Complex interactions of in-stream dissolved organic matter and nutrient spiralling unravelled by Bayesian regression analysis

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Abstract. Uptake and release patterns of dissolved organic matter (DOM) compounds and co-transported nutrients are entangled, and the current literature does not provide a consistent picture of the interactions between the retention processes of DOM fractions. We performed plateau addition experiments with five different complex DOM leachates in a small experimental stream impacted by diffuse agricultural pollution. The study used a wide range of DOM qualities by including leachates of cow dung, pig dung, corn leaves, leaves from trees, and whole nettle plants. We measured changes in nutrient and dissolved organic carbon (DOC) concentrations along the stream course and determined DOM fractions by fluorescence measurements and parallel factor (PARAFAC) decomposition. To assess the influences of hydrological transport processes, we used a 1D hydrodynamic model.

We developed a non-linear Bayesian approach based on the nutrient spiralling concept, which we named the “interactions in nutrient spirals using Bayesian regression” (INSBIRE) approach. This approach can disentangle complex interactions of biotic and abiotic drivers of reactive solutes’ uptake in multi-component DOM sources. It can show the variability of the uptake velocities and quantify their uncertainty distributions. Furthermore, previous knowledge of nutrient spiralling can be included in the model using prior probability distributions. We used INSBIRE to assess interactions of compound-specific DOM and nutrient spiralling metrics in our experiment.

Bulk DOC uptake varied among sources, showing decreasing uptake velocities in the following order: corn > pig dung > leaves > nettles > cow dung. We found no correlations between bulk DOC uptake and the amounts of protein-like compounds or co-leached soluble reactive phosphorus (SRP). The fastest uptake was observed for SRP and the tryptophan-like component, while the other DOM components’ uptake velocities more or less resembled that of the bulk DOC. Almost all DOM components showed a negative relationship between uptake and concentration, known as efficiency loss. Furthermore, we observed a few negative and (weak) positive interactions between the uptake and the concentration of different components, such as a decreased uptake of protein-like compounds at high concentrations of a high-molecular-weight humic-like compound. We also found an influence of the wetted width on the uptake of SRP and a microbially derived humic substance, which indicates the importance of the sediment–water interface for P and humic C cycling in the studied stream.

Overall, we show that bulk DOC is a weak predictor of DOC uptake behaviour for complex DOM leachates. Individual DOM compound uptake, including co-leached nutrients, is controlled by both internal (quality-related) and external (environmental) factors within the same aquatic ecosystem. We conclude that the cycling of different C fractions and their
mutual interaction with N and P uptake in streams is a complex, non-linear problem, which can only be assessed with advanced non-linear approaches, such as the presented INS-BIRE approach.

1 Introduction

Dissolved organic matter (DOM) from terrestrial sources plays a key role in the metabolism and the ecological state of streams and rivers by controlling the activity and the composition of microbial communities (e.g. Freixa et al., 2016) and influencing a variety of aquatic biogeochemical processes (Tank et al., 2010). The quantity and the quality of DOM affect the aquatic bacterial respiration (e.g. Besemer et al., 2009; Niño-García et al., 2016), change the ratio between autotrophy and heterotrophy (Lutz et al., 2012; Martínez et al., 2017), and influence the microbial uptake of dissolved inorganic nitrogen (DIN; e.g. Bernhardt and Likens, 2002; Taylor and Townsend, 2010; Wymore et al., 2016) and soluble reactive phosphorus (SRP; Gibson and O’Reilly, 2012; Stutter et al., 2020; Weigelhofer et al., 2020), amongst others.

The influence of DOM on nutrients is mutual (e.g. Mineau et al., 2013; Stutter et al., 2020; Weigelhofer et al., 2020) due to the demand of microbes for carbon (C), nitrogen (N), and phosphorus (P) in a molar ratio that approximates their C : N : P ratio (Small et al., 2009; Stutter et al., 2018; Welti et al., 2017; Godwin and Cotner, 2018). The importance of this stoichiometric control of nutrient and organic carbon uptake in streams and rivers has been known for some time (Cross et al., 2005; Dodds et al., 2004) but has increasingly gained attention over the last few years with the discovery of anthropogenic impacts on pristine C : N : P ratios and DOM compositions (Stutter et al., 2018; Xenopoulos et al., 2021). Intensive land use has changed the origin, amount, and transport of terrestrial DOM to streams, thereby probably altering the DOM in-stream processing (Weigelhofer et al., 2020).

According to ecological stoichiometry, in-stream DOM uptake and retention is primarily related to the availability of inorganic nutrients, whether they already exist in the freshwater ecosystem (environmental control of DOM uptake) or are provided by the DOM source itself (intrinsic control through the DOM quality; e.g. Bernhardt and McDowell, 2008; Graeber et al., 2015; Gücker et al., 2016; Wickland et al., 2012). Field and laboratory studies show that DOC uptake can be positively affected by N and P concentrations in the water column (Catalán et al., 2018; Mineau et al., 2013). DOC uptake is also high in N- and P-rich DOM sources, such as leaves from fertilized trees or agricultural areas (Mineau et al., 2013; Mutschlecner et al., 2018; Weigelhofer et al., 2020).

In addition to nutrient interactions, DOM uptake depends on the structure and the bioavailability of the individual DOM compounds (Guillemette and Giorgio, 2012; Mineau et al., 2016). High uptake rates have been observed for protein-rich, low-molecular-weight DOM sources such as leachates of fresh leaf litter, macrophytes, and periphyton (Berggren et al., 2010; Koehler et al., 2012). In contrast, the soil leachates’ biodegradability has been described as generally low (e.g. Fellman et al., 2009b; Hansen et al., 2016). Finally, the in-stream uptake of DOM may be influenced by environmental factors other than nutrient concentrations, such as the hydrology and morphology of the respective reach or the biofilms’ composition (Casas-Ruiz et al., 2017; Romani et al., 2004; Weigelhofer et al., 2020).

Thus, unravelling the underlying mechanisms of in-stream DOM uptake is complex. While a considerable part of the reactive N and P exists as small and simple molecules, dissolved organic carbon (DOC) is bound in a mixture of differently structured organic molecules, with retention times varying by several orders of magnitude (Cory and Kaplan, 2012; Mineau et al., 2016). The production of new compounds during DOM decomposition may further complicate an accurate assessment of the DOM uptake (Stevenson and He, 1990; Tsutsuki and Kuwatsuka, 1979). Mass balance approaches or calculations of first-order decay curves from addition experiments have been successfully used in numerous studies to estimate in-stream uptake of DIN, SRP, and bulk DOC (e.g. Bernhardt and McDowell, 2008; Catalán et al., 2017; Covino, 2012; Ensign and Doyle, 2005; Mineau et al., 2013; Schiller et al., 2011; Stream Solute Workshop, 1990; Weigelhofer et al., 2018b). However, these methods are often limited in quantifying the uptake of individual DOM components. This limitation restricts and complicates the analyses of interactions among different DOM components and their role in the overall DOC uptake (Mineau et al., 2013; Stream Solute Workshop, 1990; Weigelhofer, 2017; Weigelhofer et al., 2018b).

