# Geogenic organic carbon in terrestrial sediments and its contribution to total soil carbon

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# 15 Abstract

16 Geogenic organic carbon (GOC) from sedimentary rocks is an overlooked fraction in soils that has not

17 yet been quantified, but influences the composition, age and stability of total organic carbon (OC) in

18 soils. In this context, GOC is the OC in bedrocks deposited during sedimentation. However, the

19 contribution of GOC to total soil OC varies depending on the type of bedrock. As yet, no far studies

20 have investigated the contribution of GOC derived from different terrestrial sedimentary rocks to soil

21 OC contents.

22 In order to fill this knowledge gap, 10-m long sediment cores from three sites recovered from Pleistocene 23 Loess, Miocene Sand and Triassic Red Sandstone were analysed and the amount of GOC calculated based on <sup>14</sup>C measurements. The <sup>14</sup>C ages of bulk sedimentary OC revealed that OC is comprised of 24 25 both biogenic and geogenic components. The biogenic component relates to OC that recently entered 26 the sediments from plant sources. All the sediments contained considerable amounts of GOC (median amounts of 0.10 g kg<sup>-1</sup> in Miocene Sand, 0.27 g kg<sup>-1</sup> in Pleistocene Loess and 0.17 in Red Sandstone) 27 compared with subsoil OC contents (between 0.53-15.21 g kg<sup>-1</sup>). Long-term incubation experiments 28 29 revealed that the GOC appeared comparatively stable against biodegradation. Its possible contribution 30 to subsoil OC stocks (0.3-1.5 m depth) ranged from 1 to 26 % in soil developed in the Miocene Sand, from 16 to 21 % in the Loess soil and from 6 to 36 % at the Red Sandstone site. Thus, GOC with no 31 detectable <sup>14</sup>C content influenced the <sup>14</sup>C ages of subsoil OC, and may partly explain the strong increase 32 in <sup>14</sup>C ages observed in many subsoils. This could be particularly important in young soils on terrestrial 33 sediments with comparatively low amounts of OC, where GOC can make a large contribution to total 34 35 OC stocks. 36 37

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Keywords Geogenic organic carbon; sedimentary organic carbon; <sup>14</sup>C; terrestrial sediments; incubation
 experiment

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# 48 1. Introduction

On average, the world's soils store more than 50 % of OC in the subsoil below 30 cm depth (Batjes, 49 2014). This type of carbon is considered a highly stable carbon pool due to its apparently high <sup>14</sup>C ages 50 (Mathieu et al., 2015, Schrumpf et al., 2013). However, another explanation for this could be the 51 52 contribution of geogenic organic carbon (GOC), which is defined here as OC that originates from deposition during sedimentation and rock formation, and may increasingly influence subsoil OC with 53 depth (Graz et al., 2010, Kögel-Knabner et al., 2008, Schrumpf et al., 2013, Trumbore, 2009). GOC in 54 most cases is devoid of <sup>14</sup>C and thus may lead to an overestimation of ancient OC sources although a 55 56 number of studies showed the importance of root derived, young OC inputs to subsoils (Angst et al., 2016, Crow et al., 2009). Therefore GOC may significantly influence and affect the overall <sup>14</sup>C signal, 57 particularly in OC-poo subsoils. Vindušková et al. (2015) investigated the contribution of GOC to soils 58 in reclaimed mine soils, and found GOC contributions to total soil OC of between 26 and 99 %. 59 Furthermore OC-rich sediments with contents of 2-7 g kg<sup>-1</sup> (Hemingway et al., 2018) and 28-105 g kg<sup>-1</sup> 60 <sup>1</sup>(Frouz et al., 2011) have been investigated with regard to the stability of OC in these sediments, but no 61 62 conclusion reached about GOC contributions in soils. However, the impact of GOC on soils derived 63 from sediments or sedimentary rocks with lower OC contents has not yet been investigated. Considering that approximately 65 % of the continental earth's surface is covered with sediments and sedimentary 64 rocks (Amiotte Suchet et al., 2003), a potentially large fraction of soils could contain GOC that 65 contributes to soil OC stocks, even though a large portion might be derived from recent sedimentation 66 processes. There is not vet much literature about sediments containing only low amounts of OC. There 67 68 are estimations that assume sandstones to be free of GOC(van der Voort et al., 2018) or, in contrast, a storage model that assumes generally high GOC amounts of 2.4 g kg<sup>-1</sup> for all sandy deposits (Copard et 69 al., 2007). Therefore, more information about the amounts of GOC in sediments is needed. 70

71 To estimate the possible contribution of GOC to subsoil OC stocks, it is important to establish the 72 amount of OC in sediments that comes from sedimentation (GOC) and to distinguish it from OC derived 73 from current vegetation (biogenic OC). There are many soil- and substrate-specific factors that might 74 influence the OC contribution from current vegetation to sedimentary OC, such as potential rooting 75 depth or pore distribution. No method has yet been established to allow a direct quantification of GOC 76 in different soils or sediments, apart from promising methods to quantify the graphitic part of GOC in 77 soils (Zethof et al., 2019). The only reliable approach to distinguish between both sources is the use of <sup>14</sup>C. Since deposition of sediments mostly took place > 50,000 years BP, they do not contain <sup>14</sup>C, which 78 has a mean half-life time of 5,730 years (Libby, 1952). In addition, the  $\delta^{13}$ C values of OC in the 79 sediments allow carbonaceous sources with  $\delta^{13}$ C values around 0 % to be distinguished from organic 80 sources with  $\delta^{13}$ C values < -22 ‰. Thus, the use of both carbon isotopes could reveal whether the OC 81 is a mixture of GOC and OC from vegetation that is less than 50,000 years old. 82

One important question about the possible contribution of GOC in soils is whether the GOC is 83 mineralised when it becomes part of the soil. As GOC resists degradation once it has been deposited, it 84 can be assumed that it already exhibits a strong inherent recalcitrance. Nevertheless, this could also be 85 86 due to a physical protection that prevents microbial accessibility. However, when it becomes part of the subsoil during soil development, this OC pool could be degraded by the infiltration of water, oxygen, 87 fresh nutrients and microorganisms. Direct microbial coal degradation has already been observed in 88 89 incubation experiments on mine soils (Rumpel and Kögel-Knabner, 2002, Waschkies and Huttl, 1999) 90 or shale bedrocks directly exposed to the surface (Soulet et al., 2017). There has been no study to 91 establish whether GOC is degradable in OC-poor sediments or sedimentary rocks has not been 92 investigated, but it could differ since the amount of available OC can also drive microbial respiration 93 (Colman and Schimel, 2013). Therefore, these sediments might contain fewer microorganisms that can be spatially separated from the GOC, which may hamper its respiration. 94

95 To the best of the author's knowledge, there has only been one study by van der Voort et al. (2018) that 96 has investigated the amount of GOC in soils. They estimated that GOC makes up around 80 % of soil 97 OC in a moraine-derived soil, suggesting that GOC's contribution to soil OC is large. However, apart 98 from this study by van der Voort et al. (2018) on a very specific sediment, there have been no direct 99 calculations of the amount of GOC in soils.

The aim of this study was to quantify GOC in different terrestrial sediments and a sedimentary rock, and investigate its stability in incubation experiments in order to make assumptions about its possible contribution to soil OC stocks in soil profiles at the same site. The main research questions were: i) What is the relationship between sedimentary and subsoil OC contents? ii) Is OC in sediments <sup>14</sup>C-free and how much is really geogenic? iii) Will sedimentary GOC be degraded? and iv) How much does GOC contribute to soil OC?

