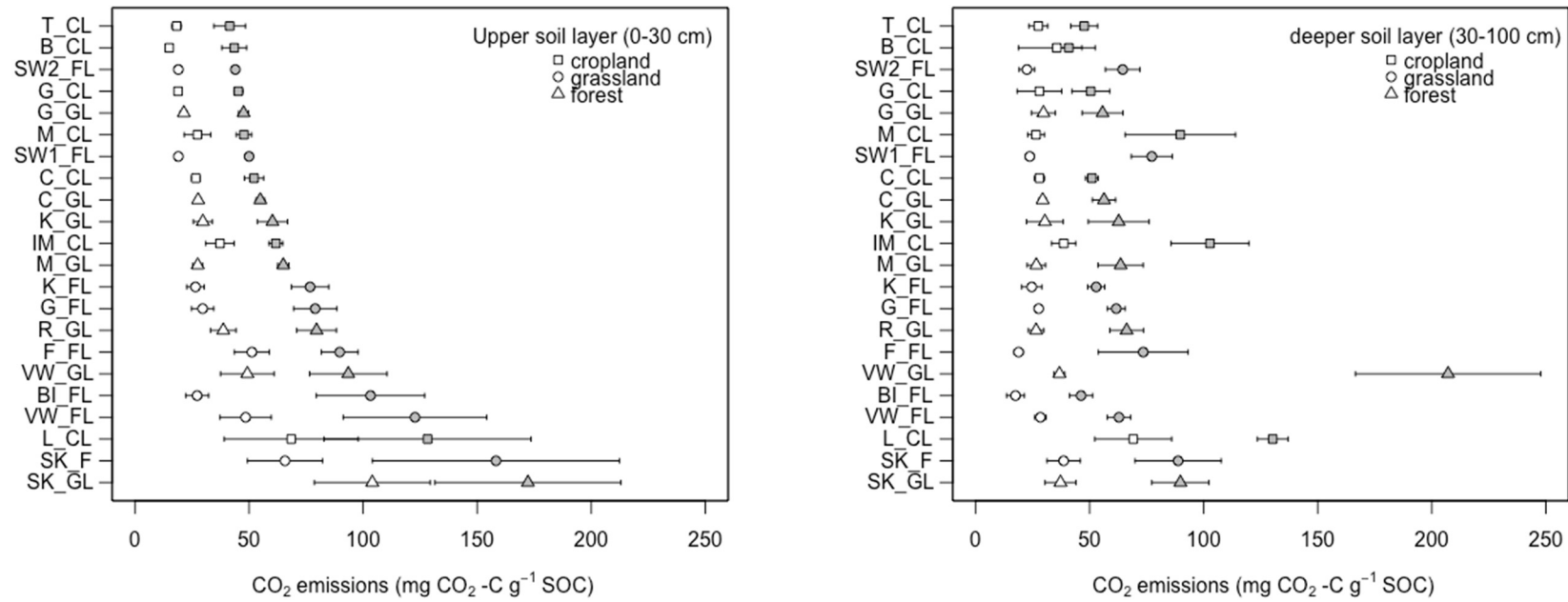


Fig. 3. CO<sub>2</sub> emissions at 10 (open symbols) and 20°C (grey symbols) displayed for upper soil layers (0-30cm) and bottom layers (30-100cm) of all sampling locations. Sampling locations are sorted from the lowest to highest CO<sub>2</sub> emissions in the left diagram. Same order of sites was taken for the diagram on the right. Error bars represent the standard error of the mean. If a symbol lacks error bars, the standard error was smaller than the symbol size or as in the cases of upper soil layer SW1 and SW2 n= 2.



