Dynamic transfer of soil bacteria and dissolved organic carbon into small streams during hydrological events

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Abstract
Small headwater streams interlink catchment soils with the river network. As water makes its way from the hillslopes to the stream, it may transport dissolved organic carbon (DOC) and potentially soil microbes into stream water. In this study, we aimed at quantifying the dynamic transfer of DOC and microbial life, namely bacteria from catchment soils into streams. We hypothesized that increased soil saturation enhances the lateral inflow of bacteria and DOC into streams. To address this hypothesis, we sampled six first order streams and three soil transects at two different depths located within the pre-alpine Oberer Seebach (OSB) catchment in Austria over a duration of 2 years. We found a strong variation in DOC concentrations (range 0.4–5.6 mg L\(^{-1}\)) and bacterial abundances (range < 500,000–3,863,000 cells mL\(^{-1}\)) measured by flow-cytometry. The highest values of DOC and bacterial cells occurred during high flow events. DOC concentration and bacterial abundance were correlated across all streams and seasons. In soils, DOC ranges were higher and were also correlated with bacterial abundance, while DOC concentrations were ~ 10 times higher per bacterial cell than in streams. Overall we show that soils provide a dynamic inflow of bacteria and DOC to first order streams. Most probably, this results in a dynamic and reoccurring inoculation of small streams from catchment soils during runoff events. We propose that this dynamic microbial inoculation of small streams is potentially relevant for microbial community dynamics of downstream receiving waters.

Keywords
Soil inoculation · Soil bacteria · Dissolved organic matter · Hydrological events

Introduction
Small headwater streams are dynamic systems that interlink the terrestrial landscape with the fluvial network (Battin et al. 2008; Richardson and Danehy 2007). As they receive water from hillslopes, they have been proposed as ‘mirrors of the landscape’, integrating the catchment heterogeneity of soils, geology and land-use into a combined biogeochemical signature (Bishop et al. 2008; Wipfli et al. 2007). Typically, headwater streams encompass 60–80% of stream networks flow length (Benda et al. 2005; Schumm 1956). As they provide nutrients, woody debris, and dissolved and particulate organic carbon (DOC and POC, respectively) to downstream water bodies (Gomi et al. 2002; Wipfli et al. 2007), they play a crucial role for the ecology of downstream river networks. Also, through matter transport, headwater streams link the terrestrial and aquatic carbon cycle (e.g., Fasching et al. 2014; Raymond et al. 2013; Schelker et al. 2016). It is therefore critical to assess the ecology of headwater streams to understand the functioning of the rest of the river network.

Despite the important role headwaters play in aquatic networks, they remain understudied. Most first order streams are not fully represented on current maps (Benda et al. 2005). Also, headwater streams are influenced by environmental change (Wipfli et al. 2007). The strong link they share with terrestrial environment affect in particular stream DOC concentrations that show fluctuations throughout the year—being lower during winter and higher during summer (Dawson et al. 2008; Laudon et al. 2011). These seasonal variations are influenced by discharge and temperature as secondary drivers that are also sensitive to climate change (Köhler et al. 2009; Sidle et al. 2000). Similar to DOC, bacterial population dynamics, such as for example the bacterial
abundances in streams show recurring seasonal patterns (Hullar et al. 2006).

DOC provides carbon and energy for bacterial heterotrophs (Williams and Del Giorgio 2005). Its variation influences bacterial biogeography (Findlay et al. 2008) and stream metabolism (Berggren and del Giorgio 2015). In some aquatic systems, the prevalence of major bacterial clades has been shown to be significantly modified by the presence and composition of DOC (Amaral et al. 2016). Thus, variations in DOC and bacterial abundances throughout the year support the dynamic nature of headwater streams. However, the origin of these variations is hitherto not well understood.

