Responses to Reviewers – Point-by-point response to reviewer comments on "Technical Note: Evaluation of a low-cost evaporation protection method for portable water samplers" by Jana von Freyberg et al.

We would like to thank Nils Michelsen and one anonymous referee for reviewing our manuscript and for providing helpful comments. The point-by-point reply to the comments is given below. The comments provided by the reviewers are shown in regular font, and our responses in blue and bold.

1) Authors' response to the interactive comment by Nils Michelsen (Referee)

Dear colleagues,

With great interest, I have read your manuscript on the design of an evaporation reduction method for automatic water samplers, facilitating their use in isotope hydrology studies. Given the popularity of these samplers, the topic is, in my opinion, relevant for the HESS community. The simple concept is described in detail and since the required parts are low-cost and readily available, the method is easy to apply (if the reader has access to an automatic sampler). The experimental designs are well-thought-out and the corresponding results indicate that the suggested mechanism is indeed capable of reducing post-sampling evaporation, although there are limitations (certain climatic conditions, storage times).

The manuscript is also well structured and written.

I do have a few remarks (see below), but overall I think the manuscript should be published as a Technical Note in HESS, after minor revisions.

We thank Nils Michelsen for this positive evaluation of your work and his comments, which were very helpful for improving our manuscript. We have addressed all comments in detail below (in blue bold font).

Introduction: I think Williams et al. (2018) should be cited somewhere in the introduction. They have tackled the topic of post-sampling evaporation from automatic samplers before and should be given credit for their efforts. As their approach was rather different from yours (other evaporation reduction methods; experiments in insulated boxes instead of real automatic samplers), mentioning their work will provide additional context (and justification) for your work.

Thank you. We will add this reference to the introduction.

Line 47: That the tube dips into the collected water is obviously the most important aspect of this mechanism, but for sake of completeness, you could also mention the complementing aspect of the Gröning et al. (2012) design, i.e. the pressure equilibration tube (e.g. "Tube-dip-in-water collector with pressure equilibration", IAEA 2014). To additionally cite the pioneer work by IAEA (2002) is a good idea, but the reference is missing in the reference list.

We will include IAEA (2002) in the reference list. We will also add a statement to the methods section: "<u>Because a small gap remains between the syringe housing and the inner rim of the sampler bottle opening</u> (i.e., not air-tight), pressure differences due to water flowing into the bottle will equilibrate with the outside conditions. Thus our system does not require an external tube for pressure equilibration such as the "tubedip-in-water collector" proposed by Gröning et al. (2012)."

Lines 61-63: This sentence is a bit misleading. It suggests that only changes in air temperature and humidity would cause problems, but evaporation and vapor mixing would also occur if temperature and humidity remained constant.

We will re-formulate the sentence: "However, because the sample bottles remain open during the sampling period, <u>vapor exchange may occur between the sample water and the atmosphere inside the sampler</u> <u>housing, which may alter</u> the isotopic compositions of the water samples in the bottles."

Line 78: I suggest to refer to Figure 1a directly after "9 cm in length"

We will add the reference to Figure 1a after the end of the suggested sentence and change the sentence to: "On the Luer tip, we fit a 1-mm inner diameter silicone tube approximately 9 cm in length, to reach the bottom of the sample bottle (Figure 1 <u>a</u>, b)." Line 80: The word "tightly" is misleading here. If the syringe housing is really plugged tightly into the bottle opening (i.e. air-tight), the whole mechanism would not work anymore. When additional water is supposed to flow into the bottle, air needs to be displaced and has to leave the bottle. You do mention this aspect later (Section 4, line 412-413), but this should be made clear here. You could even explicitly mention that you skip the "pressure equilibration tube" of the original concept (Gröning et al., 2012; IAEA 2002, 2014; see comment above).

This is a good point. We will remove the term "<u>tightly</u>" from the sentence and clarify the setup by adding the explanation proposed above: "<u>Because there remains a small gap between the syringe housing and the</u> <u>inner rim of the sampler bottle opening (i.e., not air-tight), pressure differences due to water flowing into</u> <u>the bottle will equilibrate with the outside conditions. In this way, our system does not require an external</u> <u>tube for pressure equilibration such as the "tube-dip-in-water collector" proposed by Gröning et al. (2012)</u>."