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## 107 **2. Material and methods**

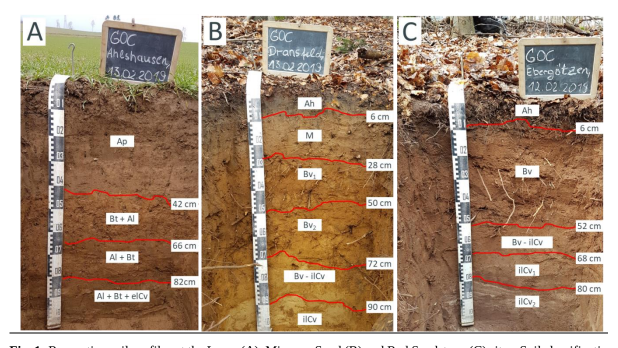
#### 108 2.1 Site description

109 Three sites were selected with different sedimentary bedrocks derived from a single geologic substrate that can be found close to the surface and is homogeneous down to 10 m depth. The sites represented 110 one sedimentary rock and two soft sediments. The sedimentary rock was a sandstone (Solling Formation, 111 112 Triassic) under European beech forest (Fagus sylvatica) 11.5 km north-east of Göttingen (51°35.012' N; 10°3.960' O), referred to below as "Red Sandstone". The soil is classified as a Folic Brunic Arenosol 113 according to the World Reference Base for Soil Resources (WRB, 2006). The sediments were loessic 114 115 deposits (Weichselian Glacial) that have been under agricultural land use for the past decades, 30 km north of Göttingen (51°48.101 N; 9°58.002' O), referred to as "Loess", and terrestrial sandy deposits 116 117 from the Miocene (Neogene formerly named Tertiary) in a European beech forest 13 km south-west of Göttingen (51°28.673 N; 9°45.323' O) referred to as "Miocene Sand". The respective soils have been 118 119 classified as a Haplic Luvisol and a Dystric Chromic Arenosol accordingly. The sediments at the Loess 120 site were deposited between the last glacial and interglacial periods between 115,000 and 400,000 years 121 BP, according to Jordan and Schwartau (1993). To the best of the author's knowledge, these forest sites have never been under agricultural use. The associated soils are classified as a Luvisol and a Cambisol 122 respectively. Mean annual air temperature and precipitation were 9.2 °C and 647 mm (1981-2010) 123 124 according to the nearby weather station that covers all three sites.

# 125 2.2 Sampling and sample preparation

Two 10-m long sediment cores with a diameter of 15 cm were drilled at each site in April 2017. For the soft sediments (Loess and Miocene Sand), drilling was conducted as percussion drilling, and for the hard sediment (Red Sandstone) as cable core drilling with water as the flushing solution. The replicates per site were drilled approximately 10 m apart. The sampled cores were subdivided into 1-m increments. For further chemical analysis, material was taken from a depth of 85-95 cm for each 1-m increment, and the outer 5 cm removed to avoid possible contamination. Thus the increment from 1-2 m for example, was represented by a sample from a 1.85-1.95 m depth.

One sample from each depth increment (1-2 m, 2-3 m, 3-4 m, 4-5 m, 5-6 m, 6-7 m, 7-8 m, 8-9 m, 9-10 m) for chemical analysis was oven dried at 60 °C and sieved to pass 2 mm. The Red Sandstone samples were crushed with a hammer before being dried and sieved. Additionally, approximately 1-m deep soil profiles were dug and soil samples taken from the different classified layers to obtain corresponding soil parameters (Fig. 1).



**Fig. 1:** Respective soil profiles at the Loess (A), Miocene Sand (B) and Red Sandstone (C) sites. Soil classification

140 was conducted using the German classification system for soil horizons (Eckelmann et al., 2006) and later

transferred to the WRB (2006). Depth transitions to the starting sediment were 82 cm for Loess (Al + Bt + elCv),

142 72 cm for Miocene Sand (Bv - ilCv) and 68 cm for Red Sandstone (ilCv<sub>1</sub>).

Samples were oven dried at 60°C and stored and sieved to pass 2 mm. For the determination of OC and
 <sup>13</sup>C, samples were ground in a planetary ball mill.

# 145 2.3 Chemical analysis and calculations

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146 Three aliquots of each sieved sample were analysed by dry combustion for total C and total N content 147 (TruMac CN LECO, St. Joseph, MI, USA). Samples with a pH value of > 6 (measured in 0,01 mol L<sup>-1</sup> 148 CaCl) were analysed for carbonates after the sample was ignited for 16 h in a muffle kiln at 450 °C 149 according to Nelson and Summers (1983), to remove the organic part of total C. The OC concentration 150 was calculated by subtracting carbonaceous C from total C, expressed as g OC kg<sup>-1</sup> dry matter. Homogenised samples were further analysed for  $\delta^{13}$ C values after carbonate removal in an isotope ratio 151 152 mass spectrometer (Delta Plus, Thermo Fisher, Waltham, MA, USA) coupled to an elemental analyser (FLASH EA 1122 NA 1500; Wigan, United Kingdom). Resulting  $\delta^{13}$ C values (‰) were expressed 153 154 relative to the international standard of Vienna Pee Dee Belemnite. The bulk densities for the soil samples were obtained with 250 cm<sup>3</sup> sampling rings from each layer of the soil profile. For the 155 sedimentary Loess and Miocene Sand samples, the bulk density of the deepest soil sample (94 and 100 156 157 cm respectively) was used. The bulk densities and densities without pore space of the intact Red Sandstone cores were determined on four subsamples (from 1.6, 3.6, 7 and 9 m depth) with a Dryflow-158 pycnometer (GeoPyc 1360) and a gas pycnometer (AccuPyc 1330) respectively. Bulk densities for the 159 missing depth increments were linearly interpolated. For radiocarbon (<sup>14</sup>C) analysis, the sediment 160 161 samples were first treated with 1% HCl to remove inorganic C, and then transferred into pre-combusted

- 162 quartz ampoules containing copper oxide and silver wool. The ampoules were evacuated, flame-sealed,
- 163 then combusted at 900  $^{\circ}$ C, and the CO<sub>2</sub> evolved was purified on a vacuum ring (Rethemeyer et al.,
- 164 2019). The  ${}^{14}C$  contents were measured with the MICADAS accelerator mass spectrometry (AMS)
- system at ETH Zürich, Switzerland. Where possible, one sedimentary sample per depth increment and
- site and one sample per soil layer was analysed. Due to the very low OC contents in some sediment
- samples, <sup>14</sup>C contents could only be determined for three samples from the Miocene Sand (from 1.9, 4.9
- and 7.9 m depth) and four from the Red Sandstone (1.9, 4.9, 7.9 and 9.9 m depth). For the Loess,  ${}^{14}C$  of
- bulk OC was measured in all depth intervals (1.9-9.9 m).
- 170 Total OC stocks (Mg ha<sup>-1</sup>) were calculated according to Eq. 1:

$$OC \ stock = \ OC \ \cdot BD \ \cdot (1 - stone \ content) \cdot depth \cdot 0.1$$
 Eq. 1

where *OC* is the weight-based OC content, either in the fine soil <2-mm fraction of the soil profiles or in the sediments (g kg<sup>-1</sup>), *BD* is the bulk density of the fine soil (g cm<sup>-3</sup>), stone content is the volumebased proportion of stones (cm<sup>3</sup> cm<sup>-3</sup>) and depth is the thickness of the depth increment (cm). To be able to compare OC stocks and contributions from GOC, it was decided to set the borders between the topsoils- and subsoils at 0.3 m and the transition from subsoils to the sediments at 1.5 m. According to Richter and Markewitz (1995) this represents a common border for the transition from soil to sediment. The sediments where further subdivided into an upper and a lower part at adepth of 4 m.