Soils constitute one of the most important microbial habitats, containing a great diversity of bacteria. The microbes have crucial roles in nutrient cycling, maintaining soil fertility and soil carbon sequestration (Fierer 2017; Philippot et al. 2013; Wagg et al. 2014). Soil microbes have a significant effect on biological processes as they produce CO₂ while respiring soil organic matter, influence soil acidity and regulate soil dynamics (Fierer 2017). Within soils, microbial densities vary greatly. Microbial biomass is commonly one to two orders of magnitude lower in deeper soil horizons than at the soil surface (Federle et al. 1986; Fierer et al. 2003; Taylor et al. 2002).

Soils are connected to river networks by water flows that enter small streams through the riparian zone (Richardson and Danehy 2007; Sidle et al. 2000). The relationship between soils and streams has been extensively studied regarding the seasonal variation in DOC concentrations. In forested catchments, snowmelt and storms mobilize DOC from the surrounding soils that reaches the streams through subsurface flow-paths and induce an increase in stream DOC concentration (Fasching et al. 2015; Laudon et al. 2004). At the same time, soils have been proposed to serve as critical reservoirs and thus source areas of microbes for surface waters (Crump et al. 2012; Hullar et al. 2006). The transport of microbes from soils to small streams was previously shown using DNA sequencing techniques and this transfer has been hypothesized to alter downstream microbial communities (Crump et al. 2012; Ruiz-González et al. 2015; Savio et al. 2015). Bacterial transfer to streams was also shown to vary throughout the year and in response to changes in temperature and precipitation (Hassell et al. 2018; Hermans et al. 2019; Teachey et al. 2019). However, DNA sequencing data do not provide quantitative information and, to this day, little is known about the dynamics and seasonality of bacterial abundances in small headwater streams (Richardson 2019).

Here we propose that the dynamic connectivity between soils and streams acts as a principle origin of variation of bacterial abundances in headwater streams. To our knowledge, no previous study has analysed bacterial abundances in soil water from catchment soils in order to compare it to nearby headwater streams and shed light on their transient connectivity. Thus we aim to explore the dynamic link between soils and headwater streams in terms of DOC and bacterial abundances through time and at different flow conditions. We pose the following questions for this study: What is the soil contribution in terms of providing DOC and microbial life to small headwater streams? What are the temporal dynamics of these provisions? We hypothesize that DOC concentrations co-vary with bacterial abundances, as both are mobilized from the soil simultaneously. In addition, we propose that varying flow conditions and soil saturation influence the degree of connectivity between soils and streams and thus control the DOC concentrations and bacterial abundances in small streams.

Methods

Study site

Sampling was carried out on six small headwater streams draining into Lake Lunz located in the eastern Alps near Lunz am See, Austria (Fig. 1). Lake Lunz receives most of its water from the Oberer Seebach Catchment. This catchment is approximately 25 km² and is largely pristine. Vegetation is dominated by Fraxinus excelsior, Acer pseudoplatanus, Fagus sylvatica, Salix caprea and Picea abies (Battin 1999; Fasching et al. 2015). During 2018, air temperature averaged 8.96 °C and precipitation averaged 1500 mm. Snow cover can extend from November to May, but in some winters little snow is present. Between May 2010 and August 2013, the Oberer Seebach had an average DOC concentration of 1.56 ± 0.3 mg L⁻¹ during autumn and of 1.79 ± 0.4 mg L⁻¹ during summer (Fasching et al. 2015). The catchment is characterized by glacial alluvial deposits, which are underlain by a low-permeability layer of ancient lake sediment and calcareous rock (Battin 1999).

Three hillslopes around Lake Lunz were selected for this study (Rehberg, RBG; Schlögelberg, SBG; and the WasserCluster Lunz slope, WCL; Fig. 1). On each hillslope, two headwater streams were chosen, referred to as RBG2 and 3; SBG1 and 2; and WCL1 and 2. The streams are of first stream order, small in size (width between 0.1 and 0.3 m) and enter Lake Lunz within 800 m of their flow length. All hillslopes are steep; the typical slope of the hillslopes is 29%.

