New on-line method for water isotope analysis of speleothem fluid inclusions using laser absorption spectroscopy (WS-CRDS)

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Abstract

A new online method to analyse water isotopes of speleothem fluid inclusions using a wavelength scanned cavity ring down spectroscopy (WS-CRDS) instrument is presented. This novel technique allows us to simultaneously measure hydrogen and oxygen isotopes for a released aliquot of water. To do so, we designed a new simple line that allows the on-line water extraction and isotope analysis of speleothem samples. The specificity of the method lies in the fact that fluid inclusions release is made on a standard water background, which mainly improves the $\delta D$ reliability.

To saturate the line, a peristaltic pump continuously injects standard water into the line that is permanently heated to 140°C and flushed with dry nitrogen gas. This permits instantaneous and complete vaporisation of the standard water resulting in an artificial water background with well-known $\delta D$ and $\delta^{18}O$ values. The speleothem sample is placed into a copper tube, attached to the line and after system stabilisation is crushed using a simple hydraulic device to liberate speleothem fluid inclusions water. The released water is carried by the nitrogen/standard water gas stream directly to a Picarro L1102-i for isotope determination. To test the accuracy and reproducibility of the line and to measure standard water during speleothem measurements a syringe injection unit was added to the line.

Peak evaluation is done similarly as in gas chromatography to obtain $\delta D$ and $\delta^{18}O$ isotopic composition of measured water aliquots. Precision is better than 1.5‰ for $\delta D$ and 0.4‰ for $\delta^{18}O$ for water measurement for an extended range ($-210$ to 0‰ for $\delta D$ and $-27$ to 0‰ for $\delta^{18}O$) primarily dependent on the amount of water released from speleothem fluid inclusions and secondarily on the isotopic composition of the sample. The results show that WS-CRDS technology is suitable for speleothem fluid inclusion measurements and gives results that are comparable to Isotope Ratio Mass Spectrometry (IRMS) technique.
1 Introduction

Speleothems are being more and more used for paleoclimate reconstructions due to their widespread occurrence and potential to deliver precisely dated and highly resolved climatic records covering very long time intervals (e.g. Cheng et al., 2009; Badertscher et al., 2011). To date, the vast majority of speleothem-based paleoclimate reconstructions are based on oxygen isotope ($\delta^{18}O$) measurements on calcite. However, their interpretation remains difficult as they can be influenced by several and sometimes competing climatic and non-climatic factors (Lachniet, 2009), such as changes in seasonality of rainfall, varying cave air temperatures and evaporation. Fluid inclusions are common in speleothems (0.05 to 0.5 weight %) and are natural repositories of cave drip water (Scheidegger et al., 2010). Therefore, the hydrogen ($\delta D$) and oxygen ($\delta^{18}O$) isotopic composition of fluid inclusions holds direct information on the isotopic composition of paleoprecipitation (Dennis, 1997; Baker and Bradley, 2010) and, when combined with isotopic analyses of speleothem calcite, it can be used to calculate either paleotemperatures (McGarry et al., 2004; Tremaine et al., 2011) or to reveal changes in the source of moisture (Fleitmann et al., 2003). Though the potential of isotope measurements of fluid inclusion water is now fully recognized, various analytical limitations (e.g. sample size restrictions, time consuming analysis, fractionation processes during extraction of fluid inclusion water) made a wider application of this crucial climate proxy difficult.

The relationship between cave drip water trapped in fluid inclusions and precipitations was first established by Harmon et al. (1979). Later, it was also shown that cave temperature generally reflects the mean temperature at the surface (Fairchild et al., 2006), so fluid inclusions are consequently a key parameter to reconstruct continental paleotemperatures. More recently, experiments have shown that fluid inclusions preserve the isotopic composition of parent drip water (Dublyansky and Spötl, 2009).

To extract water from speleothems fluid inclusions, two different techniques have been used in the past three decades: (i) the extraction by crushing the sample and...
(ii) the extraction by thermal decrepitation. Originally, investigation of inclusions in speleothem using the crushing extraction procedure was initiated by pioneering work in the late seventies (Schwarz et al., 1976; Harmon et al., 1978, 1979). Dennis et al. (1997) improved the precision for hydrogen isotopes of fluid inclusions using the zinc reduction method and a vacuum crushing device coupled to a cold trap cell. He was the first to introduce a combined oxygen and hydrogen isotope analysis on fluid inclusions (Dennis et al., 2001). Since then considerable advances have been made, in particular, in applying on-line techniques either for the isotope analyses of calcite (Smalley et al., 1989) or of water released from different archives such as ice cores, speleothems, etc. (Huber and Leuenberger, 2003; Leuenberger and Huber, 2002; Morrison et al., 2001). Analyses of both water isotopes in a unique measurement were achieved using the pyrolysis coupled to IRMS technique and gas chromatography (Gehre et al., 2004; Sharp et al., 2001). More recently, the same method was adapted and modified to successfully perform water isotopes analysis of speleothem fluid inclusions (Vonhof et al., 2006; Dublyansky and Spötl, 2009).

The second technique is the use of thermal decrepitation at temperatures of around 550 °C to extract water (Yonge, 1982). Several studies show a large fractionation of up to 30 ‰ for δD in comparison to parent cave drip waters (Yonge, 1982; Matthews et al., 2000; McGarry et al., 2004). No fractionation is observed when the speleothem sample is heated to a maximal temperature of 400 °C (Verheyden et al., 2008).

