Rapid soil organic carbon decomposition in river systems: effects of the aquatic microbial community and hydrodynamical disturbance

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Abstract. Different erosion processes deliver large amounts of terrestrial soil organic carbon (SOC) to rivers. Mounting evidence indicates that a significant fraction of this SOC, which displays a wide range of ages, is rapidly decomposed after entering the river system. The mechanisms explaining this rapid decomposition of previously stable SOC still remain unclear. In this study, we investigated the relative importance of two mechanisms that possibly control SOC decomposition rates in aquatic systems: (i) in the river water SOC is exposed to the aquatic microbial community which is able to metabolize SOC much more quickly than the soil microbial community and (ii) SOC decomposition in rivers is facilitated due to the hydrodynamic disturbance of suspended sediment particles. We performed different series of short-term (168 h) incubations quantifying the rates of SOC decomposition in an aquatic system under controlled conditions. Organic carbon decomposition was measured continuously through monitoring dissolved O$_2$ (DO) concentration using a fiber-optic sensor (FireStingO$_2$, PyroScience). Under both shaking and standing conditions, we found a significant difference in decomposition rate between SOC with aquatic microbial organisms added (SOC + AMO) and without aquatic microbial organisms (SOC − AMO). The presence of an aquatic microbial community enhanced the SOC decomposition process by 70 %–128 % depending on the soil type and shaking–standing conditions. While some recent studies suggested that aquatic respiration rates may have been substantially underestimated by performing measurement under stationary conditions, our results indicate that the effect of hydrodynamic disturbance is relatively minor, under the temperature conditions, for the soil type, and for the suspended matter concentration range used in our experiments. We propose a simple conceptual model explaining these contrasting results.

1 Introduction

Rivers play an important role in the global carbon cycle by linking terrestrial and aquatic ecosystems. Each year, rivers receive and deliver large amounts of terrestrial organic carbon to the oceans (Raymond and Bauer, 2001; Ward et al., 2017). However, rivers do not just transport OC. In 2007, Cole et al. (2007) published the idea of “rivers as an active pipe”, which highlights the fact that rivers not only transport but also process large amounts of OC. Mounting evidence indicates that a significant fraction of the OC processed in rivers is soil organic carbon (SOC), including dissolved (DOC) and particulate (POC) forms. The mobilized SOC can display a very wide range of ages among rivers depending on the carbon sources and retention time. Many studies have demonstrated that when SOC is delivered to a river, part of this SOC can be rapidly mineralized and emitted back to the atmosphere (Lapierre et al., 2013; Wilkinson et al., 2013; Mayorga et al., 2005). Understanding the mechanisms that contribute to this active mineralization process of SOC is essential for understanding the role of rivers in the global carbon cycle and for assessing how the metabolism of rivers may respond to environmental perturbations such as an increase or decrease in terrestrial carbon delivery to the river system and/or changes in hydrology and climate. Indeed, the amount of SOC imported into riverine ecosystems is often large compared to the autochthonous within-river primary...
production (Cole and Caraco, 2001). It is therefore important to understand the fate of this terrestrial SOC as this not only affects the global carbon cycle but also strongly regulates the ecological functioning of river ecosystems (Kling, 1995). A key question is then which factors control the decomposition of terrestrial SOC, which has been stable in soil for decades to centuries, when it enters into a river system.

In recent years, the mechanisms controlling this rapid riverine mineralization process have gained increasing attention (Aufdenkampe et al., 2011; Guenet et al., 2014; Ward et al., 2017). While in transit, SOC can be degraded by microbial mineralization (Ward et al., 2013) and photochemical oxidation (Spencer et al., 2009). As can be expected, these processes are closely associated with a suite of factors such as temperature (Lapierre et al., 2013; Gudasz et al., 2015), the availability of oxygen (Koehler et al., 2012), and the presence and composition of microbial communities (Ward et al., 2019a), along with physical river properties, such as river velocity and hydrodynamic conditions (Ward et al., 2018). For example, Ward et al. (2018) incubated river water and sediment under three rotation regimes to mimic river flow and found that respiration rates in rotating samples were 1.4 (under 0.22 m s$^{-1}$) and 2.4 times (under 0.66 m s$^{-1}$) higher compared with stationary conditions. Mechanical disturbance can be expected to enhance mineralization because the physical breakup of large particles induced by river water disturbance may increase the accessibility of microbial enzymes to SOC (Lal, 2003; Richardson et al., 2013).

