Continual in-situ monitoring of pore water stable isotopes in the subsurface

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Received: 13 September 2013 – Accepted: 18 September 2013 – Published: 5 November 2013

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Published by Copernicus Publications on behalf of the European Geosciences Union.
Abstract

The stable isotope signature of pore water provides an integral fingerprint of water origin, flow path, transport processes, and residence times and can thus serve as a powerful tracer of hydrological processes in the unsaturated and saturated zone. However, the full potential of stable isotopes to quantitatively characterize subsurface water dynamics is yet unfolded due to the difficulty in obtaining extensive detailed and continual measurements of spatiotemporally variable pore water signatures. With the development of field-deployable laser-based isotope analyzers, such measurements are now becoming feasible. This study presents the development and application of a functional, automatable, and cost-efficient system for non-destructive continual in-situ monitoring of pore water stable isotope signatures with high resolution. The monitoring system uses automatic-controllable valve arrays to continuously extract diluted soil air water vapor via a branching network of multiple small microporous probes into a commercial isotope analyzer. Soil temperature observations are used to convert obtained vapor phase into liquid phase water isotope signatures, but these can also be obtained based on vapor concentration measurements. In-situ sampling was conducted at six depths for each of three plots planted with varying vegetation on an experimental site in SW Germany. Two different methods based on advective and diffusive soil water vapor probing were employed suitable under unsaturated and all (including saturated) moisture conditions, respectively. The advective sampling method was applied using multiple permanently installed probes (continual mode) and using a single probe subsequently inserted to sample the various locations (push-in mode), while the diffusive sampling method was applied in push-in mode only. Using a specific identical treatment onsite calibration approach along with basic corrections for instrument bias and temperature dependent free water-vapor isotopic equilibrium fractionation, the monitoring system facilitated inference of normalized liquid pore water isotopic composition with sufficiently high accuracy and precision at sampling intervals of less than four minutes and resolved the isotopic variability along natural depth profiles. Comparison indicated
that the presented in-situ approaches may be used interchangeably with each other and with concurrent laboratory-based direct equilibration measurements of destructively collected samples, such that the choice of method will depend upon the task and anticipated conditions of sampling. The introduced sampling techniques provide powerful tools towards a detailed quantitative understanding of dynamic and heterogeneous shallow subsurface and vadose zone processes.

1 Introduction

Stable isotope ratios of hydrogen (\(^{2}\text{H}/^{1}\text{H}\)) and oxygen (\(^{18}\text{O}/^{16}\text{O}\)) provide powerful tracers of water in the hydrologic cycle (Gat, 1996). The usefulness of the stable isotopes of hydrogen and oxygen as tracers for hydrological processes lies in their (mostly) conservative nature as intrinsic part of the water molecule and the measurable natural variations of their ratios in precipitated and other natural waters imposed by large and variable fractionations during physicochemical processes and reactions such as, and most importantly, phase changes (Craig, 1961; Dansgaard, 1964; Friedman et al., 1964; Ehhalt and Knott, 1965; Majoube, 1971). The isotopically distinct nature of input waters to a system in space and time provides for a continuously and naturally imprinted signature that can be traced to reveal information on transit times, origin, flow pathways, and processes such as phase transitions, transport, and exchange.

As such, the analysis of stable water isotopes is being widely employed in hydrology and related disciplines and has brought major advances to our understanding of natural systems (see reviews by Ehleringer and Dawson, 1992; Kendall and McDonnell, 1998; Yakir and Sternberg, 2000; Vitvar et al., 2005; Aggarwal et al., 2007; West et al., 2010b; Soderberg et al., 2012). In particular, stable isotope signatures of pore water have provided insight into soil evaporation (Zimmermann et al., 1967; Allison, 1982; Allison et al., 1983; Barnes and Allison, 1988; Walker et al., 1988), recharge rates (Dincer et al., 1974; Saxena, 1984; Darling and Bath, 1988), unsaturated and saturated subsurface flow processes, mixing, and residence times (Sklash and Farvolden,
Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler

However, the full potential of stable water isotopes to quantitatively characterize water dynamics is yet to be unfolded. This holds particularly with regard to complex and heterogeneous processes, such as infiltration and shallow subsurface flow in the vadose zone subject to strong gradients and fluxes of energy and matter, demanding stable isotope data with high spatial and temporal resolution to complement traditional observations. Instead, a major limitation to the extent and scope of stable isotope applications was imposed by the available techniques for sampling and analysis (e.g. Kerstel and Meijer, 2005; Helliker and Noone, 2010; Munksgaard et al., 2011). Conventionally, measurement of water stable isotopic composition was relatively labor-, time-, and cost-intensive and constrained to laboratory-bound analysis of previously collected and processed discrete samples based on gas source isotope ratio mass spectrometry (IRMS) (Horita and Kendall, 2004). Pore water samples were most commonly obtained from destructive collection of geologic material and subsequent extraction using various methods (e.g. Buttle and Sami, 1990; Ingraham and Shadel, 1992; Walker et al., 1994; West et al., 2006) or, less frequently, using suction or gravity lysimeters (e.g. Stewart and McDonnell, 1991; Wels et al., 1991; Landon et al., 1999; Figueroa-Johnson et al., 2007), however, with considerable uncertainty and unclear implications of each method for the representativeness of obtained pore water isotopic signatures.

In recent years, major advances in analyzing techniques have come with the advent of commercially available, portable, and field-deployable laser spectroscopic instruments for isotopic analysis of water vapor (see reviews by Kerstel and Meijer, 2005; Kerstel and Gianfrani, 2008; Helliker and Noone, 2010). Available instruments based
Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler

Building upon the previous developments, this paper presents the development and application of a novel field system for continual unattended in-situ monitoring of liquid pore water stable isotopic composition by delivering diluted soil water vapor via small microporous soil water isotope probes (SWIPs) from an arbitrary number of subsurface sources into a commercial WS-CRDS instrument. Representing a trade-off between...
expenditure and universality in application, two different implementations of the monitoring system were investigated, which are henceforth referred to as the advection dilution sampling (ADS) method and the diffusion dilution sampling (DDS) method, according to the respective dominant vapor transport mechanism across the microporous probing tube (as in Munksgaard et al., 2011). The aim of this work was to develop a functional monitoring system suitable for investigating dynamic and heterogeneous shallow subsurface and vadose zone processes at the relevant temporal and spatial scales that facilitates accurate and precise in-situ measurements while meeting the following desirable attributes:

- inexpensive in terms of both acquisition and maintenance and expandable to a large number of probing locations as prerequisite to broad and locally dense application and thus, high spatial resolution;
- short sampling interval to facilitate frequent measurements and thus, high temporal resolution;
- unattended applicability under most conditions encountered in the subsurface throughout the day and year as a prerequisite for continuous application;
- compact (subsurface components), easy and quick to install imposing little disturbance and obstruction to the natural subsurface system and processes under investigation;
- mechanically robust and largely chemically inert components to facilitate safe insertion and duration in natural subsurface environments and long term weathering resistance.

