Spatial and Temporal Variability in Concentration-Discharge Relationships at the Event Scale

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Abstract  The analysis of concentration-discharge (C-Q) relationships from low-frequency observations is commonly used to assess solute sources, mobilization, and reactive transport processes at the catchment scale. High-frequency concentration measurements are increasingly available and offer additional insights into event-scale export dynamics. However, only few studies have integrated inter-annual and event-scale C-Q relationships. Here, we analyze high-frequency measurements of specific conductance (EC), nitrate (NO$_3^-$-N) concentrations and spectral absorbance at 254 nm (SAC$_{254}$, as a proxy for dissolved organic carbon) over a two year period for four neighboring catchments in Germany ranging from more pristine forested to agriculturally managed settings. We apply an integrated method that adds a hysteresis term to the established power law C-Q model so that concentration intercept, C-Q slope and hysteresis can be characterized simultaneously. We found that inter-event variability in C-Q hysteresis and slope were most pronounced for SAC$_{254}$ in all catchments and for NO$_3^-$-N in forested catchments. SAC$_{254}$ and NO$_3^-$-N event responses in the smallest forested catchment were closely coupled and explainable by antecedent conditions that hint to a common near-stream source. In contrast, the event-scale C-Q patterns of EC in all catchments and of NO$_3^-$-N in the agricultural catchment without buffer zones around streams were less variable and similar to the inter-annual C-Q relationship indicating a homogeneity of mobilization processes over time. Event-scale C-Q analysis thus added key insights into catchment functioning whenever the inter-annual C-Q relationship contrasted with event-scale responses. Analyzing long-term and event-scale behavior in one coherent framework helps to disentangle these scattered C-Q patterns.

1. Introduction

Anthropogenic activities such as intensive agriculture and waste water discharge deteriorate the quality of water resources. More specifically, anthropogenic sources increase the aquatic concentration and fluxes of nutrients like nitrogen, phosphorus and organic carbon, leading to eutrophication in rivers, lakes and coastal waters (Carpenter et al., 2011; Schlesinger, 2009). This can pose a threat to water security and downstream aquatic ecosystem health and functioning (Foley et al., 2005). For effective catchment-scale water quality management, knowledge on sources, pathways and reaction processes of critical constituents is needed. Reactive transport at the catchment scale, is however, complex and tends to span a large range of spatial and temporal scales (Gall et al., 2013; Kirchner, 2003; Sivapalan, 2006). This often hinders establishing a unique cause-effect relationship.

One way of approaching this complexity involves the analysis of the integrated response of concentration (C) of a constituent and discharge (Q) at a given point in the stream to identify underlying processes and their hierarchy (Basu et al., 2011; Godsey et al., 2009; Musolff et al., 2015). More specifically, characterizing the relationship between concentration and discharge proved valuable to link temporal patterns in the data to dominant processes at the catchment scale (Sivapalan, 2006). This link may however not always be fully
identifiable as several processes can lead to similar observable patterns (Musolff et al., 2017a). The processes that shape C-Q relationships are a consequence of solute availability and hydrological connectivity in source zones (Dupas et al., 2015; Werner et al., 2019; Winterdahl et al., 2011), source zone distribution in the catchment (Ameli et al., 2017; Dupas et al., 2019; Musolff et al., 2017a; Seibert et al., 2009; Zhi et al., 2019), reactions along subsurface flow paths (Musolff et al., 2017a; van der Velde et al., 2010; Yang et al., 2018) and in the stream network (Koenig et al., 2017; Moatar et al., 2017; Wollheim et al., 2017). C-Q relationships have been used to identify processes at different temporal scales: long-term average behavior (Basu et al., 2010; Musolff et al., 2015; Zarnetske et al., 2018), long-term changes of sources (Ehrhardt et al., 2019; Moatar et al., 2017; Westphal et al., 2020), seasonal changes (Koenig et al., 2017; Minaudo et al., 2019; Winterdahl et al., 2011) and changes at the runoff event time scale (Fovet et al., 2018; Knapp et al., 2020; Strohmeier et al., 2013; Vaughan et al., 2017; Werner et al., 2019). Runoff events are of special interest for catchment-scale assessments as they are critical moments of mobilization and export (Knapp et al., 2020; Werner et al., 2019) and often dominate the total load (Birkel et al., 2017).

C-Q relationships are most often described by power law models in the form $C = aQ^b$ with $a$ defining the concentration level and exponent $b$ defining the direction and strength of concentration changes with Q. For the latter, we distinguish dilution ($b < 0$) from enrichment ($b > 0$) and neutral ($b \sim 0$) C-Q patterns (Godsey et al., 2009; Musolff et al., 2015). However, at the scale of runoff events, hysteresis between concentration and discharge is often observed (Bieroza & Heathwaite, 2015; Evans & Davies, 1998) that cannot be captured by the power law formulation. Hysteresis results from systematic differences of concentration for the same discharge depending on the position on the event hydrograph (rising or falling limb). This behavior is explained by the mixing of different runoff sources with a different timing (e.g., base flow, interflow and event flow (Evans & Davies, 1998)) or an early or late activation of sources during the event. For dissolved constituents this results from the spatial arrangement and connectivity of the sources (Strohmeier et al., 2013), biogeochemical reactions in the source zone during the course of an event (Bao et al., 2017) or the exhaustion of a source during the event (Duncan et al., 2017; Sebestyen et al., 2008). By that, hysteresis and event-scale C-Q responses are shaped by land use and anthropogenic impacts as well as by the interplaying hydroclimatic-conditions (Burns et al., 2019; Butturini et al., 2008; Knapp et al., 2020; Vaughan et al., 2017). Hysteretic C-Q relationships are classified in terms of their direction into clockwise (CW), counterclockwise (CCW) and complex (e.g., eight-shaped) patterns (Burns et al., 2019; Evans & Davies, 1998). This type of analyses needs high frequency measurements of concentration and discharge and allow insights into the availability and the release of solutes from catchments (Kirchner et al., 2004). Here, deriving metrics to characterize single C-Q events and classifying the response such as hysteresis and flushing indices are a common approaches for this type of analysis (Lloyd et al., 2016; Vaughan et al., 2017).

High-frequency water chemistry data measured in-situ have been previously the bottleneck of extensive event-scale analyses (Kirchner et al., 2004) but now have become more available (Rode, Wade, et al., 2016; von Freyberg et al., 2017). Today, a growing number of studies (Burns et al., 2019) utilize high-frequency concentration measurements to investigate event-scale solute mobilization under different degrees of human impacts, unveiling source availability and limitation, source location and source connectivity, and reactive transport processes. These processes, however, can also be approached by the C-Q analysis of conventional low-frequency monitoring at inter-annual scale (Godsey et al., 2009; Moatar et al., 2017; Musolff et al., 2015; Thompson et al., 2011). But how do both types of analyses align and in which cases may event-analysis provide a deeper insight into reactive solute transport at catchment scale compared to inter-annual C-Q analysis? This concurs with the question of how event-scale, often hysteretic C-Q relationships aggregate to form an inter-annual C-Q relationship. There is evidence for diverging C-Q behavior comparing different time scales. For instance, intense agricultural management was shown to homogenize water flow (Boland-Brien et al., 2014) and nitrate export behavior reflected in well identifiable inter-annual C-Q relationships (Marinos et al., 2020; Musolff et al., 2017a; Thompson et al., 2011) while event-scale C-Q was found to be temporally highly variable in some cases (Lloyd et al., 2016; Vaughan et al., 2017). In case of a forested catchment Knapp et al. (2020) showed that positive C-Q relationships for nitrate at the event scale can stack up to an inter-annual dilution pattern. Similarly, DOC responses at the event scale were found to be highly variable as a result of interacting biogeochemical and hydrological processes (Blaen et al., 2017; Fovet et al., 2018; Koenig et al., 2017; Werner et al., 2019) but revealed a consistent positive inter-annual C-Q pattern. In contrast, geogenic solutes and specific conductance showed highly consistent patterns at
event-scale and inter-annual scale (Knapp et al., 2020). Although many studies used event-scale C–Q analysis and inter-annual C–Q relationships, so far only few studies integrate both types of C–Q relationships into a coherent framework (Dupas et al., 2016; Knapp et al., 2020; Minaudo et al., 2017).