Lines 85-91: I appreciate that this potential pitfall is highlighted and that you provide a maximum filling rate. I can imagine that debris (e.g. sediment, insects) may further reduce the tubing diameter (1 mm is quite small) in a field setting. Hence, it is probably a good idea to use a screen at the intake (e.g. a funnel screen) and maybe you want to mention this explicitly somewhere.

Good point. We will include a sentence after Line 86: "<u>In order to prevent debris (e.g. sediment, insects,</u> <u>leaves) from clogging the evaporation protection system, the streamwater intake or precipitation funnel can</u> <u>additionally be equipped with a screen."</u>

Line 104: For sake of consistency, this part should read ". . .during 62 two-to-three week cycles. . ." (as in lines 179 and 382)

We will change this as suggested.

Line 111: Please provide some details on the climate-controlled chamber. If this was a commercial chamber, please mention model and manufacturer. If it was a custom-made chamber, please indicate this. We will clarify this: "The ISCO autosampler was placed <u>on a heater inside a ventilated chamber</u> where the conditions were kept at approximately 35 °C air temperature and 11 % relative humidity." We will also change the term "climate-controlled chamber" to "<u>ventilated chamber</u>" throughout the text (four instances in total).

Line 113: Please also mention details on the temperature and humidity loggers (model, manufacturer, precision). Sometimes, it is these technical details that matter for the reader, particularly if they want to conduct similar experiments (as indirectly suggested in line 452). Knowing which logger can cope with such conditions (rel. humidity of up to 100 %), can be valuable.

The specifications of the temperature and humidity loggers are: RHT30 humidity-temperature data logger; EXTECH Instruments, FLIR Commercial Systems Inc., Nashua, NH, USA; measurement accuracy ±1 % rel. humidity and 0.5 °C temperature). We will include this information in the methods section of the revised manuscript.

Line 217: Wouldn't $\Delta\delta$ 18O and $\Delta\delta$ 2H be more accurate? We decided to avoid double-delta notations for the sake of easier readability.

Line 230: Please specify what the reference water is here (tightly sealed bottles). We will add this information to the sentence to be more specific: "...of the reference water (i.e., $\delta^2 H_{reference}$, $\delta^{18}O_{reference}$ of the water in the tightly sealed bottles) for each sampling period."

Lines 237-239: The 1°C temperature difference between the climate chamber and the ISCO (inside the climate chamber) is a bit surprising, particularly because the temperatures matched quite well before the logger as moved "to a more representative position".

We will add a better explanation: "Air temperature outside the ISCO sampler was around 35 ± 1 °C with distinct diurnal variations (<u>a</u> 1.2 °C temperature drop at the beginning of the 4th day was caused by moving the <u>humidity</u>-temperature <u>logger from a position close to the heater</u> to a <u>higher</u> position <u>near the sampler's</u> <u>control unit to better represent the conditions</u> inside the <u>ventilated</u> chamber).

Table 1: Probably, this should read "... relative to the reference water...".

Correct, we will change this accordingly.

Lines 264-267: The isotopic shifts observed for days 10 and 12 are indeed remarkable and I wonder if it is a coincidence that this was about the time when a humidity of about 100 % was reached. Could condensing water play a role here (i.e. liquid water drops flowing into the bottles)? Concerning the possible explanations that are given: I do not understand how spillage could explain the phenomenon.

To check whether condensation might have played a role here, we calculated the dew point temperatures based on the Magnus formula and the parameter set from Sonntag (1990). As can be seen in the figure below, the dew point was reached inside the ISCO sampler after day 12 of the experiment. Even though this coincides with the day 100% relative humidity was reached, this would suggest that all samples filled on and after day 12 would also have been affected by condensation, which was not the case. In addition, because of our sampling design, all previous samples (days 0-10) were exposed to the dew point conditions as well but did not show such large isotopic differences as those samples from days 10 and 12.

We therefore can only speculate about the larger isotope differences in the samples of days 10 and 12. We removed the possible explanation of spilling and kept measurement errors during isotope analysis.

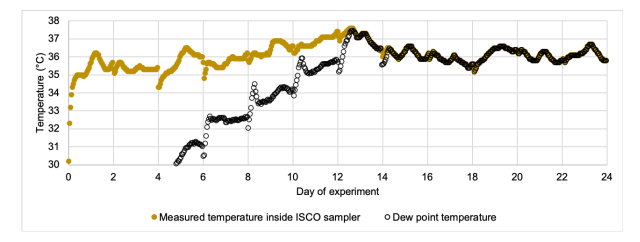


Fig. 2: I am a bit confused about the x-axis in Fig. 2c. I was able to match all blue data points in 2b and 2c and conclude that some of the blue points in 2c represent the first days of the experiment (e.g. days 0 and 2). During this time, humidity was around 50 % according to 2a, but in 2c the x-axis only shows humidities above 87.5 %.