Our study aimed at investigating the effects of DOM quality on the in-stream DOM uptake. We developed an approach to quantify complex interactions between individual DOM compounds, including co-leached N and P, and to elucidate their combined role in the overall DOM retention. For this purpose, we performed several short-term plateau additions with different DOM sources in an agriculturally influenced headwater stream according to the nutrient spiralling concept (Stream Solute Workshop, 1990). We used leachates from natural and human sources (e.g. leaves, manure) to see how in-stream DOM processing may be altered due to anthropic land use changes. Because of the diverse composition of DOM, we decided to extend the equations from the nutrient spiralling concept (Stream Solute Workshop, 1990) and use a Bayesian approach to analyse interactions between and influences of different DOM components and nutrients, including uncertainty propagation. Bayesian statistics is a suitable tool for ecological and biogeochemical questions, allowing users to assess the natural variability and assign degrees of belief in hypotheses based on measured data (Arhonditsis et al., 2008; Berger and Berry, 1988; Cox, 1946; Ellison,
2 Methods

2.1 Site description

The experiment was carried out in the Hydrological Open Air Laboratory (HOAL; Fig. 1) in Petzenkirchen, Austria (Blöschl et al., 2016; for further information and a detailed map, see https://hoal.hydrology.at/the-hoal, last access: 3 April 2021). The HOAL is a small catchment that was transformed into a hydrologic observatory to foster scientific research. It features several permanently installed sensors measuring discharge, different water parameters, and the weather. Many studies on surface and subsurface flow paths, evaporation, soil erosion, sediment transport, and nutrient dynamics have already been performed in the HOAL (Blöschl et al., 2016). The first-order stream has several inflows, including two natural springs, six drainage pipes, and one site with groundwater infiltration from a small wetland. The stream flows through a deciduous forest with two short open sections in the middle (points 4–5 and 7 in Fig. 2) dominated by herbaceous vegetation and grass on the banks. All inflows as well as the stream discharge are continuously monitored regarding water quantity and quality. Sediments are dominated by clay, which is washed in from the adjacent fields during storm events. Table 1 shows the extent and basic environmental characteristics of the stream.

For the experiments, we chose a study reach of 215 m situated between two lateral inflows which was divided into 10 subsections of 16 to 26 m lengths, depending on the accessibility (Fig. 2). The first sampling point (1) was located 19.4 m downstream of the injection point (0) to ensure complete mixing of the solutes in the water column. Full mixing was determined by measuring the electrical conductivity across the channel transect during a salt addition before the experiments. The study reach was characterized by a meandering stream course with frequent pools (up to 24 cm in depth) between sampling points 8 and 11 (Fig. 2). Between point 4 and point 5, Equisetum palustre and Juncus sp. were growing in the open channel. At point 7, the patchy canopy cover facilitated the growth of algae on the stream bed. During the experiment, the median temperature was 16.7 °C (interquartile range = 2.4), and the median conductivity was 633 µS cm⁻¹ (interquartile range = 23).

2.2 Experimental design

The experiment was performed over six consecutive weeks in July and August 2018. No major rain event occurred in the study area, and the average discharge was between 0.38 and 0.93 L s⁻¹. Ten additions with DOM leachates from five different sources were injected into the study reach using short-term plateau additions according to the Stream Solute Workshop protocol (Stream Solute Workshop, 1990; Weigelhofer et al., 2012). The respective leachate plus a NaCl solution acting as a conservative tracer were pumped into the stream at point 0 over 2 to 2.5 h via a peristaltic pump (Fig. 2). We used a mobile conductivity meter to identify plateau conditions in the stream at each sampling point. Approximately 10 min after reaching stable plateau conditions, water samples were taken at each sampling point to analyse nutrient concentrations, organic carbon concentrations, and DOM composition. Thus, the sampling time exceeded the average water travel time by far, ensuring proper mixing and a stable

Table 1. Extent and environmental characteristics of the HOAL.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length</td>
<td>620</td>
<td>m</td>
</tr>
<tr>
<td>Catchment size</td>
<td>0.66</td>
<td>km²</td>
</tr>
<tr>
<td>Arable land coverage</td>
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<td>%</td>
</tr>
<tr>
<td>Mean annual discharge</td>
<td>0.004</td>
<td>m³ s⁻¹</td>
</tr>
<tr>
<td>Peak discharge</td>
<td>2</td>
<td>m³ s⁻¹</td>
</tr>
<tr>
<td>Mean annual temperature</td>
<td>9.5</td>
<td>°C</td>
</tr>
<tr>
<td>Mean annual precipitation</td>
<td>820</td>
<td>mm yr⁻¹</td>
</tr>
</tbody>
</table>

2004; Jaynes, 2003; McCarthy, 2007). We incorporated non-linear nutrient uptake models observed in previous studies, such as the Michaelis–Menten kinetics or the nutrient efficiency loss model (Dodds et al., 2002; O’Brien et al., 2007), in our approach to describe relationships between concentrations and uptake velocities mathematically. Our approach enabled us to (1) analyse how uptake processes of different components influence each other; (2) test our mathematically pre-formulated assumptions with the measured data, including the remaining errors; (3) consider the natural variability of each parameter; and (4) include knowledge of nutrient and DOC uptake kinetics from previous studies in our models. We called our approach “interactions in nutrient spirals using Bayesian regression” (INSBIRE). With INSBIRE, we addressed the following questions:

- What are the differences in the bulk DOC uptake velocity of different leachates?
- How do selected DOM components behave in comparison to the bulk DOC uptake velocity?
- Which factors and interactions influence the uptake velocity of the bulk DOC, the individual DOM components, and the co-leached nutrients N and P?

We expected nutrient- and protein-rich leachates to show higher uptake velocities than the other leachates. Low-molecular-weight protein-like compounds would show a faster uptake, while high-molecular-weight, aromatic compounds would show a slower uptake than the bulk DOC. We also expected to find positive influences of co-transported nutrients on the bulk DOC uptake and negative influences of low-molecular-weight protein-like compounds on the uptake of high-molecular-weight, aromatic compounds.
Figure 1. Hydrologic Open Air Laboratory (HOAL): catchment, stream, sampling points, and location within Austria. A satellite image of the site also shows the inflows before and after the sampling stretch (catchment outlet: MW; inlet: SYS4; tile drains: FRAU1, FRAU2, SYS1, SYS2, and SYS3; erosion gullies: E1 and E2; springs: Q1 and K1; wetland run-off: A1 and A2). GIS data and aerial provided by the Bundesamt für Wasserwirtschaft in Petzenkirchen, Austria.