Current research focuses on the extraction of fluid inclusion water at low temperatures (∼ 140 °C) with the use of laser spectroscopy as a valuable alternative to the traditional IRMS detection technique (Arienzo et al., 2013). The use of laser spectroscopy is growing rapidly and is now being recognised as a reliable, precise and easy-to-use technique for water isotopes analysis as mass spectrometry (Brand et al., 2009) and using it for speleothem fluid inclusion measurement constitutes a logical extension for this technique. Water isotope measurements with laser spectroscopy is nowadays mainly used for continuous atmospheric isotope measurements (Aemisegger et al., 2012) as well as for discrete water measurements (Brand et al., 2009).
We present one of the first applications of the laser absorption technique to measure fluid inclusion water extracted from the speleothem calcite. The particularity of the method is that the line is always kept under humid conditions as a water background is continuously generated. To generate the water background, a procedure similar to the calibration protocol for atmospheric measurement is used (Iannone et al., 2009; Sturm and Knohl, 2010) and it requires the vaporisation of water standard in a vaporisation system. Compared to the crushing device of the Amsterdam line (Vonhof et al., 2006), our line is used to investigate whether the removal of any cryogenic traps prior to the analysis is an advantage.

This paper discusses the technical aspects of the newly designed line, the calibration of the laser instrument and the evaluation of the data. Furthermore, it documents the accuracy and precision of the method for standard and real sample measurements.

2 Method

2.1 Line description

The new line consists of basically three units: (i) a water background generator for improved instrumental performance; (ii) a syringe injection unit to allow single injections of small aliquots of standard water and (iii) a simple home-made crushing device.

The whole line, shown in Fig. 1, is flushed with nitrogen gas to transfer the water sample from the crushing device to the Picarro L1102-i analyser (Picarro Inc., Sunnyvale, CA, USA). The delivery of nitrogen gas in the line is controlled by an Analyt-MTC pressure regulator (Analyt-MTC GMBH, Müllheim, Germany), which is set to regulate a constant pressure of 1.015 bars over atmosphere. Detailed information of all three units is provided below.

A vaporisation unit (Water Background Generator) was designed to generate a stable water vapour mixing ratio and isotopic background via a constant supply of internal standard water (ST-08) with well-known $\delta^D$ and $\delta^{18}O$ isotopic values into a nitrogen
stream using a Union Tee connector as a simple dripping device. Droplets of standard water is supplied by an Ismatec IP High Precision Multichannel Pump (IDEX Health & Science GmbH, Wertheim, Germany), via a 0.13 mm inside diameter TYGON R3607 capillary extended with a heat resistant 1/32″ stainless steel capillary. The peristaltic pump is well-suited for continuous injections of standard water at a rate of ∼4 µL min⁻¹ (depending on the capillary diameter and the pump speed) into the dripping device. Performance of the pump was tested by monitoring the supply of standard water for 20 h at intervals of 1 s using a Sensirion Mass Flow Controller (Sensirion AG, Staefa, Switzerland) that was placed right after the pump. This test showed that the average delivered amount of water is 4.8 ± 1 µL min⁻¹, whereas short-term fluctuations (over five seconds) range between 2.72 and 7.38 µL min⁻¹. However, these are smoothed out by the mixing volume, the crusher device as well as the whole line. The dripping device is an integral part of the Water Background Generator where dry nitrogen and standard water are injected into the line via a capillary that slightly touches the wall of the stainless steel line (connecting lines are all made with Swagelok 1/8″ stainless steel tubes). The advantages of this design are that the droplets are vaporised instantaneously and formation of large bubbles at the tip of the capillary is minimized to prevent a delay in the release of water. The temperature (140 °C) and the pressure are kept constant inside the line to ensure total and instantaneous vaporisation of the water without isotopic fractionation. The vaporised water is carried along the line by nitrogen gas. The pump speed can be adjusted in a range of water vapour mixing ratio ranging from ∼6700 ppmv to the detection limit of the laser spectrometer (including the purge unit). The nitrogen pressure has been selected such that the stability of the water background is enhanced. The water vapour and dry nitrogen are mixed homogeneously in the mixing cavity, which has a volume of ∼2 cm³ and is placed immediately after the injection unit, in order to minimise fluctuations caused by the peristaltic pump and to improve the stability of the water background.

As the injected water vapour mixing ratio would be too high for the sensitivity of the Picarro, a 10 cm stainless steel capillary purge has been added after the mixing cavity,
which releases $\sim 350 \text{ mL min}^{-1}$ dry nitrogen/water vapour mixture. The remaining fraction ($\sim 65 \text{ mL min}^{-1}$) passes through the line and yields a stable isotopic background for $\delta D$ and $\delta^{18}O$ (see Sect. 2.4 for further details). Moreover, this setup allows us to maintain a quite stable water vapour mixing ratio over time.

A syringe injection unit has been placed prior to the crushing unit to perform reproducibility tests using various standard waters and to make calibration peaks during speleothem fluid inclusion measurement. It is made of a customised Union Tee connector with an added 7/16’’ Marathon septum to seal it and a piece to drive the needle inside the line. The septum should be replaced regularly as variations of pressure trough the septum can lead to a loss of stability when the septum is deteriorated. Liquid water samples ranging from 1 to 2 µL were injected using a 10 µL Hamilton syringe, whereas for lower amounts we used a 1 µL SGE Analytical Science syringe.