178 In a second step, the amount of GOC and biogenic OC in the sediments was calculated, considering 179 GOC as one carbon pool free of <sup>14</sup>C. For the sediments, the proportion of biogenic OC ( $f_{biogenic}$ ) of the 180 total amount of OC was calculated in a two pool model (Eq. 2) used by Cerri et al. (1985):

$$f_{biogenic} (\%) = \frac{F_{biogenic \, oC} - F_{GOC}}{F_{sample} - F_{GOC}} \cdot 100$$
 Eq. 2

181 where *F* represents the <sup>14</sup>C content in the fraction modern carbon ( $F^{14}C$ ) from a source compared with 182 the <sup>14</sup>C content of an oxalic standard (Stuiver and Polach, 1977, Torn et al., 2009). The sources were the 183 GOC fraction (*F<sub>GOC</sub>*), the sample (*F<sub>sample</sub>*) and the biogenic OC fraction (*F<sub>biogenic OC</sub>*). Since the <sup>14</sup>C content 184 of the GOC fraction can be set to zero, this equation could be simplified to:

$$f_{biogenic} (\%) = \frac{F_{biogenic \ OC}}{F_{sample}} \cdot 100$$
 Eq. 3

For the biogenic OC in the sediments, an average <sup>14</sup>C age ranging from 1,000-4,000 years BP was
assumed, based on published <sup>14</sup>C results of dissolved OC reaching greater depths (Artinger et al., 1996,
Schiff et al., 1997). The <sup>14</sup>C contents in the sediment from 2 to 4 m depth of the loess led to ages < 3,000</li>
years BP, and were therefore even younger than at 74 cm depth (4,413 years BP). Thus, they were treated

189 like the soil part for the calculation below of a GOC fraction. Respective times were converted into <sup>14</sup>C 190 contents ( $F_{biogenic OC}$ ) according to Torn et al. (2009):

$$F_{biogenic \, OC} = e^{\left(\frac{t}{-8033}\right)}$$
 Eq. 4

where *t* represents the <sup>14</sup>C age (1,000 or 4,000 years BP respectively) and 8033 represents the mean life of radiocarbon in years. The proportion of GOC in the sediments ( $f_{GOC}$ ) is therefore the portion left (Eq. 5):

$$f_{GOC} = 100 \% - f_{biogenic}$$
Eq. 5

For the depth increments without measured <sup>14</sup>C ages (Tab. S1), the calculated amounts were linearly 194 195 interpolated with measured <sup>14</sup>C ages from samples above and below. This was done by assuming a depthdependent correlation and using the adjacent values. To calculate the amount of GOC in the soil profiles, 196 the weight-based amount of GOC in the sediments was first calculated by multiplying its fraction ( $f_{GOC}$ ) 197 by the respective OC content (in g OC per kg<sup>-1</sup> dry mass). The median amount of GOC (g GOC per kg<sup>-1</sup> 198 <sup>1</sup> dry mass) of these sedimentary values was then taken and its proportion of soil OC content calculated 199 (g OC per kg<sup>-1</sup> fine soil) in the soil profile. This was done for the proportion of GOC in the sediments 200 calculated with a 1,000 and a 4,000-year-old biogenic OC fraction ( $F_{biogenic OC}$  in Eq. 3) to obtain a range 201 202 of GOC contributions. It was assumed that the GOC fraction resisted degradation during soil formation. 203 Therefore, this proportion represents the largest possible amount of GOC that could contribute to soil 204 OC stocks. On this assumption, it was also possible to define the influence of GOC in the soil profile on the resulting <sup>14</sup>C ages. Since the calculated <sup>14</sup>C ages represent a mixture of the <sup>14</sup>C content from GOC 205 and the biogenic fraction (Eq. 5), the GOC fraction has the same influence on the soil <sup>14</sup>C age as on bulk 206 207 OC (according to Eq. 3). Reducing the age by this fraction would therefore represent an "unbiased" age 208 of soil-derived-OC.

# 209 2.4 Incubation experiment

To assess the potential stability of OC in the sediments against microbial decay, two laboratory 210 incubation experiments were conducted at 20 °C for 50 and 63 days respectively. This was done to 211 reveal the potential degradation of OC from the sediments. A temperature of 20 °C was chosen because 212 only very low degradation rates were expected at lower temperatures. The first, 50-day experiment was 213 conducted with intact Red Sandstone core samples, while the second experiment was performed after 214 215 the Red Sandstone was crushed to pieces < 2 mm to simulate the process of weathering when the intact 216 sediment or sedimentary rock becomes part of the (sub)soil. For the incubations, four subsamples of 1,340-6,890 g per sample from different depth intervals were used. The sample material was stored at 217 218 room temperature until the start of the incubation experiment. Four samples were taken from each 219 sediment and from each of four depth ranges for Miocene Sand (1.2-2.8, 3.2-4.8, 6.1-7.8 and 8.2-9.9 m), 220 Red Sandstone (2-3, 4-5.8, 7-8 and 8.4-9.8 m) and Loess (1.4-2, 2.7-3, 4.8-6 and 9.1-10 m). Water

content was adjusted to correspond to 40 % of water-holding capacity based on the poured bulk density, 221 determined by filling the loose material into a defined volume and measuring its weight. The material 222 223 for the four repetitions was mixed in large plastic vats and water added before the respective four 224 subsamples were transferred to incubation vessels. There were also four blank samples with no material. 225 Based on preliminary tests and their calculated bulk density and porosity, the intact Red Sandstone 226 samples were kept in a barrel with pure water for 14 hours to reach a water content of nearly 40 %. 227 Samples were placed in polycarbonate vessels with a volume of 7069 cm<sup>3</sup> and made airtight. The lids contained two tube connectors so that the samples could be flushed with ambient air. After flushing, 228 229 samples were set to a starting pressure of about 1,300 mbar and kept closed until the end of the 230 incubation. Nine gas samples were taken in evacuated glass vials (20 mL), 0, 3, 7, 13, 21, 30, 59 and 63 231 days after the start of incubation. Samples were analysed for CO<sub>2</sub> concentrations by gas chromatography (Agilent 7890A, GC, Agilent Technologies, Santa Clara, USA) to account for the amount of 232 233 accumulated CO<sub>2</sub>. Three additional gas samples were taken 0, 30 and 63 days after the incubation started and analysed with an isotope ratio mass spectrometer (Delta Plus XP, Thermo Fisher Scientific, Bremen, 234 Germany) to account for the development of  $\delta^{13}C$  of CO<sub>2</sub> during the respiration. The corresponding 235 pressure was measured on each sampling date. When the over-pressure of a vessel was lost due to 236 237 leakages, it was removed from the sampling because contamination with ambient air could no longer be 238 ruled out. This happened for a third of all the samples.

239 The amount of respired CO<sub>2</sub>-C (mg CO<sub>2</sub>-C  $d^{-1}$ ) was calculated using Eq 6.

$$CO_2 - C = \frac{0.1 \cdot p \cdot x_i \cdot M \cdot V}{R \cdot T \cdot t}$$
 Eq. 6

where *p* is the pressure (mbar),  $x_i$  is the difference in CO<sub>2</sub> concentration between the samplings (ppm), *M* is the molar mass of C (g mol<sup>-1</sup>), *V* is the air volume of the sample (m<sup>3</sup>), *R* is the molar gas constant (J kmol<sup>-1</sup> K<sup>-1</sup>), *T* is the incubation temperature (K) and *t* is the elapsed time (d) between the samplings. Based on the  $\delta^{13}$ C values of CO<sub>2</sub> the proportion of CO<sub>2</sub> derived from carbonates was subtracted where necessary according to Bertrand et al. (2007). This respiration rate was related to the OC content of the samples (called "OC-normalised respiration") by being divided by the total amount of OC in g in the sample.