Figure 4

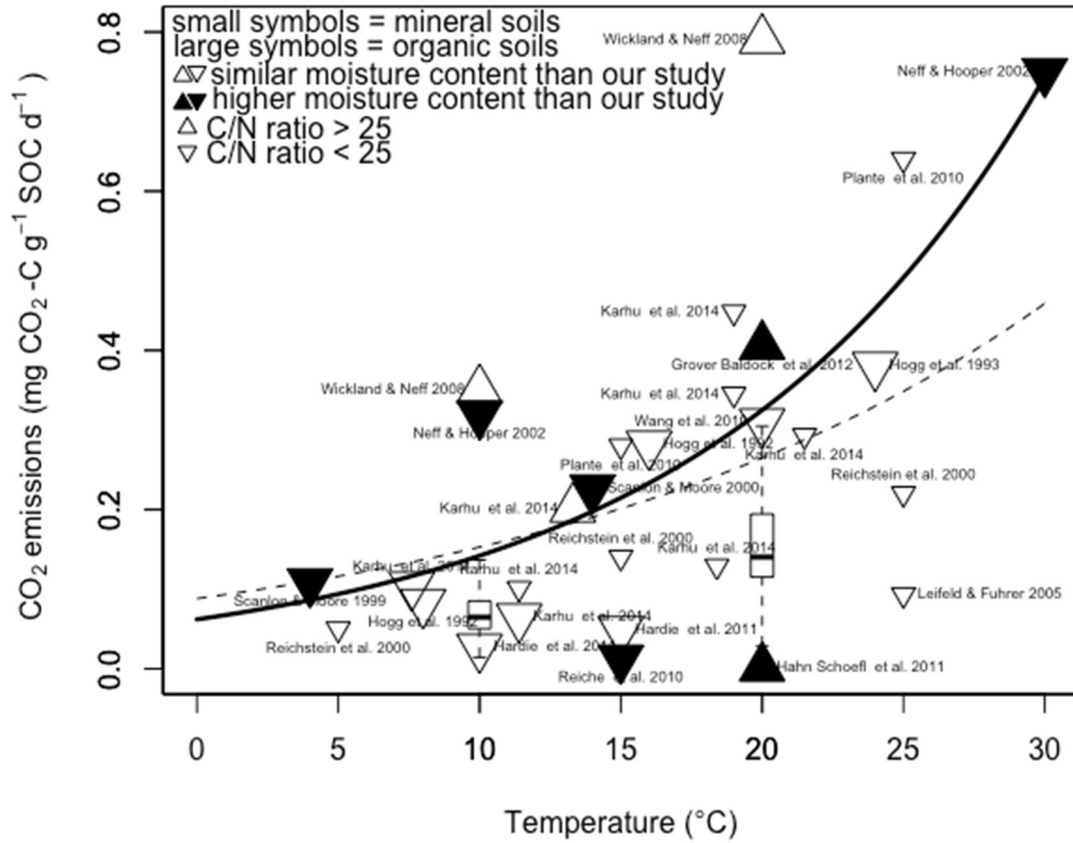


Fig. 3: Comparison of daily CO<sub>2</sub> emission rates from this study (boxplots) with rates found during other incubation studies (organic soils and mineral soils). The curves represent the modelled CO<sub>2</sub> emission rates for organic soils from other studies (solid thick line) rate=0.06e<sup>0.08t</sup> and mineral soils (dashed thin line) rate=0.09e<sup>0.05t</sup>, for temperatures between 0-30 °C.

**Table 1**

Tab. 1: List of sampling locations, including information on the land-use type, peat thickness, approximate time since drainage onset, elevation (MASL), mean annual temperature (MAT) and mean annual precipitation (MAP) of each site.