The crushing device has a simple design and consists of a Power Team P12 hydraulic press (Power Team, Rockford, IL, USA) that exerts a pressure of 220 bars onto the copper tube, which corresponds to a force of approximately 10 000 Newton. A 7 µm Swagelock inline filter was placed after the copper tube to avoid that calcite particles travel to the laser spectrometer, which may damage the laser instrument. Crushing of the sample generates no leaks or cracks to the copper tube. To estimate the crushing efficiency, a grain size analysis has been made on one sample that led to the following size distribution: 20 % for the fraction $0 < x < 0.149 \text{ mm}$, 30 % for $0.149 < x < 0.495$ and 50 % for $x > 0.495 \text{ mm}$ with possible individual particles of up to 1 mm size.

After syringe injection or sample crushing, the released water is mixed with the nitrogen/standard vapour mixture to form a three components gas that is then flushed to the laser spectrometer through a stainless steel capillary.

The entire line is uniformly heated to $140^\circ\text{C}$ by two heating sources without any cold spots to completely vaporise a very large quantity of water and to avoid condensation and accumulation of water in dead volumes. Compared to a line at room temperature, a heated line minimises the water adsorption on its wall. As shown in Fig. 1, the vapourisation unit, including the water pump inlet, the mixing cavity and the purge, are heated.
to 140°C using a digitally controlled heating band that allows a precise temperature control over time. The injection unit and the speleothem crushing unit are placed into an oven where the temperature is also kept at 140°C.

Once the three components (nitrogen/standard water/sample water) mixture reaches the laser spectrometer, it passes through the vaporiser chamber, which is equally heated to 140°C, into the optical cavity cell where it is measured at precisely controlled pressure and temperature conditions of 35 Torr and 80°C, respectively. The WS-CRDS operation procedure is well established and further details are described in Crosson (2008) and Gupta et al. (2009).

Results are given in parts per million by volume (ppmv) for the water vapour mixing ratio and in permil (‰) for the isotopic compositions. They are normalised and expressed against the common Vienna Standard Mean Ocean Water (VSMOW) standard.

### 2.2 Sample preparation

Firstly, a parallelepiped of calcite with a width of 25 mm, a height of 5 mm and a variable length taken along the growth axis of the stalagmite is extracted (preferably with apex in the middle). This piece of calcite is then cut in 50 mm long pieces and fixed with glue on both sides of a glass thin section to avoid disintegration of the sample during the cutting process (Fig. 2). This also allows to follow a marker layer by using a commercially available 3032 Precision Horizontal Diamond Wire Saw (Well Diamond Wire Saw SA, Le Locle, Switzerland) equipped with a 0.3 µm wire. This set up allows cutting the sample without any significant loss of material and, moreover, cutting the calcite by making curves using a potentiometer.

In addition, a 200 µm thin section to evaluate the distribution and volume of the fluid inclusions is prepared. One of the advantages of such a preparation is that once the abundance of fluid inclusions is accessed, the thickness of the sample can be adjusted. After the samples are cut, they are placed into a bath of acetone for a couple of hours to remove the residual glue that may have remained on the sample.
The sample varying between 0.1 and 1 g is then placed into a ∼ 16 cm long high purity copper tube with a diameter of 3/8” inch, which has been prepared and cleaned with acetone and flushed with dry air to remove any residual dust. The copper tube has a volume of ∼ 11 cm³ and samples of up to 20 mm in length can be loaded. To avoid powder dissemination into the system, quartz wool is placed at both ends of the tube. The loaded tube is then quickly attached to the line using 1/4” and 3/8” Swagelok connections.

### 2.3 Desorption tests

Atmospheric water vapour desorption of the sample can be a source of error in the water isotope analysis of speleothem fluid inclusions (Dennis et al., 2001) as the adsorbed atmospheric water present on the calcite surfaces can contaminate the released water after the crushing. It was recommended to operate fluid inclusion extraction at a temperature of 150 °C, more recent studies (Vonhof et al., 2006; Dublyansky and Spötl, 2009) showed that temperatures of 120 to 130 °C are sufficient to avoid desorption and adsorption problems. For systems working with a dry background, it is not only necessary to remove water adsorbed on the sample prior to crushing but also on all surfaces of the whole measuring system. With our analytical setup we only need to test the desorption efficiency from the speleothem sample surface. However, desorption tests were made in two steps. Firstly, we evaluate desorption from a stalagmite sample under dry nitrogen background conditions to precisely characterise the phenomenon and to estimate the total released water content. Secondly, we adapted the procedure to our normal measuring conditions in a humidified mode.

Desorption tests have been performed in the oven in which the loaded copper tube is positioned. The speleothem sample is placed at room temperature and flushed with dry nitrogen to the laser instrument for direct determination of desorbed water. With our setup, the nitrogen gas stream contains approximately 15 ppmv of residual water. After the stabilisation and recovering of a dry background, the system is heated to 140 °C to evaluate how heating affects water desorption from the calcite (Fig. 3a). We performed
five speleothem sample desorption measurements. Results of these tests show that the released amount of water varies between 0.06 and 0.13 microliter per gram of calcite. Dublyansky and Spötl (2009) showed that decrepitation of single inclusions during the heating of calcite can occur which is then clearly visible on a mass spectrometer spectrogram. So far we have not noticed such effects with our system.

The water vapour mixing ratio and the isotopic composition were monitored under measuring conditions (Fig. 3b). To minimize the migration of atmospheric water vapour into the system when a sample is loaded, the copper tube should be rapidly fixed to the heated line (140°). To ensure complete desorption of water adsorbed on the sample surface, the sample is heated until the background conditions are stable to a high degree ±0.25 and ±0.90 ‰ (if not specified standard deviation of single values is given) for δ18O and δD, respectively. This procedure typically takes 90 min before another sample measurement is processed. The time interval for the background determination is about 20 min (100 values), whereas uncertainties of the mean background is in the order of ±0.03 and ±0.1 ‰ for δ18O and δD, respectively. The background amount of water is fairly constant at ±20 ppmv for single values and ±2 ppmv for its mean. Such stable background conditions enhance the sample precision as discussed in detail in Sect. 2.4.2.