Hysteresis has been characterized and quantified by comparing the concentrations on the rising and the falling limb of the event hydrograph based on a hysteresis index (Lloyd et al., 2016; Vaughan et al., 2017). As this method relies on comparing similar discharge on both sides of the hydrograph, partial events where discharge is not falling back to initial levels are difficult to analyze. Although the value in the combined information on C–Q pattern (direction) and hysteresis have been acknowledged (Dupas et al., 2016; Vaughan et al., 2017), metrics of hysteresis and patterns were mostly derived independently from each other. However, approaches to integrate hysteresis into power-law C–Q analysis at the event scale exist (House & Warwick, 1998; Krueger et al., 2009; Minaudo et al., 2017; Winter et al., 2020) but have, to the best of our knowledge, not yet been fully exploited. Advancing event-scale C–Q analysis to a mathematical framework comparable to the established inter-annual power-law approach provides an opportunity for unifying C–Q relationships across temporal scales.

In this study we hypothesize that (1) land use and its spatial arrangement in catchments determines solute-specific source location and thus lead to systematic differences in C–Q relationships. We further hypothesize that (2) antecedent hydro-climatic conditions determine variability of concentration responses to discharge at the event-scale and lead to event C–Q relationships that contrast the inter-annual behavior for biogeochemically reactive solutes, but not for geogenic solutes. To this end we analyze inter-annual and event-scale C–Q relationships in measured high-frequency in-situ time series of three water quality parameters in four neighboring catchments along a gradient of hydroclimate and anthropogenic impact. We focus on specific conductance representing geogenic solutes, and the biogeochemically reactive solutes dissolved organic carbon being of biogenic origin and nitrate being also of anthropogenic origin. To do so we utilize a coherent mathematical framework that extends the established power law C–Q analysis by integrating a hysteresis term at the event scale. We classify and compare the results across catchments, constituents, and seasons. By testing the strength of the relationships between event metrics and antecedent conditions we discuss sources and mobilization mechanisms under contrasting land use settings. Finally, we discuss the strengths and potential weaknesses of event-scale C–Q analysis compared to the established analysis at the inter-annual scale.

2. Materials and Methods

2.1. Adding Hysteresis to C–Q Analysis

The relationship of concentration $C$ of a given solute or particulate in the stream to discharge $Q$ is commonly described by a power law:

$$C(t) = aQ(t)^b$$  \hspace{1cm} (1)

Here, parameter $a$ is related to the concentration level (intercept in log-log space, concentration when $Q = 1$) while exponent $b$ (slope in the log-log space) describes the strength and direction of the response of $C$ to $Q$ (Godsey et al., 2009; Jawitz & Mitchell, 2011): a positive $b$ indicates an enrichment pattern while a negative $b$ indicates a dilution pattern.

Several authors suggest to add a hysteresis term to this form of the C–Q relationship to be able to capture concentration deviations on the rising and falling limb (House & Warwick, 1998; Krueger et al., 2009; Minaudo et al., 2017; Winter et al., 2020). They propose the use of the derivative of discharge $dQ(t)/dt$ that allows to distinguish the rising and falling limbs of the hydrograph and thus creates a hysteresis loop around the log-log linear C–Q relationship:

$$C(t) = aQ(t)^b + c\frac{dQ(t)}{dt}$$  \hspace{1cm} (2)

with parameter $c$ describing the size and the direction of the hysteresis by its absolute value and its sign respectively: A positive $c$ indicates a clockwise hysteresis while a negative $c$ indicates a counterclockwise hysteresis. Figure 1 gives an overview over the meaning of the parameters in Equation 2. This equation can be applied on storm event time scales as well as longer timescales. However, as hysteresis patterns can change
between events (Fovet et al., 2018; Lloyd et al., 2016; Vaughan et al., 2017), applications of Equation 2 at the event scale are most appropriate.

We note that parameters $a$ and $c$ scale with discharge and concentration (see also Jawitz and Mitchell (2011)), while parameter $b$ is independent of both and thus easily transferable between catchments and solutes. That means that the event magnitude largely influences the variability of hysteresis parameter $c$ between events. More specifically, an event with higher discharge but similar concentration would lead to a lower absolute value of $c$. However, with $c$ we aim to describe the extent of the hysteresis and how it deviates from the log-log linear C-Q relationship. To allow for a better comparability of C-Q metrics between events, others have suggested to scale concentration and discharge prior to analysis (Lloyd et al., 2016; Vaughan et al., 2017). This affects both parameters $a$ and $c$. Here we scaled $dQ(t)/dt$ by its own range such that parameter $c$ is independent of the size of the specific discharge event, but we did not scale $Q(t)$ to the average event discharge or to the average inter-annual discharge. In this way, $c$ describes the loop direction and relative width of hysteresis independent of the magnitude of the discharge event. Parameter $a$ in Equation 2 remains equivalent to parameter $a$ in the simple power law C-Q analysis (Equation 1) and captures the vertical offset of individual events as shown in Figure 1a. However, depending on the purpose of the analysis, options to scale or normalize concentration and discharge should be carefully selected prior to analysis.

### 2.2. Study Area

We chose four sub-catchments of the Bode river catchment in central Germany to test our approach (Figure 2). The Bode catchment is an experimental Observatory for terrestrial and hydrological research within the TERENO initiative (Wollschlager et al., 2017). The chosen sub-catchments have been used in several studies on solute export regimes (Dupas et al., 2017; Musolff et al., 2015), instream processes (Rode, Halbedel, et al., 2016), nitrogen retention in reservoirs (Kong et al., 2019), DOC concentration and DOC quality dynamics (Raeke et al., 2017; Werner et al., 2019) and emerging effects of sub-catchment mixing (Winter et al., 2020). The catchments span a gradient in elevation which is linked to gradients in precipitation and land use (Table 1). While the more pristine and mainly forested catchments Warme Bode (FOR1) and upper Rappbode (FOR2) are higher in elevation and generally wetter, the catchments Hassel (AGR1) and upper Selke (AGR2) are dryer and characterized by a substantial share of arable land and grassland (Table 1). The AGR1 catchment drains into a drinking water reservoir and therefore fertilizer application in buffer zones around the streams is prohibited. All catchments are situated in the Harz Mountains and mainly underlain by metamorphic and partially igneous rocks. Note that FOR2 is a small first order catchment (2.3 km$^2$), while stream orders and sizes for FOR1, AGR1 and AGR2 range from 2 to 3 and 43–97 km$^2$, respectively.

### 2.3. Data Base of Concentrations and Discharge

This study mainly relies on previously published concentration and discharge data (Kong et al., 2019; Rode, Halbedel, et al., 2016; Werner et al., 2019; Winter et al., 2020). Water quality data and discharge were
measured every 15 min. We focused on the period between January 2013 and December 2014 where data was most consistently available for all four sites. The water quality parameters we considered in this study were nitrate-N (NO$_3$-N), spectral absorbance at a wavelength of 254 nm (SAC$_{254}$) and specific conductance (EC). Both, NO$_3$-N and SAC$_{254}$ were measured using in-situ UV-VIS probes (TRIOS ProPS, Trios Germany in FOR1, AGR1 and AGR2; s::can spectrolyser, scan Austria in FOR2). We used SAC$_{254}$ as a proxy for the part of DOC that absorbs light at 254 nm instead of estimated DOC concentrations to avoid a possible bias induced by differing internal routines of the probes to derive DOC concentration. The use of SAC$_{254}$ as a proxy for DOC is supported by the strong relationship between SAC$_{254}$ and lab-measured DOC concentration (see Table S1, $r_{\text{Pearson}}$ between 0.58 and 0.96). EC was measured using in-situ probes (YSI6800, YSI, USA for FOR1, AGR1 and AGR2; CTD Diver, Van Essen, Canada for FOR2). Discharge measurements relied on an established stage-discharge relationship and were provided by state authorities (LHW, 2018) (for FOR1, AGR1 and AGR2) and by Werner et al. (2019) (for FOR2). Monitoring stations were maintained every two weeks, including manual cleaning of the UV-VIS probes and grab sampling for subsequent calibration and validation with NO$_3$-N and DOC concentrations analyzed in the lab. Details on these procedures can be found in Rode, Halbedel, et al. (2016), Kong et al. (2019) and Werner et al. (2019). During winter times probes had to be partially removed for four weeks due to harsh weather conditions and limited site accessibility. Water temperature was recorded with the YSI and CTD probes.

Briefly, data preparation included five steps: drift corrections, outlier detection, gap filling, calibration, and moving averaging. Time series of UV-VIS based parameters tended to drift due to biofouling between the probe maintenances. This was corrected by distributing the offset between mean values one hour before and after the maintenance. Table 1 provides an overview of the catchment size, land use, climate and topography metrics for the four studied catchments.