Location name	Abbreviation of Location with Land-use (CL: Cropland, GL: Grassland, FL: Forest)	Co-ordinates (WGS 1984)	MASL (m)	Area size (ha)	Peat thickness <sup>1</sup> (cm)	Drainage history <sup>2</sup>	Drainage class <sup>3</sup>	MAT <sup>4</sup> (°C)	MAP <sup>5</sup> (mm)
Birmensdorf	BI_FL	8.454,47.357	560	2.6	95	unclear; peat excavation nearby	s	9.2	1122
Brüttelen	B_CL	7.175,47.033	438	3.1	290	drained by 1864	d	9.9	1003
Chreienriet	F_FL	8.486,47.434	440	6.9	330	Unclear; peat excavation site nearby until 1940	d	9.4	1040
Cressier	C_CL, C_GL	7.047,47.041	430	1.6, 1.6	120	drained by 1864	d	10.0	1145
Gals	G_CL, G_GL, G_FL	7.065,47.040	430	1.2, 0.8, 1.0	<100	drained by 1864	d	10.0	1145
Im Moos	IM_CL	9.573,47.379	414	5.6	400	drained by 1860; intensive drainage between 1942 -1962	d	10.1	1297
Katzensee	K_FL	8.495,47.433	440	1.9	230	unclear; peat excavation site nearby until 1940	s	9.4	1040
Kirchenthurnen	K_GL	7.523,46.821	540	9.9	302	drained after 1860	d	8.9	1136
Lüchingen	L_CL	9.574,47.378	414	4.4	400	drained by 1860; intensive drainage between 1942 -1962	d	10.1	1297
Mühlethurnen	M_CL, M_GL	7.523,46.821 7.523,46.817	540	8.2, 7.6	400	drained after 1860; intensive drainage in 1942	d	8.9	1136
Rüthi	R_GL	9.536,47.283	435	13.3	>700	drained by 1970	d	10.1	1533
Staatswald 1+2	SW1_FL, SW2_FL	7.092,46.984	431	30.0, 48.4	142	drained by 1864; intensive drainage in 1942	d	10.1	990
Summerigchopf	SK_GL, SK_FL	9.399,47.212	1300	11.5, 2.2	147-202	drain established between 1935 and 1960	s	6.0	1731
Treiten	T_CL	7.145,47.010	439	29.3	238	drained by 1864	d	9.9	1033
Vorderwengi	VW_GL, VW_FL	9.098,47.196	1070	1.1, 0.9	100 -146	grassland drained by 1935	s	6.2	2240

<sup>1</sup>Peat thickness was determined by excavation an additional peat core down to the underlying sediment layer; <sup>2</sup>Information on drainage was gained by viewing Siegfried topographical maps (1870-1949), considering information on Swiss organic soils by (Lüdi, 1935), as well as aerial photographs; <sup>3</sup>shallow drainage < 0.5 m s, deep drainage > 0.5 m d; <sup>4</sup>MAT is the average for the years 1981-2010; <sup>5</sup>MAP is the average of the years 1971-1991 derived from original data of MeteoSchweiz

**Table 2**

Tab. 2: Results of land-use effect analysis for the whole soil profile as well as specifically in the topsoil (0-30 cm) and bottom layers (>30 cm), displayed for SOC concentration, C-stocks, bulk density C/N, H/C and O/C ratios, CO<sub>2</sub> emissions at 10 and 20 °C, and the resulting Q10 values.