As the system is kept under humidified conditions, line walls are constantly conditioned with water molecules, and the probability of isotope exchange with adsorbed water at the walls is rather low since the mean free path length is small under our conditions. Therefore, it takes about 50 min to stabilise the background conditions. For speleothem samples measurements, 90 min were taken to ensure that no atmospheric water vapour contamination will affect the measurement.

Though there are slightly different stability conditions between empty, loaded and crushed tube conditions, these different states may slightly affect water background levels (as shown on Fig. 3b).
2.4 Laser spectrometer calibration

The use of the WS-CRDS system is new for this kind of application and the calibration and the precision of the Picarro L1102-i analyser has been determined carefully. The precision of the data will be discussed in Sect. 4. In this section, we present tests that are relevant for optimising the continuous measurements using our online system (coupling of preconditioning/crusher/laser instrument). Table 1 gives the isotopic composition of standard waters used for the calibration.

2.4.1 Isotopic calibration for water vapour measurements

Three standards (Meerwasser, ST-08 and DYE-III) have been used to perform the isotopic calibration of the instrument against the VSMOW scale. Their isotopic values were precisely measured in our institute by mass spectrometry and they ranged from 0 and $-27\,\%$ for $\delta^{18}O$ and from 0 to $-220\,\%$ for $\delta D$. These waters fit with the expected range of values of fluid inclusion water in speleothems. Injection of each standard lasts at least three hours to ensure that the system reaches an equilibrium state. The correlation between the measured ($x$ value) and the assigned value ($y$ value) is used as the instrument correction equation. For this study the corrected isotopic value for $\delta^{18}O$ is $y = 0.989x \pm 0.001 + 0.112 \pm 0.021 (r^2 = 1)$ and for $\delta D$ is $y = 0.965x \pm 0.002 - 0.574 \pm 0.299 (r^2 = 1)$. These equations needs to be updated depending on the instrument performance and they are applied to all isotopic data recorded by the laser instrument in order to get the most accurate isotopic composition of the sample.

2.4.2 Optimising a stable water background

Laser spectroscopy under dry conditions (water background 10 to 100 ppmv) results in highly variable $\delta D$ and $\delta^{18}O$ isotopic background values of $\pm 1000\,\%$ for $\delta D$ and $\pm 250\,\%$ for $\delta^{18}O$, which hampers obtaining reliable data. Furthermore, when working
close to dry conditions water molecules have a tendency to stick to the line surface and fractionation may occur. This is particularly important for small amounts of water as it is the case in our application. To avoid such an effect, we decided to continuously admix standard water with a known isotopic composition to generate an artificial water background that is close to the best operational domain of the analyser (17 000 to 23 000 ppmv). This implies that during the entire measuring procedure, the surfaces of the device are saturated with standard water molecules to avoid this volatility problem that is well-known for dry conditions.

As working reference for the speleothem fluid inclusion measurement we use an internal standard water (ST-08). To evaluate the stability of the system over a workday, a 13 h continuous admixture of ST-08 has been established. Results show that the water vapour mixing ratio in the optical cavity cell reached 13 462 ± 32 ppmv (1σ). Similarly, the isotopic stability reaches −77.31 ± 0.58 ‰ for δD and −10.93 ± 0.12 ‰ for δ18O.

The dependence of isotope ratios on the mixing ratio was evaluated by varying the mixing ratio for eleven individual levels between 6700 and 20 500 ppmv. Data were recorded for each level for ~ 40 min and data analyses were made during the last 20 min (Table 2). Standard deviation is robust for δ18O (±0.10 ‰) between 13 000 and 19 000 ppmv and only slightly enhanced below and above this range (Fig. 4). For δD, standard deviation decreases continuously from 1.1 to 0.5 ‰ between 7000 and 13 000 ppmv and then stabilises below 0.5 ‰ up to 22 000 ppmv. The stability of the water background is ±10 ppmv below 14 000 ppmv and increases up to ±30 ppmv above it. The reproducibility of the individual isotope measurements for δD increases with the increasing mixing ratios, but remains stable for δ18O (Fig. 5). The absolute values for δD show a decrease with increasing mixing ratios and then a constant behaviour over 13 000 ppmv, whereas δ18O values increases with rising mixing ratios up to 12 000 ppmv and are rather stable above it. Based on these observations we decided to perform the measurements at a background mixing ratio of ~ 13 000 ± 1000 ppmv.
2.4.3 Time needed for system stabilisation

The stabilisation time after replacing an empty copper tube by a loaded one can be evaluated by monitoring both isotope values and water vapour mixing ratios. To do so, three experiments were conducted: (i) with an empty tube, (ii) with a speleothem sample loaded tube and finally (iii) with a crushed sample. Figure 6 shows the response time of the system after switching between two standards (from ST-08 to DOME-C). For the three experiments, the system behaves in the same way as documented by the three resulting isotope traces. There are no significant offsets in the delta values as a function of time, suggesting that the presence of material in the tube has no significant influence on the time needed for the system to stabilise when switching between two isotopically different waters. Approximately 45 and 75 min are necessary to stabilise $\delta^{18}$O and $\delta$D, respectively.