A bottle that was filled on day 1 of the experiment stayed inside the ISCO for 23 days and experienced the full evolution of the relative humidity. The values shown in Figure 2c represent the average relative humidity values experienced during the full storage duration of each bottle.

We will edit the caption of Figure 2 to better explain the aggregation of humidity values: "Isotopic change in water samples relative to the reference water as a function of <u>the mean relative humidity, which represents</u> <u>the average of relative humidity values during the full storage duration of each bottle.</u>"

Line 282: Please make clear that the "low relative humidity" refers to the chamber conditions (not the conditions inside the ISCO).

We will clarify this by adding "... and low relative humidity inside the ISCO autosampler,..."

Lines 296-298: If these temperatures and humidities are not shown anywhere, please indicate this to avoid confusion.

We will edit this sentence: "Outside the ISCO autosamplers, air temperature (mean ± standard deviation) was 13.3±6.2°C in the outdoor setting and 18.6±4.7 °C indoors, while relative humidity was 73.5±23.0% outdoors with distinct daily fluctuations and 44.8±8.9% indoors (values of temperature and relative humidity measured outside the ISCO autosamplers are not shown but are provided in the Supplement)."

Lines 312-314: If wind plays a role strongly depends on the geometry of the setup. A possible scenario that is not mentioned here so far is temperature-triggered gas volume changes. In contrast to Experiment 1 (climate chamber), Experiment 2 was characterized by significant daily temperature fluctuations (20°C and more inside

the ISCO; outdoor setting; Fig. 4a). Upon heating (daytime), the air inside the ISCO expanded and some of the (moist) air was pushed out of the device. When temperatures dropped (at night), the opposite happened, i.e. the air in the ISCO contracted and sucked in fresh air from outside. This "sampler breathing" probably happened on a daily basis, resulting in a greater air exchange, which in turn (apparently) caused lower humidities inside the ISCO (and more evaporation). Maybe this effect could have played a role here as well. We will include in Line 313: "... is less important for causing isotope effects than the magnitude of the temperature fluctuations that may trigger gas volume exchanges. In contrast to Experiment 1 (ventilated chamber), Experiment 2 was characterized by significant daily temperature fluctuations (20 °C and more inside the ISCO that was situated outdoors; Figure 4a). During the daytime when air temperature increased, the air inside the ISCO expanded and some of the (moist) air was pushed out of the device; when temperatures dropped (at night), the opposite happened, i.e. the air in the ISCO contracted and sucked in fresh air from outside. This "sampler breathing" probably happened on a daily basis, resulting in a greater air exchange with the outside, which in turn may have reduced the humidity inside the ISCO and resulted in stronger evaporation (Figure 4a). In addition, the greater potential for wind-driven ventilation of the outdoor ISCO may also have enhanced evaporative fractionation effects in the open sample bottles."

Lines 328-329: This sentence is confusing. Do you mean "...exchange with the heavier RefA water..."? Correct, this was a typo. We will correct this in the revised manuscript.

Lines 370-372: I am not sure if this statistical summary is sufficient. The mean value is close to zero, but I think you should at least mention that the isotopic shifts showed a fairly large scatter and ranged from about -2 (i.e. isotopic depletion) to about +3 % (isotopic enrichment) in case of δ 2H (Fig. 7c and 7d). In case of δ 18O, the maximum deviations are remarkable as well. Here, $\Delta\delta$ 18O scatters between about 0.7 and about +1.2 % (Fig. S5c and S5d). Both values are clearly beyond the analytical precision.

We will change the text to: "deviated <u>statistically</u> significantly from zero." We will also add a note to point out the large scatter in Figure 7 c,d: "<u>The isotopic differences of the samples in open bottles relative to the</u> <u>reference water exhibited substantial scatter, with values between -2 ‰ and +3.5 ‰</u>..."

Practical implications: Here, the wind issue is stressed again. Maybe the "sampler breathing" (see above) also deserves to be mentioned here. If it really played a role, a practical consequence may be the need of a thermal insulation of the ISCO (reducing the temperature fluctuations inside the device).