Microbial organisms influence SOC cycling not only via decomposition but also because microbial products are themselves important components of soil organic matter (Kögel-Knabner, 2002). The interactions of microbial communities and SOC decomposition have been extensively studied in terrestrial ecosystems (Cleveland et al., 2007; Hu et al., 2014; Tian et al., 2016). The interactions between microbes and SOC in aquatic systems had received little attention until recently. Recently, Ward et al. (2019a) illustrated the potential importance of the composition of the aquatic microbial community by showing that mixing the water of two lowland tributaries (the Tapajós and Xingu rivers) and the main Amazon river resulted in enhanced respiration rates, which they attributed to the fact that such mixing resulted in a more diverse microbial community capable of digesting more OC. While these recent studies clearly indicate that different mechanisms do indeed control OC processing in rivers, important knowledge gaps do remain with respect to the specific controls of SOC decomposition by aquatic microbes. One of the reasons for the latter is that, hitherto, very few experiments were carried out with a factorial design allowing the investigation of the relative importance and potential interaction of different individual factors.

The main objective of this study is to shed light on the fate of SOC in river systems by investigating the relative importance of two key mechanisms that were previously suggested to potentially contribute to the rapid decomposition of previously stable SOC in aquatic systems: (i) in the river water, SOC is exposed to an aquatic microbial community which may be able to metabolize SOC much more quickly than the soil microbial community and (ii) SOC decomposition in rivers may be facilitated due to the hydrodynamic disturbance of sediment. We base our analysis on continuous measurements of DO consumption during a series of lab incubation experiments and the characterization of the evolution of SOC, POC and DOC characteristics throughout the incubation period. By doing so we were able to quantify the importance of these mechanisms both in absolute and relative terms.

2 Material and methods

2.1 Site description

We sampled the Dijle River located ca. 2 km upstream of the city of Leuven, which is situated in the central Belgian loess belt (Fig. 1). The relief of the 700 km$^2$ large Dijle River catchment upstream of Leuven varies from about 25 m above sea level (a.s.l.) in the north to ca. 165 m a.s.l. in the south. The catchment is characterized by an undulating plateau in which several brooks and rivers are incised. Slope gradients are usually less than 5%, although maximum slopes of ca. 50% can be found along the valleys. The majority of the soils are Luvisols, developed in the loess deposits (FAO, 1998). The land use in the catchment is mainly cropland, which is particularly vulnerable to erosion during spring and early summer when vegetation cover is low and rainfall intensity is high (Takken et al., 1999). The contemporary land use in the floodplain is dominated by grassland, plantation forests, and built-up area (Broothaerts et al., 2014).
2.2 Experimental approach

2.2.1 Experimental setup

Our measurement approach involved collecting a large volume of river water, subsequently treated by removing all suspended particles via filtration and dispensing the filtrate into a series of 320 mL glass incubation bottles. The treatments included (1) SOC with rotation, (2) SOC under stationary conditions, (3) DOC control with rotation, (4) DOC control under stationary conditions, (5) SOC with rotation without AMO, and (6) SOC under stationary conditions without AMO (see Table 1 for details). Mechanical breakup was simulated by a custom-designed swing system (Fig. 2), which kept the particles in suspension during the incubation. Incubation bottles were fixed in the PVC clip brackets and rotated at about seven rounds per minute. Since the bottles were connected with FireStingO₂ fiber, the swing system was first rotated 180° counterclockwise and then 180° clockwise back, which together was counted as one round. In order to get a higher disturbance intensity, 8 g of glass beads (diameter: 2 mm) were added to the bottle before adding soil and water samples. The glass beads were pre-sterilized at 450 °C for 1 h to avoid contamination.

2.2.2 Collection and incubation

River water was collected from the Dijle River (details in Table 2). The water was filtered on 0.7 μm glass fiber filters to remove all suspended particles. For treatments without aquatic microbial communities, river water was then filtered through 0.2 μm syringe filters. Since filtration at 0.7 μm already removed the majority of microbial organisms, 4 mL unfiltered river water (ratio of inoculum to the incubated water sample: 1 : 79) was added to serve as an inoculum for treatments with aquatic microbial organisms. Soil samples were collected from the Belgian loess belt near Leuven: one from arable land (50°48′31.6″N, 4°35′16.9″E), which was collected from a depth of 0–20 cm, and the other from Bertem forest (50°52′59.8″N, 4°38′24.2″E). The top litter layer was first removed, and then the soil sample was collected from a depth of 0–20 cm. These soils were oven-dried at ~ 55 °C and then sieved through 2 mm to remove all roots and stones. In order to obtain a detectable rate of oxygen consumption, the POC concentration was controlled at 10–12 mg L⁻¹ by adding 160 mg arable soil and 60 mg forest soil in 320 mL river water (details in Table 3). While the sediment and POC concentrations we used in our experiments are relatively high, they are not unrealistic: during high flood we observed POC concentrations exceeding 10 mg L⁻¹ in ca. 5 % of our samples. The bottles were subsequently closed without headspace. For the suspended condition treatments, bottles were fixed on the swing systems (Fig. 2). The remaining bottles were placed on a shelf to keep particles settled in each bottle. To avoid the influence of temperature and light (both potentially influencing OC mineralization), all bottles were wrapped with aluminum foil and incubated in a temperature-controlled room at ~ 20 °C. For each experiment, four incubation bottles with the same treatment were incubated, which allowed us to sample for POC and DOC during the course of the experiment (at t = 24, 48, 96, 168 h). Six experimental runs for each soil type were conducted to investigate the effect of water disturbance and the presence of an aquatic microbial community on SOC decomposition.