2.5 Measurement protocol

To achieve precise and reproducible measurements, several tests have been made to optimise the measurement protocol (Fig. 7). Once a stable background is reached, a first 0.5 µL internal standard (ST-08) is injected manually (peak 1 in Fig. 7). After 40 min, the empty tube is then quickly exchanged (within 30 s) with the copper tube containing a sample (peak 2) to minimise ambient water vapour contamination. We wait for approximately 90 min to be sure to restore the original conditions including the removal of adsorbed water from the sample. Insufficient time could lead to a mixture of background standard water with adsorbed atmospheric water vapour. Once a stable background is reached again (water vapour mixing ratio and isotopic composition), a second standard ST-08 reference peak is injected (peak 3, Fig. 7). After 40 min, the speleothem sample is crushed (peak 4, Fig. 7) and the released fluid inclusions water reaches the Picarro analyser approximately 40 s later. After up to 60 min (depending on the water amount released from the sample) a last standard water aliquot is injected on the crushed calcite (peak 5, Fig. 7). 40 min later the measurement sequence...
is terminated. These three standard peaks allow us to control the stability of the measurement. The overall time needed to run one speleothem sample using the described method (Fig. 7) is approximately five hours of which the sample peak itself lasts only around 20 min. In a time optimized procedure a minimal time of 2 h for one sample is necessary to guarantee the stated precision.

3 Raw data evaluation

The analyser software records all measured parameters at intervals of $\sim 12.2$ s. The water vapour mixing ratio value can be used as it is recorded, whereas $\delta D$ and $\delta^{18}O$ need to be corrected using a linear correlation that is obtained by measuring various standard waters with known isotopic values as shown in Sect. 2.4.1.

3.1 Estimation of sample water concentration

The released amount of sample water is determined as follows: various aliquots of water ranging from 0.02 to 2 µL were injected on a wet background to determine the associated signal area $S_A$ (Eq. 1a) or maximal signal amplitude $S_M$ (Eq. 1b). They are then used to determine the amount of water released through crushing the sample with a precision of $\pm 0.08$ µL for $S_A$ and $\pm 0.10$ µL for $S_M$ (Fig. 9):

$$H_2O[\mu L] = 7.16 \times 10^{-7} \cdot S_A + 0.0244 \quad (1a)$$

$$H_2O[\mu L] = 9.50 \times 10^{-9} \cdot S_M^2 + 0.000152 \cdot S_M + 0.0187. \quad (1b)$$

The sample water concentration ($c$) is obtained by dividing Eq. (1a) by the sample weight ($m$).

$$c = \frac{H_2O[\mu L]}{m[g]} \quad (2)$$
3.2 Calculation of $\delta$D and $\delta^{18}$O isotopic values

When the sample is crushed, standard and sample water with different isotopic compositions are mixed in the line. To determine the isotopic composition of the sample we integrate the product of the water amount and its isotopic value over the entire peak. However, we need to take into account the background (Fig. 10).

As the software delivered with the Picarro cannot be used for this measurement procedure, we developed a simple data evaluation protocol to calculate isotopic composition of the standard or speleothem fluid inclusion water. A criterion for peak integration is determined on the basis of the water content. The peak inflection point is well defined as the released (or injected) water leads to a rapid increase of the water vapour mixing ratio, more generally the criterion for the peak starting time ($t_0$) is defined by $dH_2O(t)/dt \geq 2$ ppmvs$^{-1}$.

Fractionation occurs right after the peak inflection point and after the peak maximum the mixing ratio will decrease slowly with time. As mentioned before, with this setup a peak lasts $\sim$20 min. Criterion used to determine the peak end time ($t_1$) is $dH_2O(t)/dt \geq -0.15$ ppmvs$^{-1}$. Precise peak definition is especially crucial when small water amounts are being measured.

The general equation for the water mixture amount is:

$$H_{2O_{mix}} = H_{2O_s} + H_{2O_b}$$  \hspace{1cm} (3)

where $s$ and $b$ stands for the sample and the background, respectively. The same can be applied to heavy-oxygen water:

$$H_{2^{18}O_{mix}} = H_{2^{18}O_s} + H_{2^{18}O_b}.$$  \hspace{1cm} (4)

With Eqs. (4)/(3) and approximating $H_{2O_{mix}} = H_{2^{16}O_{mix}} + H_{2^{18}O_{mix}}$ with $H_{2^{16}O_{mix}}$ ($H_{2^{18}O_{mix}}$ contributes only 0.2 %), we have $H_{2^{18}O_{mix}}/H_{2O_{mix}} \approx H_{2^{18}O_{mix}}/H_{2^{16}O_{mix}}$.
The isotope values in Delta-notation can be calculated as follows:

\[
\delta^{18}O_j = \frac{\int_{t_0}^{t_1} \delta^{18}O_j(t) \cdot H_2O_j(t) dt}{\int_{t_0}^{t_1} H_2O_j(t) dt} \quad (5a)
\]

\[
H_2O_j = \frac{\int_{t_0}^{t_1} H_2O_j(t) dt}{\int_{t_0}^{t_1} dt} = \frac{\int_{t_0}^{t_1} H_2O_j(t) dt}{t_1 - t_0} \quad (5b)
\]

where \( j \) is either the mixture, the background or the sample.

To calculate the isotopic value of a sample \( \delta_s \) of the injected or released water we use the following relationship.

\[
\delta_s = \frac{\delta^{18}O_{mix} \cdot H_2O_{mix} - \delta^{18}O_{b} \cdot H_2O_{b}}{H_2O_{mix} - H_2O_{b}} \quad (6)
\]

For \( \delta D \), we used a correspondent evaluation.