- Since we observed an almost linear respiration behaviour was observed in both incubation experiments, a simple linear regression model was fitted to describe the mineralisation rate per time (mineralised OC (%) =  $k \cdot t$ ).
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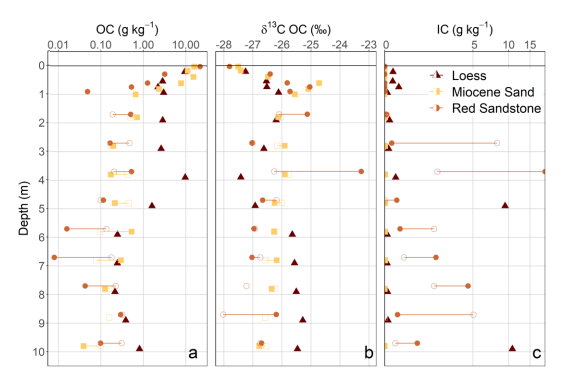
252 Statistical analyses were conducted using the statistical environment R (R Core Team (2018)) including the function "lm" to fit linear models and the package ggplot2 (Wickham, 2016) for graphical 253 presentation. The models were tested for deviations from homoscedasticity, normality of residuals and 254 255 absence of collinearity. The tests revealed heteroscedasticity of the residuals, which can be explained 256 by an increasing standard deviation with time. It subsequently became apparent that the residuals were 257 not normally distributed since the dependent variable, representing the proportion of OC being 258 mineralised, only allows values between 0 and 1. Bearing this in mind, the results should be treated as 259 an indicator of differences between the samples and a scale for the mineralisation of the OC pool. 260 Therefore, calculations of standard errors and significance of the parameters were omitted since this 261 would not produce reasonable results with the model used.

#### 262 **3. Results**

# 263 3.1 Relationship between sedimentary and subsoil organic carbon

264 Organic carbon was detectable in all the sediments analysed using three laboratory replicates per sample 265 and comparing them with the muffled samples, measurable. The mean relative standard deviation of the 266 laboratory replicates was 9.5 %. In terms of the detection limit, the sample with the lowest total C content (mean of 0.04 g C kg<sup>-1</sup> soil) showed values between 0.00 and 0.01 g C kg<sup>-1</sup> soil after removal of OC at 267 450 °C. Thus the range of 0.00 to 0.01 g C kg<sup>-1</sup> soil was assumed to be the mean standard error from the 268 measurement. Despite having the same material down to 10 m depth at each site, there were still some 269 270 inhomogeneities that were visible and measurable. This was especially true for Loess and Red Sandstone. The amount of OC in the sediments from 1 to 10 m depth was comparatively low in Miocene 271 Sand and Red Sandstone (0.04-0.71 g kg<sup>-1</sup> and 0.01-0.53 g kg<sup>-1</sup>, respectively), while much higher OC 272 273 contents of 0.21-9.71 g C kg<sup>-1</sup> were found in Loess (Fig. 2 a). The median OC content of the sediments 274 was within a range comparable to that of the respective deepest subsoil horizon. This deepest horizon 275 was a Cv horizon at 94 cm depth for Loess, 100 cm depth for Miocene Sand and 74 cm depth for Red 276 Sandstone. In detail, the median OC content in the sediments, compared with the respective Cv horizons, 277 corresponded to 27 % for Loess, 29 % for Miocene Sand and 39 % for Red Sandstone. The Loess OC 278 contents were highly variable, highlighting the changing sedimentary conditions during the past glacial 279 and interglacial periods (Jordan and Schwartau, 1993). At 4-5 m depth, the OC contents of Loess were even higher (9.7 g kg<sup>-1</sup>) than in the subsoil (3 g kg<sup>-1</sup>). This was not an outlier because the high OC content 280 could be visually confirmed by the very dark colour of the sample. In Miocene Sand and Red Sandstone, 281 no clear depth gradient of OC was found at 2-10 m depth (Fig. 2 a). Even though the OC content in the 282 sediments were low, the OC stocks could be very large. A comparison of OC stocks in topsoils (0-0.3 283 m), subsoils (0.3-1.5 m) and the sediments down to 10 m depth revealed quite high OC stocks in the 284 sediments. For Loess, OC in the sediment contributed up to 71 % of the total OC amount while it was 285 286 51 % for Red Sandstone and 21 % for Miocene Sand (Table 1).

The distribution of the  $\delta^{13}$ C values of OC in the soil and sediment profiles showed an increase in  $\delta^{13}$ C 287 with depth in the soil down to 1 m (Fig. 2 b). In contrast, the  $\delta^{13}$ C values of OC in the sediments showed 288 no clear trend with increasing depth, but they all were within the range of C<sub>3</sub> plant material. A value 289 290 above -25 ‰ for Red Sandstone at 4 m depth could be explained by the high values of inorganic carbon (IC) at this depth. It can be assumed that decarbonisation of this sample was not completely successful. 291 Unexpectedly high amounts of IC were found in parts of Red Sandstone, indicating the presence of 292 293 calcareous deposits in this terrestrial material (Fig. 2 c). The Loess also included some distinct calcareous layers at 5 and 10 m depth, while there were only small amounts of around 0.1 mg IC g<sup>-1</sup> soil 294 at the other depths. This could be due to the fact that the investigated loess deposits belong to the "Leine 295 296 Ilme Basin", a region with aoelian loamy loess that has been decalcified during weathering and soil 297 genesis (Wagner, 2011), while the contents in the soil profile can be explained by liming. No IC was 298 present in Miocene Sand.



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**Fig. 2:** Depth distribution of different bulk properties of the soil profiles and deep drilling cores. Presented parameters include the log scale organic carbon (OC) and inorganic carbon (IC) content (a and c) and the  $\delta^{13}$ C values of the organic carbon (b) related to the amount of fine soil or dry mass respectively. Filled and unfilled symbols represent the two different cores. For the Loess, only one core could be analysed.

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Table 1: OC stocks and proportions for the three sites down to 10 m depth. Proportions of biogenic and geogenic
 OC were calculated based on <sup>14</sup>C results and assumptions described in the material and methods section.
 Represented ranges are calculated based on the assumption of a 1,000 or 4,000-year-old biogenic OC fraction
 reaching the sediments.

			TOC		OC stoc	OC stocks (Mg ha <sup>-1</sup> )		Proportion of OC (%) <sup>b</sup>		
Substrate	Layer	Depth (m)			geogenic	biogenic	geogenic	biogenic		
		-	(Mg·ha <sup>-1</sup> )	(%) <sup>a</sup>	4,000 1,000 yrs. yrs.	1,000 4,000 yrs. yrs.	4,000 1,000 yrs. yrs.	1,000 4,000 yrs. yrs.		
	Topsoil	0.0 - 0.3	40.6	11	1.1 - 1.2	40 - 39	3 - 3	97 - 97		
Loess	Subsoil	0.3 - 1.5	66.0	17	5.1 - 5.3	61 - 61	8 - 8	92 - 92		
LUESS	Upper Sediment	1.5 - 4.0	218.6	57	10.6 - 11.0	208 - 208	5 - 5	95 - 95		
<u> </u>	Lower Sediment	4.0 - 10.0	55.1	15	42.9 - 46.7	8 - 12	78 - 85	15 - 22		
	Topsoil	0.0 - 0.3	22.2	30	0.6 - 0.7	22 - 22	3 - 3	97 - 97		
Red	Subsoil	0.3 - 1.5	13.7	19	1.6 - 1.7	12 - 12	12 - 12	88 - 88		
Sandstone	Upper Sediment	1.5 - 4.0	18.7	25	13.1 - 14.9	4 - 6	70 - 80	20 - 30		
	Lower Sediment	4.0 - 10.0	19.3	26	14.1 - 15.7	4 - 5	73 - 82	18 - 27		
	Topsoil	0.0 - 0.3	39.1	31	0.3 - 0.3	39 - 39	1 - 1	99 - 99		
Miocene	Subsoil	0.3 - 1.5	60.8	48	1.3 - 1.6	59 - 60	2 - 3	97 - 98		
Sand	Upper Sediment	1.5 - 4.0	10.6	8	3.7 - 5.8	5 - 7	34 - 55	45 - 66		
l	Lower Sediment	4.0 - 10.0	16.3	13	8.3 - 10.8	6 - 8	51 - 66	34 - 49		
30	<sup>a</sup> % of total O	OC stock from 0-	-10 m							