Attribute	Land-use interaction <sup>1</sup>		P values between spec. land-uses <sup>2</sup>			
	$\chi^2$ Value	P value	CL vs FL	CL vs GL	FL vs GL	
<b>Soil pH</b>	$\chi^2(2)=3.7$	0.16	-			a
Soil pH (0-30 cm)	$\chi^2(2)=14.9$	0.0006	0.0003	0.9	0.0021	a
Soil pH (>30 cm)	$\chi^2(2)=0.7$	0.7	-			a
<b>SOC</b>	$\chi^2(2)=10.7$	0.005	0.14	0.28	0.002	b n +bd
SOC (0-30 cm)	$\chi^2(2)=14.5$	0.0001	0.0001	0.5	0.009	a
SOC (>30 cm)	$\chi^2(2)=3.0$	0.2	-			a
<b>Cumul. CStock</b>						a
CStock (0-30 cm)	$\chi^2(2)=5.4$	0.07	0.06	0.4	0.6	a
CStock (0-100 cm)	$\chi^2(2)=5.4$	0.06	0.2	0.06	0.8	a
<b>Bulk density</b>	$\chi^2(2)=3.4$	0.2	-			a
Bulk density (0-30 cm)	$\chi^2(2)=10.3$	0.06	0.02	0.09	0.4	a
Bulk density (>30 cm)	$\chi^2(2)=2.0$	0.4	-			a
<b>C/N ratio</b>	$\chi^2(2)=5.9$	0.05	0.06	0.9	0.1	a
C/N ratio (0-30 cm)	$\chi^2(2)=15.0$	0.0005	0.0002	0.8	0.003	a
C/N ratio (>30 cm)	$\chi^2(2)=2.2$	0.3	-			a
<b>H/C ratio</b>	$\chi^2(2)=6.7$	0.04	0.5	0.4	0.02	a
H/C ratio (0-30 cm)	$\chi^2(2)=6.3$	0.04	0.6	0.4	0.03	a
H/C ratio (>30 cm)	$\chi^2(2)=3.5$	0.2	-			a
<b>O/C ratio</b>	$\chi^2(2)=11.5$	0.003	0.4	0.7	0.001	c bd+h
O/C ratio (0-30 cm)	$\chi^2(2)=10.5$	0.005	0.06	0.7	0.003	a
O/C ratio (>30 cm)	$\chi^2(2)=8.5$	0.014	0.008	1.0	0.003	a
<b>CO<sub>2</sub> 10°C</b>	$\chi^2(2)=2.4$	0.3	-		-	d soc+oorg
CO <sub>2</sub> 10°C (0-30 cm)	$\chi^2(2)=2.9$	0.2	-			oorg
CO <sub>2</sub> 10°C (30-60 cm)	$\chi^2(2)=7.17$	0.03	0.023	0.38	0.34	SOC oorg
CO <sub>2</sub> 10°C (>60 cm)	$\chi^2(2)=1.6$	0.4	-			SOc oorg h
<b>CO<sub>2</sub> 20°C</b>	$\chi^2(2)=1.4$	0.5	-			a
CO <sub>2</sub> 20°C (0-30 cm)	$\chi^2(2)=6.5$	0.04	0.03	0.2	0.7	a
CO <sub>2</sub> 20°C (30-60 cm)	$\chi^2(2)=1.7$	0.4	-			a
CO <sub>2</sub> 20°C (>60 cm)	$\chi^2(2)=1.2$	0.5	-			a
<b>Q10</b>	$\chi^2(2)=3.5$	0.2	-			a
Q10 (0-30 cm)	$\chi^2(2)=0.4$	0.8	-			a
Q10 (30-60 cm)	$\chi^2(2)=1.1$	0.6	-			a
Q10 (> 60 cm)	$\chi^2(2)=1.0$	0.6	-			oorg

<sup>1</sup> P value of ANOVA comparing linear mixed models with and without the factor “land-use type”.

<sup>2</sup> P value emitted using least square means between “land-use types”

**Table 3**

Tab. 3: Results of the depth influence analysis displayed for Q10 values , CO<sub>2</sub> emissions at 10 and 20°C, SOC contents, bulk densities, C/N ratios, H/C ratios and O/C ratios. Ea values (not shown) had similar significances as Q10 values.