4 Reproducibility tests

4.1 Water injections

To test the new setup three different internal standards were analysed to ensure good accuracy and precision of the line. A total of 46 manual injections of standard water of 1, 1.5 and 2 \( \mu L \) were performed. The three known isotopic standards ranged between
0.63 and $-429.39 \, \%$ for $\delta D$ and between $-0.11$ and $-54.23 \, \%$ for $\delta^{18}O$. Reproducibility test of these standard waters (Meerwasser, DYE-III and DOME-C) are summarised in Table 3 and Fig. 11.

For $\delta^{18}O$, measurements are rather accurate and precise over the entire range between 0 and $-55 \, \%$. However, standard deviations are slightly influenced by the water amount and the Delta value, both resulting in a change of the spectral signal strength of the laser system. The lower the water amount and the lower Delta value the smaller the spectral signal for the less abundant isotope gets. This is clearly visible for samples with very negative $\delta D$ values (DOME-C water) for which we receive a clear decrease in precision but retaining accuracy. Therefore, the standards used for calibrating the whole setup should be representing the expected range of sample Delta-values.

4.2 Speleothem samples

For fluid inclusion analysis, a total of twelve samples from four different layers of a stalagmite from a cave in northern Borneo in the West Pacific (Meckler et al., 2012) were analysed (Fig. 12). Each sample was split into several pieces of between 100 and 200 milligrams each to run three replicates. The results of the $\delta D$ and $\delta^{18}O$ measurements and water amounts are summarized in Table 4. The released water amounts range from 0.4 to 2.2 µL and correspond to water contents between 2 to 19 microliter per gram of calcite (average of 10.1 µLg$^{-1}$). The calculated water concentration between the three replicates is not constant (except for sample 387). This is most probably due to the unequally distributed fluid inclusions in the stalagmite and we expect the influence of the crushing to be minimal since the grain size distribution is rather constant. The released water amounts do not to have a strong influence on the isotope values measured. Nevertheless, the sample weight should be selected so that the released water amount is at least 0.5 µL or higher. In addition, two fluid inclusion analyses were made for a recent speleothem samples from Milandre cave (Switzerland) where the corresponding modern drip water is also monitored. Samples around 800 to 900 mg
were crushed releasing a constant water amount of 0.6 µL (average water concentration of 0.68 µL g⁻¹). The observed offset between drip water and fluid inclusion for δ¹⁸O of one permil might indicate an exchange between calcite and inclusion water after its formation since the isotopic composition of hydrogen matches well. Results are given in Table 5.

5 Conclusions

We present a new system for measuring hydrogen and oxygen isotope ratios of fluid inclusions water extracted from speleothems. The measuring principle is based on the wavelength-scanned cavity ring-down spectroscopy technology that allows simultaneous monitoring of hydrogen and oxygen isotopes. The main advantages of the WS-CRDS technique in relation to our study are: (i) this instrument is only dedicated to water isotopes measurement, (ii) the operational costs of the analysis are low and (iii) the speleothem fluid inclusion measurement procedure is simplified by avoiding any water treatment prior to their isotopic determination, such as the reduction of H₂O in a pyrolysis reactor into CO and H₂ molecules as it is required for the combination of gas chromatography (cleaning of the products) and IRMS method (isotope detection unit). Moreover, the line avoids the step of the freezing treatment using a cryo-focusing cell.

The new simple and easy-to-build line allows extracting and analysing the isotopic composition of water released from speleothem fluid inclusions by crushing the sample with a hydraulic press. This line could be even used for air inclusions, not discussed here.

With this setup, we achieve standard deviations smaller than 1.5‰ for δD (between 0 to −210‰ at least) and 0.4‰ for δ¹⁸O (between 0 to −27‰ at least) for sample sizes from and above 1 µL, comparable with traditional IRMS measurements. These isotopic intervals are wide enough for our speleothem target measurement. For more negative values, measurements are still possible, but with lower precision but retained accuracy.
(especially for $\delta^D$). It is best to adjust the isotope value of background water and the calibration waters to the target range (expected sample values). The reproducibility for real stalagmite samples was in the same range even for small-sized sample amounts in the order of 0.5 $\mu$L.

The sample throughput could be increased by placing two or more copper tubes in parallel to optimise the stabilisation time between speleothem crushing, or the measurement protocol could also be shortened leading to substantial gain of time and allowing the crushing of more than two samples per day.

**Acknowledgements.** We warmly thank Kim Cobb, Jess Adkins and Nele Meckler to have let us measure one of their stalagmites. We would also like to thank Yves Krüger for his active involvement in sample preparation and for making the fluid inclusions photos available and Peter Nyfeler for his helpful assistance and involvement in line maintenance. This study is part of “STALCLIM”, a Sinergia project financed through the Swiss National Science Foundation (grant no. CRSI22-132646/1).

**References**


Table 1. Isotopic composition of internal standard waters used for calibration determined independently by IRMS.

<table>
<thead>
<tr>
<th>Standard</th>
<th>δ^{18}O (VSMOW) (%)</th>
<th>δD (VSMOW) (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meerwasser</td>
<td>−0.044 ± 0.05</td>
<td>1.24 ± 0.5</td>
</tr>
<tr>
<td>DOME_C</td>
<td>−54.18 ± 0.05</td>
<td>−428.26 ± 0.5</td>
</tr>
<tr>
<td>DYE-III</td>
<td>−27.21 ± 0.05</td>
<td>−210.23 ± 0.5</td>
</tr>
<tr>
<td>ST08</td>
<td>−10.79 ± 0.05</td>
<td>−77.46 ± 0.5</td>
</tr>
</tbody>
</table>
Table 2. Effect of the water vapour mixing ratio on δD and δ^{18}O measurement vs. VSMOW based on 40 min measuring intervals per water level.