<sup>b</sup> % of OC stock in the respective depth increment 310

#### 311 3.2 Ages of organic carbon in soils and sediments and contributions from geogenic organic carbon

312 The ages of OC in the Loess soil profiles revealed a modern carbon signature (0 years BP) at 0.3 m

313 depth, with a sharp increase up to  $4,413 \pm 51$  years BP at 0.7 m depth (Fig. 3 a). For the Red Sandstone 314 soil profile there was only an increase in the ages from a modern signature at 0.04 m to  $532 \pm 41$  years 315 BP at 0.3 m depth. The Miocene Sand soil profile at 0.4 m depth showed an increase from  $1,277 \pm 41$ 316 to  $1,771 \pm 44$  years BP at 0.6 m depth. Thus, OC of the subsoil (around 0.6 m depth) in Loess was more

317 than twice as old as in Miocene Sand. In contrast, Loess had a modern signature at 0.3 m, while the soil

318 developed in Red Sandstone showed an average age of  $532 \pm 41$  years BP at 0.3 m depth.

319 The ages of OC in the sediments ranged from 2,200-30,730 years BP, with respective mean ages of

 $9,077 \pm 3,234$  years BP for Miocene Sand,  $13,674 \pm 9,632$  years BP for Loess, and  $14,463 \pm 1,992$  years 320

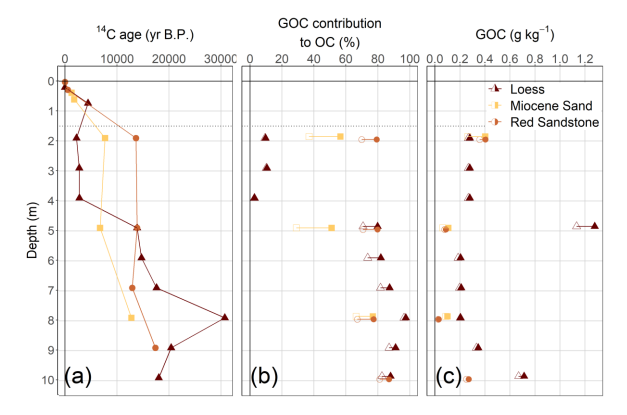
BP for Red Sandstone. For all sediments, 11 out of 16 samples had a <sup>14</sup>C content that led to an <sup>14</sup>C age 321

322 older than 11,600 years BP, which we assumed to be the time after the latest glacial period when soil

development started (Litt et al., 2007). Therefore, the sediments contained a mixture of geogenic (<sup>14</sup>C-323 324 free) and biogenic (with <sup>14</sup>C) OC. Despite being the youngest sediment, Loess partly revealed the highest

- apparent <sup>14</sup>C ages up to  $30,730 \pm 631$  years BP (Fig. 3 a). The ages of OC in the sediment of Red 325
- 326 Sandstone and Miocene Sand ranged from  $12,940 \pm 132$  to  $17,390 \pm 206$  years BP and from  $6,750 \pm 86$
- 327 to  $12,770 \pm 151$  years BP respectively, revealing no depth trend with age in the deeper sediment. The
- 328 calculated GOC fraction in the sediments was highest for Red Sandstone, ranging from 67 to 87 %, with
- 329 a mean of 77 % (Fig. 3 b). For the three Miocene Sand samples, the GOC fraction ranged from 29 to 77

- % with a mean of 53 %. Loess showed a sharp increase at a depth of five metres where GOC contribution
  rising to between 71 to 98 % while it was only 5 to 19 % at the 2 to 4 m depth.
- 332 The calculated weight-based content of GOC in the sediment revealed a comparatively uniform
- distribution for all the sediments with depth, except the extremely high contents of Loess at 5 m depth
- (Fig. 3 c). The investigated sediment depths revealed quite a narrow range of GOC contents with 0.10
- $\pm 0.03$  g kg<sup>-1</sup> for Miocene Sand,  $0.17 \pm 0.12$  g kg<sup>-1</sup> for Red Sandstone and  $0.27 \pm 0.08$  g kg<sup>-1</sup> for Loess.



336

Fig. 3: Depth distribution of apparent <sup>14</sup>C ages (in years BP) (a), GOC contribution to OC contents in the sediments (b) and
resulting weight based amounts of GOC (c) in the sediments. The range in the contribution from GOC is due to the assumption
of an average biogenic OC age of 4,000 (empty shapes) and 1,000 years (filled shapes).

#### 340 *3.3 Biodegradability of sedimentary derived organic carbon*

The incubation experiment revealed a potential, but low biodegradability of OC for all samples, but 341 without a clear depth gradient from 1 to 10 m (Fig. 4). To compare the effect of crushing on the 342 343 respiration of Red Sandstone samples, the mineralisation rates from the first incubation experiment were compared with those of the second incubation experiment (Table 2). While Red Sandstone showed very 344 low mineralisation when the samples were incubated as intact cores (0.3-1.0 mg CO<sub>2</sub>-C g<sup>-1</sup> OC y<sup>-1</sup>), the 345 mineralisation rate constants were up to five times higher when the samples were crushed (1.0-2.1 mg 346 CO<sub>2</sub>-C g<sup>-1</sup>OC y<sup>-1</sup>). In a comparison of Loess and Miocene Sand, both revealed large differences between 347 samples with the lowest (1.2 and 1.8 mg CO<sub>2</sub>-C g<sup>-1</sup> OC y<sup>-1</sup>) and highest (34.2 and 12 mg CO<sub>2</sub>-C g<sup>-1</sup> OC 348 y<sup>-1</sup>) respiration rate constants. Interestingly, there was no depth gradient for the different substrates, but 349

samples from 7 and 9 m depth of Miocene Sand tended to have up to 6.8 times lower respiration rate
constants than samples from the 2 and 4 m depth. There was also a very wide variation between samples
from the same depth (see Fig. A1) as revealed by the high standard deviations. Assuming a constant
mineralisation rate over time, the results of the incubation experiment would result in mean residence
times of between 29 and 135 years for the Loess, 83 and 556 years for the Miocene Sand and between
476 and 1,000 years for the crushed Red Sandstone. Since the CH<sub>4</sub> levels of the samples remained at a

low level (Fig. A3) there were no indications of oxygen-limited conditions during the incubation.

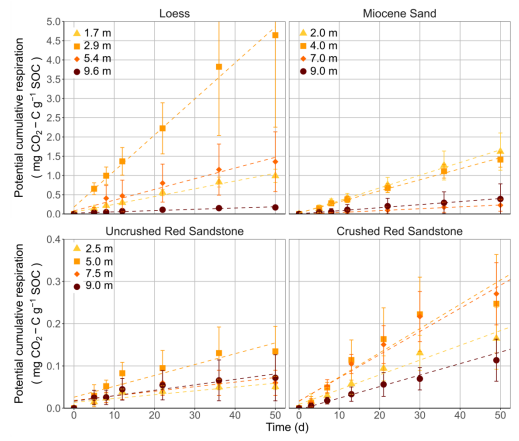
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Table 2: Comparison of mineralisation rate constants between the first (intact Red Sandstone samples) and second incubation
 experiments (crushed Red Sandstone samples) calculated using linear models.