Given that the soil samples were oven-dried at ~ 55 °C before the incubation, a certain amount of soil microbial organisms might have been eliminated. In order to test this, we carried out a series of supplementary experiments whereby air-dried and oven-dried soil material was used under stationary and rotation conditions. Soil material was collected from the same arable land (OC%: 1.5 %). The soil was divided into two parts and dried with two methods: one part was oven-dried at ~ 55 °C, and the other was air-dried. The incubations followed the same procedures mentioned above.

2.2.3 DO, POC, and DOC measurements

DO was measured every 10 s for 168 h using an optical oxygen meter (FireStingO₂), and we used the DO data to derive the total amount of C mineralized in each incubation series, using a procedure similar to the one used in previous studies (Berggren et al., 2012; Richardson et al., 2013). It is known that when OC is decomposed, the O₂ : CO₂ ratio is not constant but depends on the composition of the substrate and the characteristics of the bacterial community (Berggren et al., 2012). Richardson et al. (2013) took the elemental composition of organic matter into account and reported an O₂ : CO₂ ratio of 1.04–1.2 for detrital organic matter and 1.0 for DOC. In this study, we used an O₂ : CO₂ ratio equal to 1.0 for both SOC and DOC, thus assuming that the mineralization of 1 mol C consumed 1 mol O₂. Using a respiration ratio of
Table 1. Experimental setup for incubations under controlled laboratory conditions.

| Treatment number | Treatments          | Number of repetitions | Ingredients                                      |
|------------------|---------------------|-----------------------|--------------------------------------------------|
| 1                | SOC, rotation       | Six                   | soil + 0.7 µm filtered river water + inoculum    |
| 2                | SOC, stationary     | Six                   | soil + 0.7 µm filtered river water + inoculum    |
| 3                | DOC control, rotation | Six                 | 0.7 µm filtered river water + inoculum          |
| 4                | DOC control, stationary | Six                 | 0.7 µm filtered river water + inoculum          |
| 5                | SOC without AMO, rotation | Six                 | soil + 0.2 µm filtered river water              |
| 6                | SOC without AMO, stationary | Six                 | soil + 0.2 µm filtered river water              |

Table 2. Nutrient concentrations in the Dijle River water (data from the Flanders Environment Agency: https://www.vmm.be/data/wATERKwaliteit, last access: 13 November 2020).

| Year | Number of measurements | NH$_4^+$ (µmol L$^{-1}$) | NO$_3^-$ (µmol L$^{-1}$) | PO$_4^{3-}$ (µmol L$^{-1}$) |
|------|------------------------|--------------------------|--------------------------|----------------------------|
| 2020 | 3                      | 24 ± 7                   | 717 ± 55                 | no data                   |
| 2016 | 6                      | 48 ± 30                  | 410 ± 15                 | 6 ± 1                     |
| 2014 | 6                      | 26 ± 11                  | 537 ± 94                 | 6 ± 1                     |

Note: values are mean ± SD.

1.0 might result in an overestimate of the total amount of C mineralized, but since all treatments were calculated with the same respiration ratio and the same soil samples were used throughout the experiments, the relative variations will not be affected by this choice.

After the incubation period, the entire volume of ∼ 320 mL water was filtered for later determination of total POC, particulate nitrogen (PN), and the stable C isotope ratios ($\delta^{13}$C) of POC on pre-combusted 25 mm Whatman GF/F filters (pore size: 0.7 µm). Due to the high concentrations of soil material in each bottle, each water sample was filtered on two to five filter papers. For arable soil incubations, water samples were filtered on five filter papers. We measured all filter papers for two repetitions and two of five filter papers for the other four repetitions. Then we used the suspended sediment weight and POC concentrations measured on this subsample to calculate the POC and PN content as well as the $\delta^{13}$C signature of the entire sample. For forest soil incubations, water samples were filtered on two to five filter papers. All of the filter papers were oven-dried at 50°C and preserved for POC, PN, and $\delta^{13}$C measurements. Inorganic C was removed from the filters by exposing them to HCl fumes overnight in a desiccator. Subsequently, the dried filters were packed in Ag cups for analysis on an elemental analyzer–isotope ratio mass spectrometer (EA–IRMS, Thermo Finnigan Flash HT and Delta V Advantage). Certified (IAEA-600, caffeine) and in-house laboratory standards (leucine and tuna tissue) were analyzed throughout each run.

