The potamochemical symphony: new progress in the high-frequency acquisition of stream chemical data

Paul Floury1,2, Jérôme Gaillardet1, Eric Gayer1, Julien Bouchez1, Gaëlle Tallec2, Patrick Ansart2, Frédéric Koch3, Caroline Gorge1, Arnaud Blanchouin2, and Jean-Louis Roubaty1

1Institut de Physique du Globe de Paris (IPGP), CNRS and Université Sorbonne Paris-Cité, 1 rue Jussieu, 75238 Paris, France
2UR HBAN, Institut national de recherche en sciences et technologies pour l’environnement et l’agriculture, Antony (IRSTEA), France
3Endress+Hauser SAS, Huningue, France

Correspondence to: Paul Floury (floury@ipgp.fr) and Jérôme Gaillardet (gaillardet@ipgp.fr)

Received: 10 January 2017 – Discussion started: 13 January 2017
Revised: 12 September 2017 – Accepted: 29 October 2017 – Published: 7 December 2017

Abstract. Our understanding of hydrological and chemical processes at the catchment scale is limited by our capacity to record the full breadth of the information carried by river chemistry, both in terms of sampling frequency and precision. Here, we present a proof-of-concept study of a “lab in the field” called the “River Lab” (RL), based on the idea of permanently installing a suite of laboratory instruments in the field next to a river. Housed in a small shed, this set of instruments performs analyses at a frequency of one every 40 min for major dissolved species (Na+ , K+ , Mg2+ , Ca2+ , Cl− , SO4 2− , NO3 − ) through continuous sampling and filtration of the river water using automated ion chromatographs. The RL was deployed in the Orgeval Critical Zone Observatory, France for over a year of continuous analyses. Results show that the RL is able to capture long-term fine chemical variations with no drift and a precision significantly better than conventionally achieved in the laboratory (up to ±0.5 % for all major species for over a day and up to 1.7 % over 2 months). The RL is able to capture the abrupt changes in dissolved species concentrations during a typical 6-day rain event, as well as daily oscillations during a hydrological low-flow period of summer drought. Using the measured signals as a benchmark, we numerically assess the effects of a lower sampling frequency (typical of conventional field sampling campaigns) and of a lower precision (typically reached in the laboratory) on the hydrochemical signal. The high-resolution, high-precision measurements made possible by the RL open new perspectives for understanding critical zone hydro-bio-geochemical cycles. Finally, the RL also offers a solution for management agencies to monitor water quality in quasi-real time.

1 Introduction

Rivers are messengers from the critical zone. The chemical composition of rivers offers a window into the multiple processes that operate among water, organic matter, primary and secondary minerals and living organisms at the Earth’s surface (Calmels et al., 2011; Feng et al., 2004; Kirchner et al., 2000, 2001; Neal et al., 2012, 2013). Understanding the parameters that control the composition of river water is not only a scientific challenge but also one of the major challenges for humanity to access and preserve drinkable water (Bain et al., 2012; Banna et al., 2014; Bartam and Ballance, 1996). A limit in our understanding of water geochemistry at the Earth’s surface is limited by the temporal resolution at which sampling can be operated (Whitehead et al., 2009). As summarized by J. Kirchner: “If we want to understand the full symphony of catchment hydrochemical behaviour, then we need to be able to hear every note.” (Kirchner et al., 2004, p. 1358). Yet, taking high-frequency sample sets back to the laboratory, filtering and analysing them for several elements is limited by the requirement of considerable human resources (Chapman et al., 1996; Danielsen et al., 2008; Halliday et al., 2015; Neal et al., 2013; Rozemeijer et al., 2014; Strobl and Robillard, 2008; Telci et al., 2009).
A significant number of studies have reported high-frequency chemical measurements in watersheds. Thus far, these data have been mostly acquired during limited periods of time such as single storm events or a day (Beck et al., 2009; Brick and Moore, 1996; Chapman et al., 1997; Gammons et al., 2007; Kurz et al., 2013; Liu et al., 2008; Morel et al., 2009; de Montety et al., 2011; Neal et al., 2002; Nimick et al., 2011, 2005; Takagi, 2015; Tercier-Weaber et al., 2009). Although these studies clearly highlighted the wealth of information provided by sampling rivers at sub-hourly frequency, they underestimate the legacy of past hydrological episodes (Kirchner, 2006; Jasechko et al., 2016; Rode et al., 2016) and are of limited use when mass budgets are to be calculated for a typical hydrological cycle.