2 Methods

In this section, the basic set-up, working principle, operation, and data processing of the monitoring system are first described, followed by the field evaluation approach.
Throughout this paper, the δ-notation (e.g. δ\(^{18}\)O or δ\(^2\)H) is used to express the isotopic composition of a sample in units of parts per thousand (‰), which, for a given element (e.g. oxygen or hydrogen), is written as

\[
\delta = \left( \frac{R_{\text{sample}}}{R_{\text{reference}}} - 1 \right) \cdot 1000 \text{‰} \tag{1}
\]

where \(R_{\text{sample}}\) and \(R_{\text{reference}}\) denote the atomic ratios of the rare to the most abundant isotope of the respective element (e.g. \(^{18}\)O/\(^{16}\)O or \(^2\)H/\(^1\)H) of the sample and of the employed international reference standard Vienna Standard Mean Ocean Water (VSMOW). Subscripts l and v are used to indicate reference to liquid and vapor phase water isotopic composition, respectively.

2.1 In-situ monitoring system

2.1.1 Monitoring system set-up

A schematic representation of the monitoring system set-up is provided in Fig. 1. All components were selected in compliance with the above stated desirable attributes. The system is built up modularly and can simply be modified or augmented to accommodate a certain number of probes with relatively little extra cost. A single set of central analyzing and supply devices is sufficient to operate a multitude of SWIP sampling locations via a branching tubing network and valve control system. A commercial laser spectroscopic instrument (Picarro L2120-i WS-CRDS, Santa Clara, CA, USA) is used, analyzing an air stream drawn through the attached main sample line by a downstream vacuum pump at a frequency of 0.5 Hz for δ\(^{18}\)O\(_v\), δ\(^2\)H\(_v\), and vapor concentration. Further, a nitrogen gas (N\(_2\)) supply installation is employed, comprising a regulated pressurized source to the main dilution line with a flow rate adjustable to a target value sequence imposed by a programmable digital mass flow controller (Analyt-MTC, Müllheim, Germany). In case of the diffusion dilution sampling (DDS)
method, the N\textsubscript{2} supply is split upstream of the first mass flow controller and a second mass flow controller is used to regulate the flow rate through the main throughflow line. The CRDS instrument and N\textsubscript{2} supply installations are housed in a customized trailer for field deployment. The trailer comprises a voltage transformer and high capacity accumulators to enable stable and gap-free operation using an electric generator power source in remote field applications and is vented to prevent instrument overheating.

Each of the main gas transport lines is branched in two steps to facilitate operation of a multitude of probes at varying locations with minimum total tubing length. The main gas transport lines are guided from the trailer to a central distribution box where radial multi-port connectors (Upchurch Scientific, Oak Harbor, WA, USA) are used to distribute the main lines towards local control boxes associated with a cluster of probes installed in the subsurface (e.g. at varying depths beneath a specific surface location). Arrays of two-way normally-closed electromechanically actuated solenoid valves (Clippard Minimatic, Cincinnati, OH, USA) mounted on manifold blocks (custom manufactured from polyvinylidene fluoride (PVDF) by Fischer Plastics, Gundelfingen, Germany) are used to further split and control the incoming gas transport lines towards individual probes. The valves are actuated either manually or automatically by a microcontroller switching unit. The valves controlling the ports receiving gas transport lines of a specific probe on each manifold are coupled and operated simultaneously.

All tubing used for the monitoring system gas transport lines consists of fluorinated ethylene propylene (FEP) and is protected by mesh hoses. All sample lines and all lines immediately attached to the probes have an outer diameter (OD) of 1/16 in and an inner diameter (ID) of 1 mm, while the remaining N\textsubscript{2} supply lines have an OD of 1/8 in and an ID of 1/16 in. The total effective length of each gas transport line (from each probe to each end device) depends on the study design and field situation and amounted to approx. 10 m in the present study. Outside the trailer, all tubing connections can quickly and tool less be finger tightened using removable and reusable fittings (Upchurch Scientific), while stainless steel fittings (Swagelok, Solon, OH, USA) are used inside the trailer.
Each soil water isotope probe (Fig. 2) is composed of three major custom parts connectable via threads. The core of the probe is a rigid hydrophobic microporous polyethylene (PE) probing tube (Porex Technologies, Aachen, Germany) with a length of 50 mm and an OD of 10 mm. The hydrophobic material allows water vapor contained in the soil air to pass while prohibiting liquid water penetration at sufficiently low differential pressures. The microporous tube is sealed on one end and connected on the other end to a central element manufactured from PVDF (Fischer Plastics). The central element contains a mixing chamber and accepts a line retainer with two (ADS; Fig. 2a) or three (DDS; Fig. 2b) gas transport lines attached. The central element can be connected to a rigid insertion shaft of variable length (here, 100 to 700 mm) made of anodized aluminum, guiding and protecting the gas transport lines in the subsurface. This design facilitates convenient probe installation from the soil surface with little disturbance of the subsurface environment. The probes are installed into pre-drilled holes using a steel tube with an OD of 10 mm, thus favoring immediate close contact of the subsequently inserted probes with and preventing compaction of the surrounding soil matrix.