To determine the DOC concentration and its stable isotope composition, 40 mL water samples (filtered at 0.2 µm) were collected and stored in glass vials with Teflon-coated screw caps, and 100 µL of H$_3$PO$_4$ was added for preservation. Analysis of DOC and $\delta^{13}$CDOC was performed on a wet oxidation TOC analyzer (IO Analytical Aurora 1030W) coupled with an isotope ratio mass spectrometer (Thermo Finnigan Delta V Advantage). Quantification and calibration were performed with IAEA-C6 ($\delta^{13}$C = −10.4‰) and an internal sucrose standard ($\delta^{13}$C = −26.99 ± 0.04‰).

The two soil samples used were analyzed for their $^{14}$C content on a MICADAS accelerator mass spectrometry system at the Royal Institute for Cultural Heritage (KIK/IRPA, Brussels). Data are reported in years BP, whereby 1950 is taken as the present.

2.3 Statistical analysis

Statistical tests were performed in R (version 3.5.1, R Core team). The normality of data was tested with the Shapiro–Wilk test. The paired sample t test was used to test for differences in DO consumption rates, total amount of C mineralized, and POC loss between treatments under rotation and stationary conditions as well as to test for differences between treatments with and without the presence of AMO. To identify the combined effect of AMO and physical disturbance on the C decomposition rates, we used two-way ANOVA with the presence of AMO and mechanical rotation as the main factors. ANOVA was carried out separately for the two soil types. Average values are given ± the standard deviation.

3 Results

3.1 Effect of water disturbance and AMO on DO consumption rates

The initial DO concentration in the river water varied from 7.58 to 10.67 mg L$^{-1}$. In all treatments, DO concentrations followed a decreasing trend, and the DO consumption rates were relatively constant over time (Fig. 3). For incubations where soil was present, cumulative DO consumption ranged between 0.7 and 2.6 mg L$^{-1}$ during the 168 h incubation period. The highest DO consumption occurred in the SOC + AMO treatments with average con-
sumption rates of 0.015 ± 0.005 mg O$_2$ L$^{-1}$ h$^{-1}$ (arable soil) and 0.010 ± 0.002 mg O$_2$ L$^{-1}$ h$^{-1}$ (forest soil). DO consumption was lowest for treatments where soil was present without AMO with an average of 0.007 ± 0.002 mg O$_2$ L$^{-1}$ h$^{-1}$ (arable soil) and 0.004 ± 0.001 mg O$_2$ L$^{-1}$ h$^{-1}$ (forest soil). We found that keeping soil particles in suspension resulted in a relatively small acceleration of the OC decomposition process. With the presence of aquatic microbial organisms (SOC+AMO), DO consumption rate was increased on average by 13 % (p < 0.05) for the arable soil, while no significant effect was found for forest soil (p > 0.05) (Table 4).

The addition of AMO, on the other hand, had a much stronger stimulation effect on OC decomposition. Compared with SOC−AMO treatments, DO consumption rates were approximately doubled by the presence of AMO for both soil types. This increase was present under both rotation (arable soil: 0.015 ± 0.005 vs. 0.007 ± 0.002 mg O$_2$ L$^{-1}$ h$^{-1}$, p < 0.01; forest soil: 0.010 ± 0.002 vs. 0.005 ± 0.002 mg O$_2$ L$^{-1}$ h$^{-1}$, p < 0.01) and stationary conditions (arable soil: 0.014 ± 0.003 vs. 0.007 ± 0.002 mg O$_2$ L$^{-1}$ h$^{-1}$, p < 0.01; forest soil: 0.009 ± 0.002 vs. 0.004 ± 0.001 mg O$_2$ L$^{-1}$ h$^{-1}$, p < 0.01) (Table 4). In addition, the presence of AMO and rotation had no significant combined effect on the C decomposition rates for both soil types (p > 0.05).

### 3.2 Total amount of C mineralized

We calculated the total amount of C mineralized in each treatment, using an O$_2$ : CO$_2$ ratio of 1.0, and expressed mineralization rates on a per carbon basis. Obviously, trends in total C mineralization are similar to those in oxygen consumption. For SOC+AMO incubations with arable soil, keeping particles in suspension increased the total amount of C mineralized by 13 % (p < 0.05, Fig. 4). Also, for forest soil a small increase was noted (11 %), but this was not statistically significant (p > 0.05). Compared with SOC−AMO, the presence of aquatic microbial organisms (SOC+AMO) – regardless of soil type and rotating–stationary conditions – significantly stimulated OC decomposition (Fig. 4). For arable soil, AMO addition led to 91 % (under rotation conditions, p < 0.01) to 128 % (under stationary conditions, p < 0.01) more C mineralized by the end of the incubation and 116 % (under rotation conditions, p < 0.01) to 118 % (under stationary conditions, p < 0.01) with forest soil, depending on rotation and stationary conditions.