Figure 2. Sampling scheme and general parameters of the stream. Point 0 marks the location of the solute addition, and points 1–11 the sampling sites.

state during sampling (see Fig. 2). After shutting the addition off, the change in conductivity was recorded until salt concentrations had returned to ambient levels. Additions were limited to a maximum of two times per week with at least 48 h between two consecutive samplings, allowing the system time to recover. Each leachate was added twice to the stream with an interval of 5–7 d to minimize adaptions of the microbial community and interferences among leachates. The additions created concentration peaks equal to or below local rain events. Each Monday, we sampled ambient concentrations to interpolate background conditions for the days with addition experiments. All samples were taken between 10:00 and 14:00 CEST to ensure comparability. As the environment changes naturally over time (e.g. discharge, temperature), different additions cannot be compared if the interval between them is too long. However, extremely short intervals and/or long addition times may lead to adaptions of the microbial community. Thus, the sampling schedule represents a compromise based on our experiences in nutrient addition experiments (Weigelhofer, 2017; Weigelhofer et al., 2012, 2018b) and on the long-term weather and discharge data of the stream. During our experiments, environmental changes were negligible due to highly stable weather conditions and the lack of human activities in the experimental area. We also observed no systematic changes of the DOM, N-NO$_3$, and SRP uptake over time, indicating that any potential adaptions or responses of the microbial community to these short and low pulses did not affect the results of the study significantly.

2.3 Preparation of the leachates

The leachates were prepared from 50 g L$^{-1}$ dry matter of cow and pig dung, foliage from local trees (Acer platanoides, Acer pseudoplatanus, Lonicera xylosteum, Pteridium aquilinum, Sambucus nigra), nettles (Urtica dioica), and corn plant (Zea mays) leaves. We leached the substrates with nutrient-poor water from a local well under aerated condi-
flections in a barrel over 24 h. The leachates were filtered in steps of 2 and 0.5 mm using stainless-steel sieves and 50 μm using a 25 cm spun filter cartridge (PureOne PS-10). The end volume was between 40 and 60 L. To avoid post-leaching changes in DOM, the leachates were prepared freshly for each addition.

Average DOC concentrations in the stream water were about 1.3 mg L$^{-1}$. We aimed to achieve an increase by about 3 mg L$^{-1}$ DOC in the experiments. Some sources proved difficult to leach in sufficient amounts, and parts of the leached DOC were degraded even during short storage. Thus, the DOC increase achieved during the experiments was between 0.2 and 2.3 mg L$^{-1}$. Even within the same source, leached amounts varied in concentration and composition between different additions. We consider these variations neglectable since we defined the leachates by their measured composition and not solely by their source. However, the fluctuations broadened the distributions of the measured values, providing more stable models and a more general picture of the uptake processes.

2.4 Analyses

Before the lab analyses, samples were filtered through pre-combusted Whatman glass microfiber filters, Grade GF/F (0.7 μm). We measured inorganic nitrogen as N-NO$_3^-$, nitrite (N-NO$_2^-$), and ammonium (N-NH$_4^+$) as well as SRP with a continuous-flow analyser (accuracy ±0.1 μg L$^{-1}$). DOC was measured with a Sievers® 900 portable total organic carbon analyser (accuracy ±2 %). We measured the DOM quality (excitation–emission matrices, or EEMs) via fluorescence spectroscopy with a Hitachi F-7000 fluorescence spectrophotometer and DOM absorbance with a Shimadzu UV-1700 spectrophotometer.

We analysed the data using R software version 3.5 (R Development Core Team, 2019) and tidyverse (Wickham et al., 2019). The DOM EEMs (11 sampling points, 16 sampling dates, 176 samples in total) were pre-processed using eemR (Massicotte, 2019); the fractions by fluorescence measurements and parallel factor (PARAFAC) analysis was done with staRdom (Pucher et al., 2019). The measured fluorescence EEMs were corrected for inner-filter effects, samples of ultra-pure water were subtracted, scatter bands were removed and interpolated, and the samples were normalized to Raman units. Samples were screened visually, and no unusual noise was found. After obtaining the first models, three outliers were identified using the samples’ leverages and excluded from the model. The components’ spectra were visually checked for plausibility. After that, a suitable model was validated using a split-half analysis. The final model did not express any problems related to those criteria. The outliers were included again to calculate loadings under the already-fixed components. To calculate the PARAFAC models and the split-half validation, we used 256 random initializations, a tolerance of 10$^{-11}$, and staRdom’s standard way of splitting the data (Pucher et al., 2019). We used Open-Fluor (Murphy et al., 2014) to compare and link the found components with other studies (Table 2).

2.5 Hydrodynamic modelling

A hydrodynamic 1D model was used to calculate the necessary hydraulic parameters using the software package HEC-RAS (Hydrologic Engineering Center’s River Analysis System). To create the digital terrain model, a cross-sectional approach was applied, where 64 cross sections were recorded at a distance of 0.8 to 6.8 m depending on structural variations and accessibility. A total of 251 points were measured in the stream with a theodolite (Leica TC805) and then merged with a 1 × 1 m floodplain area model (based on the official laser scan data of the province of Lower Austria) using the software package Surface-water Modeling System (Aquaveo, LLC). The model was calibrated with the discharge data recorded at the HOAL site by comparing the measured water surface elevation with the modelled one. The calibrated 1D model was used to calculate the hydraulic parameters flow velocity, water depth, wetted width, and water travel time at each sampling point for each sampling day.

2.6 Calculating interactions in nutrient spirals using Bayesian regression (INSBIRE)

The uptake was calculated using a Bayesian non-linear model and solved with a Markov chain Monte Carlo (MCMC) algorithm as provided in the R package brms (Bürkner, 2017) relying on stan (Carpenter et al., 2017). We used the Bayes factor (BF; Goodman, 1999a, b) for hypothesis testing and model comparisons. The BF is the ratio of the marginal likelihood of two competing hypotheses or models. A BF of 10 favouring a particular hypothesis or model means that this model is 10 times more likely to explain the measured data. The interpretation of the BF was conducted according to Kass and Raftery (1995). There, the evidence is “barely noteworthy” (1 < BF < 3.2), “substantial” (3.2 < BF < 10), “strong” (10 < BF < 100) or “decisive” (BF > 100). A BF < 1 corresponds to the inverse of the BF but in favour of the other hypothesis. Model selection using the BF also allows removing models prone to collinearity problems (Ghosh and Ghattas, 2015). The Bayes $R^2$ (Gelman et al., 2019) for each model was calculated to demonstrate the accuracy of the analysis.

We used the equations of the nutrient spiralling concept provided by the Stream Solute Workshop (1990) to develop our solute spiralling model, INSBIRE. All equations providing the basis of the model from the Stream Solute Workshop (1990) and all equations derived, transformed, and developed from these basic equations for the model development are shown and explained in detail in Sect. S1 in the Supplement. For a straightforward solving scheme like INSBIRE, a single-step analysis is necessary to determine all in-
terdependent parameters’ posterior distributions at once. Interactions, model weaknesses, collinearity (Ghosh and Ghat-
tas, 2015), and the variation of parameters can then be as-
essed and interpreted consistently.