<table>
<thead>
<tr>
<th>Water vapour mixing ratio (ppmv)</th>
<th>Standard deviation (ppmv)</th>
<th>δ^{18}O (%)</th>
<th>Standard deviation (%)</th>
<th>δD (%)</th>
<th>Standard deviation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6717</td>
<td>5</td>
<td>−11.44</td>
<td>0.13</td>
<td>−77.41</td>
<td>1.01</td>
</tr>
<tr>
<td>8000</td>
<td>5</td>
<td>−11.36</td>
<td>0.11</td>
<td>−77.48</td>
<td>0.78</td>
</tr>
<tr>
<td>9520</td>
<td>6</td>
<td>−11.23</td>
<td>0.11</td>
<td>−77.83</td>
<td>0.67</td>
</tr>
<tr>
<td>10 810</td>
<td>5</td>
<td>−11.18</td>
<td>0.11</td>
<td>−78.08</td>
<td>0.62</td>
</tr>
<tr>
<td>12 093</td>
<td>6</td>
<td>−11.16</td>
<td>0.11</td>
<td>−78.04</td>
<td>0.54</td>
</tr>
<tr>
<td>13 414</td>
<td>8</td>
<td>−11.13</td>
<td>0.10</td>
<td>−78.32</td>
<td>0.46</td>
</tr>
<tr>
<td>15 016</td>
<td>29</td>
<td>−11.12</td>
<td>0.09</td>
<td>−78.34</td>
<td>0.50</td>
</tr>
<tr>
<td>16 365</td>
<td>28</td>
<td>−11.06</td>
<td>0.10</td>
<td>−78.40</td>
<td>0.51</td>
</tr>
<tr>
<td>17 698</td>
<td>22</td>
<td>−11.07</td>
<td>0.10</td>
<td>−78.45</td>
<td>0.40</td>
</tr>
<tr>
<td>18 968</td>
<td>15</td>
<td>−11.04</td>
<td>0.10</td>
<td>−78.44</td>
<td>0.48</td>
</tr>
<tr>
<td>20 476</td>
<td>35</td>
<td>−11.12</td>
<td>0.11</td>
<td>−78.45</td>
<td>0.45</td>
</tr>
</tbody>
</table>
**Table 3.** Experimental results with injected aliquots of standard water (refer to Table 1 for uncertainties of the expected values). Note that the aliquots slightly deviate from values given in Table 1.

<table>
<thead>
<tr>
<th>Standard</th>
<th>Injected amount (µL)</th>
<th>Number of injections (n)</th>
<th>( \delta^{18} \text{O} ) average (‰)</th>
<th>Standard deviation (‰)</th>
<th>( \delta \text{D} ) average (‰)</th>
<th>Standard deviation (‰)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meerwasser</td>
<td>2</td>
<td>6</td>
<td>0.3</td>
<td>0.1</td>
<td>-1.0</td>
<td>0.3</td>
</tr>
<tr>
<td></td>
<td>1.5</td>
<td>5</td>
<td>0.3</td>
<td>0.3</td>
<td>-1.1</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>5</td>
<td>0.2</td>
<td>0.2</td>
<td>-0.8</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>expected values</td>
<td></td>
<td>-0.1</td>
<td></td>
<td>0.6</td>
<td></td>
</tr>
<tr>
<td>DYE-III</td>
<td>2</td>
<td>5</td>
<td>-27.1</td>
<td>0.2</td>
<td>-212.6</td>
<td>0.4</td>
</tr>
<tr>
<td></td>
<td>1.5</td>
<td>5</td>
<td>-27.1</td>
<td>0.3</td>
<td>-211.5</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>4</td>
<td>-27.1</td>
<td>0.1</td>
<td>-212.1</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td>expected values</td>
<td></td>
<td>-27.3</td>
<td></td>
<td>-212.8</td>
<td></td>
</tr>
<tr>
<td>DOME-C</td>
<td>2</td>
<td>5</td>
<td>-54.3</td>
<td>0.3</td>
<td>-429.7</td>
<td>2.0</td>
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<tr>
<td></td>
<td>1.5</td>
<td>5</td>
<td>-54.4</td>
<td>0.2</td>
<td>-430.1</td>
<td>2.1</td>
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<tr>
<td></td>
<td>1</td>
<td>6</td>
<td>-54.2</td>
<td>0.7</td>
<td>-428.7</td>
<td>4.2</td>
</tr>
<tr>
<td></td>
<td>expected values</td>
<td></td>
<td>-54.2</td>
<td></td>
<td>-429.4</td>
<td></td>
</tr>
</tbody>
</table>
### Table 4. Experimental results with crushed speleothem samples from Borneo.