In addition, when comparing the mineralization of oven-dried and air-dried soil without AMO addition, oven-dried soil incubation showed a 40 % lower C loss in comparison to air-dried soil incubation at the end of the incubation (Fig. 5, p < 0.05). However, the addition of AMO resulted in a significantly higher C loss with both treatments (air drying: 70 %, p < 0.05; oven drying: 165 %, p < 0.05).

### 3.3 Particulate and dissolved organic carbon concentrations and δ$_{13}$C$_{POC}$ and δ$_{13}$C$_{DOC}$ values

Measurements of final POC concentrations showed a reduction of POC at the end of the incubations, where 5 %–13 % of POC was mineralized for incubations with arable soil and 1 %–11 % for incubations with forest soil (Table 4). Keeping particles in suspension resulted in ca. 4 % more POC loss in SOC+AMO incubation series for both soil types, although the increase was only significant for arable soil (arable soil: 13 % vs. 9 %, p < 0.05; forest soil: 11 % vs. 7 %, p > 0.05, Table 4). Conversely, keeping particles in suspension in SOC−AMO incubation series showed negative effects with 1 % (arable soil) to 4 % (forest soil) less POC loss, but this difference was not statistically significant (p > 0.05). For both soils, the presence of AMO led to ca. 8 %–10 % more POC loss under rotation conditions (p < 0.05). This effect was smaller (2 %–3 %) and non-significant (p > 0.05) under stationary conditions (Table 4).

These trends are generally consistent with the patterns derived from the oxygen consumption measurements, i.e., a larger reduction in POC when AMO is present and a limited effect of hydrodynamic disturbance on POC decomposition. However, it is also clear that variations in residual POC are less consistent than those observed from oxygen consumption rates. In 7 of 12 incubation series, the combined POC

### Table 3. Characteristics of the two soils used in this study. Texture was determined using laser diffraction (Coulter LS 13 320).

| Soil samples | Soil texture | OC content (%) | C/N (weight/weight) | δ$_{13}$C (%) | $^{14}$C age (year BP) |
|--------------|-------------|----------------|---------------------|---------------|----------------------|
| Arable soil  | Sand % 23   | 2.40           | 9.9                 | -29.4         | 267 ± 21             |
|              | Loam % 69   |                |                     |               |                      |
|              | Clay % 8    |                |                     |               |                      |
| Forest soil  | Sand % 39   | 5.20           | 17.6                | -28.6         | 334 ± 22             |
|              | Loam % 56   |                |                     |               |                      |
|              | Clay % 5    |                |                     |               |                      |

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Figure 3. DO consumption of the SOC + AMO treatment (red), the SOC − AMO treatment (blue), and the DOC control treatment (green) during the incubations, under rotating (full lines) and stationary (dashed lines) conditions: (a) incubation with arable soil, (b) incubation with forest soil. Standard errors were calculated based upon six repetitions with different river water samples.

Figure 4. Total amount of C mineralized during the incubations with and without AMO under rotation and stationary conditions with (a) arable soil and (b) forest soil. Different capital letters indicate significant differences between treatments under rotation and stationary conditions at $p < 0.05$; different lowercase letters indicate significant differences between treatments with and without AMO at $p < 0.01$.

Figure 5. Cumulative amount of C mineralized during incubation with oven-dried soil (red) and air-dried soil (black), with addition of AMO (full line) and without addition of AMO (dashed line). Standard errors were calculated based upon three repetitions with different river water samples.

Variations in $\delta^{13}$C_POC values showed a similar pattern for both soil types (Fig. 6): firstly, an increase in $\delta^{13}$C_POC occurred during the first 24 h (for forest soil) or 48 h (arable soil). The increase in $\delta^{13}$C_POC appears to be more important for forest soil (0.4 ‰–0.6 ‰) in comparison to arable soil (0.2 ‰–0.4 ‰). After this initial period, $\delta^{13}$C_POC values stabilized.

DOC concentrations were relatively stable during the incubation (Fig. 7; Table 4). The initial $\delta^{13}$C_Doc value was lower in arable soil experiments where no AMO was present. When AMO was present the initially higher $\delta^{13}$C_Doc values declined to similar values as those observed in experiments without AMO within the first 48 h. In the experiments with forest soil, initial $\delta^{13}$C_Doc values were slightly but not significantly higher (0.6 ‰, $p > 0.05$) than the initial $\delta^{13}$C_Doc values for arable soil without AMO. $\delta^{13}$C_Doc values were stable throughout the incubation period for all forest soil experiments.