To date, the best combination of high-frequency and long-term monitoring ever reported for river chemistry is a 7 h frequency sampling over 18 months (Neal et al., 2012). In this study, the authors demonstrate the “act of discovery” permitted by such sampling scheme, by showing that the high sampling frequency of river hydrochemistry over sufficiently long time spans reveals patterns related to hydrological and biological drivers that are imperceptible at lower sampling frequency. Automated approaches, developed using probes installed directly in the river (Rozemeijer et al., 2010a; Macintosh et al., 2011; Cassidy and Jordan 2011; Dábakk et al., 1999; Glasgow et al., 2004; Zhu et al., 2010; Yang et al., 2008), or using online instrumental devices in which continuously pumped water is injected (Rozemeijer et al., 2010b; Zabiegala et al., 2010; Jordan and Cassidy, 2011), are alternatives to sampling methods requiring human intervention. Several papers have been published over the last decade reporting existing devices mostly focused on monitoring dissolved N or P and organic matter (Clough et al., 2007; Kunz et al., 2012; Aubert et al., 2013b; Escoffier et al., 2016). A recent overview of the potential of available conductivity, dissolved oxygen and carbon dioxide, nutrients, dissolved organic matter and chlorophyll in situ probes is given by Rode et al. (2016). A new solution for high-frequency measurement of river chemistry is offered by bringing the laboratory’s measuring devices to the field (the “lab in the field” concept). A Swiss group has recently developed such a system (von Freyberg et al., 2017) by installing ionic chromatography devices in a hut next to a stream. In this paper, we present a parallel initiative named the River Lab (RL) and funded by the French programme CRITEX: “Innovative sensors for the temporal and spatial Exploration of the Critical Zone at the catchment scale” (https://www.critex.fr). This approach, like the previously published one, overcomes traditional limitations on the number of samples and avoids several issues related to sample transport, filtration and storage. The RL is able to perform a complete chemical analysis of all inorganic major anionic and cationic species in the dissolved load of river water using ion chromatography (IC), with a frequency of up to one complete measurement every 40 min.

This article is a proof-of-concept paper that describes the analytical design of the RL and its performance by evaluating the precision, reproducibility and accuracy of concentration measurements. The first results from the RL reveal a significant improvement in reproducibility compared to conventional sampling and analysis techniques. Leveraging these optimal analytical conditions, the RL is able to reveal temporal patterns of river chemistry, such as daily concentration variations. The RL opens thus new opportunities in the field of river chemistry research and environmental monitoring.

2 Monitoring site

The RL was installed in the Orgeval Critical Zone Observatory, located 70 km eastward from Paris, France (https://gisoracle.irstea.fr/), a temperate agricultural catchment, within the Seine River watershed, and part of the French Critical Zone Research Infrastructure OZCAR (“Observatoires de la Zone Critique, Applications et Recherche”). The Orgeval catchment is one of the most instrumented and documented river observatories in France, with 50 years of hydrological data (Garnier et al., 2014). Catchment hydrologic data are available on the ORACLE website (https://bdoh.irstea.fr/ORACLE/).

The RL was installed at the outlet of the Avenelles River, a sub-catchment in the Orgeval watershed. The Avenelles River drains an area of 45 km². The climate is temperate and oceanic, with cool winters (mean temperature 3 °C), warm summers (20 °C on average) and an annual precipitation rate of ~ 650 mm on average. The Avenelles sub-catchment sits within the sedimentary carbonate-dominated Paris Basin. The river is perennial, supplied by groundwater from the Brie aquifer, with water chemistry dominated by Ca²⁺, SO₄²⁻, HCO₃⁻ and NO₃⁻ ions. The water level at the Avenelles gauging station shows an average daily volumetric flow rate of 0.2 m³ s⁻¹ (from 1962 to 2016), with low water period in summer (0.1 m³ s⁻¹) and flash flood events reaching 10.4 m³ s⁻¹ in spring.

3 Design of the River Lab

The concept of the RL is to pump river water and feed it to a set of physico-chemical probes and ion chromatography (IC) instruments for a complete analysis of major dissolved species continuously at high frequency (40 min is needed for a complete analysis). All the instruments of the RL fit into a sub-catchment in the Orgeval watershed. The Avenelles River drains an area of 45 km². The climate is temperate and oceanic, with cool winters (mean temperature 3 °C), warm summers (20 °C on average) and an annual precipitation rate of ~ 650 mm on average. The Avenelles sub-catchment sits within the sedimentary carbonate-dominated Paris Basin. The river is perennial, supplied by groundwater from the Brie aquifer, with water chemistry dominated by Ca²⁺, SO₄²⁻, HCO₃⁻ and NO₃⁻ ions. The water level at the Avenelles gauging station shows an average daily volumetric flow rate of 0.2 m³ s⁻¹ (from 1962 to 2016), with low water period in summer (0.1 m³ s⁻¹) and flash flood events reaching 10.4 m³ s⁻¹ in spring.
The RL has been designed around a primary circuit, which pumps the river water at 700 L per hour. First, the unfiltered river water sampled in the middle of the stream (Fig. 1) continuously supplies an overflow tank where six parameters are measured: pH, conductivity, dissolved O$_2$, dissolved organic carbon (DOC), turbidity and temperature. The water is then released into the river downstream from the RL. The turnover time of water in this primary circuit is 2 min. The turbidity probe is installed upstream of the overflow tank in a pipe perpendicular to the flow to provide more accurate measurements. The turbidity and DOC probes benefit from an automatic self-cleaning every 5 min using compressed air. For all probes, the frequency of acquisition is one measurement per minute. The tank and each probe are hand-cleaned weekly. All probes are developed and provided by Endress & Hauser (E+H®).