Commonly, uptake length ($s_w$), uptake velocity ($v_f$), and areal uptake rate ($U$) are used to describe nutrient uptake (Dodds et al., 2002; O’Brien et al., 2007; Trentman et al., 2015; Weigelhofer et al., 2018b). We fitted all three param-
ters to the equations. The uptake length $s_w$ is known to change with different discharges, while $v_f$ is independent
of discharge (Dodds et al., 2002) and $U$ is related to the solvent concentration. We used $v_f$ to address our research
questions because the exclusion of hydrology provides the
best conditions for analysing biogeochemical relationships.
In alignment with Bayesian statistics (e.g. McCarthy, 2007),
we defined prior distributions for each parameter based on
the knowledge from other studies (e.g. Mineau et al., 2016)
to keep the parameters (e.g. $v_f$) within realistic ranges and
foster a stable fitting procedure. We provide an exemplary
R script that demonstrates INSBIRE (Pucher, 2020). A de-
tailed mathematical description of the INSBIRE approach
can be found in Sect. S1.

We used data from all experiments combined to perform
the parameter estimation, thus increasing the number of
points in our model. By doing that, we got a better insight
into processes and interactions underlying uptake that can
only be observed with different nutrient and DOM ratios.
In contrast to a fitting algorithm, which determines only the
most suitable value, the result of a Bayesian fitting is a dis-
tribution of probable parameter values showing the variabil-
ity in the stream and between experiments. Furthermore, we
set a threshold for complete retention of the added solutes,
at which the difference between plateau and ambient condi-
tions was equal to 2 times the accuracy of the lab analyses.
Measured values below this threshold were removed from the
analyses. Since the fluorescence of DOM increases linearly
with concentration (Kothawala et al., 2013), we used $F_{\text{max}}$
of the PARAFAC components analogues to concentrations
in these models.

We tested trends in the longitudinal nutrient concentra-
tions by comparing the assumption of constant concentra-
tion with exponential decay as proposed in the nutrient spiralling
concept (Stream Solute Workshop, 1990). The BF between
those two models was calculated to show which one is more
likely. For the analysis of the uptake of bulk DOM and indi-
vidual components (research questions 1 and 2), we used the
equations from the nutrient spiralling concept (Stream So-
lute Workshop, 1990) to calculate $v_f$ via a one-step fitting
procedure (for details, see Sect. S1 and Eq. S4). By adding
the leachate source to the uptake models as a random fac-
tor, we could determine quality-related differences between
the leachate sources. We also added the sampling date as a
random effect to our models to see if there are differences
between the two sampling dates of the same leachate source
(e.g. Ohno and Bro, 2006). A systematic change with the
sampling date for all or at least most nutrients and compo-
nents can also reveal experimental or analytical problems in
the execution process but was not observed in this study.

The analyses of influencing factors in bulk DOM and
component-specific uptake (research question 3) required the
adaptation of the original concept by including relationships
among different DOM components and co-leached nutrients.
This step was motivated by nutrient addition studies showing
different uptake models such as linear functions, power func-
tions (efficiency loss model), and Michaelis–Menten kinetics
(Dodds et al., 2002; O’Brien et al., 2007; Trentman et al.,
2015). We additionally tested an exponential function and an
asymptotic regression function. We decided to present only
the results of the power function (Eq. 1) because its inclusion
in the models showed the highest BFs (highest probability
to explain the observed data) in most cases. A considerable
advantage of the power function is that there is only one pa-
rameter to fit, making it less prone to over-fitting in complex
models (e.g. discussed in McElreath, 2016, chap. 7). Also, in
our experiment, the concentrations of the DOM components
and the co-leached nutrients did not reach uptake limits. In
such cases, uptake rate curves often exhibit a power func-
tion, probably representing the lower unsaturated part of a
saturation model within a concentration range naturally met
in stream systems. O’Brien et al. (2007) limited the exponent
of the power function ($m_i$ in Eq. 1) to negative values ($m_i < 1$
for $U$, but $m_i < 0$ for $v_f$) to describe the efficiency loss
behaviour. However, positive $m_i$ can also be used in the models
to describe situations where a substance improves the uptake
of another (e.g. Stutter et al., 2020). In our study, we looked
for both positive and negative interactions among compo-
nents, and thus we did not constrain the sign of $m_i$. For pos-
itive exponents $m_i$ in Eq. (1), the function would pass through
zero, which means that the absence of a stimulating compo-
nent automatically leads to a total collapse of DOM or nu-
trient uptake. However, such a total collapse is not expected
for DOM fractions since microbes can use other C sources.
Thus, we incorporated an added value $l$ in Eq. (1) as a degree
of freedom, whose relevance was tested during the model se-
lection process.

From a modelling point of view, any available variable can
be included and tested at that point. We decided to include
the wetted width in the formula because we expected an influ-
ence of the available benthic surface on the uptake processes.
Furthermore, the stream showed a fluctuation of the wetted
width by a factor of 2.8 between different dates and cross
sections. The considerations made in this chapter resulted in
the following equation:

$$v_f = kw\left(l + \prod_{i} C_{i,x,t}^{mi}\right),$$  \hspace{4cm} (1)

where $v_f$ is nutrient uptake velocity; $k$ is the uptake rate
factor (model parameter); $w$ is wetted width, constant at 1
to represent no influence (calculated by HEC-RAS and then
fixed); $l$ is the additive value (model parameter); $i$ is the index

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of DOM component or nutrient; $C_{i,x,t}$ is the concentration of compound $i$ at point $x$ and date $t$ (measured variable); and $m_i$ is an exponent determining the strength of the relations (model parameter).

The same fitting algorithm as for research questions 1 and 2 was used to derive the parameters $k$, $m_i$, and $l$ in Eq. (1). Additional information is provided in Sect. S1 and Eq. (S8). Relationships among components (including co-leached nutrients) were tested individually and in different combinations by adding factors of power functions according to the single factors in Eq. (1). The different combinations were compared to the initial model and to the next simpler models and were rated according to their BFs. When models with specific variables did not improve the predictability of the observed data, they were rejected and are not presented in the Results section below. By doing that, we determined models with meaningful component relationships and derived BFs for each variable included, representing the strength of evidence to support this inclusion.

For comparisons of uptake velocities of all co-leached nutrients and DOM fractions, we transformed the equation from the nutrient spiralling concept (Stream Solute Workshop, 1990) to derive $v_f$ for each nutrient and DOM component and between all pairs of sequent sampling points directly. Uptake velocities between nutrients and DOM fractions were compared using a Bayesian test for linear correlation (Jeffreys, 1998; Ly et al., 2016) implemented in the R package BayesFactor (Morey et al., 2018) to be aware of and avoid the effects of collinearity on the models calculated. Furthermore, the distributions of differences between $v_f$ of different DOM components and nutrients were calculated using a Monte Carlo simulation. In this way, we gained a probability distribution of differences that can be used to measure the evidence favouring a difference (motivated by the posterior distribution of the difference in means; Kruschke, 2013).

3 Results

3.1 PARAFAC components

We could successfully fit a six-component PARAFAC model (Fig. 3, Table 2). Leachates of pig and cow dung exhibited high levels of tryptophan-like (Trp, C5) and tyrosine-like (Tyr, C6) compounds. Leaf leachate showed high peaks of microbiologically produced humic-like (Hum-mic, C1) fluorescence, which is assumed to represent low-molecular-weight, aliphatic DOM originating from microbial degradation. Ambient water was characterized by humic-like material from terrestrial sources (Hum-ter, C2) and microbiologically processed terrestrial DOM associated with agriculture (Hum-micer, C3). Another humic-like fluorophore, which showed a similar fluorescence to pure quinone, was identified in all sources (Qui, C4). The ambient DOM composition resembled the leachate of pig dung.