Substrate	Depth (m)	Mineralisation rate constant (mg CO <sub>2</sub> -C g <sup>-1</sup> OC y <sup>-1</sup> ) <sup>a</sup>
	2	12.0
Maria Card	4	10.5
Miocene Sand	7	1.8
	9	2.9
	1.7	7.4
T	2.9	34.2
Loess	5.4	10.2
	9.6	1.3
	2.5	0.3
Uncrushed	5	1.0
Red Sandstone	7.5	0.4
	9	0.4
	2.5	1.3
Crushed	5	2.1
Red Sandstone	7.5	2.0
	9	1.0

360 <sup>a</sup> represents the slope of the fitted linear model for respiration during the incubation experiment

361



362

Fig. 4: Potential degradability of sedimentary OC from three sites. Results represent cumulative respiration from
 Loess and Miocene Sand samples and both Red Sandstone incubation experiments with uncrushed and crushed
 samples, with respective standard deviations (n = 4). Dashed lines represent a fitted linear model to the respiration
 data.

#### 367 *3.4 Possible contribution from geogenic organic carbon to soil organic carbon*

As bedrock weathers, it becomes part of the soil and GOC also becomes soil OC. The potential amount 368 369 of GOC in the subsoil was dependent on the sedimentary bedrock (Fig. 5). In the subsoil of Miocene 370 Sand the proportion of GOC amounted to 2-3 %, at the Loess site it was 8 %, and at the Red Sandstone 371 site it was 12 % (Table 1). For the defined topsoils (0-30 cm depth), contributions of GOC to soil OC were smaller with 0.7-0.9 % for Miocene Sand, 2.8-2.9 % for Loess and 2.8-3.0 % for Red Sandstone 372 373 The possible contribution of GOC to soil OC was also calculated under the assumption that the biogenic fraction had an unrealistically high average age of 10,000 years (Fig. A5). This resulted in the GOC 374 375 fraction falling to 0 for Miocene Sand. For Loess the geogenic fraction fell by  $\sim 25$  %, and for Red 376 Sandstone by  $\sim 16$  %.

- The presence of <sup>14</sup>C-free GOC in soil OC reduced the mean bulk soil OC <sup>14</sup>C ages, depending on its proportion in soil OC content. Topsoils that developed in Loess and Red Sandstone had a modern <sup>14</sup>C content of 1.029 and 1.035 F<sup>14</sup>C, similar to the atmospheric <sup>14</sup>C content in 1950. Due to the large proportion of biogenic OC, no influence of a geogenic fraction was detected in the topsoil of these sites.
- 381 No <sup>14</sup>C data were available for the topsoil of Miocene Sand. For all subsoils, the influence of GOC on

bulk soil OC <sup>14</sup>C contents depended greatly on the depth and corresponding OC contents. For Loess, the 382 383 possible influence on <sup>14</sup>C ages in the subsoils was quite high, with an average of 10 % reduction in mean apparent <sup>14</sup>C ages in the subsoil. Thus it would reduce the measured age of 4,413 years BP in 74 cm 384 depth by 532-555 years BP. Geogenic OC potentially reduced the mean apparent radiocarbon age of 385 1,277 years BP at 0.39 m depth in Miocene Sand by about 7-9 years, and the radiocarbon age of 1,771 386 years BP at 0.61 m depth by 20-24 years. These reductions are below the respective standard deviations 387 388 of the measurement. Nevertheless, at 1 m depth a given possible proportion of 13.1-16.3 % would reduce an interpolated <sup>14</sup>C age of 3,053 years BP by 399-497 years For Red Sandstone the influence of GOC 389 390 on <sup>14</sup>C ages would be highest in the subsoil. At 74 cm depth it would increase an age of 1,453 years BP 391 by 451-490 years BP. Due to the low amounts of soil OC at 90 cm depth at the Red Sandstone site, the 392 weight-based median amount of GOC in the sediments was four times higher than the biogenic amount 393 of soil OC.

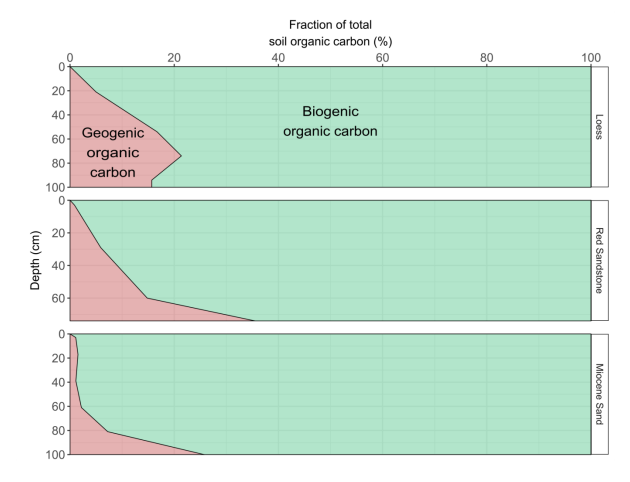


Fig. 5: Largest possible contribution of GOC to OC (red area) in relation to bulk OC content, taking into account the median
 GOC contents of the sediments for the respective horizontal weight based OC contents. The contribution of GOC is the mean
 amount based on the assumption of an average biogenic OC age of between 4,000 and 1,000 years

# 398 4. Discussion

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#### 399 *4.1 Geogenic OC in the sediments*

Generally, our calculations on the GOC fraction in the sediments are based on the assumption that 400 biogenic OC in the sediments is not older than 4,000 yrs BP on average. And we also excluded the 401 402 influence of a biogenic OC fraction that derives from soils that developed before the latest glacial period. 403 Thus, there is uncertainty of a biogenic OC fraction in the sediments since it is unknown when biogenic 404 OC entered the sediments. We assumed a mean age of 1000 to 4000 years based on DO<sup>14</sup>C data that was 405 leached from the soil. Nevertheless, even with an assumed age of 10,000 years for the biogenic OC 406 fraction, the highest possible contribution was 15 % for the Loss (94 cm depth) and 22 % for the Red 407 Sandstone (74 cm depth) (Fig. A5). A mean age of 10.000 years would be an unrealistic assumption 408 since sediments are open systems and may receive OC input throughout the pedogenic period if 409 vegetation is present and not only at the start of pedogenesis.

#### 410 *4.1.1 Site dependent contents of GOC*

411 Regarding the calculated contribution of GOC to OC in the sediments, the assumed range of biogenic 412 <sup>14</sup>C ages from 1,000-4,000 years BP was within the typical range for ages of dissolved OC leaching from 413 soils (Artinger et al., 1996, Jia et al., 2019). Nevertheless, the range of 1,000-4,000 years BP did not 414 greatly influence the range of calculated sedimentary contribution from GOC, especially for Loess (~ 6 % difference) and Red Sandstone (~ 9 % difference). The calculated GOC contribution for Miocene 415 416 Sand was comparatively low (29-77 %) compared with the contribution for Red Sandstone and Loess, 417 especially at 5 m depth. Compared with the Red Sandstone, this could be due to deep biogenic carbon 418 inputs, such as roots and root exudates from the trees (Angst et al., 2016, John et al., 2016, Kirfel et al., 419 2017, Tückmantel et al., 2017), since the loosely bedded Miocene Sand allows for deep infiltration 420 compared with the Red Sandstone site with its shallow bedrock as a root-restricting layer (Schneider 421 and Don, 2019). Although it was not possible to define the exact rooting depth for Miocene Sand, this 422 depth probably does not exceed 4 m, according to Schenk and Jackson (2005). Therefore, the increase 423 in sedimentary contribution from 5 to 8 m depth of Miocene Sand could be due to the decreasing 424 influence of roots and root exudates at 8 m depth.