The system can be used to sample the distribution of pore water stable isotopic composition in two different modes. On the one hand, a multitude of soil probes can be permanently installed in clusters to continually monitor, e.g., depth profiles of pore water stable isotopic composition below various surface locations (i.e. continual sampling mode). On the other hand, a single probe can be used to subsequently sample spatial pore water isotope distributions by repeated insertion at various locations and depths (i.e. push-in mode). Of course, continuous sampling using a single permanently installed probe is also possible; however, this will seldom present a relevant mode of field application.

A specific calibration approach following the principle of identical treatment is used to facilitate normalization of measured isotope signatures to the VSMOW reference scale. Therefore, four polyvinyl chloride (PVC) boxes of 20 cm × 30 cm × 30 cm were equipped with a centrally placed SWIP, closely neighbored by two 5TE probes and one
MPS-1 probe (both Decagon Devices, Pullman, WA, USA) sensing soil water content, temperature, electrical conductivity, and matric potential, and filled with oven-dried soil material from the upper 60 cm of the study site. Subsequently, water of known isotopic composition was added to generate a soil water content of approx. 20 vol% at the depth of the probes and the boxes were immediately sealed. The waters used included an isotopically light and a heavy laboratory standard, serving for normalization during calibration, and an intermediate laboratory standard used twice (i.e. for two boxes) and serving as quality control standard. The respective boxes are henceforth referred to as calibration and control standard boxes. During field sampling, three of the boxes were placed into a thermo-insulated housing to reduce temperature variations, while one control standard box was placed outside the housing, thus being subject to stronger temperature variations.

### 2.1.2 Monitoring system operation

An individual probe is operated by simultaneously opening the associated valve on each of the manifolds, while all other valves are closed. Once the valves are open, sample air is drawn through the sample line at a rate of 30–35 mL min\(^{-1}\) imposed by the vacuum pump of the CRDS instrument. At the same time, \(N_2\) is supplied via the dilution line into the mixing chamber with a controllable rate. If the \(N_2\) supply rate is lower than the sample intake rate, soil air saturated with water vapor at the local temperature advects through the walls of the microporous probing tube in response to the applied absolute pressure gradient. The soil air flows into the mixing chamber, where it is diluted to a lower vapor concentration as determined by the ratio of \(N_2\) supply rate to sample intake rate. This diluted vapor sample stream enters the sample line and flows to the CRDS instrument (ADS method). By varying the \(N_2\) supply rate, the vapor mixing ratio can be controlled from close to zero to saturation at soil source temperature. The primary purpose of the vapor dilution is to preclude any condensation of water due to varying temperature along the sample line while avoiding any sample line heating system.
In case of the DDS method, an additional third gas transport line (throughflow line) supplies N\textsubscript{2} to the bottom of the probe at a rate controlled by a second mass flow controller to equal the difference between sample intake rate and dilution rate, such that no absolute pressure gradients result across the microporous wall. Instead, driven by a partial pressure gradient, vapor diffuses into the probing tube to saturate the through flowing N\textsubscript{2} carrier gas. As for the ADS method, the sample stream is transported to the analyzer and diluted in the process. While this method raises the expenses for monitoring system acquisition and operation, it provides the advantage that it can be applied under saturated conditions as long as the water entry pressure of the employed microporous material is not exceeded.

The monitoring system allows for manual or automated sampling using programmed probe and dilution rate sequences. In a typical probe sequence, all SWIPs are subsequently sampled, moving from one cluster to the next, occasionally intermitted by sampling the standard boxes. A given location is typically probed using a dilution rate sequence starting with a short flushing phase with high dilution, thus clearing the mixing chamber, sample lines, and cavity (i.e. the entire path for the new sample vapor) from vapor of previous samples. This is followed by a sampling phase with lower dilution. In the present application, each in-situ measurement was conducted using a sequence of 60 s flushing and 150 s sampling. N\textsubscript{2} flow rates of 36 and 18 mL min\textsuperscript{-1} were used during flushing and sampling, respectively. The former slightly exceeds the nominal sample intake rate, while the latter presents the minimal dilution rate necessary to avoid condensation during > 95 % of the time of a day and year in the region of application, as computed from multi-year temperature records.

### 2.1.3 Signal processing

Processing of the acquired time series signals is fully automated into a MATLAB\textsuperscript{®} (The MathWorks, Inc., Natick, MA, USA) toolbox. Standard processing for the CRDS-based measurements includes derivation of raw sample values from the observed high-frequency signals, subsequent corrections to account for instrument-specific vapor...
concentration biases, drift, and deviations from the internationally accepted reference scale (e.g. Schmidt et al., 2010; Sturm and Knohl, 2010; Gröning, 2011; Aemisegger et al., 2012), as well as conversion of the corrected vapor to respective liquid isotopic compositions.

Inspecting the acquired time series of water vapor concentration (Fig. 3a), individual sample phases can be easily identified as step input responses with a vapor concentration asymptotically approaching a level depending on the soil temperature and the imposed dilution rate. The asymptotic behavior can be primarily attributed to advective-diffusive-reactive transport through the soil pore space and the various segments of the probing system as well as storage effects such as cavity reservoir gas exchange. The preceding flush phase is characterized by an initial peak followed by declining vapor concentration, emptying the sample line for the next sample intake. Note also that a short time lag (approx. 20 s) exists between observed responses and switching of valves and dilution rates due to the bulk transport through the sample lines.

To determine the vapor concentration and raw isotopic composition of each sample, the acquired time series of vapor concentration is first scanned to identify characteristic times based on differentiation of the signal (i.e., using a multi-point moving slope to reduce the impact of noise). These include the defined start time of sample arrival, start time of asymptotic behavior, and the end time of sample arrival, identified as the stationary point, first inflection point (from convex to concave downwards), and second inflection point (from concave to convex downwards) following a declining flush phase, respectively. Second, sample values are obtained as the steady-state solution of simple exponential function least-squares-fits to the asymptotic period of sample arrival. Using the root mean squared error (RMSE) as quality criterion, it was found that exponential models of type

\[ \delta(t) = \delta_\infty + (\delta_0 - \delta_\infty) e^{-t/\tau} \]  