3.2 Ambient concentrations and introduced material

Peak DOC concentrations were highest for cow dung leachate, followed by corn and leaves, and lowest in nettles and pig dung (Fig. 4). Leachates of cow dung, pig dung, and leaves showed the highest concentrations of SRP. The overall background concentrations of N-NO$_3$ were highly fluctuating, high in concentration, and hardly influenced by leachate additions. Most components declined during downstream travel, while Hum-ter (C2) and Hum-micer (C3) increased during corn and leaf additions. Concentrations and fluorescence tended to return to ambient conditions while travelling downstream. The BFs for an exponential decay during downstream travel compared to a conservative behaviour (no concentration change along the stream course) were 3.3 for DOC and larger than 10$^7$ for all DOM components and SRP. Thus, the evidence for an exponential decay curve was strong to decisive for these components. N-NO$_3$ only exhibited a BF of 0.03, which means that there is strong evidence that the concentration was constant along the stream course during each sampling. The correlations of DOC, N-NO$_3$, SRP concentrations, and the fluorescence-based concentrations of the DOM fractions can be found in Table 3.

3.3 Uptake velocities of bulk DOC and DOM components

Probability densities of DOC uptake velocities from corn leachate, leaf leachate, and cow dung leachate were narrow (Fig. 5), allowing for a clear distinction of $v_f$ between these three leachates (Table 4). Here, corn leachate was taken up fastest, followed by leaf and cow dung leachate. During nettle and pig dung leachate additions, the DOC peaks were lower (Fig. 4) and measurement errors had a stronger relative influence, leading to broader posterior probability densities (Fig. 5). This hampered a clear separation of the $v_f$ of nettle and pig dung leachates from the other leachates’ $v_f$ probabilities. Nevertheless, we can assume that the uptake velocities of nettles and pig dung leachates did not exceed 6 mm min$^{-1}$ and that they were faster than the $v_f$ of cow dung leachate.

Modelling $v_f$ of the different components and nutrients without considering interactions or influencing factors showed that bulk DOC reflected the average uptake of the different DOM components. N-NO$_3$ and Hum-mic (C1) were taken up slower, whereas SRP, Hum-micer (C3), and Trp (C5) were taken up faster than the bulk DOC and all other components (Fig. 6, Table 5).

3.4 Influencing factors and component interactions

Differences between samplings using the same source can be caused by day-dependent characteristics, such as discharge and weather, or by differences in the leachate composition due to, e.g., degradation during storage. Hence, we tested whether the date of the different experiments and/or the
**Table 2.** PARAFAC components and their comparison to other studies. The abbreviations and symbols used stand for the following: a: ambient; m: corn; c: cow dung; l: leaves; n: nettles; p: pig dung; ▲: high; ─: intermediate; and ▼: low.

<table>
<thead>
<tr>
<th>Component</th>
<th>Similar components in other studies</th>
<th>Interpretation</th>
<th>Relative share in leachates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hum-mic (C1)</td>
<td>G2 (Murphy et al., 2011), C2 (Lambert et al., 2016b), D2 (Shutova et al., 2014)</td>
<td>Microbial humic-like, DOM produced during the microbial degradation of terrestrial DOM within freshwaters</td>
<td>▼ ▼ ▲ ▼</td>
</tr>
<tr>
<td>Hum-ter (C2)</td>
<td>C2 (Lambert et al., 2016a), F3 (Heibati et al., 2017)</td>
<td>Terrestrial humic-like, high-molecular-weight and aromatic compounds of terrestrial origin</td>
<td>▲ ▼ ▲ ▼ ▼ ▲</td>
</tr>
<tr>
<td>Hum-micter (C3)</td>
<td>C5 (Lambert et al., 2017), C4 (Williams et al., 2010), C5 (Williams et al., 2013)</td>
<td>Microbial humic-like, positively correlated with bacterial activity and crop-lands in the catchment, associated with microbial transformation of terrestrial organic matter</td>
<td>▲ ▼ ▼ ▼ ▼ ▲</td>
</tr>
<tr>
<td>Qui (C4)</td>
<td>C2 (Yamashita et al., 2011), C2 (Garcia et al., 2015)</td>
<td>Humic-like, A and C peaks, terrestrial origin, with an aromatic chemical nature, may be derived from old soil organic matter, some similarity to pure quinone</td>
<td>─ ▲ ▲ ─ ─</td>
</tr>
<tr>
<td>Trp (C5)</td>
<td>C7 (Stedmon and Markager, 2005), C6 (Murphy et al., 2011)</td>
<td>Tryptophan-like fluorescence, peak almost identical to free tryptophan, derived from autochthonous processes, correlated to terrestrial fluorescent material in forested catchments</td>
<td>─ ─ ▲ ─ ─</td>
</tr>
<tr>
<td>Tyr (C6)</td>
<td>G7 (Murphy et al., 2011), C3 (Yamashita et al., 2013), J3 (Wünsch et al., 2015)</td>
<td>Tyrosine-like, suggested as degradation products of peptides/proteins</td>
<td>▼ ─ ▲ ▲ ▲ ─</td>
</tr>
</tbody>
</table>
Figure 4. Measured concentrations of DOC, SRP, and N-NO$_3$, and Raman units (RUs) of DOM fractions along the stream course for the different samplings. The values are corrected for dilution effects (see also Table S1 in the Supplement). Arrows show the general trend of the concentrations/RUs from up- to downstream. The ambient concentrations were interpolated from measurements taken between leachate additions and are visualized as grey ribbons (see Table S1 for ambient conditions and additional amounts from leachate additions at the upstream station).

Table 3. Linear correlation of nutrient concentrations and DOM fraction fluorescence; BFs in brackets; only shown if BF > 1.

<table>
<thead>
<tr>
<th></th>
<th>Hum-mic (C1)</th>
<th>Hum-ter (C2)</th>
<th>Hum-micter (C3)</th>
<th>Qui (C4)</th>
<th>Trp (C5)</th>
<th>Tyr (C6)</th>
<th>DOC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hum-micter (C3)</td>
<td>0.87 (5.47)</td>
<td>0.62 (3.24)</td>
<td>0.59 (1.46)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Qui (C4)</td>
<td></td>
<td>0.86 (3.25)</td>
<td>0.73 (2.45)</td>
<td>0.58 (1.03)</td>
<td>0.56 (1.38)</td>
<td>0.47 (1.18)</td>
<td></td>
</tr>
<tr>
<td>Trp (C5)</td>
<td></td>
<td></td>
<td>0.87 (8.22)</td>
<td>0.80 (12.62)</td>
<td>0.91 (8.83)</td>
<td>0.69 (4.74)</td>
<td>0.37 (1.35)</td>
</tr>
<tr>
<td>Tyr (C6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DOC</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4. Comparison of $v_f$ of DOC depending on leachate source by the BF of one $v_f$ being lower than the other one. Additionally, the table presents median values of the $v_f$ distributions in millimetres per minute.