Second, a fraction of water pumped through the primary circuit feeds another circuit directed toward two IC instruments for the measurement of major dissolved species concentrations. A filtration system is deployed between the primary circuit and the IC instruments, consisting of a tangential filter with a 2 µm pore size, followed by a 0.2 µm frontal filtration system through cellulose acetate filters (Fig. 1), crucial for the IC instruments. Cation and anion chromatographs, connected in series, are fed simultaneously every 40 min from the filtered water circuit through a injection valve. Between two injections, the water in the filtered circuit is constantly renewed (1 L per hour). Our tests show that the frequency for a complete and uncontaminated analysis of cation and anion is actually limited by the filtration device (see Sect. 4.3).

The IC analysis is performed using two Dionex® ICS-2100 (Thermo Fisher Scientific®) instruments using eluent produced with concentrated eluent cartridges and ultra-pure water (Fig. 1). The cationic species measured are Na$^+$, K$^+$, Mg$^{2+}$ and Ca$^{2+}$, and anionic species are Cl$^-$, NO$_3^-$ and SO$_4^{2-}$. The chosen analysis time is 30 min (40 min if Sr$^{2+}$ concentration measurements are included; see details in Supplement “Ion Chromatographs characteristics”). The multi-port valve installed upstream of the ICs allows us to check the drift of the instruments and the background signal by regular introduction of calibration solutions and pure distilled water (see Sect. 4). Pure distilled water is regularly (every 2 weeks) introduced to check the residual noise. Both cationic and anionic chromatographs are calibrated every 2 months using synthetic solutions mimicking the river chemistry, made from 1000 ppm mono-elemental standard solutions. Two sets of calibration solutions are prepared, one for anions and the second for cations. The first solution (called “River ×1”) is prepared based on concentrations of the river water during summer, i.e. with the highest measured concentrations for most species. In the second solution, these concentrations are doubled (called “River ×2”). Further solutions are produced out of River ×1 and ×2 through dilution by up to tenfold to achieve lower concentrations (“River ×0.5; ×0.25; ×0.1”). The resulting five calibration solutions cover the entire range of possible natural variability of each species observed for the Orgeval River, including flood events.
Table 1. Assessment of the RL accuracy and instrumental drift based on concentration measurements made after several injections of the standard solution “River ×1”. The uncertainty on the calibration solution is the quadratic sum of the uncertainty on the standard solutions (provided by the manufacturer) and the overall uncertainty for weighing during solution preparation. Measurement errors over 1 week and over 2 months are expressed as the relative standard deviation (RSD) calculated for repeated injections of the solution “River ×1” directly into the IC instruments via the multiport valve (see Fig. 1).

<table>
<thead>
<tr>
<th>Species</th>
<th>Calibration concentration</th>
<th>Uncertainty (mg L(^{-1}))</th>
<th>Uncertainty (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg(^{2+})</td>
<td>10.0</td>
<td>0.03</td>
<td>0.3</td>
</tr>
<tr>
<td>K(^{+})</td>
<td>3.0</td>
<td>0.01</td>
<td>0.45</td>
</tr>
<tr>
<td>Ca(^{2+})</td>
<td>130.0</td>
<td>0.39</td>
<td>0.3</td>
</tr>
<tr>
<td>Na(^{+})</td>
<td>10.0</td>
<td>0.03</td>
<td>0.3</td>
</tr>
<tr>
<td>SO(_4^{2-})</td>
<td>70.0</td>
<td>0.84</td>
<td>1.2</td>
</tr>
<tr>
<td>NO(_3^-)</td>
<td>60.0</td>
<td>0.84</td>
<td>1.4</td>
</tr>
<tr>
<td>Cl(^-)</td>
<td>40.0</td>
<td>0.28</td>
<td>0.7</td>
</tr>
</tbody>
</table>

One measurement (injection of “River ×1” solution 4 times successively)

<table>
<thead>
<tr>
<th>Number of measurements</th>
<th>Average (mg L(^{-1}))</th>
<th>SD (mg L(^{-1}))</th>
<th>RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(4)</td>
<td>10.08 3.00 129.86 9.98</td>
<td>0.02 0.16 0.21 0.21</td>
<td>0.16 0.27 0.12 0.16</td>
</tr>
</tbody>
</table>