Attributes	Depth interaction		P-values between specific depth classes			
	$\chi^2$ Values	P-value	0-30 vs 30-60	0-30 vs >60	30-60 vs >60	
<b>Q10 Values</b>	$\chi^2(2)=46.2$	$9.56 \times 10^{-11}$	0.05	<0.0001	<0.0001	a
Q10 Cropland	$\chi^2(2)=16.1$	0.0003	0.8	0.0002	0.002	a
Q10 Forest	$\chi^2(2)=5.2$	0.08	-	-	-	a
Q10 Grassland	$\chi^2(2)=29.5$	$3.9 \times 10^{-7}$	0.06	<0.0001	0.009	a
<b>CO<sub>2</sub> emiss.(10°C)</b>	$\chi^2(2)=6.1$	<0.05	0.03	0.7	0.2	a
Cropland (10°C)	$\chi^2(2)=1.5$	0.5	-	-	-	a
Forest (10°C)	$\chi^2(2)=17.3$	0.0001	0.0001	0.01	0.5	a
Grassland (10°C)	$\chi^2(2)=7.9$	0.02	0.01	0.3	0.5	a
<b>CO<sub>2</sub> emiss.(20°C)</b>	$\chi^2(2)=0.9$	0.6	-	-	-	a
Cropland (20°C)	$\chi^2(2)=8.4$	0.015	0.02	1.0	0.09	a
Forest (20°C)	$\chi^2(2)=13.2$	0.0001	0.0007	<0.05	0.6	a
Grassland (20°C)	$\chi^2(2)=3.5$	0.17	-	-	-	a
<b>pH</b>	$\chi^2(2)=6.0$					SOC+Oorg+H+N1+BD
Cropland	$\chi^2(2)=19.4$	$6.2 \times 10^{-5}$	<0.02	<0.0001	0.09	
Forest	$\chi^2(2)=36.8$	$1 \times 10^{-8}$	0.004	<0.0001	0.001	emittCO2
Grassland	$\chi^2(2)=27.4$	$1.1 \times 10^{-6}$	0.0001	<0.0001	0.9	emittCO2.2+SOC+Oorg+H
<b>SOC</b>	$\chi^2(2)=157.7$	$<2.2 \times 10^{-16}$	<0.0001	<0.0001	0.0001	N+BD
Cropland	$\chi^2(2)=158.2$	$<2.2 \times 10^{-16}$	<0.0001	<0.0001	<0.0001	N+BD
Forest	$\chi^2(2)=3.8$	0.15	0.2	1.0	0.3	N+BD
Grassland	$\chi^2(2)=143.2$	$<2.2 \times 10^{-16}$	<0.0001	<0.0001	<0.0001	N+BD
<b>Bulk Density</b>	$\chi^2(2)=57.6$	$<3.1 \times 10^{-13}$	<0.0001	<0.0001	0.9	corg
Cropland	$\chi^2(2)=312.6$	$<2.2 \times 10^{-16}$	<0.0001	<0.0001	<0.0001	a
Forest	$\chi^2(2)=31.6$	$1.4 \times 10^{-7}$	0.7	<0.0001	<0.0001	a
Grassland	$\chi^2(2)=49.9$	$1.4 \times 10^{-11}$	<0.0001	<0.0001	0.08	corg
<b>C/N ratio</b>	$\chi^2(2)=325.5$	$<2.2 \times 10^{-16}$	<0.0001	<0.001	<0.0001	bd
Cropland	$\chi^2(2)=199$	$<2.2 \times 10^{-16}$	<0.0001	<0.001	<0.0001	bd
Forest	$\chi^2(2)=41.2$	$1.5 \times 10^{-9}$	0.4	<0.001	<0.0001	bd
Grassland	$\chi^2(2)=152.8$	$<2.2 \times 10^{-16}$	<0.0001	<0.001	<0.0001	bd
<b>H/C ratio</b>	$\chi^2(2)=19.9$	$<4.7 \times 10^{-5}$	<0.0001	<0.0002	0.7	Oorg und BD und N
Cropland	$\chi^2(2)=46.7$	$7.3 \times 10^{-11}$	<0.0001	<0.0001	0.3	a
Forest	$\chi^2(2)=1.9$	0.38	-	-	-	a
Grassland	$\chi^2(2)=52.6$	$3.7 \times 10^{-12}$	<0.0001	<0.0001	0.07	Oorg und BD und N und CO2
<b>O/C ratio</b>	$\chi^2(2)=22.0$	0.0005	<0.06	0.03	0.9	H
Cropland	$\chi^2(2)=0.01$	1.0	-	-	-	H
Forest	$\chi^2(2)=2.6$	0.3	-	-	-	CO2
Grassland	$\chi^2(2)=6.3$	0.04	0.03	0.5	0.5	H

**Table 4**

Tab. 4: Coefficients of determination and correlation for CO<sub>2</sub> emissions measured at 20°C and different soil attributes as explanatory variables (profile depth, SOC content, bulk density, C/N, O/C and H/C ratio). Ea values (not shown) behaved similar as Q10 values.