We will change the respective sentence to "The larger isotopic change observed during the field deployment of Experiment 3 may be attributed to more variable climatic conditions (e.g., due to diurnal temperate variations) <u>causing "sampler breathing"</u>, ...". And we will also add "sampler breathing" at a later point in the practical implications Section: "Larger temperature and humidity contrasts due to diurnal fluctuations in outdoor conditions may have resulted in repeated evaporation and condensation inside the ISCO housing, and in enhanced vapor exchange between the sample bottles and the outside atmosphere <u>("sampler breathing"</u>)."

Practical implications/Conclusions: The partly significant δ 2H and δ 18O values observed in Experiment 3 (see above) underscore the relevance of such "field controls". These pre-filled bottles (known isotopic composition and mass), placed into the automatic sampler upon field installation, allow for a hindsight evaluation of the samples' isotopic integrity. Although their advantage is somewhat obvious, it may be a good idea to explicitly recommend this technique to the reader.

This is a good suggestion. We will add a sentence to the revised manuscript near the end of the "Practical implications" section: "<u>Control samples with known isotopic compositions in open, retrofitted and closed bottles placed in the autosampler for the entire storage duration, should be used to monitor composite isotope effects and to allow a retrospective quality assessment of the automatically collected samples."</u>

References

Gröning, M., Lutz, H. O., Roller-Lutz, Z., Kralik, M., Gourcy, L., and Pöltenstein, L.: A simple rain collector preventing water re-evaporation dedicated for δ18O and δ2H analysis of cumulative precipitation samples, Journal of Hydrology, 448-449, 195-200, 2012.

Sonntag D.: Important New Values of the Physical Constants of 1986, Vapour Pressure Formulations based on the IST-90 and Psychrometer Formulae; Z. Meteorol., 70 (5), pp. 340-344, 1990.

Williams, M.R., Lartey, J.L., Sanders, L.L., 2018. Isotopic (δ 18O and δ 2H) Integrity of Water Samples Collected and Stored by Automatic Samplers. Agricultural & Environmental Letters, 3(1), 1-5, 2018

2) Authors' response to the interactive comment by Anonymous Referee #2

General comments

The manuscript entitled 'Evaluation of a low-cost evaporation protection method for portable water samplers' by von Freyberg et al. describes the development of a robust and inexpensive method for an evaporation reduction method for automatic water samplers that are often used in hydrology. In order to evaluate their developed setup, laboratory and field tests were conducted to simulate extremely dry and warm conditions, to test for vapor transfer between samples and to quantify the isotopic change during 3-week storage periods. It could be shown that the method efficiently protects the collected water samples from undergoing isotopic changes due to evaporative fractionation and vapor mixing and that the protection method significantly reduced isotopic fractionation over the 3-week periods under ambient climatic conditions in the field. The manuscript is well structured and nicely written. The topic of this promising approach fits well to the scope of the journal and appears to be of interest for isotope hydrologists. Most of my editing comments match those of Referee 1 and have already been addressed by the authors; therefore I only suggest minor revisions prior to acceptance and publication in Hydrology and Earth System Sciences.

We thank the anonymous referee for the positive evaluation of our manuscript. We address his/her two specific comments below (in blue bold font).

Specific comments

Introduction, L. 45-47: I suggest mentioning styrofoam beads as an additional mechanical protection method, because this is commonly used as an evaporation protection method in ISCO-bottles.

We will add the Styrofoam beads to the introduction: "Alternative mechanical evaporation protection modifications have been suggested, like *covering the water surface with Styrofoam beads (Angermann et al., 2017)* or placing a table tennis ball in the collection funnel..."

L. 75, 81, 84: Please consider replacing 'Our...' by 'The...:' at the beginning of these sentences, otherwise it sounds a bit like the conclusion section.

In the revised manuscript, we will change "our evaporation protection" to "*the* presented evaporation protection" (line 75), "our setup" to "*the described system*" (line 81, sentence changed from original based on comments from Referee #1), and "our design" to "*the presented* design" (line 81).

References:

Angermann, L., Jackisch, C., Allroggen, N., Sprenger, M., Zehe, E., Tronicke, J., Weiler, M., Blume, T., 2017. Form and function in hillslope hydrology: characterization of subsurface flow based on response observations. Hydrol. Earth Syst. Sci. 21, 3727–3748. http://dx.doi.org/10.5194/hess-21-3727-2017.