One week (injection of “River ×1” solution every 8 h)

<table>
<thead>
<tr>
<th>Number of measurements</th>
<th>Average (mg L(^{-1}))</th>
<th>SD (mg L(^{-1}))</th>
<th>RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(19)</td>
<td>10.13 3.02 130.64 10.01</td>
<td>0.03 0.39 0.22 0.22</td>
<td>0.28 0.32 0.30 0.22</td>
</tr>
</tbody>
</table>

Two months (injection of “River ×1” solution every 2 days)

<table>
<thead>
<tr>
<th>Number of measurements</th>
<th>Average (mg L(^{-1}))</th>
<th>SD (mg L(^{-1}))</th>
<th>RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(28)</td>
<td>10.33 3.14 134.34 10.05</td>
<td>0.06 0.80 0.50 0.50</td>
<td>0.54 1.34 0.59 0.50</td>
</tr>
</tbody>
</table>

4 Analytical performances of the River Lab

RL data acquisition started on 12 June 2015. The reliability of the system was assessed through five different tests involving IC measurements and the sampling procedure (accuracy, drift, precision of the whole system, cross-contamination and reproducibility). We refer to the third edition of JCGM 200-2012 (Joint Committee for Guides in Metrology) (JCGM, 2012) for the terminology used in assessing the performance criteria.

4.1 Accuracy and instrumental drift

The aim of the RL is to achieve very high-frequency measurements of river chemistry over long periods of time (pluriannual). To compensate for any long-term drift in the IC calibration, instruments are calibrated with a new set of solutions every 2 months or after each maintenance operation on the IC instruments. However, calibration drift can occur over timescales shorter than 2 months, resulting in systematic and/or random errors in concentration measurements. We evaluated this effect using a set of injections of the “River ×1” solutions, over 1 week and over 2 months (Table 1). For all species measured, no systematic variation was observed in the measured concentration of the solution “River ×1”, showing that at the two timescales, instrumental drift does not induce any systematic bias on concentration measurements, and that most of the error is of random nature. Therefore, the standard deviation of the concentration measurements of a given solution can be used as a reliable measure of the error due to instrumental drift. The measurement error over 1 week is calculated as the standard deviation of concentration measurements over 19 injections of solution “River ×1” performed every 8 h during 1 week (from 5 to 12 November 2015). The measurement error over 2 months is calculated as the standard deviation of concent-
The agreement between the calculated concentrations of the "River \times 1" solution and the RL measurements also demonstrates the accuracy of the prototype (Table 1).

4.2 Precision of the whole system

In order to estimate the precision of the whole system (IC instruments combined with the sampling device including the primary circuit, the pump and the filtration units), we performed a "closed-loop experiment" over the course of 1 day by connecting the inlet and the outlet of the primary circuit to a 300 L tank containing river water. The test was performed 3 times over two different seasons (on 20 July 2015, 28 August 2015 and 17 April 2016). The conductivity probe (one measurement every minute) was used to check the stability of the water chemistry during the course of the experiment (Fig. S2). Our results show that a lapse of 2 h at least is necessary for the system to stabilize, corresponding to the homogenization time of the water within the closed loop (Fig. 2). After 2 h, major anion and cation concentrations show a remarkable stability, indicating the absence of drift over 24 h time lapse despite the temperature variations in the river water, and allowing us to estimate the precision of the whole system over 1 day using the standard deviation of the measurements performed during the test. The results of the test are presented in Table 2. The precision reached is better than 0.5 % for all species except for potassium, for which it is better than 1.2 %.

4.3 Cross-contamination

The ability of the RL to detect rapid variations in river chemistry (typically expected during storm events) depends on (1) the response time of the RL to a perturbation in the river and (2) the potential cross-contamination from one sample to the next one. We assessed these two effects by a tracer injection experiment. After establishing a closed-loop experiment (on 29 August 2015) and allowing for the period of stabilization, we introduced a known amount of NaCl (200 g previously dissolved in a small amount of river water) into the 300 L tank of river water in order to simulate a "spike" in the river chemistry. The monitoring of conductivity in the primary circuit allowed us to follow the propagation of the spike injection into the primary circuit while Cl⁻ concentrations measured by the IC every 40 min allowed us to follow its propagation through the filtration devices and IC instruments (Fig. 3). The conductivity probe shows that the salinity spike is detected very quickly and stabilized after 5 min. This indicates that the water in the primary circuit is quickly homogenized (in agreement with the high flow rate of the primary circuit: 700 L h⁻¹). Conversely, the Cl⁻ and Na⁺ concentrations only reach the expected concentration at the second IC measurement, i.e. after 80 min.