For both soil types, the equilibrium $\delta^{13}$C_Doc values appear to be somewhat lower than the equilibrium $\delta^{13}$C_POC val-
ues: this difference was more pronounced for the forest soil (ca. 1 ‰ in comparison to ca. 0.5 ‰ on average), but non-
significant in both cases ($p > 0.05$).

4 Discussion

Our results show that a fraction of the terrestrial SOC can indeed be mineralized relatively quickly when introduced in an aquatic environment. With the presence of aquatic microfloral organisms (SOC + AMO), up to 0.58 (forest soil) or 0.97 (arable soil) mg CL$^{-1}$ was mineralized within the 168 h incubation period, equivalent to 83–139 µg CL$^{-1}$ d$^{-1}$ or 9–12 µg C (mg SOC$^{-1}$) d$^{-1}$. Comparable respiration rates of 20–80 µg CL$^{-1}$ d$^{-1}$ were reported under similar temperature conditions by Berggren et al. (2010). Similar incubation experiments with water samples collected from northern temperate lakes and streams reported respiration rates from 16 to 54 µg CL$^{-1}$ d$^{-1}$ (McCallister and Del Giorgio, 2012). We found that C mineralization rates were quite significant: in the presence of an AMO (which is always the case in natural conditions), 4% to 6% of the OC present at the initiation of the experiment was mineralized during the experiment (Fig. 4). Typical soil incubation experiments show a loss of max. 2% of the SOC in the first 2 weeks: Gillabel et al. (2010) incubated soil samples under 25° and found ca. 2% SOC was respired in 20 d with a mineralization rate at around 1 µg C mg C$^{-1}$ d$^{-1}$; Angst et al. (2019) conducted a similar soil incubation and found a cumulative respiration of 20 mg CO$_2$ (g C soil)$^{-1}$ in 14 d, equivalent to around 0.4 µg C mg C$^{-1}$ d$^{-1}$. Li et al. (2018) incubated soil with fertilization and straw application and found highest cumulative respiration at around 800 mg CO$_2$-C kg$^{-1}$ soil in 20 d, equivalent to around 2 µg C mg C$^{-1}$d$^{-1}$. Thus mineralization rates observed in soil incubation experiment (0.4–2 µg C mg C$^{-1}$ d$^{-1}$) are much lower than those observed in our study, where 6–9 µg C is mineralized per milligram of C per day. This suggests that SOC indeed decomposes 5–10 times more rapidly in aquatic systems than in the terrestrial environment and that SOC is not as recalcitrant as previously thought in aquatic systems (Mayorga et al., 2005; McCallister and Del Giorgio, 2012). Several studies already suggested that the transition from terrestrial to aquatic conditions likely facilitated SOC decomposition rates because of shifts in environmental conditions (Butman and Raymond, 2011; McCallister and Del Giorgio, 2012). In soils, sorption of OC to mineral surfaces and encapsulation of C within soil aggregates may protect SOC from complete mineralization (Bianchi et al., 2011; Schmidt, et al., 2011). This results in
the accumulation of older SOC in pools that are less accessible to decomposers and their extracellular enzymes (Marín-Spiotta et al., 2014). When SOC enters aquatic systems, a disruption of the mechanisms protecting C from mineralization, such as a physical disturbance due to the physical action of transport in water but also due to aggregate slaking (Le Bissonnais, 1996), may lead to the exposure of these protected pools to decomposers and therefore to an increase in the SOC decomposition rate. Alternatively, SOC decomposition may be accelerated due to the fact that, in an aquatic environment, a population of possible consumers is present that is different from that in the soil and that may be capable of rapidly mineralizing SOC that is otherwise preserved over long timescales in a soil environment (Wu et al., 2018; Ward et al., 2019b).

Our results indicate that physical soil disturbance, which was simulated by rotation in our experiments, had a relatively minor effect on SOC decomposition (arable soil: \( p < 0.05 \); forest soil: \( p > 0.05 \)) (Figs. 3, 4). This result is different from that reported in two previous studies with similar incubation approaches. Richardson et al. (2013) and Ward et al. (2018) found that keeping sediment particles in suspension increased river-borne organic matter decomposition by 40% to 140%, depending on the rotation speed. The differences in experimental settings, such as POC composition, POC concentration, and microbial community may influence the POC decomposition rates (Weyhenmeyer et al., 2012; Ward et al., 2019a). However, these differences as such do not explain the minor effect of physical disturbance observed in this study. Thus, our data do indicate that the relative role of physical disturbance vs. that of exposure to an aquatic microbial community may vary considerably between different ecosystems. We propose two possible reasons as to why we did not find a strong effect of physical disturbance on SOC decomposition. It is hypothetically possible that the level of physical protection of (a large part of) the SOC that is present is such that it is not disrupted by the physical disturbance that we imposed. Alternatively, the level of physical protection was so weak that the mere immersion of soil particles in water was sufficient to destroy most of it. Given the fact that loess soils are known for their very low structural stability (Le Bissonnais, 1996) and the high POC mineralization rates we observed, we propose that the latter is more likely in our experiments. The results of our last series of experiments (Fig. 5) also show that when SOC is introduced into an aquatic environment where no aquatic microbial commu-
Table 4. Total amount of C mineralized and POC and DOC loss at the end of the incubation series calculated as the difference in the mass of carbon that was introduced and the mass of carbon recovered from the final sample.