In Loess, the low  $^{14}$ C ages at 2-4 m depth (2,200-2,770 years BP) in contrast to a  $^{14}$ C age of 4,413 years 425 426 BP at 0.74 m depth were surprising. This might indicate past anthropogenic activities or erosion-driven 427 material movement that might have led to a mixing of the upper part of the profile. Furthermore, the 428 modern <sup>14</sup>C signature at 21 cm depth could be due to the plough layer at the Loess site mixing the upper 30 cm. Nevertheless, at depths below 4 m the high <sup>14</sup>C ages of OC in the sediments indicated a large 429 430 proportion of GOC. This could be due to different sedimentation periods, soil forming and also soil burial processes (Chaopricha and Marin-Spiotta, 2014) that took place during the Pleistocene. These 431 432 processes can lead to the presence of buried layers in Loess with varying amounts of rather recalcitrant 433 OC, as shown by Marin-Spiotta et al. (2014). For the investigated Loess site, different sedimentation 434 and soil forming processes can be expected due to the presence of completely different material in the 435 cores in terms of colour and measured OC contents. For example, the very dark Loess at 4 m depth with

its high OC content support the assumption of sedimentation circumstances that favoured the
accumulation and preservation of OC. This is in accordance with Jordan and Schwartau (1993) who
investigated the same site and assigned the different layers to specific Pleistocene sedimentation periods.

In summary, the contribution of GOC to sedimentary OC was substrate dependent. A loosely bedded sediment like Miocene Sand with extremely low concentrations of OC could be more prone to infiltration of biogenic OC and dilution of GOC. This resulted in contributions of biogenic OC to the sediments of about 50 %. In contrast, the Loess site with comparatively low infiltration rates or the Red Sandstone site with reduced possibilities for deep rooting seemed to contain relative constant contributions from GOC of around 80 %.

445 The GOC contribution within the sediments was not found to increase with soil depth. This is in contrast 446 to the results of Frouz et al. (2011) which showed that different sediment types from a Miocene clay 447 sediment had higher weight-based carbon contents at 150 m compared with 30 m depth. However, a 448 comparison with the present study is difficult since Frouz et al. (2011) did not distinguish between the 449 geogenic and biogenic OC fractions, and OC contents were much higher (28-112 g kg<sup>-1</sup> dry mass compared to 0.008-10 g kg<sup>-1</sup> dry mass in our results). However, it also underlines the importance of 450 451 different sedimentation processes for the amount and depth distribution of OC in sediments and 452 sedimentary rocks. 4.1.2 Geogenic OC in Miocene Sand and Red Sandstone compared with other studies

It is hard to compare the weight-based amounts of GOC in terrestrial sediments since most studies in 453 454 this field rarely determine the amounts of OC in terrestrial sediments, or presume that sandy sediments, 455 for example, do not contain large amounts of OC (Artinger et al., 1996). Quite high amounts of OC have been found in the skeleton part of different soils on sandstones with 0.61-1.97 t ha<sup>-1</sup> (Corti et 456 457 al.,2002). Nevertheless, they mentioned the possible strong influence of organic substances from the 458 soil solution without quantifying it, and did not directly investigate OC in the sediments. Additionally, Copard et al. (2007) assumed an OC amount of 2.4 g kg<sup>-1</sup> from an unknown source for all sandy 459 460 sediments in a global storage modelling approach for the first metre of sediments. This would fit with the amounts of around 1 and 5 mg OC g<sup>-1</sup> found by Krummholz et al. (1997) for Dakota sandstone layers 461 at > 180 m depth, but is much greater than the median GOC amount in Red Sandstone (0.2 g kg<sup>-1</sup>) and 462 463 Miocene Sand  $(0.1 \text{ g kg}^{-1})$  in the present study.

# 464 *4.1.3 Geogenic OC in Loess compared with other studies*

Loess deposits are relatively well investigated because they provide a record of paleoenvironmental conditions (Hatté et al., 1998, Head et al., 1989, Murton et al., 2015, Wang et al., 1996). The median amount of 0.27 g kg<sup>-1</sup> from the present study was low compared with studies by Hatté et al. (1998), Wang et al. (1996) and Strauss et al. (2012). Hatté et al. (1998) investigated 20-m deep loess deposits in the Rhine valley and found OC contents between 1.0-8.6 g kg<sup>-1</sup>, Wang et al. (1996) investigated 12-

470 m deep loess deposits in China and found OC contents of  $31.2 \pm 30.5$  g kg<sup>-1</sup> and Strauss et al. (2012)

found OC contents of  $15 \pm 14$  g kg<sup>-1</sup> in Yedoma loess deposits in Siberia. This shows that the deposits from the site in the present study stored comparatively low GOC contents, although most of the abovementioned studies do not distinguish between a biogenic and geogenic OC pool. Nevertheless, as also shown in the present study, Loess had a high OC content compared with other sediments. This was in line with highest OC contents in subsoils at the Loess site and may indicate the importance and contribution of bedrock OC to subsoil OC.

#### 477 *4.2 Is sedimentary derived organic carbon biodegradable?*

478 The incubation experiment revealed a mineralisation of OC within the sediments with values between 479 0.1 and 3.4 % of total OC being mineralised after one year, assuming a constant mineralisation rate. The incubation temperature of 20 °C is above the typical mean temperatures in the subsoils underlining 480 incubation derived mineralisation rates to be potential rates. For subsoils with comparable climatic 481 482 conditions, Wordell-Dietrich et al. (2019) found temperatures at 150 cm depth ranging from 4 to 14.4°C 483 over a two-year period. Assuming a  $Q_{10}$  value of around two for the assumable difference (Hamdi et al., 484 2013), the respiration rate at typical subsoil temperatures would be roughly half that. A direct 485 mineralisation of OC from sediments is in agreement with several studies investigating the direct 486 mineralisation from outcrops (Copard et al., 2007, Horan et al., 2017, Petsch et al., 2000, Soulet et al., 487 2017) Seifert et al., 2001). The difference to the present study is, that those studies observed this 488 mineralisation when the sediments were directly exposed to the surface or/and were part of a very rapidly 489 eroding area with GOC from the sediments getting inputs of recent vegetation. However, Frouz et al. 490 (2011) conducted an incubation experiment with sedimentary samples from OC-rich Miocene clay sediments and found quite high respiration rate, with values between 0.4-1.2 % OC loss per year during 491 492 a 91 day incubation experiment.. Kieft and Rosacker (1991) also found high respiration rates of 493 sedimentary samples, with values between 0.1-1 % OC loss per year. Those results are in fairly good 494 agreement with the respiration rate constants observed in the present study. Compared with subsoil 495 incubation experiments, the mineralisation we found for sediments was quite low. For example, in a subsoil incubation experiments at 20 °C, Wordell-Dietrich et al. (2017) found that between 5-9.5 mg 496 CO<sub>2</sub>-C g<sup>-1</sup> OC of OC are mineralised after incubation for 63 days, Wang et al. (2013) report values 497 between 5-15 mg CO<sub>2</sub>-C g<sup>-1</sup> OC after 28 days, and Soucemarianadin et al. (2018) report values between 498 10- and 12.5 mg CO<sub>2</sub>-C g<sup>-1</sup> OC after 70 days. The difference between the respiration rates observed in 499 500 the present incubation experiment and the results from sedimentary and subsoil incubation experiments could be due to different microbial communities, OC quality and the physical disconnection between 501 502 OC and potential decomposers.

503 Our study indicated that there was a considerable portion of biogenic OC in the sediments. The observed 504 mineralisation could therefore be mainly due to the consumption of this biogenic OC part. Furthermore, 505 the low mineralisation rate of Red Sandstone during the incubation as intact cores (Fig. A1) promoted 506 the stability of GOC when it is part of the sediments. The low mineralisation rates of the OC in the sediments might be caused by a lack of fresh substrates and/or microorganisms that could enhance the degradation of OC (Fontaine et al., 2007). Seifert et al. (2011) have shown that microorganisms are able to degrade sedimentary OC after the addition of glucose in black slate outcrops. Nevertheless, mineralisation of OC could only be observed with the addition of water, which indicates the widely recognised fact of the presence of an active microbial community in the sediments (Bomberg et al., 2017, Joergensen and Wichern, 2018, Magnabosco et al., 2018).