Attribute	0-30 cm				30-100 cm			
	Intercept	cor	P value	R <sup>2</sup>	Intercept	cor	P value	R <sup>2</sup>
CO <sub>2</sub> at 20°C								
Depth (cm)	<2.0x10 <sup>-16</sup>	-0.23	0.01	0.05	0.0001	0.11	0.2	0.01
SOC (g kg <sup>-1</sup> )	8.06x10 <sup>-6</sup>	0.31	0.001	0.09	7.6x10 <sup>-6</sup>	-0.01	0.9	4.8x10 <sup>-5</sup>
Bulk density (g kg <sup>-1</sup> )	<2.0x10 <sup>-16</sup>	-0.27	0.003	0.07	<2x10 <sup>-16</sup>	-0.19	0.02	0.04
C/N ratio	0.002	0.28	0.002	0.08	3.5x10 <sup>-6</sup>	-0.1	0.5	0.004
O/C ratio	1.33x10 <sup>-6</sup>	-0.02	0.9	0.0002	2.1x10 <sup>-5</sup>	0.06	0.4	0.01
H/C ratio	0.046	0.03	0.8	0.0005	0.292	0.07	0.4	0.01
pH	2.3x10 <sup>-8</sup>	-0.29	0.001	0.09	2.3x10 <sup>-8</sup>	-0.25	0.003	0.06
CO <sub>2</sub> at 10°C								
Depth (cm)	1.7x10 <sup>-13</sup>	0.19	0.2	0.02	0.0002	0.10	0.2	0.02
SOC (g kg <sup>-1</sup> )	1.7x10 <sup>-5</sup>	-0.13	0.04	0.03	4.8x10 <sup>-16</sup>	-0.24	0.002	0.06
Bulk density (g kg <sup>-1</sup> )	<2.0x10 <sup>-16</sup>	-0.22	0.02	0.04	<2.0x10 <sup>-16</sup>	-0.01	0.9	9.1x10 <sup>-5</sup>
C/N ratio	0.0007	0.31	0.0007	0.09	1.05x10 <sup>-5</sup>	-0.03	0.7	0.0008
O/C ratio	3.8x10 <sup>-5</sup>	0.01	1.0	3.4x10 <sup>-5</sup>	3.4x10 <sup>-5</sup>	0.10	0.2	0.01
H/C ratio	0.5	0.14	0.1	0.2	0.3	0.24	0.002	0.06
pH	6.6x10 <sup>-5</sup>	-0.14	0.1	0.02	0.0001	-0.09	0.3	0.007
Q10 values								
Depth (cm)	<2.0x10 <sup>-16</sup>	0.18	0.12	0.01	0.0001	-0.30	0.0001	0.08
SOC (g kg <sup>-1</sup> )	<2.0x10 <sup>-16</sup>	-0.16	0.03	0.03	<2.0x10 <sup>-16</sup>	0.12	0.06	0.02
Bulk density (g kg <sup>-1</sup> )	<2.0x10 <sup>-16</sup>	-0.02	0.6	-0.006	<2.0x10 <sup>-16</sup>	-0.02	0.4	-0.002
C/N ratio	8.1x10 <sup>-16</sup>	-0.10	0.15	0.01	1.2x10 <sup>-15</sup>	-0.08	0.4	-0.002
O/C ratio	5.3x10 <sup>-15</sup>	0.20	0.07	0.02	<2.0x10 <sup>-16</sup>	-0.13	0.3	0.0002
H/C ratio	0.0001	0.06	0.2	0.004	2.9x10 <sup>-8</sup>	-0.13	0.06	0.02
pH	1.02x10 <sup>-9</sup>	-0.03	0.8	0.0006	2.7x10 <sup>-8</sup>	-0.02	0.8	0.0003