![Figure 2. Land use and topography for the study catchments. Blue lines show the stream network. Inlet figure displays the location within Germany. Land use is based on Corine Land Cover (CLC, 2016).](image)

<table>
<thead>
<tr>
<th>Catchment</th>
<th>Abbr</th>
<th>Area [km$^2$]</th>
<th>SO</th>
<th>P [mm a$^{-1}$]</th>
<th>Gr</th>
<th>LC$_{ara}$ [%]</th>
<th>LC$_{gras}$ [%]</th>
<th>LC$_{for}$ [%]</th>
<th>Elevation [m asl]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Warme Bode</td>
<td>FOR1</td>
<td>97.15</td>
<td>2</td>
<td>1,184</td>
<td>4.32</td>
<td>2.7</td>
<td>4.9</td>
<td>89.6</td>
<td>585</td>
</tr>
<tr>
<td>upper Rappbode</td>
<td>FOR2</td>
<td>2.33</td>
<td>1</td>
<td>1,061</td>
<td>3.06</td>
<td>0.0</td>
<td>1.2</td>
<td>98.8</td>
<td>581</td>
</tr>
<tr>
<td>Hassel</td>
<td>AGR1</td>
<td>43.40</td>
<td>2</td>
<td>865</td>
<td>2.78</td>
<td>41.9</td>
<td>20.0</td>
<td>35.0</td>
<td>505</td>
</tr>
<tr>
<td>upper Selke</td>
<td>AGR2</td>
<td>94.53</td>
<td>3</td>
<td>698</td>
<td>3.91</td>
<td>25.7</td>
<td>10.3</td>
<td>61.4</td>
<td>456</td>
</tr>
</tbody>
</table>

*Note. SO-stream order. P-mean annual precipitation (1986–2015, Cornes et al. (2018)), Gr-topographic gradient (based on 25 m DEM, EEA (2013)), LC-land cover based on CORINE Land Cover (CLC, 2016): ara-cropland, pas-pastures, for-forest. Elevation refers to the catchment mean based on 25m DEM.*
after cleaning equally among the two weeks maintenance interval assuming an exponential growth. Outliers were detected with a two-step procedure. First, values outside a physically likely range were removed (see Table S2). Second, Grubbs test (Grubbs, 1950) was applied to a moving window of 100 values to detect and remove outliers. Data gaps smaller than two hours were filled using cubic spline interpolation (similar to Vaughan et al. (2017)). The resulting time series were globally calibrated against the lab measured concentration of NO$_3$-N (all stations) and against SAC$_{254}$ (all stations but AGR2). Here, probe values averaged one hour before and one hour after sampling were used. EC was calibrated against field values obtained with a handheld WTW probe (WTW Multi 430, Xylem Analytics Germany) for FOR2, while YSI-probes for FOR1, AGR1 and AGR2 were regularly calibrated in field making later corrections obsolete. Parameters of the linear global calibration can be found in Table S3 and indicate a good fit with $R^2 > 0.59$ (mean $R^2 = 0.83$).

In a final step, noise in the signal of both discharge and water quality was reduced by a moving average between 2.5 and 6 h (see Table S4). The length of the moving average window was adjusted visually to the variability of the time series and in general larger for the slower reacting larger catchments (see also Rode, Halbedel, et al., 2016).

At the AGR2 station no SAC$_{254}$ lab measurements were available to calibrate the high-frequency SAC$_{254}$ time series. To validate this time series we explored the relationship of SAC$_{254}$ to DOC. At station AGR2 the slope of SAC$_{254}$-DOC regression is low (Table S1, 0.08 mg m/L) compared to the other catchments (mean 0.18 mg m/L) indicating a potential offset in probe measurements of SAC$_{254}$. Assuming the SAC$_{254}$-DOC regression slope (which equals 1/SUVA$_{254}$) at AGR2 to be similar to the other stations would result in average SAC$_{254}$ values smaller by a factor of 2 and thus very similar to the neighboring AGR1 catchment. We prefer to leave SAC$_{254}$ at AGR2 uncalibrated noting that C–Q relationships and hysteresis patterns remain unaffected by that.

### 2.4. Data Analysis

All data analysis was performed using the R statistical software environment and is available as a script in an open repository (Musolff, 2020b).

To test our hypotheses we first detected discharge events using the methodology of Dupas et al. (2016). Details on the manually adjusted parameters can be found in Table S4. The parameters aimed to encompass the full range of discharge in the event to capture the C-Q relationship including the hysteresis. The start of an event was defined in a moving window (5–12 h, Table S4) based on a discharge increase of 10%. The event end was defined when discharge decreased less than 0.3%–0.5% within the moving window. Multiple peaks were considered as one event if discharge dropped less than 50% after discharge peak. Only events above a defined threshold were included (Table S4). Number of separated events ranged from 32 (AGR2) to 56 (AGR1) (Table S4).

After discharge-based event separation, the simple power-law C–Q (Equation 1) as well as the C–Q model accounting for hysteresis (Equation 2) were fitted (minimizing the sum of the square of the errors) separately for each event and constituent. To assess if the addition of the hysteresis parameter is justified, the performance of both models was compared using the Akaike information criterion (AIC). AIC indicate a better performance of Equation 2 when the AIC improvement was larger than two units. We classified the event response from Equation 2 plotting parameter $c$ against parameter $b$ as suggested by Vaughan et al. (2017). To specifically address hypothesis (1) we tested differences of all model parameters across the catchment using the Kruskal-Wallis rank sum test for significance ($\alpha = 0.05$). In case of a significant difference we used the Wilcoxon rank sum test for pairwise comparison with Bonferroni correction for the p-values.

To compare the parameter $c$ from Equation 2 to an established method, we derived the hysteresis index $H_I$ (Lloyd et al., 2016; Vaughan et al., 2017), based on the range-normalized discharge ($Q_{\text{norm}}$) and range-normalized concentration ($C_{\text{norm}}$) of a given event. $C_{\text{norm}}$ was linearly interpolated between adjacent measurements at 2% intervals of $Q_{\text{norm}}$ to allow comparison for the $Q_{\text{norm}}$ intervals that exists on both, rising and falling limb:

$$H_I = \text{mean}\left(C_{\text{norm, rising}} - C_{\text{norm, falling}}\right)$$

(S3)
To quantify time lags between concentration and discharge as a potential main control of hysteresis we utilized cross correlation. For the comparison between parameter \( c \), \( H_I \) and the time lag Spearman’s rank correlation (significance level of \( p < 0.01 \)) was used.

Hypothesis (2) was addressed using two approaches. First, to allow for a discussion on the similarities and differences in event-scale C-Q responses of biogeochemically reactive (\( \text{NO}_3^-\text{N} \) and \( \text{SAC}_{254} \)) and geogenic constituents (represented by \( \text{EC} \)) we used Spearman’s rank correlation between model parameters \( a \), \( b \), and \( c \) within each catchments (significance level of \( p < 0.01 \)). Second, to quantify and characterize the role of antecedent conditions for the C-Q response we applied partial least squares regression (PLSR, Wold et al. (2001)), being a robust method that accounts for collinearities among the predicting variables. As predictors we used the 7-, 15- and 30-days mean of the pre-event discharge and water temperature and took event characteristics such as minimum, maximum, cumulative discharge as well as concentration and discharge ranges into account. Similar to Musolff et al. (2015) we used the variable influence on projection (VIP, Wold et al. (2001)) to rank the predictors and the regression coefficient to characterize the direction of influence on the response variable. Only models with \( R^2 > 0.5 \) are considered and only predictors with a \( \text{VIP} > 1 \) are reported (Onderka et al., 2012).

3. Results

3.1. Observed Discharge, Concentrations and Inter-Annual C–Q Relationships

Between 56,000 and 70,000 measurements of discharge, EC, \( \text{NO}_3^-\text{N} \) and \( \text{SAC}_{254} \) (equating 580–730 days) were available for analysis. Here, we report on the basic statistics of the time series (Table 2) and on the inter-annual C-Q relationships from applying Equation 1 to the entire time series.