Technical Note: Evaluation of a low-cost evaporation protection method for portable water

samplers

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Abstract Automated field sampling of streamwater or precipitation for subsequent analysis of stable water isotopes (²H and ¹⁸O) is often conducted with off-the-shelf automated samplers. However, <u>when</u> water samples <u>are</u> stored in the field for days and weeks in open bottles inside autosamplers, <u>their isotopic signatures</u> can be altered by evaporative <u>-undergo isotopic</u> fractionation and vapor mixing, <u>thus altering their isotopic signature</u>. We therefore designed an evaporation protection method which modifies autosampler bottles using a syringe housing and silicone tube, and tested whether this method reduces evaporative fractionation and vapor mixing in water samples stored for up to 24 days in <u>6712 Full-size Portable</u> <u>Samplers ISCO autosamplers</u> (Teledyne ISCO-, Lincoln, US). Laboratory and field tests under different temperature and humidity conditions showed that water samples in bottles with evaporation protection were far less altered by evaporative fractionation and vapor mixing than samples in conventional open bottles. Our design is a cost-efficient approach to upgrade the 1-litre sample bottles of <u>the ISCO-autosamplers</u>6712 Full size Portable Samplers, allowing secure water sample collection in warm and dry environments. Our design can be readily adapted (e.g., by using a different syringe size) to fit the bottles used by many other field autosamplers.

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landscape, and thus provide important insights into water sources, flowpaths, and travel times in hydrologic systems (e.g., Gat et al., 2001; Kendall and McDonnell, 1998; Klaus and McDonnell, 2013; McGuire and McDonnell, 2008). Furthermore, deuterium and oxygen-18 signatures in precipitation and/or streamwater can help to track the movement of atmospheric air

The stable water isotopes deuterium (²H) and oxygen-18 (¹⁸O) are used as natural tracers for water flow through the

masses (Fischer et al., 2017), identify the water sources of plants (Dawson and Ehleringer, 1991), and reconstruct climate records (Shanley et al., 1998). Long-term data sets of stable water isotopes in precipitation and streamwater are available

35 from global monitoring networks (the Global Network of Isotopes in Precipitation, GNIP, and the Global Network of Isotopes in Rivers, GNIR) and various national monitoring networks (e.g., the ISOT monitoring program of the Swiss Federal Office for the Environment).

Streamwater is usually collected through instantaneous grab sampling, after which the sample containers are sealed and 40 cooled until laboratory analysis. In contrast, precipitation is usually collected over periods of weeks to months with open buckets or funnels mounted onto sample bottles. To prevent evaporative fractionation of the precipitation sample during the sampling period, paraffin oil can be used to that forms a protective layer of oil floating on the water sample, thus preventing evaporative fractionation (IAEA, 2014; Williams et al., 2018). However, residual oil in the water sample can alter subsequent laser spectroscopy measurements (Gröning et al., 2012). The contamination risk is particularly high if the 45 sample volume is small, so the addition of oil is only suitable for longer sampling durations (weekly or monthly), but not recommended for daily or sub-daily sampling. Alternative mechanical evaporation protection modifications have been suggested, like covering the water surface with Styrofoam beads (Angermann et al., 2017), or placing a table tennis ball in the collection funnel ("ball-in-funnel") to seal the inflow during times without precipitation (Prechsl et al., 2014). Another widely used collector modification is the "tube-dip-in-water" collector (Gröning et al., 2012; IAEA, 2002), where the 50 collection bottle is sealed except for a small-diameter tube that reaches from the bottom outlet of the funnel into the water sample. This setup substantially reduces the contact area between the water sample and the atmosphere. While some of these modifications may substantially reduce evaporative fractionation of the water sample in the bottle, others were found to be less effective (Michelsen et al., 2018; Terzer et al., 2016).