The first IC measurement following the spike injection indicates that only 93 % of the final steady-state concentration is reached, revealing a contamination of the (n)th sample by 7 % of the (n − 1)th sample. In practice, such a contamination will only be significant if the instantaneous derivative of river concentration with time is important. In the case of the Orgeval River, where the RL is deployed, the relative derivative of the concentration with respect to time is lower than 1 % per hour for 90 % of the time for all species. In this case, the cross-contamination induces an error of 0.07 % compared to the true concentration, which means that the effect of cross-contamination is negligible compared to the precision of the RL (see Sect. 4.2). However, in the case of flood events, when the stream flow increases quickly, the derivative of concentration can change by more than 10 % per hour. In such cases, cross-contamination will induce an error of 1 % or more. The injection test shows that the time resolution of the RL is limited by the transfer time of the water between sampling and injection into the IC instruments. This transfer time of the water in the RL is mainly due to the design of the filtration system, which may be improved in the future.

4.4 Reproducibility: RL vs. laboratory

As a final test for assessing the ability of the RL to record fine natural variations of river chemistry in comparison to conventional techniques of filtration and analyses in the laboratory, we focused on 2 days in the summer of 2015 following long periods without rain (21 July 2015 for cations and 19 April 2016 for anions), which showed very high resolution diurnal variations (< 5 % relative) in chemical compo-

<table>
<thead>
<tr>
<th>Date</th>
<th>Number of measurements</th>
<th>Mg²⁺ RSD (%)</th>
<th>K⁺ RSD (%)</th>
<th>Ca²⁺ RSD (%)</th>
<th>Na⁺ RSD (%)</th>
<th>SO₄²⁻ RSD (%)</th>
<th>NO₃⁻ RSD (%)</th>
<th>Cl⁻ RSD (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 July 2015</td>
<td>(22)</td>
<td>0.17</td>
<td>0.90</td>
<td>0.21</td>
<td>0.22</td>
<td>0.39</td>
<td>0.47</td>
<td>0.24</td>
</tr>
<tr>
<td>28 August 2015</td>
<td>(20)</td>
<td>0.32</td>
<td>0.63</td>
<td>0.31</td>
<td>0.36</td>
<td>0.20</td>
<td>0.25</td>
<td>0.19</td>
</tr>
<tr>
<td>17 April 2016</td>
<td>(35)</td>
<td>0.38</td>
<td>1.20</td>
<td>0.17</td>
<td>0.31</td>
<td>0.31</td>
<td>0.38</td>
<td>0.30</td>
</tr>
</tbody>
</table>
Stabilization

24

4

22

SO

20

12

8

16

(C - C)

Measured

Mean

ately using a Teflon

by the RL every 40 min, we conducted hourly sampling of

sition of the Orgeval River. In addition to the analyses made

injection. The conductivity measurement frequency is 1 per minute,

the time between two measurements of chloride concentra-

tion is 40 min. Error bars for conductivity and Cl\(^-\)

corresponding to the homogenization of water in the circuit and tank (see conductivity measurements in Fig. S2) for the calculation of

Figure 2. Assessment of the precision (in deviation from the mean for four dissolved species) of the whole RL system including the primary circuit, filtration systems and IC instruments (17 April 2016). A closed system is established on the primary circuit of the RL by connecting the inlet and the outlet through a 300 L tank of river water. The system is then run for a period of 24 h. The time between two IC analyses is 40 min. The purple curve represents data of temperature of the water in the tank. We do not consider the first 2 h (three first measurements), whereas the time between two IC analyses is within symbols’ size. Results are normalized to initial and final values

Figure 3. Cross-contamination assessment and response time of the RL system after a spike injection of 200 g of NaCl. A closed system is established on the primary circuit of the RL by connecting the inlet and outlet through a 300 L tank of river water prior to the injection. The conductivity measurement frequency is 1 per minute, whereas the time between two measurements of chloride concentration is 40 min. Error bars for conductivity and Cl\(^-\) concentration measurements are within symbols’ size. Results are normalized to the difference between the minimum value, before the tracer injection (0 %), and the maximum value, at the end of the experiment (100 %).