Mean specific discharge [mm/d] nearly halved from FOR1 to AGR2. This is in line with the decrease in precipitation along the elevation gradient from west to east (Table 1, Figure 2). The variability of discharge was high in all catchments (\( CV_Q > 1 \)) being highest in the small FOR2 catchment (\( CV_Q > 2 \)). Mean EC, in turn, was twice as high in AGR2 compared to FOR1. \( CV_{EC}/CV_Q \) clearly indicated chemostatic conditions for EC (\( CV_{EC}/CV_Q < 0.15 \)) with an overall dilution C–Q pattern (\( b_{\text{glob}} < 0 \)). The strongest dilution pattern coincided

<table>
<thead>
<tr>
<th>Catchment</th>
<th>Constituent</th>
<th>( n )</th>
<th>Mean</th>
<th>Median</th>
<th>CV</th>
<th>Min</th>
<th>Max</th>
<th>( CV_C/CV_Q )</th>
<th>( b_{\text{glob}} )</th>
<th>SE ( b_{\text{glob}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>FOR1</td>
<td>Q [mm/d]</td>
<td>70,080</td>
<td>1.56</td>
<td>0.88</td>
<td>1.28</td>
<td>0.16</td>
<td>27.48</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>EC [µS/cm]</td>
<td>68,680</td>
<td>150.85</td>
<td>148.16</td>
<td>0.12</td>
<td>93.79</td>
<td>310.79</td>
<td>0.10</td>
<td>−0.06</td>
<td>4.1E−04</td>
</tr>
<tr>
<td></td>
<td>( \text{NO}_3^-\text{N} ) [mg/L]</td>
<td>67,447</td>
<td>0.66</td>
<td>0.64</td>
<td>0.33</td>
<td>0.10</td>
<td>2.11</td>
<td>0.26</td>
<td>0.15</td>
<td>1.2E−03</td>
</tr>
<tr>
<td></td>
<td>( \text{SAC}_{254} ) [1/m]</td>
<td>66,348</td>
<td>16.27</td>
<td>14.50</td>
<td>0.55</td>
<td>3.73</td>
<td>88.66</td>
<td>0.43</td>
<td>0.11</td>
<td>2.1E−03</td>
</tr>
<tr>
<td>FOR2</td>
<td>Q [mm/d]</td>
<td>64,660</td>
<td>0.90</td>
<td>0.45</td>
<td>2.19</td>
<td>0.02</td>
<td>71.12</td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>EC [µS/cm]</td>
<td>63,126</td>
<td>191.46</td>
<td>187.78</td>
<td>0.13</td>
<td>126.79</td>
<td>253.61</td>
<td>0.06</td>
<td>−0.09</td>
<td>3.3E−04</td>
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<tr>
<td></td>
<td>( \text{NO}_3^-\text{N} ) [mg/L]</td>
<td>55,852</td>
<td>0.37</td>
<td>0.36</td>
<td>0.35</td>
<td>0.12</td>
<td>1.29</td>
<td>0.16</td>
<td>0.08</td>
<td>1.3E−03</td>
</tr>
<tr>
<td></td>
<td>( \text{SAC}_{254} ) [1/m]</td>
<td>55,895</td>
<td>20.76</td>
<td>18.35</td>
<td>0.61</td>
<td>1.15</td>
<td>73.48</td>
<td>0.28</td>
<td>0.14</td>
<td>2.6E−03</td>
</tr>
<tr>
<td>AGR1</td>
<td>Q [mm/d]</td>
<td>62,511</td>
<td>1.12</td>
<td>0.61</td>
<td>1.39</td>
<td>0.05</td>
<td>16.83</td>
<td></td>
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<td></td>
</tr>
<tr>
<td></td>
<td>EC [µS/cm]</td>
<td>68,002</td>
<td>203.07</td>
<td>193.96</td>
<td>0.20</td>
<td>123.16</td>
<td>366.39</td>
<td>0.15</td>
<td>−0.12</td>
<td>4.1E−04</td>
</tr>
<tr>
<td></td>
<td>( \text{NO}_3^-\text{N} ) [mg/L]</td>
<td>64,498</td>
<td>2.60</td>
<td>2.49</td>
<td>0.51</td>
<td>0.20</td>
<td>6.91</td>
<td>0.37</td>
<td>0.36</td>
<td>1.7E−03</td>
</tr>
<tr>
<td></td>
<td>( \text{SAC}_{254} ) [1/m]</td>
<td>64,378</td>
<td>21.54</td>
<td>20.29</td>
<td>0.26</td>
<td>11.75</td>
<td>79.92</td>
<td>0.19</td>
<td>0.03</td>
<td>9.1E−04</td>
</tr>
<tr>
<td>AGR2</td>
<td>Q [mm/d]</td>
<td>70,080</td>
<td>0.79</td>
<td>0.79</td>
<td>1.26</td>
<td>0.11</td>
<td>12.81</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>EC [µS/cm]</td>
<td>69,872</td>
<td>326.41</td>
<td>321.80</td>
<td>0.19</td>
<td>179.22</td>
<td>484.35</td>
<td>0.15</td>
<td>−0.21</td>
<td>2.8E−04</td>
</tr>
<tr>
<td></td>
<td>( \text{NO}_3^-\text{N} ) [mg/L]</td>
<td>69,430</td>
<td>1.76</td>
<td>1.48</td>
<td>0.65</td>
<td>0.01</td>
<td>4.73</td>
<td>0.51</td>
<td>0.73</td>
<td>1.8E−03</td>
</tr>
<tr>
<td></td>
<td>( \text{SAC}_{254} ) [1/m]</td>
<td>65,284</td>
<td>36.36</td>
<td>36.48</td>
<td>0.17</td>
<td>26.26</td>
<td>62.04</td>
<td>0.14</td>
<td>−0.02</td>
<td>7.9E−04</td>
</tr>
</tbody>
</table>

Note. \( n \)-Number of Observations, CV-Coefficient of Variation, \( b_{\text{glob}} \)-Slope of the Inter-annual (Global) logC-logQ relationship. All \( b_{\text{glob}} \) values are significant (\( p < 0.01 \)). SE \( b_{\text{glob}} \)-standard error of \( b_{\text{glob}} \).
with the highest concentrations in the AGR2 catchment. Among the three constituents standard errors for EC $b_{glob}$ were lowest, indicating only few scatter in the C–Q relationship across all data with decreasing values from FOR1 to AGR2. Note that in the AGR2 catchment, EC was strongly influenced by drainage from an abandoned mine, affecting both the concentration level and the dynamic (Musolf et al., 2015). Concentrations of NO$_3$-N were consistently low in the more pristine FOR1 and FOR2 catchments (mean $\leq 0.66$ mg/L) and high in the more agricultural AGR1 and AGR2 catchments (mean $\geq 1.76$ mg/L). $CV_{c}/CV_{q}$ indicates a higher variability in the AGR1 and AGR2 catchments compared to FOR1 and FOR2, with a chemodynamic regime in AGR2 ($CV_{c}/CV_{q} > 0.5$). The global logC-logQ slope $b$ of NO$_3$-N indicates enrichment patterns ($b_{glob} > 0$) for all catchments with lowest values in FOR2 and highest in AGR2. Standard errors of NO$_3$-N $b_{glob}$ were higher in the forested compared to the agricultural catchments. Mean SAC$_{254}$ increased from FOR1 to AGR2 by a factor of 2 (see chapter 2.4 on issues in data calibration and validation for station AGR2) while $CV_{SAC}/CV_{Q}$ clearly decreased. Slope $b_{glob}$ of SAC$_{254}$ was near zero in AGR1 and AGR2 and indicates enrichment patterns in FOR1 and FOR2 ($b_{glob} > 0.10$). Standard errors of $b_{glob}$ for SAC$_{254}$ were higher as for EC and NO$_3$-N indicating a more scattered C–Q relationship. This was more pronounced for the agricultural than for the forested catchments.

3.2. Storm Event Analysis and Parameter Differences Between Catchments

From the time series of discharge and concentrations events have been separated and event-scale C–Q models have been fitted using Equation 2. In the following separated events are described in terms of number and length. The C–Q model are described in comparison to the simple C–Q models without hysteresis (Equation 1) and in terms of parameter variability between catchments and constituents. Finally, event-scale C–Q model parameters are compared to the inter-annual behavior and to other hysteresis metrics.

The event analyses resulted in a larger number of runoff events in FOR1 ($n = 50$, Table S7), FOR2 ($n = 55$) and AGR1 ($n = 56$) compared to AGR2 ($n = 32$). Events were considerably shorter in the small FOR2 catchments (mean $= 1.8$ d) compared to the three larger catchments (mean $\geq 3.6$ d) covering only 14% of the total time (Table S4). In contrast, in the AGR2 catchment events were longest (mean 6.8 d) and covered 30% of the total time series. The highest number of events with a mean discharge exceeding the 95th percentile of inter-annual discharge was found in FOR2 (25.5% of events) followed by FOR1 (14.0%), AGR2 (9.4%) and AGR1 (8.9%).

The observed continuous time series, global C–Q and selected event models are displayed in Figure 3 and Figures S1–S3. The event-scale C–Q models including the hysteresis parameter (Equation 2) were compared to the models based on the simple power-law model (Equation 1) using the Akaike information criterion (AIC). AIC indicated a better performance (AIC improvement larger than two units) for the models including hysteresis for 538 out of 559 analyzed events (Table S7) thus justifying the addition of a third model parameter. Comparing both models we note that the addition of the hysteresis term to the simple power law formulation left the slope $b$ unaffected (linear regression slope between $b$ from Equations 1 and 2 of 0.99, $R^2 = 0.9$, Figure S4).