Stream sample collection

All six streams were sampled for DOC every two weeks in spring, summer and fall, respectively and approx. every four weeks in wintertime from July 2017 to end of June.
2019. Analyses of bacterial abundances started in May 2018. Additional samplings were performed during and following high flow events in order to characterize those events and compare the dynamics of the soil/stream connectivity at different flow conditions. At each sampling, the same four to five locations along each stream, with an approximate distance of 35 m between sites were sampled (Fig. 1).

The three hillslopes differed in orientation, aspect and land use (Fig. 1). WCL and SBG are located on a north-facing forested mountainside, whereas the RGB streams face towards the southwest. Land use of the SBG streams is mixed deciduous forest, with little forestry activity during the last two decades. The WCL hillslope is similarly forested, but WCL1 enters an open meadow towards the lake. Also, both WCL streams were temporarily affected by forestry machinery crossing the stream in summer 2017. Land cover of the RGB streams is also forest, except an open meadow uphill of RGB3, which is used for low intensity cattle grazing during summer.

**Soil runoff collection**

Soil water was collected at two different soil depths (approximately 30 and 50 cm, referred to A and B, respectively in the following) from each hillslope (RGB, SBG, and WCL). These samples were drawn from soil runoff samplers, which were permanently installed in wet locations in close proximity (<15 m) to at least one of the two streams on each hillslope (Fig. 1). Each soil runoff sampler consisted of two stainless steel soil water collectors of 2 m width. The collectors were pressed laterally (angle approx. 10° to allow water flow) into the hillslope (30–40 cm) at the respective depth and the outflow covered by a thick plastic tarp. The samplers intercept soil water from the unsaturated zone of the soil profile as it drains downwards along the hillslope.
The sampling of the soil samplers was then performed by placing acid washed and pre-combusted (4 h at ~450 °C) 2 L Schott bottles below the outflow of each sampler the day before sampling. Sampling was then done by homogenizing the content of the Schott bottles and filtering samples of soil water, similar to stream samples. During many occasions, Schott bottles were overflowing at the time of sampling, while at the driest summer conditions, some soil samplers also remained dry.

**DOC sampling**

Stream water samples were filtered in the field on 0.7 µm pre-combusted glass-fiber filters (Whatmann GF/F) into 40 mL glass vials using a syringe and syringe-type filter holders. Vials were prepared ‘organic carbon free’ by acid-washing and subsequent combustion (450 °C, 4 h). The filtrate was analysed for DOC concentration within 24 h following sampling using a Sievers 900 TOC Analyser (GE Analytical Instruments, Boulder, CO, USA) operated with an inorganic removal unit.

**Bacterial cells counting**

Unfiltered water samples were fixed for 2 h with paraformaldehyde (1% final sample concentration) within 6 h after sampling. Fixed samples were then diluted 1:2 for stream water and 1:10 for soil runoff water using deionised and sterile ultrapure water. In order to separate bacteria from larger inorganic particles, samples were sonicated in a bath for 5 min and immediately filtered using sterile bolting cloth of 20 µm mesh-size. This method yields the total number of bacteria in the water column, independent of their potential bond to a particle. The filtrate was kept at 4 °C until counting, which was done within 48 h.

SYBR Green I was prepared by diluting a 10,000 × stock solution (prod. nr. 7563; Invitrogen, Life Technologies, Paisley, UK) with sterile Tris EDTA buffer (AMRESCO Inc., Solon, OH, USA) to a concentration of 100× and was used to stain the samples at a final concentration of 1×. Dye was stored at −20 °C. The staining was performed in the dark for at least 20 min prior to analysis. All measurements were performed with a Beckman Coulter CytoFLEX flow cytometer (Beckman Coulter GmbH, Krefeld, Germany) equipped with a laser providing 50 mW at 488 nm and five different fluorescent channels. The threshold to trigger events was set in green fluorescence (525 nm) as stained bacteria would emit green light when excited with blue (488 nm). Bacterial cells were discriminated from the background using previously measured unstained samples and green fluorescence versus side scatter dot plots.