| Soil type | Treatments | Total amount of C mineralized (mg) | Significant POC losses (%) | Percentage POC loss (mg) | Significant DOC losses (%) | Percentage DOC loss (mg) |
|-----------|------------|-----------------------------------|---------------------------|-------------------------|---------------------------|-------------------------|
| Arable soil | SOC + AMO/rotation | 0.31 ± 0.09 | Y | 13 ± 5 | Y | 0.54 ± 0.15 | Y | 0.32 ± 0.08 | Y |
| | SOC/rotation | 0.27 ± 0.06 | N | 9 ± 3 | N | 0.43 ± 0.14 | N | 0.24 ± 0.04 | N |
| | SOC control/rotation | 0.15 ± 0.05 | N | 6 ± 3 | N | 0.31 ± 0.12 | N | 0.14 ± 0.06 | N |
| | SOC control/stationary | 0.13 ± 0.04 | N | 6 ± 3 | N | 0.31 ± 0.12 | N | 0.14 ± 0.06 | N |
| Forest soil | SOC + AMO/rotation | 0.23 ± 0.03 | Y | 14 ± 5 | Y | 0.24 ± 0.10 | Y | 0.24 ± 0.10 | Y |
| | SOC/rotation | 0.18 ± 0.04 | N | 8 ± 3 | N | 0.24 ± 0.10 | N | 0.14 ± 0.06 | N |
| | SOC control/rotation | 0.11 ± 0.04 | N | 6 ± 3 | N | 0.31 ± 0.12 | N | 0.14 ± 0.06 | N |
| | SOC control/stationary | 0.09 ± 0.03 | N | 6 ± 3 | N | 0.31 ± 0.12 | N | 0.14 ± 0.06 | N |
| | SOC + AMO/stationary | 0.08 ± 0.02 | N | 6 ± 3 | N | 0.31 ± 0.12 | N | 0.14 ± 0.06 | N |

The higher POC consumption rates observed when an AMO is present may, of course, partly be due to the fact that more microbes were present in those experiments where AMO was introduced. We compared the initial population of bacteria with and without the addition of AMO. The addition of AMO led to 20 %–30 % more bacteria present in the water at the beginning of the experiment (arable soil: 4.30 × 10^5 vs. 3.52 × 10^5 cells mL^-1; forest soil: 3.67 × 10^5 vs. 2.86 ×
10^5 cells mL^{-1}). This larger initial population could partly explain the higher SOC decomposition rates with the addition of AMO, but the fact that the addition of AMO increases SOC decomposition rates by 70%–165% rather than 20%–30% does suggest that the aquatic microbial community is indeed capable of attacking old, stable SOC more effectively than the soil microbial community. Although the microbial community is considered to play a central role in shaping OC reactivity in both terrestrial and aquatic systems (Schmidt et al., 2011), such a strong stimulation effect of the addition of AMO on SOC has rarely been reported.

The evolution of POC and DOC concentrations during the experiments is generally in agreement with the patterns derived from the oxygen consumption measurements. However, the variations in residual POC are less consistent than those observed from oxygen consumption. Several reasons might explain the discrepancy between the total C mineralization as calculated from oxygen consumption in comparison to direct measurements of POC. Firstly, POC and DOC samples were collected from four incubation bottles for one treatment. Even though we controlled the initial conditions of each bottle as closely as possible, there might be heterogeneity between different bottles with respect to the OC content of the soil sample that was placed into the bottle. Secondly, we compared POC measurements from two to five filter papers for arable soil and five filter papers for forest soil. For each individual measurement the filter weight has to be subtracted from the gross weight of the filter plus the sediment. Given the small quantities of sediment present on the filters, small weighing errors will result in relatively large errors in the calculation of the amount of POC that is remaining. DO measurements are non-intrusive and are not subject to measurement errors related to the weighing of small quantities. We therefore believe that the oxygen consumption measurements provide us with more robust measurements of OC decomposition in comparison to direct measurements of the OC content of the remaining sample. The direct measurements suggest that, overall, POC mineralization was more important than DOC mineralization when POC was present. Indeed, DOC concentrations showed little variation during the experiments, despite significant oxygen consumption rates.