The parameters of Equation 2 (see also Figure 1 for interpretation) show distinct differences between constituents, between catchments and over time and allowed for a classification of event responses in terms of $b$ (dilution vs. enrichment) and $c$ (clockwise vs. counterclockwise hysteresis) (overview Figure 4, and in Table S7). The Kruskal-Wallis test confirmed that the distribution of all three model parameters for each constituent significantly differed across the catchments (Table S5). In the following model parameters $a$, $b$ and $c$ are described separately for each constituent focusing on differences between the catchments.

Parameter $a$ for EC significantly differed between all catchments (pairwise tests, Table S6, Figure 4) and increased from FOR1 over FOR2 to AGR1 and AGR2 (Table S7). The variability of $a$ across all events at each station was however low ($CV \leq 0.13$), meaning that concentration levels hardly shifted between events. The majority of events showed dilution patterns for EC with a median $b \leq -0.07$ with significant differences between all catchments except FOR2 and AGR1 (Table S6). The hysteresis parameter $c$ significantly differed between all catchments and indicates a similar proportion of clockwise and counterclockwise events in the catchments FOR1 and FOR2. In contrast, most events showed clockwise hysteresis in AGR1 (84%) and
counterclockwise hysteresis in AGR2 (84%). Compared to NO$_3$-N and SAC$_{254}$, parameters $b$ and $c$ for EC varied little between events (see scatter in Figure 4d compared to 4h and 4i).

Parameter $a$ for NO$_3$-N significantly differed between all catchments (Table S6). In the agricultural catchments AGR1 and AGR2 parameter $a$ was higher and more variable. Catchments FOR1, FOR2 and AGR2 dominantly showed enrichment patterns ($b > 0$) while in AGR1 NO$_3$-N dilution patterns ($b < 0$) prevailed. Here, parameter $b$ did not significantly differ between FOR1 and AGR1 and between FOR2 and AGR2 (Table S6). The hysteresis parameter $c$ indicates that counterclockwise events (>70%) dominated for FOR1, FOR2 and AGR2, while more clockwise hysteresis were observed for AGR1 (60%). Here, the pairwise Wilcoxon test revealed no significant differences between most of the catchments (Table S6).

Parameter $a$ of the SAC$_{254}$ increased from west to east (FOR1 over FOR2, AGR1 to AGR2) with significant differences between all catchments except FOR2 and AGR1 (Table S6). However, note the explanation on potentially too high SAC$_{254}$ values in AGR2 above. The majority of events in all four catchments showed enrichment patterns for SAC$_{254}$ ($b > 0$). However, the size and variability of $b$ consistently decreased from FOR1 to AGR2 (Figure 4j) reflected in significant difference between all catchments except FOR2 and AGR1. Hysteresis parameter $c$ was more variable for SAC$_{254}$ compared to EC and NO$_3$-N (Figures 4d, 4h and 4i). More counterclockwise events were identified for FOR1 and FOR2 (>60%), while a higher number of events with a clockwise hysteresis was found in the AGR1 catchment (>65%). In the AGR2 catchment

Figure 3. Time series analysis of EC, NO$_3$-N, and SAC$_{254}$ in the AGR1 catchment. Left column (a, d, g, j) - time series with separated events highlighted with shaded bars; middle column (b, e, h) - global C-Q relationships and power law model (Equation 1); right column (c, f, and i) - example of a modeled event (Equation 2) with observations as dots and the modeled time series as red line; (k) - probability distribution of discharge; (l) time series of the modeled event. Half of the separated events are also plotted as lines in the global C-Q relationships in the second column. The color gradient in the event C-Q plots shows the time from blue at the event start over red to yellow at the event end. The red star in (j) marks the selected event.
clockwise SAC$_{254}$ hysteresis was detected almost exclusively (>95%). The variability of the parameter $c$ between events decreased considerably from FOR1 to AGR2 and the pairwise Wilcoxon test confirmed significant differences between all catchments except for FOR1 with FOR2 and AGR1 (Table S6).

The inter-annual logC-logQ slope $b_{\text{glob}}$ using the entire time series was found to be very similar to the median of the slope $b$ for the event models for EC and diverging more strongly for NO$_3$-N and SAC$_{254}$ (Table 2, Figure 4). In the case of NO$_3$-N in the AGR1 catchment, the global slope $b_{\text{glob}}$ indicates an overall enrichment pattern (Figure S2e) while the majority of individual events showed dilution patterns (Figure 4f); in the other catchments the sign of $b_{\text{glob}}$ and that of event-scale C-Q slopes were both positive for NO$_3$-N.

The comparison of model parameter $c$ to the established hysteresis index $H_I$ (Lloyd et al., 2016; Vaughan et al., 2017) revealed a good agreement (Figure S5, mean Spearman's rank correlation for all events 0.79). In addition, parameter $c$ and the cross-correlation time lag between concentration and discharge in each event was found to be positively correlated (mean rank correlation for all events was 0.68). This relationship only fails for EC in the AGR2 catchment where mine drainage affected EC (Table S7).
3.3. Similarities of Event C-Q Models Across Constituents

The comparison of event parameters \( a \), \( b \) and \( c \) between constituents within a catchment provides information on similarities and differences in sources and mobilization mechanisms for EC, NO\(_3\)-N and SAC\(_{254}\) exported by the same events. Overall we found that parameters in the small FOR2 catchment were often strongly positively or negatively correlated with each other. This was not the case for the larger catchments FOR1, AGR1, and AGR2 (Figure 5). In FOR2, rank correlation between NO\(_3\)-N and SAC\(_{254}\) was highest for the hysteresis parameter \( c \), followed by parameter \( b \) and lowest for \( a \). In this catchment, the hysteresis parameter \( c \) for EC was also negatively correlated with the \( c \) of NO\(_3\)-N and SAC\(_{254}\) (\( r < -0.55 \)). In the managed catchments AGR1 and AGR2, parameters \( a \) for NO\(_3\)-N and for SAC\(_{254}\) were found to be negatively related to each other (\( r < -0.60 \)).

3.4. C-Q Model Parameter Variability Over Time

The event analysis revealed a variability of model parameters over time in each of the catchments (Figure 4). To explore the role of event- and pre-event conditions on the model parameters of individual events we systematically checked the explorative power of ambient (WT0 - mean event water temperature) and antecedent mean water temperature (WT7, WT15, WT30 - mean water temperature 7, 15, 30 days before the event start) and of ambient (Q0 - mean event discharge, Qmax - maximum event discharge, Qsum - sum of event discharge) and antecedent discharge (Q7, Q15, Q30 - mean discharge 7, 15, 30 days before the event start) using PLSR analysis (Table 3).

![Figure 5. Spearman rank correlation between model parameters for each constituent within each catchment. White tiles are not significant (p > 0.01).](image-url)
The PLSR analysis yielded 10 out of 26 models with \( R^2 > 0.5 \). Most of those regression models satisfactorily predict event model parameters for SAC\(_{254} \) (\( n = 6 \)) while only two models predict parameters for EC and NO\(_3\)-N with acceptable performances. PLSR models for EC were found acceptable for the small FOR2 catchment only. Here parameter \( a \) was well predicted by event- and antecedent water temperature indicating higher EC values with higher water temperature. Parameter \( c \) was predicted by event discharge metrics. Higher maximum discharge within an event was connected to a more pronounced clockwise hysteresis while mean event discharge and sum of event discharge had an opposite effect on parameter \( c \).

The regression analysis for NO\(_3\)-N showed that parameter \( a \) in AGR2 was controlled by event- and antecedent water temperature and discharge yielding the highest \( R^2 \) of all PLSR models. Parameter \( a \) in AGR2 thus shows a clear seasonal pattern with increasing values and consequent shifts of C-Q event relationships with colder and wetter conditions. The rank correlation analysis indicated a similar relationship for AGR1 (Figure S6) not strong enough, however, for an acceptable PLSR model. The temporal changes of parameter \( b \) of NO\(_3\)-N in FOR2 were predicted mainly by antecedent and event water temperature indicating steeper C-Q slopes under warmer conditions.

The PLSR analysis for SAC\(_{254} \) yielded similar models satisfactorily predicting parameter \( a \) in the forested catchments FOR1 and FOR2 mainly from antecedent and event water temperature. In AGR2 a high parameter \( a \) was linked to low antecedent discharge 30 days before the event and to high water temperatures.