(2)

where \( t \) indicates the time, \( \tau \) is a time constant, and subscripts 0 and \( \infty \) indicate the initial and steady-state solution, respectively, fit well (RMSE < 0.4‰ for \( \delta^{18}O_v \))
and < 3.3 ‰ for $\delta^2$H$_v$) to the observed $\delta$-value response signal. Further, exponential models of type

$$c(t) = c_\infty + (c_0 - c_\infty) e^{-t/\tau_1} + \lambda e^{-t/\tau_2}$$

(3)

where $\tau_1$ and $\tau_2$ are time constants and $\lambda$ is a parameter, fit well (RMSE < 80 ppmv) to the observed vapor concentration $c$ response signal. Note that device related corrections (e.g. vapor concentration dependency) may be applied prior to fitting a model in case they are non-linear. The 95% confidence interval (CI) of the exponential model fits is computed based on the estimated coefficient covariance matrix (on average 0.19 ‰ for $\delta^{18}$O$_v$, 1.5 ‰ for $\delta^2$H$_v$ and 11 ppmv for $c$) and illustrates the uncertainty of sample value estimation from the signal (Fig. 3). While some average value over a specified segment of the asymptotic period could actually be used without significant deterioration of the data quality, the asymptotic model makes better use of the data, can reduce the impact of noise and increase the precision, is independent of the averaging interval, less dependent on the averaging period, and provides for a statistically more valid approach. These aspects are of particular relevance when short sampling times are targeted. At the same time, we omit fitting a more complex model to the entire response function at this point, partially to avoid the impact of noisy observations at low vapor concentration.

The obtained raw sample values of $\delta^{18}$O$_v$ and $\delta^2$H$_v$ are then corrected for analyzer specific vapor concentration dependent bias relative to an arbitrary reference concentration of $10^4$ ppmv. Least-squares-fitted relations were derived prior to the field sampling based on vaporized discrete liquid water laboratory standard aliquots of three isotopic compositions and varying volume injected into a vaporizer using a liquid auto sampler and N$_2$ as a carrier gas. The sample liquid water isotopic compositions $\delta^{18}$O$_l$ and $\delta^2$H$_l$ associated with each $\delta^{18}$O$_v$ and $\delta^2$H$_v$ value is then estimated using soil temperatures measured at the respective depth of vapor extraction along with the equations for free water liquid-vapor equilibrium fractionation established by Majoube (1971). Finally, a combined calibration and drift correction is applied for normalization.
of $\delta^{18}O_l$ and $\delta^2H_l$ values to the VSMOW reference scale and elimination of temporal biases. Therefore, calibration standard measurements obtained approximately every two to three hours and processed identically to in soil measurements up to this point are linearly interpolated to the time of each other sample to conduct a linear two point calibration (e.g. Gröning, 2011).

2.2 Field application and evaluation

2.2.1 Site description and instrumentation

The in-situ monitoring system was tested under field conditions and evaluated against results from concurrent destructive sampling. Measurements were conducted during one day in September 2012 on an experimental site located at 310 m a.s.l. on a vineyard terrace on the eastern flank of the Kaiserstuhl mountain range (48°05′33.48” N; 7°42′24.06” E) in southwest Germany. The mean annual temperature is 11 °C and the mean annual precipitation amounts to 900 mm. A local meteoric water line (LMWL) was obtained based on weekly bulk samples of precipitation collected on the site from June 2011 to August 2012. The permeable slightly clayey silt soils (Luvisol) of the study site developed from a thick layer of Pleistocene aeolian loess covering the Tertiary volcanic rock forming the Kaiserstuhl (Hädrich and Stahr, 2001). Due to the low slope (< 5 %) and deep permeable soils, water fluxes are predominantly vertical.

Plots of size 50 cm × 50 cm were instrumented and planted with various species in 2011 as part of ongoing research. 5TE and MPS-1 probes installed at depths of 5, 10, 20, . . . , 60 cm deliver data at 10 min intervals. Three plots planted with grass, beech, and no vegetation (i.e., bare soil) were selected for this study. The plots had been sheltered from precipitation approximately two weeks prior to sampling such that quasi-static conditions in terms of pore water content and isotopic composition may be assumed.
2.2.2 Destructive pore water isotope sampling

For each plot, three soil cores were retrieved using a Pürckhauer drill. The cores were immediately partitioned into samples representing depth intervals of 0–5, 5–15, . . . , 75–85 cm and stored into gas-impermeable bags (Weber Packaging, Güglingen, Germany). The bags were subsequently filled with N₂, heat-sealed, and analyzed in the laboratory using the direct equilibration method as detailed by Wassenaar et al. (2008) and Garvelmann et al. (2012). The headspace water vapor isotopic composition of each sample was probed for 390 s at constant temperature using the same WS-CRDS analyzer as above. Raw sample δ¹⁸Oᵥ and δ²Hᵥ values are obtained as the arithmetic mean of the last 90 s of the period of sample arrival and consequently normalized, drift corrected, and converted to δ¹⁸O𝑙 and δ²H𝑙 values as described for the in-situ sampling. Following a similar identical treatment approach, calibration and quality control were conducted based on reference samples prepared from oven-dried soil material from the experimental site mixed with three laboratory water standards. The reference samples were prepared to have the same dry mass as the average soil samples and a water content of 20 vol%, otherwise treated identically to the soil core samples, and alternatingly analyzed between groups of three soil core samples.

2.2.3 In-situ pore water isotope sampling and evaluation

Three in-situ sampling approaches were selected, each to obtain two pore water isotope profiles for each plot with discrete measurements at depths of 10, 20, . . . , 60 cm. In-situ measurements were conducted and processed as described above and using (i) a subset of 18 ADS probes permanently installed at the respective locations and depths in July 2012 as part of ongoing research, (ii) an ADS probe in push-in mode (i.e., subsequently inserted at the respective locations and depths), and (iii) a DDS probe in push-in mode.