- The above methods and modifications were originally designed for single-sample collection using a precipitation totalizer (e.g., IAEA, 2014). For many hydrological questions, however, higher-frequency measurements of stable water isotopes are of interest, requiring daily or even sub-daily sampling of precipitation or streamwater (e.g., Knapp et al., 2019; Rücker et al., 2019; von Freyberg et al., 2018; Wang et al., 2019). This can be achieved with field-deployable automatic water samplers with programmable pump-and-distribution systems that fill and store a series of empty open bottles. Many hydrologic
 studies deploy use off-the-shelf automatic water samplers (available from, e.g., Teledyne ISCO, Lincoln (NE), USA, and Maxx GmbH, Rangendingen, Germany), because these systems are rugged, robust, versatile, and easy to program. For automatic samplers with a 24-bottle configuration, this setup reduces the manual labor of daily precipitation sampling to the collection of sample bottles only once every 24 days. However, because the sample bottles remain open during the sampling period, vapor exchangethe water samples are exposed to changes in air temperature and humidity may occur between the sample water and the atmosphere inside the autosampler housing, which may alter the isotopic compositions of the water samples in the bottles due to evaporative fractionation and mixing in the vapor phase (Williams et al., 2018).
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While attempts have been made to design more sophisticated field-deployable, programmable water samplers which reduce these isotope fractionation effects, most of these devices are not readily available (i.e., prototypes), or are technically

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complex or expensive (Ankor et al., 2019; Berman et al., 2009; Hartmann et al., 2018; Michelsen et al., 2019). We therefore designed and tested a low-cost evaporation protection modification that can be used with Teledyne ISCO's 6712 Full-size Portable Samplers and 1-litre ISCO sample bottles. We retrofitted the 1-litre ISCO sample-bottles with a simplified "tubedip-in-water" collector type that allows rapid sample flow, but reduces isotope effects due to vapor exchangeevaporative fractionation. The proposed setup is cheap, easy to handle and suitable for a wide range of sample volumes that are common in daily precipitation or streamwater sampling.

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2 Methods

2.1 Evaporation protection

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We designed an evaporation protection modification for the 1-litre sample bottles of the 6712 Full-size Portable Sampler (Teledyne ISCO₇, Lincoln, US; hereafter referred to as 'ISCO autosampler'). Our The presented evaporation protection consists of a 100-ml syringe housing (i.e., BP PlastipakTM 100 ml syringe with catheter tip, without its piston and rubber piston stopper) with attached Luer tip adapter (BP PlastipakTM). On the Luer tip, we fit a 1-mm inner diameter silicone tube of approximately 9 cm in length, to reach below the water level of the sample in the sthe bottom of the sample bottle (Figure 1a, b). The barrel flange of the syringe housing is trimmed on one side (Figure 1a) to allow the retrofitted sample bottles to properly fit into the bottom compartment of the ISCO autosampler-ensuring that they do not block the distributor arm. This 85 modified syringe housing is then plugged tightly into the opening of an ISCO sample bottle (Figure 1b). Because there remains a small gap remains between the syringe housing and the inner rim of the sampler bottle opening (i.e., not air-tight), pressure differences due to water flowing into the bottle will equilibrate with the outside conditions. In this way, Thus our the described system does not require an external tube for pressure equilibration, such as the "tube-dip-in-water collector" proposed by Gröning et al. (2012). Our setup ensures a smooth, splash-free sample flow from the syringe through the 90 silicone tube into the bottle. Because the end of the silicone tube is fully immersed in the sample liquid, only the crosssectional area of the silicone tube is exposed to the ambient atmosphere (rather than the entire cross-sectional area of the water surface), minimizing vapor exchange with the surrounding atmosphere. The presented-Our design of the evaporationprotected ISCO bottle Our setup-ensures a smooth, splash-free sample flow from the syringe through the silicone tube into the bottle when filled at a flow rate of approximately 100 ml min⁻¹. Our design of the evaporation-protected ISCO bottle 95 Furthermore, it is robust, cheap (<5 USD per /sample bottle), chemically inert, easy to disassemble and to clean, and allows

rapid sample flow of approximately 100 ml min⁻¹ into the bottle.

In field operation for the collection of streamwater samples, the autosampler should be programmed to not exceed the filling rate that can be accommodated by the narrow silicone tube. This can be accomplished by programming the autosampler to 100 deliver a series of 100--ml aliquots, allowing enough time between them (about 1 minute minimum) so that they can drain from the 100--ml syringe into the sample bottle. If possible, one should also limit the total sample volume so that the water line is somewhere in the narrow silicone tube and not in the syringe, in order to limit the water surface that is available for evaporation or condensation. In order to protect prevent debris (e.g. sediment, insects, leaves) from clogging the evaporation protection system from clogging with debris (e.g. sediment, insects, leaves), the streamwater intake or precipitation funnel 105 can additionally be equipped with a screen. To transport the filled sample bottles, the syringe housing has to be removed and the bottles have to be sealed with screw caps supplied by the manufacturer. If the bottles are transported upright and leakage is unlikely to occur, the syringe housing can also stay in place and its upper opening can be sealed with the black rubber piston stoppers that are supplied together with the syringes (BP PlastipakTM 100-ml syringe with catheter tip; Figure 1c).