<table>
<thead>
<tr>
<th>Sample</th>
<th>( \delta^{18}O ) (‰)</th>
<th>( \delta D ) (‰)</th>
<th>Released water (µL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>246-1</td>
<td>-5.6</td>
<td>-35.4</td>
<td>1.99</td>
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<tr>
<td>246-2</td>
<td>-6.1</td>
<td>-38.1</td>
<td>0.95</td>
</tr>
<tr>
<td>246-3</td>
<td>-6.0</td>
<td>-39.6</td>
<td>2.04</td>
</tr>
<tr>
<td>average</td>
<td>-5.9</td>
<td>-37.7</td>
<td></td>
</tr>
<tr>
<td>standard deviation</td>
<td>0.3</td>
<td>2.1</td>
<td></td>
</tr>
<tr>
<td>266-1</td>
<td>-8.5</td>
<td>-58.7</td>
<td>2.18</td>
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<tr>
<td>266-2</td>
<td>-8.3</td>
<td>-58.4</td>
<td>1.87</td>
</tr>
<tr>
<td>266-3</td>
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<td>-58.1</td>
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<tr>
<td>standard deviation</td>
<td>0.1</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td>277-1</td>
<td>-8.5</td>
<td>-60.2</td>
<td>1.34</td>
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<tr>
<td>277-2</td>
<td>-9.0</td>
<td>-61.1</td>
<td>0.44</td>
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<tr>
<td>277-3</td>
<td>-8.3</td>
<td>-59.1</td>
<td>0.62</td>
</tr>
<tr>
<td>average</td>
<td>-8.6</td>
<td>-60.1</td>
<td></td>
</tr>
<tr>
<td>standard deviation</td>
<td>0.4</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>387-1</td>
<td>-7.7</td>
<td>-53.6</td>
<td>1.95</td>
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<td>387-2</td>
<td>-7.9</td>
<td>-52.3</td>
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<td>387-3</td>
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<tr>
<td>average</td>
<td>-7.7</td>
<td>-52.8</td>
<td></td>
</tr>
<tr>
<td>standard deviation</td>
<td>0.1</td>
<td>0.7</td>
<td></td>
</tr>
</tbody>
</table>
Table 5. Isotopic results for two replicates of a recent sample from Milandre cave (Switzerland) and for the corresponding drip water. $\delta^D$ values of speleothem and the modern drip water are coherent and really close whereas $\delta^{18}O$ seems to be slightly fractionated.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$\delta^{18}O$ (%)</th>
<th>$\delta^D$ (%)</th>
<th>Released water (µL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milandre recent speleothem 2</td>
<td>$-7.7$</td>
<td>$-63.0$</td>
<td>$0.62$</td>
</tr>
<tr>
<td>Milandre recent speleothem 3</td>
<td>$-8.1$</td>
<td>$-61.8$</td>
<td>$0.67$</td>
</tr>
<tr>
<td>average</td>
<td>$-7.9$</td>
<td>$-62.4$</td>
<td></td>
</tr>
<tr>
<td>standard deviation</td>
<td>$0.3$</td>
<td>$0.9$</td>
<td></td>
</tr>
<tr>
<td>Drip water</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>average</td>
<td>$-8.9$</td>
<td>$-62.0$</td>
<td></td>
</tr>
<tr>
<td>standard deviation</td>
<td>$0.2$</td>
<td>$0.8$</td>
<td></td>
</tr>
</tbody>
</table>
Fig. 1. Diagram of the speleothem fluid inclusions water extraction line.
Fig. 2. Speleothem sample. (a) Conditioning of the sample before being cut with a wire saw. Growth laminae are well visible. (b) Speleothem thin sections with their laminae. Fluid inclusions are visible in both cross polarised (black) or bright field view (white). (c) Enlargement of single fluid inclusion.
Fig. 3. (a) Desorption test: the large peak corresponds to ambient water vapour migration during the opening of the line when loading the sample. After closing, the sample is flushed with a dry nitrogen stream without heating. After reaching the dry background level the system is heated to 140 °C to desorb the water from the calcite. (b) Example of desorption and stabilisation of water vapour amount and its isotopic composition after loading the sample. Desorption from the sample surface or from fluid inclusion cracks is hardly visible at this resolution.
Fig. 4. $\delta^{18}O$ and $\delta D$ reproducibility with changing water vapour amounts. (a) Water vapour mixing ratios were increased by changing the speed of the pump in steps of 0.2 % from 1 to 3 % of its capacity. (b) Evolution of the standard deviation of the water vapour isotopic measurement for $\delta D$ and $\delta^{18}O$ and water vapour amount as a function of the water vapour amount.
Fig. 5. $\delta^{18}$O and $\delta$D dependence on injected water vapour amounts presented in Fig.4.
Fig. 6. System response for switching from ST-08 to DOME-C standard water. No difference is noticeable for empty, loaded or crushed tube conditions.
Fig. 7. Water vapour evolution during a measurement sequence. Peaks 1, 3 and 5 are 0.5 µL injections of ST-08 standard water. Peak 4 corresponds to the released fluid inclusion water. The depression 2 corresponds to the exchange between the empty and the loaded copper tube.
Fig. 8. These spectrograms show the $\delta^{18}O$, $\delta D$ and $H_2O$ responses of the Picarro for a speleothem sample measurement (Fig. 7, peak 4). Both isotope ratios behave similarly. Due to diffusional fractionation of the light and heavy water isotopologues during the sample transfer by nitrogen, depleted values are observed for the very first part of the sample. Maximal isotope values are reached after the maximum of the water amount. The signals reach background stability again after about 20 min.
Fig. 9. The relationship between the sample signal and the injected water amount is used to calibrate the released water amount from fluid inclusions. Calibration was determined for amounts up to 2 µL using wet background and based on two schemes: (i) with the signal area converted into water amounts (linear regression in red) and (ii) relative to the maximal signal intensity (polynomial regression in blue).
Fig. 10. Schematic diagram illustrating the evaluation protocol (see text for details).
Fig. 11. Summary of the reproducibility tests made for different standard waters ranging from 0 to $-450 \, \permil$ for $\delta D$ (left panel) and from 0 to $-55 \, \permil$ for $\delta^{18}O$ (right panel).
Fig. 12. $\delta^D$ vs. $\delta^{18}O$ measurements with standard deviations based on three replicates for speleothem samples from Borneo. Results are close to the Global Meteoric Water Line (GMWL).