514 The incubation experiment was unable to answer the question of whether GOC is mineralised when it becomes part of the (sub-)soil. Assuming that a large part of the biogenic OC in the sediments was 515 516 mineralised during the incubation experiment, geogenic OC could still be preserved during soil 517 formation especially in the subsoils. This is in accordance with the indirect approach taken by Graz et 518 al. (2010) to determine the mineralisation of sedimentary OC when it becomes part of the subsoil. They 519 stated that 30 % of GOC resists degradation when it becomes part of the soil due to the results of a 520 quantitative palynofacies analysis of bedrock and soil samples. Hemingway et al. (2018) found that 521 sedimentary OC directly exposed to the surface in a rapidly eroding tropical mountain area exhibits 522 considerable mineralisation down to 1 m below the surface, also leading to around 30 % of GOC remaining in the soil. Based on <sup>14</sup>C measurements they found out that on average  $67 \pm 11$  % of the OC 523 524 fraction in the sediments could be lost during soil formation, but no distinction was made between biogenic and geogenic OC fractions. This indicates that a microbial mineralisation of bedrock OC takes 525 526 place but may be partly restricted to biogenic OC.

527 Regarding the depth distribution of GOC in the sediments, the amount of GOC (in  $g kg^{-1}$ ) did not 528 increase with depth, but there were clear differences. This reflects the sedimentation history with different initial amounts of OC and degradation during sedimentation, which is particularly evident in 529 530 the high amounts of GOC at 5 m depth in Loess. However, the contents of GOC, especially in Red 531 Sandstone and Miocene Sand, were within the same range for the whole depth. This might indicate that degradation of GOC is not depth-dependent within the sediments. If there were a stronger degradation 532 533 of GOC with decreasing depth, a decreasing amount of GOC could be expected due to the input of water, 534 microorganisms and fresh nutrients from above. Furthermore, there was a relatively constant 535 contribution of biogenic OC within the sediments, meaning that if biogenic OC enters the sediments 536 with possibly degrading microorganisms, this biogenic OC might also be largely mineralised. A study by Heitkötter et al. (2018) demonstrate effectively that degradation of OC in subsoils is primarily limited 537 to small hotspot areas. Also in sediments, Krummholz et al. (1997) show that microbial communities 538 539 can particularly be found in spatially discrete areas. Nevertheless, Heitkötter et al. (2018) also show that microorganisms outside the hotspots can be activated when substrate is supplied. Thus, the 540 bioavailability of GOC might be site-dependent since root channels as microbial hotspots, for example, 541 542 are less abundant and stable in sandy soils (Schneider and Don, 2019). With regard to the sites

investigated here, the solid Red Sandstone might only obtain water in preferential flow paths (Swanson, 543 544 2006) leading to a comparatively stable OC pool in the sediment. Meanwhile for weathered, poorly structured sandy soils, prevalent matrix flow conditions can be assumed (Flury et al., 1994). Thus, the 545 546 bulk GOC might be supplied with fresh substrates and water from above more often than for the well-547 structured Loess soil with more frequent and stable preferential flow paths (Schneider and Don, 2019). 548 This might lead to a lower accessibility and therefore slower turnover rates of OC (Dungait et al., 2012). 549 The conclusion can be drawn that sediment derived OC can and will be degraded when it becomes part 550 of the subsoil but probably to a comparatively limited extent and low rates with dominant mineralisation 551 of biogenic OC.

# 4.3 How much does GOC contributes to soil organic carbon?

The contribution of GOC to soil OC stocks in this study was driven by the amount of OC in the soil, the 553 554 amount of GOC in the respective sediment and also the turnover of GOC when it becomes part of the 555 soil. Our results revealed that despite differing between sediments, GOC content varied in a quite narrow 556 range between 0.1 and 0.3 g kg<sup>-1</sup>. The contribution of GOC to topsoil OC was negligible. Assuming no 557 degradation of GOC, the greatest possible contributions to total subsoil OC were found for Red 558 Sandstone ( $\sim 30\%$ ) and the lowest for Miocene Sand (0.6%). This was due to the range of OC contents 559 in the subsoils (0.53 g kg<sup>-1</sup>-15.21 g kg<sup>-1</sup>). When soil OC contents were low, the possible contributions 560 from GOC were high and vice versa For the investigated soils, the OC contents of 3 g kg<sup>-1</sup> soil allowed 561 for possible GOC contributions of between 5- and 10 %. For OC contents around 1 g kg<sup>-1</sup> soil, a GOC contribution of between 10- and 20 % seemed possible. Thus, greater contributions were made by GOC 562 rich sediments such as Loess, while smaller contributions from sandy sediments. In comparison, van der 563 564 Voort et al. (2018) estimated the contribution of GOC in a soil derived from glacial deposits (flysch) at between 80-100 cm depth to be around 40 %. For a soil developed from a poorly consolidated 565 sedimentary rock (calcareous and shaly moraine), they calculate the contribution of GOC to range from 566 567 20 % at 145 cm depth to 80 % at 310 cm depth. An attempt has also been made to fractionate subsoils to extract the most stable OC that may be derived from GOC. Paul et al. (2001) investigated a soil 568 developed on loess over till with 30 % of subsoil OC as a non-hydrolysable fraction showing a <sup>14</sup>C age 569 of 13,000 years BP. They also concluded that this high <sup>14</sup>C age can partly be explained by a GOC 570 571 fraction. These results indicate that deposits from the past glacial periods, such as flysch or till in 572 particular, have much greater potential for OC contributions from GOC, possibly due to the higher amounts of GOC in their sediments. Since only terrestrial sediments were investigated in the present 573 study, it should be noted that marine sediments or shales also contain much higher amounts of OC, up 574 575 to 250 g kg<sup>-1</sup> (Hemingway et al., 2018, Petsch et al., 2000). Thus, the amount of GOC they contain and their possible contribution to subsoil OC stocks might therefore be much higher. 576

577 Nevertheless, the <sup>14</sup>C ages of OC in the subsoil can be high in soils derived from igneous parent materials
578 without GOC (Rumpel et al., 2002), although even crystalline bedrocks contain microbial communities

(Purkamo et al., 2020). Furthermore, on a global scale, the <sup>14</sup>C ages of soil OC are primarily driven by climatic conditions, clay content and age of the soil (Mathieu et al., 2015). However, for terrestrial sediments with comparatively low amounts of GOC that started their soil development after the latest glacial period, a scale of possible contributions could be obtained when the amount of OC is known. Thus, on a global scale, the high <sup>14</sup>C age of subsoils is not driven just by the GOC fraction but the presence of GOC may greatly influence subsoil <sup>14</sup>C.

#### 585 **5.** Conclusions

This approach of estimating the GOC contribution to soil OC showed that common and abundant 586 587 terrestrial sediments with low amounts of sedimentary OC can make a considerable contribution to subsoil OC stocks. One fraction of OC in the sediments is of geogenic origin and could therefore 588 influence measured <sup>14</sup>C ages in soil, particularly in subsoils. Subsoils are known for their high <sup>14</sup>C ages 589 and slow turnover rates and slow reaction to changing environmental condition. These subsoil OC 590 properties may partly be derived from the GOC in the subsoil. The sediments at the investigated sites 591 592 contained OC in a range of 0.1-0.3 g kg<sup>-1</sup>, allowing for contributions from GOC of between 10-30 % in 593 subsoils. Incubation of sediments indicated that this geogenic contribution presents a quite stable OC 594 pool, especially for subsoils. Thus, even sediments with comparatively low amounts of OC were also 595 able to demonstrate the large contribution of GOC.

#### 596 Data availability

597 The data will be made available on request

#### 598 Author contribution

- 599 AD conceived of and designed the study, FK performed the sampling and analysis, and wrote the first
- draft. All the authors contributed to generating and reviewing the subsequent versions of the manuscript.

# 601 Competing interest

602 The authors declare that they have no conflict of interest

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612

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