<table>
<thead>
<tr>
<th></th>
<th>Leaves</th>
<th>Nettles</th>
<th>Pig dung</th>
<th>Corn</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cow dung</td>
<td>0.66</td>
<td>981</td>
<td>7.04</td>
<td>&gt; 1000</td>
</tr>
<tr>
<td>Leaves</td>
<td>2.08</td>
<td>1.40</td>
<td>7.33</td>
<td>38.7</td>
</tr>
<tr>
<td>Nettles</td>
<td>2.42</td>
<td>2.24</td>
<td>2.74</td>
<td></td>
</tr>
<tr>
<td>Pig dung</td>
<td>3.37</td>
<td></td>
<td>1.19</td>
<td></td>
</tr>
<tr>
<td>Corn</td>
<td>3.54</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

source significantly affected the uptake of the bulk DOC, the DOM components, and the leached nutrients. Hum-mic (C1) retention was substantially (BF 4.6) and Trp (C5) retention was decisively (BF 134.2) influenced by the addition date. Bulk DOC and Tyr (C6) retentions were influenced by both the DOM source and the date. However, for bulk DOC retention, the source had a stronger effect (BF 1563) than the date (BF 146), while the reverse was true for Tyr (BF 10.7 and 10$^8$ for source and date, respectively). Hum-ter (C2), Hum-micter (C3), and Qui (C4) as well as SRP and N-NO$_3$ showed conservative uptake behaviour independent of the source or the addition date (BF < 1; see also Table S3).

To further disentangle the interaction effects between nutrient and DOM component uptake, as proposed in Eq. (1),
Table 5. Comparison of \( v_f \) of DOM components and nutrients by the BF of one \( v_f \) being lower than the other one. Additionally, the table presents median values of the \( v_f \) distributions in millimetres per minute.

<table>
<thead>
<tr>
<th>Component</th>
<th>( v_f ) median in mm min(^{-1})</th>
<th>BF for ( v_f ) (row) &lt; ( v_f ) (column)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-NO(_3)</td>
<td>0.73</td>
<td>6.66</td>
</tr>
<tr>
<td>Hum-mic (C1)</td>
<td>1.60</td>
<td>133</td>
</tr>
<tr>
<td>Hum-ter (C2)</td>
<td>2.98</td>
<td>38.4</td>
</tr>
<tr>
<td>DOC</td>
<td>3.21</td>
<td></td>
</tr>
<tr>
<td>Qui (C4)</td>
<td>3.52</td>
<td></td>
</tr>
<tr>
<td>Tyr (C6)</td>
<td>3.97</td>
<td></td>
</tr>
<tr>
<td>Hum-micter (C3)</td>
<td>6.66</td>
<td></td>
</tr>
<tr>
<td>SRP</td>
<td>6.66</td>
<td></td>
</tr>
<tr>
<td>Trp (C5)</td>
<td>133</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 5. Posterior density distribution curves of uptake velocity \( v_f \) of DOC depending on the leachate source.

Figure 6. Posterior density distribution curves of uptake velocity \( v_f \) for different compounds and nutrients.

We also analysed correlations between the uptake velocities of different DOM components to check for concurrent retention, indicating interrelations among or dependencies of different microbial processes, such as the combined need of these substances in the microbial metabolism (Table S2). We found a positive correlation between the \( v_f \) of Qui (C4) and the \( v_f \) of Tyr (C6) and DOC.

3.5 Brief propagation of uncertainty

The simulated probability density of the residuals (Fig. S1) was compared to the instruments’ expected accuracy for DOC and SRP. The models depended on three measured values (\( C_{x,t} \), \( C_{amb,t} \), \( C_{x-1,t} \)). So we multiplied the instrument errors by 3 to get the effect of their uncertainty on the model results. For DOC measurements with an accuracy of 2 %, three of our concentrations at around 2000 µg L\(^{-1}\) would have an approximate effect of 120 µg L\(^{-1}\) on the model uncertainty. The 95 % probability interval of the model resid-
Table 6. Interactions between uptake velocity and concentrations of other nutrients or DOM components using the INSBIRE approach. $v_f$: uptake velocity; $k$: uptake rate factor; $w$: wetted width; $C_i$: fluorescence of PARAFAC components; $m_i$: exponent of relation; $l$: additive parameter.

<table>
<thead>
<tr>
<th>Fraction/nutrient</th>
<th>Most probable model (Eq. 1)</th>
<th>Bayes $R^2$</th>
<th>BF vs. $v_f$</th>
<th>Estimates of parameter values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hum-mic (C1)</td>
<td>$v_f = kw^C1^{mc1}$</td>
<td>0.60</td>
<td>16.74</td>
<td>$k = 2.11$ $mc1 = -0.38$</td>
</tr>
<tr>
<td>Hum-ter (C2)</td>
<td>$v_f = k(l + DOC^{mc})$</td>
<td>0.34</td>
<td>7.69</td>
<td>$K = 0.11$ $l = 3.16$ $mc = 0.32$</td>
</tr>
<tr>
<td>Hum-micter (C3)</td>
<td>$v_f = v_f$</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Qui (C4)</td>
<td>$v_f = kC1^{mc1}C4^{mc4}$</td>
<td>0.44</td>
<td>3.13</td>
<td>$K = 0.71$ $mc1 = -0.25$ $mc4 = -0.35$</td>
</tr>
<tr>
<td>Trp (C5)</td>
<td>$v_f = kC2^{mc2}C5^{mc5}$</td>
<td>0.30</td>
<td>3.87</td>
<td>$k = 0.85$ $mc2 = -0.44$ $mc5 = -0.55$</td>
</tr>
<tr>
<td>Tyr (C6)</td>
<td>$v_f = kC2^{mc2}C6^{mc6}$</td>
<td>0.45</td>
<td>$1.51 \times 10^7$</td>
<td>$k = 0.27$ $mc2 = -0.23$ $mc6 = -0.96$</td>
</tr>
<tr>
<td>DOC</td>
<td>$v_f = kC6^{mc6}$</td>
<td>0.28</td>
<td>10.50</td>
<td>$k = 0.30$ $mc6 = -0.62$</td>
</tr>
<tr>
<td>NO$_3$</td>
<td>$v_f = v_f$</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>SRP</td>
<td>$v_f = kw^{SRP^{mp}}$</td>
<td>0.63</td>
<td>$1.45 \times 10^4$</td>
<td>$k = 26.18$ $mp = -0.31$</td>
</tr>
</tbody>
</table>