### Table 3

**Partial Least Squares Regression Models to Predict the Model Parameter a, b, and c From Event- and Pre-Event Conditions**

<table>
<thead>
<tr>
<th></th>
<th>EC</th>
<th>NO(_3)-N</th>
<th>SAC(_{254} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( a )</td>
<td>FOR2 ( R^2 = 0.59 )</td>
<td>AGR2 ( R^2 = 0.83 )</td>
<td>FOR1 ( R^2 = 0.56 )</td>
</tr>
<tr>
<td>pred</td>
<td>VIP</td>
<td>dir</td>
<td>pred</td>
</tr>
<tr>
<td>WT0</td>
<td>1.40</td>
<td>+</td>
<td>WT0</td>
</tr>
<tr>
<td>WT15</td>
<td>1.35</td>
<td>+</td>
<td>WT30</td>
</tr>
<tr>
<td>WT7</td>
<td>1.34</td>
<td>+</td>
<td>WT7</td>
</tr>
<tr>
<td>WT30</td>
<td>1.26</td>
<td>+</td>
<td>WT15</td>
</tr>
<tr>
<td>Q30</td>
<td>1.11</td>
<td>+</td>
<td>Q0</td>
</tr>
</tbody>
</table>

| \( b \) | FOR2 \( R^2 = 0.52 \) | dir | FOR2 \( R^2 = 0.56 \) | dir |
| pred | VIP | dir | pred | VIP | dir |
| WT7  | 1.29 | + | Q30  | 1.40 | + |
| WT15 | 1.28 | + | Qmax | 1.30 | - |
| WT0  | 1.25 | + | Qsum | 1.30 | - |
| WT30 | 1.21 | + | Q0   | 1.19 | - |
| Q0   | 1.09 | - | Q0   | 1.06 | - |
| Qsum | 1.27 | - | Q0   | 1.00 | + |

| \( c \) | FOR2 \( R^2 = 0.55 \) | dir | FOR2 \( R^2 = 0.60 \) | dir | AGR2 \( R^2 = 0.51 \) | dir |
| pred | VIP | dir | pred | VIP | pred | VIP |
| Qmax | 2.00 | + | WT0  | 1.31 | - | Qmax | 1.59 | + |
| Q0   | 1.32 | - | WT15 | 1.09 | + | Qsum | 1.49 | + |
| Qsum | 1.27 | - | WT30 | 1.07 | + | Q0   | 1.30 | + |
| Qmax | 1.06 | - | WT7  | 1.07 | - | Q7   | 1.04 | - |
| Q0   | 1.00 | + | Q0   | 1.00 | + |

Note. Only models with \( R^2 > 0.5 \) are reported. Models are separated by constituent, catchment, and the three predicted model parameters. For each model the most relevant predictors (pred) are ranked by the VIP value. Only predictors with a VIP > are reported. dir - direction of influence of the predictors derived from the PLSR model coefficients. A full correlation matrix of model parameters with the predictors in all catchments is given in Figure S6.
as well. Consequently \( \text{SAC}_{254} \) levels of the analyzed events are characterized by a pronounced seasonality with high values under dry and warm conditions across the forested catchments and AGR2. The inter-event variability of parameter \( b \) in FOR2 was linked to antecedent and event discharge conditions. Here, wetter antecedent conditions and smaller events (low maximum, mean and sum of event discharge) were connected to steeper event-scale C-Q relationships. Hysteresis parameter \( c \) in FOR2 was mainly explained by water temperature but with mixed directions. A more pronounced (wider) counterclockwise hysteresis pattern was connected to colder event temperatures but warmer antecedent conditions. In AGR2 parameter \( c \) varied mainly with event discharge indicating more pronounced clockwise hysteresis with larger events.

4. Discussion

4.1. Controls of C-Q Relationships at the Event Scale

The exploration of model parameters fitted for each event (Equation 2) allows an interpretation on the location of solute mobilization in the catchment and changes in source with time. Our analysis considers solutes with geogenic, biogenic and anthropogenic sources in both agriculturally managed and more pristine forested catchments. The following sections addresses hypothesis (1) on the land use control on event-scale C-Q relationships by discussing the findings for EC across all catchments and the findings for \( \text{NO}_3^- \) and \( \text{SAC}_{254} \) separated by the forested and agricultural catchments.

4.1.1. Similarity of EC Dynamics Across all Catchments

Specific conductance EC is a sum parameter of dissolved ions and therefore depends on the major ion concentrations. In the studied catchments these include calcium, sulphate, magnesium and chloride (Musolff et al., 2015) with a dominant geogenic origin. The groundwater dominated base flow has the highest concentrations of these solutes and is likely increasingly diluted with an increase of younger and shallower water components during discharge events. The dependence of geogenic major ions and thus EC in the stream on the depth of the source similar to the transit time since ion mobilization has been demonstrated before using modeling and data-driven approaches (Botter et al., 2020; Musolff et al., 2017a; Zhi et al., 2019). This is reflected in the mildly negative slope \( b \) for EC across FOR1, FOR2 and AGR1 (Table 2, Figure 3) catchments which has also been observed for geogenic solutes across a wide range of catchments (Godsey et al., 2009; Knapp et al., 2020; Rose et al., 2018). The export regime of EC (\( CV_c/CV_Q \), Table 2) was overall chemostatic with only minor concentration variability compared to discharge variability (Table 2). At the event scale, C-Q parameters varied little between events which underlines the spatial and temporal stability of the mobilization mechanism described above (Figure 4). Among the four catchments, median concentrations and model parameter \( a \) systematically decreased as annual runoff increased (Figure 4). This strong hydroclimatic control on average concentrations of geogenic solutes is in line with the findings of Godsey et al. (2019) along a much wider range of different catchments. In the small FOR2 catchment the high predictive power of water temperature on parameter \( a \) (PLSR in Table 3) can be interpreted as a temperature effect on weathering and thus availability of geogenic solutes, similar to findings of Knapp et al. (2020) for magnesium. This effect seems not to be evident in the larger catchments FOR1, AGR1 and AGR2. Hysteresis loops were of minor importance for the C-Q relationship of EC (Figure 4). Among the different catchments with contrasting land use (excluding mine drainage in AGR2), differences in both parameters \( b \) and \( c \) of EC were small compared to those for \( \text{NO}_3^- \) and \( \text{SAC}_{254} \) (Figure 4). This similarity across all catchments hint to a similar source in groundwater and a similar mobilization mechanism as described above that is independent from land use and hydroclimatic settings. This is further justified by the similarity in the geological settings and the major ion chemistry (Musolff et al., 2015). Our starting hypothesis (1) on the land-use control of source arrangement and C-Q relationships does thus not hold true for EC and the group of geogenic solutes with a low reactivity.

4.1.2. Coupled \( \text{NO}_3^- \) and \( \text{SAC}_{254} \) Event Behavior in Forested Catchments

In contrast to EC the C-Q behavior of \( \text{NO}_3^- \) as well as \( \text{SAC}_{254} \) was clearly different between the forested and agriculturally managed catchments corroborating hypothesis (1). Concentration of \( \text{NO}_3^- \) was generally low in the forested catchments and the event-scale analysis revealed a dominance of counterclockwise enrichment patterns (Figure 4), indicating a delayed increase in concentration compared to discharge. Parameter \( a \) varied between events and exhibited a seasonal trend, as reflected in the high predictive power
of water temperature and antecedent discharge (FOR2) (Table 3) comparable to the findings of Knapp et al. (2020). In agreement with other studies in forested settings (Duncan et al., 2017; Knorr, 2013; Koenig et al., 2017), we found mainly NO\textsubscript{3}-N enrichment C–Q patterns. However, the same studies reported mainly clockwise hysteresis, while we found for >69% counterclockwise hysteretic patterns. Knorr (2013), Koenig et al. (2017) and Duncan et al. (2017) argue for a finite pool of NO\textsubscript{3}-N in the riparian zone and shallow groundwater that is exhausted during the events which seems not to be the case in our catchments.