so of the Orgeval River. In addition to the analyses made by the RL every 40 min, we conducted hourly sampling of the river by collecting 5 L of water and filtering it immediately using a Teflon\(^\text{®}\) frontal filtration unit (Sartorius\(^\text{®}\)) with 0.2 µm porosity polysulfonether filters. Bottles of acidified (at pH = 2) and unacidified river water were transported to the laboratory at IPGP for measurement of major cations and anions, respectively, using IC devices similar to those installed in the RL (Thermo Fisher\(^\text{®}\) ICS-2100). In the laboratory, measurements were performed using a Thermo Fisher\(^\text{®}\) ICS-5000 for cation measurements and a Dionex\(^\text{®}\) 120 from Thermo Fisher\(^\text{®}\) for anion measurements. The calibration procedure in both laboratory and RL is the same, using the same set of calibration solutions. The error measurement reached in the laboratory is estimated at 1 % through repeated injections of the standard solution “River × 1” (every five samples). Comparison between the RL and the laboratory for the seven measured species are shown in Fig. 4. First, the measurements made by the RL are more precise than those performed in the laboratory, a feature that can be primarily attributed to the greater stability of the continuously working injection system of the RL. Second, the fine variations measured by the RL are reproduced in the laboratory, validating the observed diurnal variations and supporting the reliability of the RL to detect changes of the order of a percent within a day. The third observation is that small yet systematic offsets between the two sets of data exist, up to 3 % for Mg. One possible explanation for this difference is that the filtration procedures differed between the RL and the manual sampling, which may have led to a discrepancy in the concentration measurements related to the potential for some elements to be hosted in the colloidal phase (Dupré et al., 1999). In addition, the most accurate measurements were obtained with the RL rather than with the laboratory equipment because the RL is continuously processing solutions with a similar matrix, thereby minimizing memory effects and cross-contamination that can compromise measurements if widely differing samples are run successively on the same instrument. These features of the measurement protocol, representative of most laboratory workflows for hydrochemical measurements, are likely to lead to inaccuracies. Regardless of the observed discrepancy between the two sets of measurements, we note that variations in concentration recorded by the RL and measured at the IPGP laboratory have the same amplitudes and are synchronous.
The tests performed and reported above clearly demonstrate an improvement in precision compared to the analysis of bottled samples taken back to the lab. We see three main reasons for this improvement.

1. In a given river, dissolved concentrations typically vary by less than 1 order of magnitude when water discharge changes by several orders of magnitude (Godsey et al., 2009). This constancy allows us to select a relatively narrow range of concentration for establishing specific calibration curves of the IC instruments, a condition which is rarely possible in the laboratory, where different kinds of samples are analysed.

2. While in the laboratory samples are injected discretely, in the RL river water samples are injected as a continuous flow. Thus, the primary circuit and the filtration system operate continuously at a constant pressure, which supports stable and accurate analyses.

3. The third factor is the experimental conditions in the bungalow. The temperature is maintained at 24 °C ± 2° (in addition to the 40 °C thermostatically controlled temperature in the column, precolumn and detection device of the ICs) allowing for better stability of the IC measurements. Moreover, the RL IC instruments are never stopped, which favours stability.

5.2 What is revealed by a higher sampling frequency?

To our knowledge, the high frequency of measurements (one measurement every 40 min) reached by the RL installed on the Orgeval River is the highest ever reported for stream chemistry over several months. To highlight the corresponding improvement in the recorded concentration signal, we tested the effect of sampling frequency on the concentration signal. First, we artificially subsampled the RL original signal at two lower sampling frequencies: every 7 h (starting 5 October 2015 at 10 pm) and every 24 h. The 7 h frequency was chosen to reproduce the sampling frequency of Neal et al. (2012) made in the Plynlimon watershed, Wales. The daily sampling frequency is typically what is achievable on the long term by “human grab-sampling” in the field. Second, we calculated the probability density function (PDF) of concentration measurements over a given time interval. The use of PDFs allows us to explore the structure of concentration signals beyond the mean concentration, which constitutes an important metric for river solute budget, but lacks any insight into the variations in concentrations that can be used to retrieve information on catchment processes. We describe the PDF by three statistical parameters: mean, standard deviation and skewness. Skewness indicates the distribution asymmetry, both in magnitude and direction (a positive skewness means that most values are higher than the mean). Altogether, the three parameters account, at first order, for the structure of a concentration signal. We compared these three parameters for the computed PDFs to quantify the signal degradation induced by artificial subsampling.
We applied this statistical approach to two representative periods of the hydrological cycle of the Orgeval Critical Zone Observatory: a typical 6-day rain event caused by the arrival of a wet, Atlantic meteorological front (in October 2015) and a dry summer low water stage period (July 2015), where the stream is essentially sustained by groundwater, during an apparently steady hydrological period. We first present the behaviour of calcium and sulfate concentrations as an example during the two considered periods (Figs. 5 and 6), before generalizing to all measured species (Supplement and Figs. S3, S5 and S6).