**Hydrology**

We obtained precipitation data from the records of the meteorological station at the Biological Station Lunz operated by the Zentralanstalt für Meteorologie und Geodynamik (ZAMG), Austria. Daily values represent the accumulated precipitation (in mm) for the past 24 h, measured at 7 am CET.

Variability in stream runoff was analysed using data from the Oberer Seebach station at the inlet of Lake Lunz (Fasching et al. 2015). As such, this station is located upstream of the study streams which drain directly into Lake Lunz (Fig. 1). However, the Oberer Seebach station was assumed more representative of headwater streams as compared to the station at the lake outlet. Stream discharge ($Q$ in m$^3\cdot$s$^{-1}$) was derived from 10 min water level measurements by using well established rating curves (Fasching et al. 2015). From these data, daily average discharge was calculated and used for further analysis. As some data were missing due to malfunctioning of logger systems, missing daily discharge values were estimated from the gage at the lake outlet ($R^2=0.91$ for daily $Q$’s from both stations).

We classified flow conditions by the level of stream discharge. Based on daily average discharge on the whole study period, we defined discharge higher than 90% of the time ($Q > 1.86 \text{ m}^3\text{ s}^{-1}$) as high flow. This classification based on the flow percentile is similar to the definition used in earlier work at the OSB (Fasching et al. 2015). However, we note that our threshold value is different, due to different hydrological conditions during our study period as compared to the earlier study. Similarly, low flow was defined as all discharge lower than the high flow threshold ($Q ≤ 1.86 \text{ m}^3\text{ s}^{-1}$), thus effectively representing intermediate and low flows.

**Data analysis**

DOC concentrations and bacterial abundances were tested for differences between high and low flows. In order to avoid statistical inconsistencies due to the nested sampling design, only a mean value of DOC concentrations and bacterial abundance per stream and sampling event was used for this comparison. Tests for normal distribution were conducted using the Shapiro–Wilk test. As the results indicated non-normal data, we performed non-parametric Mann–Whitney $U$ test to compare the average difference in DOC concentrations and bacterial abundances between high flow and low flow. We considered $p < 0.05$ as significant and $p < 0.001$ as highly significant. To compare the correlation between DOC and bacterial abundances from stream water with the one from soil water, we used an analysis of covariance with the type of sample as the nominal variable. Because of the large range of values measured for both DOC concentrations and bacterial
abundances, data were log-transformed ($\log_{10}$) before the analysis. When seasonal comparisons (also Mann–Whitney $U$ test) were carried out, seasons were defined similar to earlier work (Fasching et al. 2015) by the month as follows: Spring from 01.03 to 31.05; summer from 01.06 to 31.08; autumn from 01.09 to 30.11; and winter from 01.12 to 28–29.02.

**Results**

**DOC concentrations**

DOC concentrations ranged from 0.39 to 5.56 mg L$^{-1}$ with an overall mean and standard deviation of $1.16 \pm 0.56$ mg

Fig. 2  a Daily precipitation and b daily discharge in the OSB, c DOC concentrations and d bacterial abundances throughout the study years. Symbols denote the streams as follows: blue dots RBG2, blue triangles RBG3, green filled squares SBG1, green pluses SBG2, red crossed squares WCL1 and red stars WCL2. Each symbol represents a mean, bars the standard deviation of all samples taken along each stream during one sampling occasion, respectively. The dataset includes a total number of 811 DOC samples and 547 bacterial samples (color figure online)
L$^{-1}$ in stream water throughout the study period (Fig. 2). All streams presented an increase in DOC concentration of on average 17.9% between the most upstream and the most downstream sampling locations (Fig. 3a). This increase along the streams’ flow path was significant during low flow conditions (+18.1%, Mann–Whitney $U$, $p < 0.001$) but not during high flow conditions (+4.4%, $p = 1$). Further, stream DOC was found to be lower during the winter—with a mean concentration of 0.87 ± 0.2 mg L$^{-1}$—and higher during autumn and summer—with respective means of 1.16 ± 0.35 and 1.31 ± 0.53 mg L$^{-1}$. The higher values were measured during high flow events, especially summer storms (Fig. 4a). During storms, the mean DOC concentration increased significantly by 44% (Mann–Whitney $U$, $p < 0.001$), from 1.1 ± 0.36 to 1.57 ± 0.7 mg L$^{-1}$ as compared to low flow conditions.