The methodological accuracy of and the agreement between processed in-situ and destructive laboratory-based isotope measurements are assessed in terms of the
differences to known control standard values and to each other (i.e. evaluating all measurements from equal depths and similar vegetation cover against each other), respectively. Summary statistics provided include the respective root mean squared differences (RMSE), mean absolute differences (MAE), mean differences (BIAS), and 95% limits of agreement (LOA) estimated as the BIAS ± 1.96-fold standard deviation (SD) of the differences (Bland and Altman, 1986). For each method, the methodological precision and the reproducibility (or variability) of soil profile measurements are assessed in terms of the (pooled) SD of processed control standard measurements and repeated soil measurements at equal depths, respectively. Control standards placed inside and outside a thermo-insulated housing are evaluated separately to assess the impact of larger temperature and associated vapor concentration variations and differences to calibration conditions. In addition to data obtained during the day of profile sampling, a multi-day record of control standard measurements obtained as part of ongoing research is used to increase the sample size.

3 Results

3.1 Evaluation of sampling method

The monitoring system facilitates rapid signal stabilization enabling high-quality sample value estimates at short sampling intervals for the employed dilution sequence and under the field conditions encountered (Table 1; Fig. 3). The time required to reach a stable signal depends upon a variety of operational and environmental factors, such as dilution sequence and rates or magnitude of isotopic differences between successive samples. While a systematic assessment is beyond the scope of this paper and sample value estimation was based on exponential model fitting, it is noted that time constants for exponential fits to the asymptotic periods of the acquired signals, starting approx. $18 \pm 4 \text{s (arithmetic mean \pm SD)}$ after first sample arrival and with a bulk delay time of approx. $20 \text{s}$ after initiation of a sample phase, were similar for all in-situ
sampling approaches and averaged to approx. $31 \pm 12$ s for $\delta^{18}O_v$ and $34 \pm 13$ s for $\delta^2H_v$. This given, the time required for the signal to approach a steady value to within a range of e.g. $\pm 0.1 \, ^\circ$ for $\delta^{18}O_v$ and $\pm 0.5 \, ^\circ$ for $\delta^2H_v$, which corresponds to half the analyzer precision specification (Picarro, 2012), can be estimated to amount $108 \pm 45$ s for $\delta^{18}O_v$ and $128 \pm 48$ s for $\delta^2H_v$. This indicates that short sampling intervals are feasible also without employing an asymptotic model.

Evaluation of control standard measurements indicates high accuracy and precision for both the ADS and the DDS method over multiple days of field operation. Summary statistics (Table 1) show that systematic errors were small, while random errors were on the order of the precision specified for the employed CRDS instrument (Picarro, 2012) and approximately twice to thrice as high as that currently attainable with IRMS instrumentation under laboratory conditions (Brand, 2004; Horita and Kendall, 2004), however, excluding effects related to sample collection and preparation in case of the latter which often compromise the overall accuracy (Kerstel and Gianfrani, 2008). Thereby, larger temperature variations and deviations from calibration conditions (approx. $-3$ to $+4 \, ^\circ$C) did not significantly impact the control standard measurement error (two-sample $t$ test, $p < 0.001$). Further, the quality of in-situ measurements approached that attained for laboratory-based measurements conducted using the direct equilibration technique under constant temperature conditions and with higher vapor concentrations. Note that in case of the laboratory-based measurements, some variability may be associated with the preparation of individual reference samples for calibration and control of soil core sample measurements in contrast to repeated in-situ measurements of identical calibration and control standard boxes.

### 3.2 Evaluation of soil profile measurements

Depth profiles of pore water isotopic composition obtained based on the various in-situ sampling approaches and destructive sampling are physically plausible and generally in good agreement for all three plots under investigation (Fig. 4). This holds with respect
to all, the range, distribution, and magnitude of observed isotope signatures and under the conditions encountered, with observed soil water contents and matric potentials ranging from approx. 6 to 24 vol% and −270 to −27 kPa, respectively.

Repeated in soil measurements obtained for installed identical ADS probes coincide closely (SD 0.22 ‰ for δ^{18}O_l and 1.6 ‰ for δ^{2}H_l), exhibiting a reproducibility approaching that obtained for control standard measurements and indicating the value of continual in-situ multi-probe sampling. Stronger inter-profile variability exceeding the respective estimated methodological precision is observed for push-in profile sampling using the ADS (SD 0.38 ‰ for δ^{18}O_l and 2.6 ‰ for δ^{2}H_l) or DDS probes (SD 0.37 ‰ for δ^{18}O_l and 2.8 ‰ for δ^{2}H_l) and for destructive profile sampling (SD 0.34 ‰ for δ^{18}O_l and 2.4 ‰ for δ^{2}H_l). However, this can primarily be attributed to varying target soil water isotopic composition encountered at differing sampling locations.

Comparing in-situ and destructive profile sampling results, observed differences are overall similar for all in-situ sampling approaches (Table 1). Comparable agreement with destructive sampling results is obtained for spatially more proximate push-in mode sampling using either method compared to continual mode sampling, supporting the assumption of sampling distance as source of intra-method variability. Similar results could be observed when comparing in-situ sampling results obtained from different methods and modes of operation with each other (Fig. 4). In all cases, random deviations appear dominant, while biases are small. The inter-method SD and RMSE are on the order of the estimated inter-profile variability for the non-continual sampling methods, indicating that intra-method variability is the major source of inter-method disagreement. Clearly, additional sources of error and uncertainty including spatial variation of targeted pore water isotopic composition along with lack of spatial coincidence of sampling location, imprecise depth assignment, and compression during destructive sampling add to the methodological differences and errors. However, no major bias or obvious systematic dependency can be observed regarding the agreement and relative variability along the soil profiles and thus, with varying soil properties or states and gradients of soil water content, matric potential, and temperature. This indicates that
any such effect, over the range of field conditions encountered and if present, does not substantially exceed the reported level of agreement or reproducibility and may thus be deemed negligible, and is at least similar for both sampling techniques, such that the methods may be used interchangeably. Further, the spatial variations in isotopic composition obtained from destructive sampling at the measurement scale of 10 cm are well reproduced by the in-situ sampling methods, indicating that in fact a localized isotopic equilibrium between the pore water phases existed and was represented in the sampling results.