– Rain event. The Ca concentration time series recorded at a 40 min frequency shows that minimum Ca concentrations are recorded at maximum water discharge, but this relationship is invisible at lower sampling frequency (Fig. 5). Narrow peaks during the maximum of the stream flow are unresolved at a daily or 7 h frequency. The comparison of the calculated PDFs shows that a bimodal character is captured at all frequencies. The average and standard deviation are not significantly affected by the sampling frequency, with a relative difference of less than 2 % for the values of these parameters between the three distributions. However, the skewness values vary among the different records. From the 40 min frequency to the daily frequency signals, the skewness is weaker, which means that even if the overall concentration variability is well captured at the lower sampling frequencies, the concentration signal is clearly degraded. This degradation is particularly intense during the middle of the rain event, where the concentration signal evolves quickly.

– Summer event. Despite the absence of rain events during the 2015 summer, the River Lab recorded high-frequency variations revealing a diurnal structure with 7 % relative variations between day and night. Each element exhibits its own type of daily variation in terms of amplitude and regularity. Figure 6 shows that the structure of this signal is altered when the sampling frequency decreases. While these daily variations are still captured when sampling occurs every 7 h, their amplitude is somewhat altered (5 %) compared to the 40 min sampling frequency (8 %). The daily variability of the signal is absent on the daily sampling frequency. While the mean remains the same over the range of sampling frequency, the variability quantified by the relative standard deviation decreases with lower sampling frequency, by up to 50 % for the daily frequency compared to the 40 min frequency signal, indicating a significant loss of information. The skewness of the concentration distribution recorded at a subsampled daily frequency has a value that is opposite in sign compared to the other two frequencies, indicating that there is an inversion of the measured asymmetry of the PDF at lower sampling frequencies. Therefore, too coarse a sampling frequency can yield a strongly altered signal compared to higher frequencies, resulting in a biased shape of the distribution of the concentrations.

– Generalization. The resampling approach applied above is generalized and expanded to other elements for both the summer and rain events. The generalization to all species measured is presented in the Supplement. In Figs. 5 and 6, we arbitrarily chose the hour of sampling (10:00 and 14:00 for Figs. 5 and 6, respectively). In Figs. S3, S5 and S6, the subsampling is performed
at each of the possible sampling hours. This statistical analysis quantitatively demonstrates that such high-frequency measurements are able to capture the day–night chemical cycles of the Orgeval River. Given the amplitude and duration of typical rain events in the catchment, the alteration of the signal by lowering the sampling frequency is less critical but still significant during these periods (Supplement; Figs. S3, S5 and S6).

5.3 What is revealed by better analytical precision?

As shown above, the Orgeval RL not only achieves high-frequency measurements but also results in improved precision compared to conventional lab analysis following manual sampling. Therefore, any sampling procedure, even at a high frequency, involving conventional lab analysis induces a loss of precision. We demonstrate this effect through a numerically generated artificial degradation of the precision. Using the original RL concentration signal as a reference, we artificially degraded the signals by adding a normally distributed noise onto the concentration signals recorded by the RL. Noise levels of 4 and 2 % were tested; they are representative of the “standard” analytical precision reported for most laboratory IC devices. The same representative periods as in the previous section (summer and rain events) were utilized for these tests. In this section we present the example of one element for each characteristic period (Ca$^{2+}$ for rain event Fig. 7 and SO$_4^{2-}$ for summer event Fig. 8. The generalization for all elements is detailed in the Supplement (see Figs. S4, S7 and S8).

- Rain event. Figure 7 illustrates the concentration PDF obtained after degradation of the analytical precision for the Ca concentration. The narrow peaks recorded during the maximum of the stream flow are virtually invisible in the signal at a 4 % precision, and strongly smoothed in the signal at a 2 % precision. The original bimodal characteristic of the PDF is still visible in the 2 % precision signal but no longer in the 4 % precision signal. The mean and standard deviation appear to be insensitive to these changes in analytical precision, while the skewness is strongly impacted, reflecting significant alteration of the concentration PDF at lower precision.

- Summer event. Figure 8 shows how the sulfate concentration signal is affected when the precision is degraded. Day–night variations are only visible in the original RL signal because of its high analytical precision. The effect of degraded precision on the PDFs is
Figure 7. (a) Calcium concentration and stream flow in the Orgeval River during a rain event (from 1 to 25 October 2015), as recorded by RL and for two artificially degraded signals using a normally distributed noise with standard deviation of 2 and 4 %, to reflect the effect of decreased analytical precision. Black dots represent data during the rain event strictly from 5 (12:00) to 10 October 2015. The probability density functions (PDFs) of concentration are calculated and represented as histograms (b). For each PDF, the following statistical parameters are calculated: average (Ave.), standard deviation (SD) and skewness (Skew.). Grey dots represent concentration values outside of the rain event, which are not considered for the analysis presented in (b).

more important than for the rain event (Fig. 7). While the mean value is robust, the standard deviation is altered (+150 % from the RL signal to the 4 % precision signal). The skewness decreases (but keeps the same sign) by up to 90 % for the signal at 4 % precision compared to the original signal and 74 % for the signal at 2 % precision, indicating that the original RL signal asymmetry is lost as precision is worsened. These changes in the parameters of the concentration PDF show that the structure of the concentration signal in the Orgeval River would be significantly altered if the measurements were made with analytical precision lower than that of the RL prototype.