In soil runoff, measured DOC concentrations were significantly higher than in streams, ranging from 0.98 to 36.9 mg L$^{-1}$ with an overall mean of 9.37 ± 8.21 mg L$^{-1}$, being eight times higher than the overall stream average. However, in soil water, DOC concentrations were not found to differ significantly between high flow and low flow conditions (Mann–Whitney $U$, $p = 0.9$) and seasons (pairwise Mann–Whitney $U$, $p = 0.38$) (Fig. 4c).

**Bacterial abundances**

Bacterial abundances ranged from 19,300 to 3,862,800 cells mL$^{-1}$ with an overall mean of 394,000 ± 509,000 cells mL$^{-1}$ in stream water throughout the year (Fig. 2). Concentrations were found to be lower during winter—with a mean abundance of 162,900 ± 69,500 cells mL$^{-1}$—and higher during summer—with a mean abundance of 488,700 ± 514,400 cells mL$^{-1}$. In contrast to DOC, only some streams showed an increase in bacterial abundances between the most upstream and the most downstream sampling locations, while in others no consistent pattern was visible (Fig. 3b). Similar to DOC, high values were measured during high flow events (Fig. 4b), during which the mean bacterial abundance in streams...
increased significantly by 164.5% (Mann–Whitney U, $p < 0.001$), from $304,700 \pm 372,400$ to $806,100 \pm 620,700$ cells mL$^{-1}$ as compared to low flow conditions. In soil runoff, measured bacterial abundances were significantly higher than in streams ($p < 0.001$), ranging from $327,300$ to $35,412,300$ cells mL$^{-1}$ with a mean of $4,450,400 \pm 5,985,200$ cells mL$^{-1}$, being ~10 times higher than the overall stream average. Similar to DOC, bacterial abundances were not found to be significantly different between high flow and low flow conditions in soil water (Fig. 4d).

**Correlation between DOC and bacterial abundance**

We investigated the relationship between DOC concentration and bacterial abundance in stream and soil water (Fig. 5). Log$_{10}$ DOC was found to be positively correlated with log$_{10}$ bacterial abundance in stream water ($r^2 = 0.61$, $p < 0.001$) and also in soil water ($r^2 = 0.34$, $p < 0.05$). The slopes of the regression lines were significantly different (0.35 for stream and 0.47 for soil water, respectively, $p < 0.01$) as well as their $y$-intercept ($p < 0.001$).

**Discussion**

The major finding of this study is that the mobilization of bacteria from catchment soils to headwater streams is controlled by hydrological variability. Specifically, we provide evidence that high-flow events, such as rainstorms, cause an enhancement of bacterial mobilization (Fig. 4). Further we show that the timing of increase and the magnitude are similar to the mobilization of DOC from catchment soils. But whereas the responses of DOC to storm events in first order streams have been well described in the past (Fasching et al. 2015; Hinton et al. 1998), we now demonstrate that dynamics similar to those of DOC are also controlling microbial abundances in small streams.

We observed an increase in DOC concentration at all streams along their flow length from their spring to the end of our study reach (Fig. 3a). We propose that this increase is the result of intensified soil contributions along the streams flow length. Our study was located in a karst region. Most permanently draining streams in this region are groundwater fed, as they would otherwise cease during dry spells. The increase of DOC along the streams...
indicates that our study streams are likely fed with spring water with low DOC that is then enriched along the flow at the steep hillslope by high DOC soil water. This conceptual model contrasts somewhat with the traditional view of first order streams being a direct link between catchment soils and streams in other regions (Schelker et al. 2012).