The majority of samples plot closely along the global (Craig, 1961) and local meteoric water lines (GMWL and LMWL, respectively; Fig. 5), indicating coherence of obtained \( \delta^{18}O \) and \( \delta^2H \) values assuming a mostly unaltered meteoric origin of waters. Only few samples, obtained in close proximity to the surface, show potentially evaporative enrichment effects. While a more complex model based interpretation of vapor data is necessary to obtain liquid water estimates under non-equilibrium conditions of actual evaporation, the fact that all in-situ samples displayed approximately water vapor saturation within measurement precision of spatiotemporally local temperature (i.e. based on either Kelvin or Magnus Equation and taking dilution rate into account) and agree well with headspace samples from soil cores may indicate that no additional net evaporation was occurring at the time of in-situ sampling and further, that the effects are not related to surface proximity of in-situ sampling.

While a physical interpretation of the measured isotope profiles is beyond the scope of this paper, it can be noted that the last major summer rainfall reaching the ground surface had a composition of approx. \(-3.5\%\) \( \delta^{18}O \) and \(-22\%\) \( \delta^2H \), while typical winter rains are at \(-9.5\%\) \( \delta^{18}O \) and \(-66\%\) \( \delta^2H \), and thus, the obtained results may be deemed physically plausible and reflect the common perception of an attenuated seasonal meteoric variation subject to losses by transpiration and some evaporative enrichment in the top-soil (Dansgaard, 1964; Barnes and Allison, 1988; Garvelmann et al., 2012).
4 Discussion

4.1 Monitoring system implementation and functionality

The presented system was developed to allow for unattended functional and efficient minimal invasive field monitoring of pore water stable isotope distributions with high sampling rate and limited cost. The monitoring system is set-up modularly and can be easily and flexibly installed, adapted, and extended for accommodation of a multitude of probes and changing applications with relatively little extra expenses. The rigid tubular design and small dimensions of the probes (diameter of 10 mm) allow for simple installation causing little disturbance and obstruction to the subsurface flow field and processes under investigation as well as for localized vapor extraction. All parts and materials were selected to be robust and weathering resistant, allowing the system to persist in the subsurface for continuous or repeated application. Additionally, all parts that come into contact with the sample air, most importantly the FEP sample lines, have low dead volume, are chemically mostly inert, and have low water sorption, porosity, and gas permeability (e.g. Mark, 2009).

Exceeding the capabilities of a sample line heating approach, the in-situ dilution technique provides optimal control over sample mixing ratios from the subsurface sampling location to the analyzer cavity. Adequate dilution sequences allow for an efficient purging of the sample lines and clear sample detection, prevent condensation and associated fractionation effects at any point of the system, and potentially reduce the effects of sorption and storage in the various compartments of the monitoring system. Along with the low pumping rate, the physicochemical and isotopic disturbance of the natural soil water phase system in the probing environment (i.e., due to imposed absolute or partial pressure gradients and soil air losses) is reduced during actual and prolonged sampling. In this context, a rough impression can be gained by noting that the volume of liquid water extracted per sample amounts to approx. 0.4 mL (assuming 20 °C and 1013 hPa), and this can be estimated to modify the bulk water content in an isolated soil volume surrounding the probe to an arbitrary distance of 1 cm by $10^{-3}$ %vol.
or, at a low water content of e.g. 5 vol%, by 0.02 %. Thus, approx. 50 repeated local measurements could be conducted before a modification of even a dry moisture state by 1 % would be caused.

As stated above, all in-situ samples displayed approximate water vapor saturation, independent of local soil physical states or gradients. Further, it is noted that clear optima of agreement between sample vapor concentrations and saturation vapor pressures computed based on temperature measurements for various depths and times could be observed when considering the spatially and temporally closest measurement. Also, neither repeated nor prolonged extraction (i.e. > 30 min) moved the liquid-vapor system from chemical equilibrium or caused continuous modification of the observed isotopic composition. This is consistent with recent laboratory results from Rothfuss et al. (2013). Importantly, these observations also imply that actual measurement of soil temperatures is not necessarily required to derive isotopic fractionation factors for application of the monitoring system to obtain liquid pore water isotopic compositions as these may be inferred based on observed vapor concentrations, provided that their measurements are accurate (see e.g. Aemisegger et al., 2012), equilibrium conditions prevail, and effects of water potential are negligible or can be estimated. For the present application, virtually identical estimates of $\delta^{18}O_l$ and $\delta^2H_l$ could be obtained using measured and vapor-based estimates of fractionation factors (data not shown).

### 4.2 Monitoring system performance

The presented technical and operational implementation is essential in practical field applications with continual accurate, precise, and rapid sampling for at least most of the year and for different climatic regions. The presented and evaluated excerpt of a field application employing a branching network of actually more than 30 SWIPs demonstrates these capabilities for the conditions encountered. A quick stabilization of the sample signal is attained, facilitating sampling intervals of approximately four minutes. Employing a specific identical treatment approach for calibration and drift correction, along with a vapor concentration bias correction and the well established formulations
for free water liquid-vapor equilibrium fractionation by Majoube (1971), accurate and reproducible inference of normalized liquid pore water isotopic composition from vapor analysis was possible without specific prior characterization of additional sampling effects. Thereby, the two in-situ methods tested (ADS and DDS) and the two modes of employment (continual and push-in) were shown to provide similar overall results in good agreement with laboratory-based direct equilibration measurements of destructively collected samples. The comparison of in-situ and destructive sampling results, the physical plausibility and reproducibility of the measurements, and sample vapor concentrations indicate that, for the level of data quality given, the extracted vapor did in fact physicochemically and isotopically represent the spatiotemporally local liquid pore water.

The obtained precision of liquid water $\delta$ value estimates is deemed sufficient for the anticipated applications. Improvements may come from further enhanced technical and operational implementation and calibration and correction approaches. Longer sampling and integration times and higher vapor concentrations (i.e. lower dilution) may often reduce sampling errors and uncertainties due to proceeded signal stabilization, reduced impact of background short-term noise, and instrument precision dependencies on vapor concentration (e.g. Sturm and Knohl, 2010; Aemisegger et al., 2012), while a higher pumping rate could accelerate signal stabilization. However, the former and both must be traded off for lower sample generation rate and increased actual and long term disturbance of the natural system, respectively. Further, it is important to note in this context that slow signal stabilization would impede proper signal attainment when the temporal scale of changes in liquid source water or of other effects on the vapor isotopic composition (e.g. temperature changes, instrument drift) is exceeded, which also limits the maximum integration time.