Although positive C-Q patterns at the event scale prevailed for SAC\textsubscript{254}, we observed a clear difference of event behavior between forested and agricultural catchments. In the forested catchments C-Q slopes (b) were more strongly positive, while parameter a was lower and hysteresis parameter c showed a higher variability over time (Figure 4) with a high share of events revealing counterclockwise hysteresis. The counterclockwise hysteresis is in line with findings of Strohmeier et al. (2013) and Vaughan et al. (2017) for DOC in forested catchments. Both argue for a greater connectivity of near stream riparian wetland DOC source zones on the falling limb of the event hydrograph. This also agrees with a model-based assessment by Zhi et al. (2019) stating that the hysteresis links to higher rates in the reaction providing DOC from soil organic matter in the uppermost part of the soil. The generally positive (parameter b) C-Q response in catchments with well-connected wetlands corresponds with findings of Winterdahl et al. (2014), Musolff et al. (2018), and Zarrnetske et al. (2018), with analyses over a large range of catchments. The reason for this enrichment patterns were shown to be the higher availability of DOC in the shallow organic soil horizons in contrast to low DOC availability in deeper groundwater. The soil sources get increasingly activated when the catchment wets up and near-surface flow paths are activated as demonstrated by both modeling and data-driven approaches (Musolff et al., 2018; Richardson et al., 2020; Seibert et al., 2009; Zhi et al., 2019). At the small FOR2 catchment we found a strongly positive correlation between NO\textsubscript{3}-N and SAC\textsubscript{254}, hysteresis parameters c, but also between C-Q slope b and parameter a (Figure 5) pointing to a similar source of both constituents. Werner et al. (2019) could link the DOC concentration and quality dynamics in FOR2 to the hydroclimatically controlled biogeochemical conditions in the riparian zone. In the data presented here, this is reflected in the predictive power of antecedent conditions for the model parameters at the forested sites (Table 3). Warm antecedent conditions are connected to a faster decomposition of soil organic matter, a higher availability of DOC and thus SAC\textsubscript{254} as well as NO\textsubscript{3}-N in the permanently wet riparian source zone (Zhi et al., 2019) resulting in higher concentrations (parameter a), steeper C-Q slopes (b) and more pronounced hysteresis (c). In the riparian zone, nitrification and denitrification can lead to a very close spatial (vertical) proximity of nitrate and DOC source zones (Frei et al., 2012; Riedel et al., 2013). This suggests that in the studied forested catchments dominant sources for the stream export SAC\textsubscript{254} and NO\textsubscript{3}-N can be found in the top-soil horizons of the riparian zones reflected in the tight positive correlation of event-scale C-Q parameters in FOR2. This coupling was not as clear for the larger FOR1 catchment (Figure 5) and possibly blurred by the overall higher variability of C-Q parameters between events.

### 4.1.3. Agricultural Management Defines NO\textsubscript{3}-N Event Behavior

The agricultural catchments are characterized by much higher NO\textsubscript{3}-N concentration compared to the forested catchments (Table 2, Figure 4). At the event-scale we found difference between AGR1 with more clockwise dilution and AGR2 with more counterclockwise enrichment NO\textsubscript{3}-N event C-Q patterns while the seasonal temporal variation in concentration is comparable (Figure 4). The event behavior in AGR1 is similar to NO\textsubscript{3}-N event analysis in agricultural catchments by Fovet et al. (2018), Lloyd et al. (2016) and Rose et al. (2018). The observation of a positive global C-Q pattern across all data but a dilution at the event scale was described by Dupas et al. (2016) and also for 24% of 219 French catchment studied by Minaudo et al. (2019). The observed event-scale C-Q behavior of NO\textsubscript{3}-N in the AGR1 catchment mimics geogenic solutes with a higher availability in deeper groundwater compared to shallow water (Seibert et al., 2009) and a lower concentration at the falling limb of the hydrograph. In contrast the AGR2 catchment behaved very consistently across events with regards to C-Q slope b and hysteresis parameter c (Figure 4). Slope b was indicating enrichment patterns and was similar between event- and global C-Q response. This is in line with findings of Yang et al. (2018) in a subcatchment of the AGR2 and Musolff et al. (2016) who showed that higher water levels lead to a stronger connectivity between the stream and the shallow young, NO\textsubscript{3}-N rich groundwater. It was suggested that tile drains exaggerate this relationship (Musolff et al., 2015). Interestingly there is a similarity of NO\textsubscript{3}-N C-Q pattern (parameter b and c) in the AGR2 catchment with the forested catchments (Figure 4) but at a much higher concentration level. This similarity
between agricultural and undeveloped sites was also stated by (Zhi & Li, 2020). The more effective drainage of the NO$_3$-N source on the falling limb of the hydrograph in the riparian soils of the forested catchments seems to be functional similar to the role of tile drainage in the agricultural catchment. Consequently, we need to refine the starting hypothesis (1): Different land use patterns can lead to similar C-Q patterns at the event scale when source arrangement and mobilization mechanisms work in a comparable way.

The dominance of counterclockwise patterns with higher concentrations on the falling limb of the hydrograph is in line with findings of Outram et al. (2016) where it is shown that tile drains create these hysteretic patterns and stated by (Zhang et al., 2020) as well. The AGR2 catchment also exhibits a high predictive power of water temperature and antecedent discharge on model intercept $a$ (Table 3). This stresses the role of NO$_3$-N source zone availability in the cold season without plant uptake and of catchment wetness in shaping the connectivity between NO$_3$-N source zone and the stream. The question remains why two neighboring catchments with comparable land use, geology, topography and climatic characteristics exhibit diverging C-Q slopes and hysteresis at the event scale. We hypothesize that this can be related to agricultural management in the near stream zones. The AGR1 catchment drains into a drinking water reservoir which means that in a 100 m zone around the stream, the use of fertilizer is not allowed and inputs from tile drains are strongly regulated. In consequence near the stream the NO$_3$-N abundance is higher in the deeper groundwater fed from the agricultural hinterland compared to shallow ground- and soil water. We thus argue that the AGR1 catchment does not have a pronounced shallow near stream source zones of NO$_3$-N (absence of fertilization) and nor efficient connectivity of hinterland NO$_3$-N sources (absence of tile drains) exists. The lack of wide buffer zones with restricted fertilization in AGR2 lead to a high abundance of NO$_3$-N in shallow ground- and soil water while denitrification was shown to partly remove NO$_3$-N in deeper and older groundwater components in this catchment (Nguyen et al., 2021; Yang et al., 2018). The interplay of source distribution with the efficient connectivity by tile drains (Musolff et al., 2015) can explain the observed homogeneous positive event C-Q slopes and dominance of counterclockwise hysteresis in the AGR2 catchment. However, to fully disentangle the effect of spatial source heterogeneity in conjunction with potential differences in the spatial pattern of rainfall and discharge generation a spatially distributed monitoring and modelling approach in both agricultural catchments would be needed.

In both agriculturally managed catchments the event C-Q responses of SAC$_{254}$ were higher in model intercept $a$ but weaker and positive in $b$ and less variable in both $b$ and $c$. Here, clockwise C-Q patterns prevailed (Figure 4). Model parameter $a$ of SAC$_{254}$ was inversely related to parameter $a$ of NO$_3$-N for the same events (Figure 5) and exhibited a clear seasonality with positive predictive power of water temperature in AGR2 (Table 3). The inverse carbon-NO$_3$-N concentration relationship was described for larger scale datasets (Ebeling et al., 2021; Guillemot et al., 2021) and across different ecosystems globally (Taylor and Townsend, 2010). This is also in line with a better mobilization of DOC under NO$_3$-N reducing conditions in riparian zones (Musolff et al., 2017b; Strohmenger et al., 2020). The lower SAC$_{254}$ on the falling limb of the hydrograph that creates clockwise hysteresis also reflects lower DOC concentrations in the shallow NO$_3$-N source zones as discussed by Fovet et al. (2018). Fast connectivity of NO$_3$-N rich and DOC depleted source zones by tile drains may further enhance this tight coupling. Overall SAC$_{254}$ behavior shows that the near-stream source of DOC is likely similar in all catchments and therefore less controlled by land use and more by topographical settings and soil type distribution. However, biogeochemically mediated availability of DOC and hydrological connectivity can differ between catchments with similar sources.

### 4.2. How Do Event-Scale and Inter-Annual C-Q Models Align?

With regards to hypothesis (2) we found discrepancies between global and event-scale models that differed between solutes and catchments (Figure 6). In consequence, the event analysis added insights into solute mobilization in some but not in all cases. It is worth noting that for the AGR1 and AGR2 catchment a previous study provides an analysis of long-term slope $b$ for EC, NO$_3$-N and total organic carbon, (biweekly to monthly data from 1993–2009, Musolff et al. (2015)) that is nearly identical to the two-year $b_{glob}$ (2013–2014) reported here. This hints to a robustness of the C-Q slope $b_{glob}$ also for shorter time periods as well as to a persistence of C-Q relationships in time.

We found a close similarity of the inter-annual logC-logQ slope $b_{glob}$ of the entire EC time series to the event scale parameter $b$ (Figure 6 upper row). Since the event model parameters $a$, $b$ and $c$ did not vary
considerably between the events, the resulting global power law C-Q relationship was robust and allowed us to draw similar conclusions on sources and mobilization mechanisms following the idea of higher concentrations in older and deeper groundwater compared to shallower and younger soil water as described in chapter 4.1. The added value of event-scale analysis compared to inter-annual C-Q analysis is therefore rather minor when assessing geogenic solutes with low reactivity or EC.