As an alternative explanation, enrichment of DOC along the flow length could also be generated by enhanced primary production that enhances DOC concentrations, especially during spring and summer low flow (Harjung et al. 2019). However, many previous studies have shown that first order streams presented very low autochthonous primary production that accounted for less than 5% of the annual organic carbon inputs in these ecosystems (Fisher and Likens 1973; Mulholland 1997; Richardson 2019). In addition, the fact that the increase was independent of season, i.e., that it was also present during the fall and winter, makes this explanation less probable. Further, if DOC deliveries from primary production were relevant, additional variation based on light availability for stream primary producers would be expected (Dodds et al. 1996; Hall et al. 2015). Our study streams differed in their light regimes (shading by closed forest canopy at SBG1, SBG2 and RGB2, while WCL1, WCL2 and RBG3 are at least partly open, Fig. 1), as well as their aspect. However, we were unable to identify a light-induced variation in our downstream DOC pattern. Thus we conclude that increasing soil contributions along the flow length is the most probable explanation for the increasing DOC concentrations along our study streams. Interestingly, an increase in bacterial abundances along the stream flow path was also indicated for some streams (Fig. 3b) but this increase was not significant and generally weaker than the one observed for DOC concentrations. At present, we cannot explain these differences, but consider soil community dynamics or minor variations of how bacteria are mobilized from soils as compared to DOC as possible explanations. However, these minor differences appear to be overwhelmed during events resulting in synchrony in increase.

Our findings on the variation of DOC concentration and bacterial abundance with varying discharge expand our present knowledge on the dynamics of headwater streams. High flow events led to a significant increase in DOC concentration by 70.8% and in bacterial abundance by 32.4%. Previous work has described the increase of DOC concentration in streams during storms following enhanced soil contributions.
(e.g., Ågren et al. 2008; Fasching et al. 2015; Hinton et al. 1998).

The link between soils and streams in terms of microbial communities has been investigated in the past using sequencing techniques (Crump et al. 2012; Ruiz-González et al. 2015; Savio et al. 2015). However, few studies have investigated the temporal dynamics of bacterial abundances in streams (Hullar et al. 2006; Richardson 2019). Here we found that high flow conditions, as they are commonly present in our study streams during rainstorms or snowmelt, caused a general increase in bacterial abundances (Fig. 4b). This pattern may be explained by increased deliveries of microbes from the catchment soils, similar to the mobilization of DOC. Alternatively high bacterial abundances could also be caused by the remobilization of in-stream bacteria in upstream reaches. The latter would be caused by high flow velocities transiently enhancing the shear stress on benthic biofilms. The biofilm would then release bacteria during high flow (Paul et al. 2012; Stewart 1993). However, some of our observational points were located at the very source area of the first order streams (upstream length < 20 m). These sites showed the same increase in bacterial abundance during events, as those further downstream. Thus we suggest that provision of microbes from upstream reaches is small, as compared to soil contributions.

The water residence time is small in the study streams, ranging from minutes to hours at the maximum. For example, we quantified a mean flow velocity of 0.1 m s^{-1} across a distance of ~ 100 m in WCL1 at intermediate flow conditions. Assuming similar velocities for all streams suggest that the residence time in the study reaches rarely exceeds 2 h. We argue that this is too short for relevant bacterial growth in the free flowing water column, especially during high flow events. Therefore, we attribute the dynamic increases in bacteria during runoff events in our streams to enlarged soil contributions, rather than instream growth.

Following these notions, we suggest that soil inoculation of headwater streams is a dynamic process that may be largely controlled by the intensity of hydrometeorological events. Consequently, we propose that most headwater streams are subject to a dynamic and reoccurring microbial inoculation from their catchment soils during storm events. This dynamic microbial inoculation of streams by soils may then have some impact on the development of the microbial communities in the downstream river network (Besemer et al. 2013; Ruiz-González et al. 2015; Savio et al. 2015; Widder et al. 2014). However, at present this potential impact remains largely unknown, as most previous studies on the role of soil inoculation on stream microbial communities did not consider the temporal dynamics of inoculation (Besemer et al. 2013; Monard et al. 2016; Ruiz-González et al. 2015; Widder et al. 2014).