### 4.3 Assumptions and limitations

The advantage of the employed calibration approach is that any possible present constant or temporally variable sampling effects (e.g., due to fractionation, background
Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler

noise or drift) independent of variable outer conditions diverging from the reference conditions can be assumed to be removed or reflected in the estimated methodological precision. If necessary, systematic effects (e.g. due to variations of vapor concentration) equally impacting control standard and soil probing could also be corrected based on assimilated data. In principle, a range of environmental and sampling related characteristics and processes could have a potential impact on the actual and the observed gas phase isotope signatures and their relation to the local target liquid phase isotope signature in space and time. For example, physicochemical states such as chemical potential or activity of water can have specific effects on liquid-vapor equilibrium fractionation factors and equilibration times. In soils, such effects could arise due to changes in osmotic potential (see e.g. Sofer and Gat, 1972; Horita, 1989; Horita and Wesolowski, 1994) or matric potential, while experimental results of Allison et al. (1987) and Rothfuss et al. (2013) did not show observable effects. Natural and sampling induced spatiotemporal gradients causing water and vapor movement, net phase changes, and isotopic exchange may cause temporal non-equilibrium between local pore water and vapor and will require specific interpretation and modeling (see Soderberg et al., 2012, and references therein). Vapor extraction related evaporative surface cooling may need to be accounted for (Cappa et al., 2003). Organics concentrations can cause spectroscopic interference (West et al., 2010a), but can be corrected for (Schultz et al., 2011; West et al., 2011). Specific fractionation effects can also be caused for a particular porous probing material and vapor extraction technique. Initial headspace experiments comparing isotopic compositions of water vapor sampled directly and via SWIPs into the CRDS instrument suggested that isotopic fractionation associated with transport across the microporous probe wall is negligible (data not shown). However, such may arise under saturated conditions (Herbstritt et al., 2012) and depending upon concentration gradients across the microporous walls, vapor extraction rate, as well as permeability and geometry of the probing tube. In addition and as for previous sampling approaches (e.g. Landon et al., 1999; Figueroa-Johnson et al., 2007; Wassenaar et al., 2008), the information revealed about physical transport processes may be blurred by
unconsidered alteration of liquid soil water isotopic composition itself due to interactions with inorganic or organic soil constituents, or unclear contribution of water stored in different compartments of the pore space. Finally, the actual resolvability of dynamic liquid pore water isotopic composition independent of the monitoring system may be limited depending upon, e.g., the properties of geologic material (Hsieh et al., 1998b; Wassenaar et al., 2008), physicochemical states such as temperature (Ingraham and Criss, 1998), and soil water content controlled liquid surface area to volume ratio (Ingraham and Criss, 1993; Costanza-Robinson and Brusseau, 2002), determining the relevant rates of isotopic exchange between liquid and vapor phase.

The above incomplete enumeration of potential effects that may interact and vary depending upon the specific soil properties (e.g. material, texture, and pore size distribution) clarifies that further research into both, the underlying physical and sampling specific effects, is needed to attain complete confidence in the physical representativeness of results obtained using the presented or any other, particularly vapor-based pore water isotope sampling approach. While we did not observe obvious systematic impediment to multi-day measurements of control standards or deviations from methodologically similar direct equilibration measurements of soil core samples, such cannot be generally precluded but may be anticipated to emerge when more extreme and variable conditions in space and time are encountered and the level of “identity” of natural and reference conditions present for this study cannot be practically attained. As such, the presented monitoring system will certainly profit from continued laboratory experimentation and broad field application.

4.4 Outlook

Nevertheless, while subject to its own peculiarities, the presented sampling techniques drastically decrease the expenditure of time, labor, and money for pore water stable isotope monitoring while eliminating a variety of error and uncertainty sources associated with sample collection, transport, storage, and preparation. The ability of minimal-invasive repeated sample generation at a multitude of locations and at an
unprecedented rate extends the scope of stable isotope analysis for manifold applications such as investigations of vertical and lateral subsurface flow and transport processes or plant soil water interactions. According to the premise that an experimental design should provide the maximum information for the minimum expenditure, the choice of method should depend upon the processes and anticipated conditions at the site under investigation. While application of the DDS probes is more expensive in terms of acquisition and maintenance (i.e. requiring an additional carrier gas supply line including mass flow controller, valve system, tubing, and higher gas consumption), such will be necessary when saturated conditions could be encountered. In contrast, the push-in application strongly reduces acquisition cost (i.e. a single probe with only two or three valves can be used for a multitude of measurements) and will be particularly useful when spatial variations of soil water isotopic composition are of primary interest, while continuous investigation of highly dynamic processes will require permanently installed probes, and a combination of both applications with each other and/or destructive sampling can provide valuable spatiotemporal information.

5 Conclusions

This study presented the development and application of a novel system for unattended functional and efficient minimal invasive field monitoring of pore water stable isotope distributions with high sampling rate and limited cost. Along with a specific identical treatment onsite calibration approach and basic corrections for instrument bias and temperature dependent free water-vapor isotopic equilibrium fractionation, the technical and operational implementation of the system allowed for inference of normalized liquid pore water isotopic composition with sufficiently high accuracy and precision at sampling intervals of less than four minutes and resolved isotopic variability along natural depth profiles at the measurement scale of 10 cm under the conditions encountered. Comparison indicated that the presented in-situ approaches may be used interchangeably with each other and with laboratory-based direct equilibration measurements of
destructively collected samples, such that the choice of method will depend upon the task and anticipated conditions of sampling. While further research is needed to enhance the reliability and universal applicability of the monitoring system, the demonstrated capabilities of continual sample generation at a multitude of locations and at an unprecedented rate with minimal disturbance of natural processes can open new avenues towards a detailed quantitative understanding of subsurface water dynamics and fluxes in and out of the soil such as infiltration, recharge, evapotranspiration, and runoff.

Acknowledgements. This work is part of the project “Coupled soil-plant water dynamics – Environmental drivers and species effects” funded by the German Research Foundation (DFG). The authors gratefully thank the technicians, student assistants, and other helpers during preparation and realization of the field experiments, especially Emil Blattmann, Nicolai Dietermann, Benjamin Eisele, Barbara Herbstritt, Lara Kirn, and Lukas Neuhaus.