1 **Table 5**

2 Tab. 5: Incubation studies with organic and mineral soils at different moisture levels, soil depths and temperatures. If moisture level says moist, samples were incubated directly after being retrieved  
 3 from the field, while saturated samples were incubated under wetter, i.e. anaerobic conditions. Similar stands for samples whose water content was similar to our samples  
 4

	Soil	Region	Moisture level	°C	days	C/N	depths (cm)	CO <sub>2</sub> emissions (mg CO <sub>2</sub> -C g <sup>-1</sup> SOC d <sup>-1</sup> )	Q10	Ea (kJ mol <sup>-1</sup> )
This study	drained Fens	Switzerland	-10 kPa	10 20	416	17.7	5-150	0.078 0.18	2.57	69.5
Chapman and Thurlow, 1998	Drained/undrained Bogs	UK (Scotland)	Moist	10 20	unclear	-	0-20	0.051* 0.030*	3.2	80.0
Grover and Baldock (2012)	Bog	Australia	Moist	20	38	15-25	5-110	0.13-0.78	-	
Hahn-Schöfl et al. (2011)	Fen	Germany	Saturated	20	346	15.3		0.013	-	
Hardie et al. (2011)	Bog	UK	Dryer	5 10 15	6	30	0-30	0.027 0.049 0.093	3.66	86.4
Hartley and Ineson (2008)	Mineral soil	UK	Dryer	10 20	124		unclear	0.046 0.074	3.25	81.3
Hilasvuori et al. (2013)	Bog	Finland	Moist	10 20	short	83	0-44	0.016* 0.061*	2	22.7
Hogg et al. (1992)	Fen	Canada	Similar	8 16 24	120	40.6	5-40	0.083 0.282 0.381	1.9 -2.2	62.0
Karhu et al. (2014)	Organic soil	UK (Scotland)	Similar	11.4	174	28.6	0-10	0.065	-	
	Organic soil	UK (Scotland)		7.6	174	36.5	0-10	0.105	-	
	Organic soil	UK		13.3	174	18.7	0-10	0.201	-	
	Mineral soil	UK		11.4	174	13.3	0-10	0.101	-	
	Mineral soil	Spain		21.5	174	14.3	0-10	0.293	-	
	Mineral soil	Spain		19	174	13.0	0-10	0.345	-	
	Mineral soil	Spain		19	174	18.6	0-10	0.448	-	
	Mineral soil	Italy		18.4	174	13.2	0-10	0.129	-	
Karhu et al. (2010)	Mineral soil	Finland		8-25	540	-	0-30	-	3.0	45.0
Koch et al. (2007)	Organic soil	Austria	Moist	0-30	25	21.6	0-5	-	2.0	31.9
Leifeld and Fuhrer (2005)	Mineral soil	Switzerland	Similar	25	707	7.85	5-35	0.12	4.6	110.8
Neff and Hooper (2002)	Organic soil	USA (Alaska)	unclear	10 30	352	34.6	0-10	0.32 0.75	1.9	22.9
Plante et al. (2010)	Mineral soil	USA	Similar	15	56		0-20	0.28	1.36 -1.79	31.7
Reiche et al. (2010)	Fen	Germany	Saturated	15	31	30.1	0-40	0.0022	-	
Reichstein et al. (2000)	Mineral soil	Switzerland	Similar	5 15 25	104	30.3		0.05 0.14 0.22	2.5-2.7	65.9
Scanlon and Moore (2000)	Fen	Canada	Moist	4 14	12	43	5-45	0.227 0.109	2.0	45.8
Wang et al. (2010)	Organic soil	China	Similar	5-20	40	28.5	10-30	0.31	2.2	53.3
Wickland and Neff (2008)	Organic soil	Canada	Similar	10 20	57	24.7	2-30	0.35 0.79	1.7	36.6
Yavitt et al. (2000)	Bog	Canada	Moist	12-22	2		0-54	-	1.4	32.5

\* Study authors are not specific about the SOC content of peat, therefore we assumed it to be 400 gkg<sup>-1</sup>, according to the findings of undisturbed bog peat (Loisel et al., 2014).