The correlation we found between log_{10} DOC and log_{10} bacterial abundance in streams further supports our view that both constituents are mobilized concurrently from the catchment soils and that this mobilization is enhanced during high flow. The general mechanisms proposed for DOC mobilization from soils to streams include the flushing of DOC from the organic top soil (A-horizon) (Boyer et al. 1996), as well as from carbon rich riparian soils (Laudon et al. 2011; Schelker et al. 2013). Similarly, the abundance of soil bacteria has been suggested to vary with soil depth, with the highest abundance and microbial diversity present in the organic rich top soils (Fierer et al. 2003). This suggests that conceptual models developed for the mobilization of DOC from soils to streams (Boyer et al. 1996; Laudon et al. 2011; Weiler and McDonnell 2006; Zarnetske et al. 2018) could potentially also be applied to explain the variation in bacterial abundances in streams in the future. Nevertheless, the high ratio between soil-stream interfaces and water volume in headwater streams promotes transfer from soil to streams but the relationship might be weaker in larger fluvial systems. Also, pre-alpine streams have low DOC concentrations (Fasching et al. 2015) in comparison to other fluvial ecosystems, such as boreal streams (Dawson et al. 2008; Laudon et al. 2011). Further investigations must be conducted to determine if the relationship between DOC concentrations and bacterial abundances also holds true in more DOC-rich environments.

Interestingly, the strong correlation of log_{10} DOC concentration per log_{10} bacterial abundance observed in streams was also present in soil water. Although the slopes and the Y-intercepts of the linear models for soils and streams were significantly different, the general pattern of increasing log_{10} abundance with increasing log_{10} DOC remained. However, the significant differences between the relationships likely indicate fundamental differences in ecosystem structure regarding the availability of carbon versus the size of the maintained bacterial population in stream versus soil ecosystems.

Among the main distinctions between lotic and terrestrial ecosystems is the difference in the ratio between dissolved and particulate organic carbon, with a larger ratio in streams versus soils (Grimm et al. 2003). Also, primary producers show stoichiometric and lifespan differences (lower C:N and P:N ratios, more short-lived in streams) (Grimm et al. 2003; Nowlin et al. 2008). Thus one possible explanation for the observed differences in the relationships in our oligotrophic system would be that streams can maintain higher populations of primary producers during nutrient limitation. Further, the generally higher ratio of consumers versus producers in streams (Grimm et al. 2003) could then result in the difference of higher log_{10} bacterial abundance versus log_{10} DOC.
Alternatively, differences in the size of the maintained microbial population between soils and streams could also stem from differences in DOM composition between the two systems. In soils, degraded humic-like DOM components are typically dominant, while in streams labile, protein-like DOM is present along with humic substances (Fasching et al. 2015; Hutchins et al. 2017). This could also cause a general advantage in maintaining an extended microbial population in streams.

Future research should address two main aspects. First, the precise nature of the microbial communities that are delivered from soils to streams during hydrological events. At present, the community composition of these pulses of microbial life are not well understood (Monard et al. 2016; Ruiz-González et al. 2015). Second, there is currently only limited knowledge about the relevance of the dynamic and reoccurring microbial soil inoculation for stream microbial community development in the receiving waters (Besemer et al. 2013; Widder et al. 2014).

Overall we show that soils provide a dynamic inflow of bacteria and DOC to first order streams resulting in a dynamic and reoccurring inoculation of stream microbial communities from catchment soils during runoff events. We propose that this process is potentially important for the development of microbial communities of downstream river networks. We suggest future research on the community composition and the potential downstream effects of this dynamic soil inoculation process.

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