In the forested catchments we found a general agreement of parameter $b_{\text{glob}}$ for NO$_3$-N and SAC$_{254}$ with the event scale C-Q patterns leading to similar conclusions on proximal (shallow), hydrologically connected sources for both constituents (Figure 6, middle row). A considerable variability in the event model parameters $b$ and $c$ point to a large scatter in the global C-Q relationship. That is a clear indication of variable hydrological or biogeochemical conditions before and during the different events that control availability of NO$_3$-N and DOC as hypothesized and as discussed in detail in chapter 4.1 above. In consequence, the analysis of the variability between events clearly adds insight into the role of antecedent conditions on SAC$_{254}$ or NO$_3$-N mobilization and exports that cannot be captured in inter-annual C-Q models.

In the agricultural catchment AGR1 global and event-scale C-Q patterns for NO$_3$-N diverged, clearly indicating the added value of event-scale analysis for understanding the role of protected stream buffer zones on NO$_3$-N mobilization. This behavior resembles Simpson’s paradox (Simpson’s reversal) a statistical phenomenon of diverging trends within and across groups (Simpson, 1951). For the C-Q relationship of NO$_3$-N in AGR1 the means that we cannot determine the global C-Q response from responses at the event scale. An interpretations of the C-Q relationship with coarser temporal resolution may therefore lead to a false or
incomplete inference of the underlying processes. In contrast, in the agricultural catchment AGR2 without a protected buffer zone the global C-Q relationship was aligned with the event-scale patterns that only showed low variability in the model parameter $b$ and $c$ between events. This was also observed by Knapp et al. (2020) and by Rose et al. (2018) for EC and geogenic solutes such as magnesium, calcium and sodium. For NO$_3$-N this similarity in behavior hints to a homogeneity of event responses in intensively managed agricultural catchments as also stated by Thompson et al. (2011), Basu et al. (2010) and by Rose et al. (2018). The reasoning behind that lies in the strong NO$_3$-N source from fertilizers in the agricultural catchments, as well as the enhanced connectivity of source zones and stream by tile drains in the AGR2 catchment (Dupas et al., 2017; Musolff et al., 2015). This catchment can be seen as an archetype of a managed catchment where NO$_3$-N export is similar across events and dominantly controlled by hydrologically induced connectivity of NO$_3$-N source and the stream network. We also note that this functional homogeneity of hydrologically driven exports is not necessarily connected to chemostatic export regimes. Consequently we need to refine hypothesis (2): Biogeochemically reactive solutes do not necessarily exhibit contrasting C-Q hydrologically driven exports is not necessarily connected to chemostatic export regimes. Consequently we need to refine hypothesis (2): Biogeochemically reactive solutes do not necessarily exhibit contrasting C-Q relationships at event- and inter-annual scale when intensive agriculture homogenize sources and mobilization mechanisms.

In the agricultural catchments, the global C-Q behavior of SAC$_{254}$ was hardly captured by the single inter-annual power law model. The generally positive event-scale C-Q patterns shift in their level (parameter $a$) with antecedent discharge conditions to stack to a non-identifiable global C-Q pattern (see the discussion on Simpson’s paradox above) independent of the presence of protected buffers (Figure 6). Here the event analysis allowed for insights into mobilization processes driven by the interplay of biogeochemical and hydrological processes as hypothesized and as described in chapter 4.1 that would not be obtained with inter-annual low-frequency C-Q analysis.

Large scatter and high standard errors of parameter $b$ in inter-annual C-Q relationships can be an indicator for a divergence of event-scale behavior. In that case a careful evaluation is needed that may separate event from non-event samples (see e.g., Minaudo et al. (2019)) to not draw wrong conclusions on solute sources and mobilization mechanisms.

In general, we found that the variability of the event scale C-Q model parameters for the small FOR2 catchment were more often explainable by event- and antecedent conditions than for the other, larger, catchments (Table 3). This was especially true for the hysteresis parameter $c$ that was predictable for two of the three studied constituents in FOR2 but only for one constituent in all other catchments. We argue that this parameter is particularly sensitive to small variations in event timing and catchment heterogeneity while parameters $a$ and $b$ are more robust and stable across events. In a small headwater catchment a rainfall event will lead to a more homogeneous and synchronous discharge and solute mobilization response. Large catchments will exhibit stronger differences in discharge timing in different parts of the catchment when a rainfall event passes through. Hysteresis is especially sensitive to discharge and solute mobilization timing because short time shifts between both can lead to large hysteresis (see Table S7, mean time shifts between C and Q for NO$_3$-N and SAC$_{254}$ are smaller than one hour). In the integration at the catchment outlet this may lead to less traceable hysteretic patterns as also described by Vaughan et al. (2017) and by Butturini et al. (2008). This may be especially true in catchments with heterogeneous solute sources. Classifying the type of rainfall event and the discharge response in a better way (e.g., Tarasova et al. (2018)), may help resolving differences between event C-Q responses.

5. Conclusions and Outlook

In this study we analyze two-year time series of specific conductance, nitrate and spectral absorbance as a surrogate for dissolved organic carbon in four neighboring catchments by applying a model that captures concentration level (parameter $a$), C-Q slope ($b$) and hysteresis ($c$) at the same time. The model parameter $c$ describes direction and magnitude of hysteresis and compared well to the established hysteresis index. The model used here does not aim to replace the established index but rather to provide an alternative straightforward approach when the integration of event- and inter-annual C-Q patterns are of interest.

The study catchments represent a hydroclimatic gradient and range from more pristine forested to agriculturally managed conditions. As hypothesized, we found a land use control on the event-scale C–Q behavior...
and differences between geogenic and biogeochemically reactive solutes in terms of inter-event variability. However we need to refine our initial hypothesis as solute-specific differences can occur. We found that EC dynamics were well captured by a global C–Q analysis derived from inter-annual time series. The C–Q slope hardly changed between the events and hysteresis was not pronounced. The dilution behavior with chemostatic export regimes (low CV_c/CV_Q ratio) was consistent between catchments, independent of land use and resembled the typical pattern of hydrologically controlled geogenic solutes. In contrast, SAC_254 was strongly variable between events with dominantly positive C–Q patterns and more pronounced hysteresis as a result of the interplay of biogeochemical controlled availability of DOC in near stream riparian soils and hydrologically controlled exports. An inter-annual C–Q analysis across all events was able to capture the general positive C–Q pattern but not the strong variability between events. In contrast there was less variability between events for the agricultural catchments but individual positive C–Q relationships at the event scale stack to an unclear inter-annual C–Q relationship that does not allow any inference of underlying processes. NO3-N also responded differently in forested and agricultural catchments. In the forested catchments we observed a similarity of NO3-N and SAC_254 exports at the event scale that points to similar sources and export mechanisms. In one of the agricultural catchments we found a functional homogeneity of NO3-N event responses with steep positive C–Q slopes that is reflected in the similarity of event-scale and global-scale C–Q analysis. Here, exports were dominantly controlled by connectivity to the agricultural source zones. In contrast, event behavior was different in the agricultural catchment that drains into a drinking water reservoir. A more strict regulation of near-stream zones in terms of fertilizer application and the absence of tile drains may explain the negative event-scale C–Q slopes while the overall concentration level was found to be comparable to the other managed catchment.

In consequence, event-scale C–Q analysis provided more insights when the global composite C–Q relationship was not able to capture the dynamic concentration behavior especially true for biogeochemically reactive solutes. We also found that the hysteric behavior of concentrations were most consistent in the small headwater catchment. Here, we could better predict the magnitude and direction of hysteresis parameter c from antecedent and event conditions. In larger catchments, however, we found the hysteresis to be less traceable. We argue that the hysteresis is very sensitive to different timing and mixing of storm events in larger catchments. This may blur the mechanistic meaning of the hysteresis. We also note that the logC-logQ slope b is more robust against this mixing. In conclusion, the interpretation of hysteresis should be handled with care and is likely most helpful in headwater catchments.

We hope analyzing the growing availability of high-frequency data across a larger range of catchment properties will provide more insights into mechanisms of reactive solute transport at the catchment scale in the future. Here, a refinement of the typology of a storm event that can be extracted from rainfall and discharge patterns is a promising approach (Tarasova et al., 2018). Adding more advanced measures of concentration and discharge variability such as spectral analysis (Hensley et al., 2018) or using machine-learning techniques (Aguilera & Melack, 2018) will furthermore improve our understanding of catchment-scale transport processes. In turn we argue that a better understanding gained from event-scale behavior of high-frequency data will feed back into the interpretation of low-frequency C-Q analysis especially with diverging global or seasonal versus event-scale C–Q patterns as shown by Minaudo et al. (2019).

Data Availability Statement

For this research are available in these in-text data citation references: Musolff (2020a) [creative commons attribution CC BY], LHW (2018) [for scientific, non-commercial use only], Cornes et al. (2018) [observational data are strictly for use in non-commercial research and non-commercial education projects only], EEA (2013) [no limitations on public access], (CLC, 2016) [no limitations on public access].

References


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