6 Conclusions

This paper demonstrates the feasibility of deploying conventional laboratory instruments in the field to measure the concentration of major dissolved anions and cations in rivers (Na\(^+\), K\(^+\), Mg\(^{2+}\), Ca\(^{2+}\), Cl\(^-\), SO\(_4^{2-}\), NO\(_3^-\)) at a high frequency (one measurement every 40 min) and at a high analytical precision (better than 1 %) over several months. The River Lab prototype was installed in the Avenelles stream at the Orgeval Critical Zone Observatory, France. The RL features physico-chemical probes, an online 0.2 µm pore size filtration system, and two ionic chromatographic devices, all installed in a closed, air-conditioned bungalow. The RL is autonomous, remotely operable, and data can be transmitted automatically. Human intervention is required only once a week. Therefore, the RL allows for an efficient attribution of human resources, as well as considerable saving of consumables.

A suite of tests performed on the RL to assess quality measurement and to compare with more conventional “grab sampling” followed by laboratory measurements revealed only a minor drift in the instrument calibration, leading to improved precision. This precision is not easily achieved in the laboratory under standard analysis conditions, showing the benefit of transporting the laboratory devices to the field. The analytical capabilities of the RL for major dissolved elements could theoretically be extended to other elements separable by ion chromatography. Preliminary tests demonstrate that species present in trace amounts in river water (down to ppb, such as strontium or lithium) could be measured with the same gain in precision.

For this particular prototype, the measurement frequency (every 40 min) appears to be limited by the turnover time of water in the filtered water circuit, which is itself imposed by the filtration unit. However, the high frequency and high precision of the RL enabled precise and accurate observations on the fine structure in hydrochemical time series. Their interpretation is beyond the scope of the present proof-of-concept paper but the RL is able to capture the abrupt changes in dissolved species concentrations during a typical 6-day rain event, as well as daily oscillations during a hydrological steady period of summer drought.

Using the high-frequency RL signal as a benchmark, it is possible to artificially alter the sample frequency and the analytical precision and study the resulting effect on the hydrochemical distribution obtained for characteristic hydrological events. This analysis shows that in order to retrieve the fine structure of the hydrochemical signal, high sampling
frequency and improved analytical precision are both necessary conditions. To paraphrase James Kirchner’s quote: “If we want to understand the full symphony of catchment hydrochemical behaviour, then we need to be able to hear every note” (Kirchner et al., 2004). The improvements made possible by the RL here or concomitantly by von Freyberg et al. (2017) allow us to consider hearing the full potamological symphony.

Future work will explore the relationships between the desired measurement frequency and the timescales characterizing the complex interactions between primary and secondary minerals, biotic processes and hydrological processes in catchments. Recording such fine stream hydrochemical variations has the potential to offer a new perspective in critical zone science development.

**Data availability.** Data will be available in a dedicated database website after a contract accepted on behalf of all institutes.

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

**Acknowledgements.** This work was supported by the EQUIPEX CRITEX programme (grant no. ANR-11-EQPX-0011, PIs J. Gaillardet and L. Longuevergne) and funding from IRSTEA (Institut national de Recherche en Sciences et Technologies pour l’Environnement et l’Agriculture). We thank Magadalena Niska for administrative help.

We would like to thank Jérôme Laurent, Xu Zhang, Quentin Charbonnier, Damien Calmels, Pascale Louvat, James Kirchner, Jenny Druhan, Susan Brantley, Bill McDowell and Jon Chorover for their help in the field and helpful comments. Alain Guerin (IRSTEA), Sylvain Losa (Thermo Fisher), Cedric Fagot, Patrick Reignier and Matthieu Bauer from Endress+Hauser (colleagues of Frédéric Koch) are thanked for technical assistance. Paul Floury benefited from a doctorate grant from MESR, France. The Orgeval CZO River basin belongs to the French National Infrastructure OZCAR (Observatoires de la Zone Critique, Applications et Recherche).

Edited by: Laurent Pfister
Reviewed by: two anonymous referees

Figure 8. (a) Sulfate concentration in the Orgeval River recorded by the RL during 2 weeks in summer (7 to 19 July 2015), and for two artificially degraded signals, using a normally distributed noise with a standard deviation of 2 and 4 %, to reflect the effect of degraded analytical precision. The probability density functions (PDFs) of concentration are calculated and represented as histograms (b). The average (Ave.), standard deviation (SD), and skewness (Skew.) are calculated for each PDF.
References