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Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler


Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler


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Continual in-situ monitoring of pore water stable isotopes

T. H. M. Volkmann and M. Weiler


Picarro L2120-i δD/δ¹⁸O isotopic water analyzer: http://www.picarro.com/sites/default/files/L2120-i%20Datasheet.pdf (last access: 10 April 2013), 2012.


Table 1. Summary statistics of in-situ and destructive laboratory-based pore water stable isotope measurements of quality control standards with and without thermo-insulation (TI) and soil profiles.

<table>
<thead>
<tr>
<th></th>
<th>ADS Control w/TI</th>
<th>ADS Control w/o TI</th>
<th>ADS Soil Installed</th>
<th>ADS Soil Push-in</th>
<th>DDS Control w/TI</th>
<th>DDS Control w/o TI</th>
<th>DDS Soil Push-in</th>
<th>Destructive Control</th>
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<tr>
<td>$\delta^{18}$O$_i$ (%)</td>
<td>0.16 0.17</td>
<td>0.38 0.43</td>
<td>0.15 0.18</td>
<td>0.42 0.15</td>
<td>0.15 0.18</td>
<td>0.42 0.15</td>
<td>0.15 0.11</td>
<td>0.15 0.11</td>
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<td>RMSE</td>
<td>0.13 0.14</td>
<td>0.31 0.35</td>
<td>0.12 0.14</td>
<td>0.34 0.11</td>
<td>0.12 0.14</td>
<td>0.34 0.11</td>
<td>0.15 0.11</td>
<td>0.15 0.11</td>
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<tr>
<td>MAE</td>
<td>-0.02 0.02</td>
<td>0.04 -0.02</td>
<td>-0.02 0.03</td>
<td>-0.04 -0.01</td>
<td>-0.02 0.03</td>
<td>-0.04 -0.01</td>
<td>-0.01 -0.01</td>
<td>-0.01 -0.01</td>
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<tr>
<td>BIAS</td>
<td>0.16 0.17</td>
<td>0.37 0.43</td>
<td>0.15 0.18</td>
<td>0.42 0.15</td>
<td>0.15 0.18</td>
<td>0.42 0.15</td>
<td>0.15 0.11</td>
<td>0.15 0.11</td>
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<tr>
<td>SD</td>
<td>0.29 0.35</td>
<td>0.77 0.82</td>
<td>0.27 0.38</td>
<td>0.78 0.28</td>
<td>0.27 0.38</td>
<td>0.78 0.28</td>
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<td>LOA Upper</td>
<td>-0.33 -0.31</td>
<td>-0.69 -0.86</td>
<td>-0.31 -0.32</td>
<td>-0.86 -0.30</td>
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<td>-0.86 -0.30</td>
<td>0.28</td>
<td>0.28 0.28</td>
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<tr>
<td>LOA Lower</td>
<td>1.23 1.15</td>
<td>2.87 2.92</td>
<td>1.32 1.20</td>
<td>3.09 1.03</td>
<td>1.32 1.20</td>
<td>3.09 1.03</td>
<td>1.03</td>
<td>1.03 1.03</td>
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<tr>
<td>MAE</td>
<td>1.05 0.98</td>
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<td>1.10 1.02</td>
<td>2.53 0.86</td>
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<tr>
<td>BIAS</td>
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<td>0.32 -0.20</td>
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<tr>
<td>SD</td>
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<td>2.85 2.91</td>
<td>1.32 1.20</td>
<td>3.06 1.03</td>
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<tr>
<td>LOA Upper</td>
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<td>5.91 5.50</td>
<td>2.42 2.47</td>
<td>5.63 1.92</td>
<td>2.42 2.47</td>
<td>5.63 1.92</td>
<td>1.92</td>
<td>1.92 1.92</td>
</tr>
<tr>
<td>$\delta^2$H$_i$ (%)</td>
<td>13/21 11/23</td>
<td>14/19 14/19</td>
<td>14/19 12/22</td>
<td>14/19 21/22</td>
<td>14/19 12/22</td>
<td>14/19 21/22</td>
<td>21/22</td>
<td>21/22</td>
</tr>
<tr>
<td>$c_b$ (10$^3$ ppmv)</td>
<td>7/13 6/14</td>
<td>7/10 7/10</td>
<td>7/10 6/13</td>
<td>7/10 26/27</td>
<td>7/10 6/13</td>
<td>7/10 26/27</td>
<td>26/27</td>
<td>26/27</td>
</tr>
<tr>
<td>$n_c$</td>
<td>18 18</td>
<td>36 36</td>
<td>9 9</td>
<td>36 9</td>
<td>9 9</td>
<td>36 9</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a Temperature  

b Vapor concentration  

c Sample size
Fig. 1. Schematic representation of the in-situ pore water stable isotope monitoring system set-up according to the DDS method.
Fig. 2. Schematic three-dimensional representation of the in-situ soil water isotope probes according to (a) the ADS and (b) the DDS method. Labels indicate (1) microporous tube, (2) central element, (3) mixing chamber, (4) line retainer, (5) sample line, (6) dilution line, (7) through-flow line, and (8) insertion and protection shaft.
Fig. 3. Exemplary time series signal representing the acquisition of a pore water stable isotope profile using the in-situ sampling system with permanently installed ADS probes and a dilution sequence of 60 s flushing and 210 s sampling. Solid vertical lines enclose periods of sample arrival. Exponential fits use data from the asymptotic period up to 150 s after start of sampling, as enclosed by the dashed vertical lines. A moving average (15 s backward) is provided for illustrative purposes.
Fig. 4. Depth profiles of pore water $\delta^2$H and $\delta^{18}$O for a bare soil (a, d), grass (b, e) and beach (c, f) plot obtained based on in-situ (vertically shifted for visibility) and destructive sampling. The methodological accuracy associated with each symbol is indicated in terms of 95 % limits of agreement.
Fig. 5. Plot of pore water $\delta^2H$ vs. $\delta^{18}O$ obtained based on in-situ and destructive sampling of soil profiles (depths from 10 to 60 cm). The methodological accuracy associated with each symbol is indicated in terms of 95% limits of agreement.