Discussion

4.1 Uptake of bulk DOC from different sources

The uptake velocity of bulk DOC varied between leachate sources (Fig. 5), as observed in previous studies (e.g. Bernardt and McDowell, 2008; Mineau et al., 2016; Mutschlechner et al., 2018). However, in contrast to our original hypothesis, neither a high-SRP content nor an increased amount of low-molecular-weight protein-like compounds could be linked to higher $v_f$ of bulk DOC (Table 2). Cow and pig dung leachates, for example, both had high peaks of the tryptophan-like component C5, which showed the fastest uptake of all DOM fractions, indicating a high bioavailability. Nevertheless, cow dung leachate was taken up most slowly, while pig dung leachate was among the leachates with the fastest uptake. Corn leachate showed the highest uptake of all sources, while the uptake velocity of leaf leachate was intermediate. Interestingly, the same sequence of increasing uptake velocities from cow dung leachate to leaf and corn leachate was observed in a laboratory flume experiment using the same organic matter sources as this field study but different sediments (Weigelhofer et al., 2020). However, in that study, DOC uptake was positively influenced by the
SRP concentrations in the leachates. Comparisons with the literature are difficult, as other field and laboratory studies have used various organic matter sources. Among those, leaf leachates have been used most frequently so far. Leaf leachates show a wide range of biodegradability, depending on the species, the region, the pre-treatment, and the decomposition or leaching stage (e.g. Wickland et al., 2007). Long-term fertilization of trees, for example, resulted in an increased DOC uptake of leaf leachates due to their increased P content (Mutschlecner et al., 2018). Reported uptake velocities for leaf leachates range from 0.002 to 7.8 mm min\(^{-1}\), showing a high variability among leaf sources and aquatic systems (review by Mineau et al., 2016; Graeber et al., 2019). The median \(v_f\) of our leaf leachate was 2.08 mm min\(^{-1}\).

**Figure 7.** Simulated change of uptake velocity \(v_f\) with variation of one variable using the fitted models from Table 6. The colours show the 50 % (violet) and the 90 % (yellow) probability intervals.
and thus lies within this range. Regarding the other sources used in our study, we only found one addition study using cow manure (Kuserk et al., 1984). There, the median uptake velocity was slightly lower (0.31 mm min$^{-1}$; calculated in Mineau et al., 2016) than the one observed in our study (0.66 mm min$^{-1}$).

4.2 Uptake of DOM fractions and nutrients

The various DOM fluorophores were retained with different uptake velocities, whereby the velocity density curves partly overlapped (Fig. 6, Table 5). Hum-mic (C1), described as a product of microbial degradation of terrestrial organic matter, showed the lowest $v_f$ of all components. Hum-ter (C2; high molecular weight, aromatic), Qui (C4; aromatic), and Tyr (C6; tyrosine-like) showed large overlaps and exhibited uptake velocities comparable to the bulk DOC, followed by slightly higher uptake velocities for Hum-micter (C3). As expected, the fastest uptake was observed for the tryptophan-like component Trp (C5), concordant with previous studies of different amino-acid-like fractions (Findlay and Sinsabaugh, 2003). Several studies report a high biodegradability of protein-like components, while humic-like, aromatic components have proved to be much more refractory (e.g. Fellman et al., 2009a, b; Casas-Ruiz et al., 2017). However, in contrast to Trp (C5), the tyrosine-like Tyr (C6) showed only medium uptake velocities. This may have been caused by the release of Tyr (C6) as a degradation product of humic substances during the experiment (Stevenson and He, 1990; Tsutsuki and Kuwatsuka, 1979). Other studies also report the generation of protein-like components during passage through the system due to either the release of algal exudates or the decomposition of humic substances (Casas-Ruiz et al., 2017; Weigelhofer et al., 2020).

The uptake of N-NO$_3$ was the lowest of all components due to its high background concentrations in the water column exceeding even those of the ambient DOC (Fig. 4). In contrast, the co-leached SRP showed the highest uptake velocity (together with Trp, C5). An equally fast uptake was observed in flume experiments, especially in the presence of algae (Weigelhofer et al., 2020). Despite the low-to-moderate background concentrations of P in the stream water (Fig. 4), background molar C : P ratios in the water column of our study stream were usually below 80 : 1, displaying an ideal ratio for a vast number of bacterial strains (Cross et al., 2005: Godwin and Cotner, 2018). The C : P ratios were even decreased by the additions to < 30 : 1 at point 1, followed by an increase to background ratios in the downstream sections. While stoichiometry is a key factor for C, N, and P uptake (e.g. Cross et al., 2005; Gibson and O’Reilly, 2012; Stutter et al., 2020), we thus do not believe that stoichiometric control played a significant role in the P uptake in our study stream. Instead, we assume that the co-leached P was taken up faster than the DOC due to the P demand of both bacteria and algae (in contrast to the exclusive OC demand of heterotrophs; see, e.g., Oviedo-Vargas et al., 2013; Weigelhofer et al., 2020).

In general, the bioavailability of a fraction does not only depend on the chemical composition but also on the ecosystem and the involved microbial community (Kamjunke et al., 2015), the overall availability of different fractions and nutrients (Berggren and Giorgio, 2015; Bernhardt and McDowell, 2008; Mutschlecner et al., 2018), and transport characteristics (Ejarque et al., 2017). We performed the experiments in a small homogeneous stretch of a stream and already found considerable variability in DOM fluorophore-specific uptake between sampling dates. Thus, we propose that the bioavailability of DOM fractions and sources should be increasingly determined in situ under different environmental conditions to determine the effective biodegradability range of the respective component.

4.3 Relationships between uptake and concentrations of other compounds

In contrast to our assumptions, we found no influence of the co-leached SRP on the bulk DOC uptake, although there is evidence in other studies that DOC uptake can be stimulated by P, especially in P-limited systems (Mutschlecner et al., 2018; Stutter et al., 2020). However, the molar C : P ratios were low in our stream, showing no P limitation. Furthermore, most P peaks during the additions were relatively small, containing only the leached P from the DOM sources. Thus, potentially stimulating effects of SRP on the DOC retention may have remained undetected. Although the source-independent model showed a relationship between the bulk DOC uptake velocity and the Tyr (C6) fluorescence, the mixed-effects model, including the leachate source, showed a higher probability of explaining the measured values. The better performance of the mixed-effects model indicates that other, probably non-fluorescent components, which we could not detect with our methods, influenced the bulk DOC uptake apart from Tyr. Almost all DOM fractions showed a negative relationship between uptake and concentration of the same component. Decreasing uptake velocities with increasing concentrations have been previously described for nitrogen as nutrient efficiency loss (Dodds et al., 2002; O’Brien et al., 2007). This efficiency loss may occur especially in streams where the microbial community is adapted to chronic loading, showing high flexibility towards increasing nutrient (or DOC) concentrations and thus a delayed saturation (e.g. O’Brien et al., 2007; Mulholland et al., 2009; Merseburger et al., 2011). Furthermore, transport limitations between the source in the water column and the reactive sites in the sediments (e.g. due to eutrophication and sediment clogging) may lead to delays and reductions in the microbial response to short nutrient peaks (Weigelhofer et al., 2018a, b; Teissier et al., 2007; Ribot et al., 2013). Hum-mic (C2), Hum-micter (C3), and DOC retention showed no evidence of efficiency loss (BF was around 1) at the measured concentrations, indicating that the microbial community would have been able
to retain more of these substances without a decline in uptake velocity. So far, we have not found any other studies presenting efficiency loss or any other uptake–concentration relationship for DOM fractions.