The increase in δ^{13}C_{POC} values during the first 24–48 h suggests that during this period an isotopically lighter POC fraction was preferentially mineralized. This resulted in the POC in the aquatic environment becoming enriched in 13C by 0.2‰–0.6‰ compared to the POC in the original soil sample. After the initial adjustment period, δ^{13}C_{POC} remained stable, suggesting the initial adjustment is indeed due to the preferential consumption of a somewhat lighter, less recalcitrant POC fraction rather than continued preferential consumption of lighter POC. The fact that over the whole course of the experiments the δ^{13}C_{DOC} values are lower than the corresponding δ^{13}C_{POC} values is on the other hand best explained by a continuous leaching/release of DOC with a somewhat lower δ^{13}C signature from the soil POC, replacing the mineralized DOC as it is unlikely that the DOC fraction would be entirely stable while POC is continuously mineralized. Mineralization of the original DOC and its replacement with soil-derived DOC could also explain the drop in δ^{13}C_{DOC} during the initial phase of the experiments with arable soil and AMO, because the lighter DOC that was originally present in the river water is replaced by soil-derived DOC. However, this drop was not observed in the experiments with forest soil. This may be partly explained by the higher δ^{13}C value of the forest soil (−28.6‰) in comparison to the δ^{13}C signature of the arable soil (−29.4‰), causing the DOC released from the forest soil to have an isotopic signature close to that of the river water. While the patterns described above would be consistent with the preferential decomposition of isotopically lighter POC, we did not observe an increase in δ^{13}C_{POC} during our experiments as might be expected by the selective mobilization of an isotopically lighter soil fraction: this can be explained by the relatively small differences in δ^{13}C values between POC and DOC in combination with the fact that only a small fraction of the POC is ultimately mineralized, whereby most of this mineralized fraction may have been directly transformed to CO₂. If this mineralization does not selectively affect specific fractions of the POC pool, the δ^{13}C_{POC} values can be expected to remain more or less constant throughout the incubation period.

Based on our findings, we propose that the relative importance of physical disturbance on SOC decomposition upon entering river water.

Figure 8. Conceptual figure representing the effect of physical disturbance on SOC decomposition upon entering river water.
needed to expose a significant fraction of SOC to the microbial community present in water (Fig. 8). The loess-derived soils we used in our experiments do have a weak structure (Govers, 1991), resulting in rapid aggregate breakdown upon immersion in water and a relatively small effect of mechanical disturbance on mineralization rates. This also explains the insignificant interaction effect of hydrodynamic disturbance and the presence of an aquatic microbial community on the C decomposition rates for both soil types ($p > 0.05$). The mere immersion of soil particles in water is sufficient to destroy most of the soil aggregates. Therefore, further disturbance would not strongly increase the interactions between soil particles and microbial organisms. The effect of the presence of an aquatic microbial community, on the other hand, will depend on its composition and its vigor. The addition of SOC may shift aquatic microbial metabolisms and make it more prone to SOC decomposition (Lennon and Pfaff, 2005). Other factors such as water temperatures and nutrient availability may also play a role. The river water we used in our study has a high nutrient content (Table 2): the availability of a large nutrient pool may allow AMO to use the SOC more effectively and may be one of the factors explaining why we saw such an important increase in SOC decomposition when AMO was added. However, it is fair to state that our current understanding of the interaction between old SOC and different microbial communities is too limited to develop general principles describing which factors may stimulate or slow the decomposition of SOC exposed to a new microbial community.

5 Conclusions

We investigated the relative importance of physical disturbance vs. exposure to a novel microbial community for SOC decomposition rates in aquatic environments. While some recent studies found that the impact of mechanical disturbance on SOC decomposition rates was very important, we found only a very modest increase in SOC decomposition when soil particles were mechanically disturbed and kept in suspension. A simple conceptual model, whereby the effect of mechanical disturbance is assumed to depend on the initial structural stability of soil aggregates, can explain this difference in findings: mechanical disturbance is only important when soil aggregates are strong enough to withstand the disruptive forces imposed by immersion in water.

Our study also highlights the role of aquatic microbial organisms in SOC decomposition in river systems. Aquatic microbial organisms are capable of attacking SOC, leading to rapid SOC decomposition in river systems. Given the variability of aquatic microbial community composition in different aquatic systems, understanding the linkage between aquatic microbial community composition and abundance on the one hand and the resultant SOC mineralization rates on the other hand is important not only to better understand CO$_2$ outgassing from aquatic systems, but also to understand how the ecological functioning of rivers and lakes is affected by the large changes in SOC inputs that result from human activities.

Data availability. All data used and produced through this study are available with the Supplement.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/bg-18-1511-2021-supplement.

Author contributions. GG, SB, LJ, and MZ were involved in the design of the experiments, and MZ carried out the experiments. SB and MZ conducted the laboratory analysis. All authors offered advice on the data analysis and contributed to the paper preparation.

Competing interests. The authors declare that they have no conflict of interest.

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