Additionally, we observed both positive and negative interactions among different DOM fractions, with higher/lower uptake velocities of one component at increased concentrations of another component. These interactions can have various reasons and are, therefore, difficult to interpret. Positive effects on $v_f$ can arise, e.g., from the stimulation of the uptake of one substance by the presence of another through priming (but see critical discussion in Bengtsson et al., 2018). Direct negative effects can be caused by the preferential uptake of one fraction over another (Brailsford et al., 2019) or inhibitory effects between different substances (Freeman et al., 1990). Furthermore, the degradation of DOM can cause one molecule to break down into others, causing an increase of the degradation product, while the degraded component decreases (Kamjunke et al., 2017). In our study, the retention of Qui (C4) was lower at higher Hum-mic (C1) fluorescence. The molecular structures found in the literature (Stevenson and He, 1990; Tsutsuki and Kuwatsuka, 1979) suggest that Qui (C4) is a product of Hum-mic (C1) degradation, resulting in reduced net retention due to simultaneous production and degradation processes. Similarly, Trp (C5) and Tyr (C6) might have been degradation products of Hum-ter (C2). In contrast, Hum-ter (C2) degradation was stimulated by high DOC concentrations, probably due to the supply of energy in the form of carbohydrates or other essential components needed for degradation (Bengtsson et al., 2018; Bianchi et al., 2015). We also saw a weak probability that the uptake velocity of Hum-ter (C2) was stimulated by Qui (C4; BF = 1.9) and Tyr (C6; BF = 1.8). Based on the correlation of the uptake velocities, we found substantial evidence that Qui (C4) was degraded simultaneously with Tyr (C6; BF = 6.10) and bulk DOC (BF = 4.63). Good degradation conditions – such as low transport limitations (Weigelhofer et al., 2018b), ideal stoichiometric C:N:P ratios of the organic source for the microbial metabolism (Cross et al., 2005; Godwin and Cotner, 2018; Stutter et al., 2018), or stretch-wise more productive microbial communities – can foster simultaneous turnover (Guillemette and Giorgio, 2012). We consider concurrent degradation and negative and positive interactions to be essential characteristics of the complex DOM degradation processes. Using the INSBIRE approach in future experiments may help to elucidate which of the proposed mechanisms is responsible under certain environmental conditions.

Our models also revealed some hydromorphological effects on DOM fluorophore and nutrient uptake. The wetted width could partly explain the uptake of Hum-mic (C1) and SRP, probably due to adsorption to sediments or extracellular polymeric substances playing a role in retaining these compounds. The adsorption of humic substances to clay is generally strong when the ionic strength is high (Theng, 2012). The conductivity around 630 µS cm$^{-1}$, which was measured during the experiment, and the clay-dominated sediments offered good conditions for adsorption (Theng, 2012). The role of the sediment surface in the uptake of solutes is not surprising and has been observed elsewhere (Romani et al., 2004; Sabater et al., 2002; Battin et al., 2016). However, the component-specific influence of wetted width suggests that different DOM components are preferably taken up in different stream compartments. Unlike the common assumption that uptake processes are dominated by the benthic community (Battin et al., 2016; Wiegner et al., 2005), Graeber et al. (2018) and Kamjunke et al. (2015) proposed a potentially important impact of planktonic bacteria on in-stream DOM uptake processes. In our study stream, such planktonic uptake might have controlled the uptake of most DOM fractions except Hum-ter (C1), where the substantial influence of wetted width indicates the importance of the benthic community.

### 4.4 Potential and limitations of the INSBIRE approach

The INSBIRE approach was only developed after the data from the experiment were acquired. Thus, our study represents a case study for the application of INSBIRE in the analysis of DOM uptake but does not claim to be a systematic check of the developed approach. Nevertheless, we can make some statements about the potential and the limitations of INSBIRE. INSBIRE helped to reveal positive and negative interactions among different DOM fractions previously not possible in such detail. We based our model on concepts, such as nutrient spiralling (Stream Solute Workshop, 1990) and Bayesian statistics, which have been investigated and developed for at least some decades. With this available knowledge, it was possible to develop the approach on a solid theoretical basis and with existing algorithms. However, INSBIRE can be adapted to various research questions by changing the underlying equations, using different solving schemes and different types of data. While we used fluorescence measurements to determine the DOM quality, INSBIRE can incorporate any other data of solvents (e.g. toxins or pesticides) and methods (e.g. mass spectroscopy, liquid chromatography). The power function has proven useful in our study, but the approach also facilitates the use of other equations. Due to the formal description of the uptake processes, extrapolations to ambient or event-related concentrations can be performed (Payn et al., 2005).

The presented plots of the $v_f$ posterior density curves are intuitive to interpret and can support our understanding of the retention processes. The presentation in the form of probability distributions instead of single values reflects the heterogeneity of ecosystems (McCarty, 2007). For further studies, these posterior density curves can be directly used as prior information for similar models. The Bayesian nature of the analysis allows even weak relations to be evaluated, which can be tested in further experiments. We could also show the limitation of the bulk DOC retention model due to
measurement accuracy and the heterogeneity of the different molecules. When only a small number of observations are available, data from previous studies and expert knowledge can be included using non-conservative prior densities of the parameters. In this way, results can be more precise, and decisions can be based on both measured data and available knowledge (Kuhnert et al., 2010; Lemoine, 2019). Even a low number of observations may show specific trends in DOM uptake (Fig. 5), which can be especially useful for monitoring or management decisions. We want to encourage systematic tests under controlled laboratory conditions to assess the full applicability of the INSBIRE approach for modelling the uptake of complex solutes.

5 Conclusion

Human impacts, such as agricultural land use or wastewater discharges, have changed the quantity and composition of terrestrially derived DOM in stream ecosystems. Our study demonstrates that in-stream DOM uptake is source-dependent and thus influenced by DOM quality. However, bulk DOC uptake did not significantly correlate with any of the leached nutrients or individual fluorophores. One reason for this lack of correlation could be the complexity of DOM retention. DOM uptake comprises a variety of simultaneously and sequentially occurring microbial degradation and production processes. Hence, the INSBIRE approach provided evidence for positive and negative effects among the DOM components’ uptake, indicating transformations of one substance into another during processing. Furthermore, the identification of different DOM components via spectroscopic characterization may be too imprecise to reveal influences of DOM components on DOM uptake. Non-fluorescent components, not detected by spectroscopy, may also play a role in bulk DOM uptake. Thus, further DOM uptake studies are required which identify important molecular groups – such as amino acids, sugars, and humic acids – more accurately.

Our study also shows that the uptake of bulk DOC and individual DOM components may be subject to efficiency loss, so far only known from nutrient uptake. This means that the uptake efficiency declines with increasing concentration of the respective component. However, individual DOM components were not equally affected by efficiency loss and interactions. Thus, we assume that the component-specific uptake capacity of benthic biofilms depends on the respective microbial processes involved. Further studies need to look more closely into the underlying mechanisms of efficiency loss and components interactions during DOM processing in aquatic ecosystems. Our study also demonstrates that the cycling of different C fractions and their mutual interaction with N and P uptake in streams is a complex, non-linear problem, requiring advanced non-linear approaches. Here, the developed INSBIRE approach may help in finding concurrent retention and interactions of DOM components, thus providing an efficient tool for the analysis and the management of organic carbon cycling in aquatic systems.


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

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