2.2 Monitoring evaporation and fractionation

110 We conducted three experiments to assess the effects of evaporation and vapor mixing on the isotopic composition of the liquid samples in ISCO 6712 autosamplers, comparing the retrofitted ISCO bottles to un-modified ISCO bottles. In Experiment 1, we simulated a daily sampling routine under extremely dry and warm conditions to test for evaporative fractionation effects over different storage durations. In Experiment 2, we used two contrasting reference waters to test for changes in their isotopic compositions due to vapor transfer between samples, in addition to fractionation effects under 115 ambient conditions with diurnal fluctuations in temperature and humidity. Experiment 3 evaluated the performance of the retrofitted ISCO bottles during 62 two-to-three³-week cycles over a nearly 4-year deployment at two field sites in the northern Swiss pre-Alps.

Experiment 1

120 We prepared one ISCO autosampler for a 24-day test of the retrofitted bottles under controlled laboratory conditions. The autosampler contained 24 sample bottles, of which 12 were retrofitted with the modified syringe housing and the other 12 bottles remained open (i.e., as they do in normal operation). Open and retrofitted bottles were arranged alternatingly in the autosampler carousel. The ISCO autosampler was placed on a heater inside a climate controlled ventilated chamber where the conditions were kept at approximately 35 °C air temperature and 11 % relative humidity. Air temperature and 125 relative humidity were measured every hour in the elimate ventilated chamber and inside the bottom compartment of the ISCO autosampler with RHT30 humidity-temperature loggers (EXTECH Instruments, FLIR Commercial Systems Inc., Nashua, USA; measurement accuracy ± 1 % relative humidity and 0.5 °C temperature).

The bottom compartment (containing the sample bottles) and the middle compartment (containing the pump and control

unit) of the ISCO autosampler remained attached for the entire duration of the experiment. The water samples were distributed among the bottles by using the instruments' software to move the distributor arm to the desired position. The instrument's sampling tube was not threaded through the peristaltic pump, but instead was directly attached to the inlet of the distributor arm. This setup allowed <u>us to pour</u> an exact <u>amount-volume</u> of water <u>into the sampling tube</u>, with the water to flowflowing gravitationally from the inlet tube through the distributor arm into the sample bottle (we bypassed the peristaltic pump because it does not allow such exact sample dosing and might introduce air bubbles into the sample during pumping). This sampling protocol is consistent with the automated sampling of precipitation under field conditions, when the autosampler's sample inlet tube is connected directly to a precipitation collection funnel (i.e., bypassing the peristaltic pump) so that incoming precipitation flows directly through the distributor arm towards the pre-programmed bottle position (e.g.,

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Rücker et al., 2018).

To ensure that the initial isotopic compositions of all water samples were comparable, we filled a 20-litre tank with distilled reference water before the beginning of the monitoring period. This reference water tank was tightly sealed and stored at room temperature. It was only opened every second day to retrieve 801.5 ml of reference water. From this aliquot, 1.5 ml were filled into a glass vial with screw cap (screw thread vials 1.5ml, PP-screw thread caps with silicone-/PTFE-septum,

145 WICOM Germany GmbH, Heppenheim, Germany) and stored at 4 °C until isotope analysis. The purpose of these samples was to monitor the isotopic composition of the reference water and account for possible fractionation inside the storage tank. The remaining 800 ml of reference water was filled into two empty ISCO sample bottles (400 ml each into an open and a retrofitted bottle). For the open bottle, 400 ml were emptied rapidly into the inlet tube. Because of the small tubing diameter in the retrofitted sample bottles, we poured the 400 ml of reference water into the inlet tube in four steps of 100 ml min⁻¹ to prevent overflow.

Starting on day 1 and then every second day, one open and one retrofitted bottle were filled with 400 ml of reference water each following the protocol described above, and the last two sample bottles (No. 23 and 24) were filled at the start of on the 23rd day.

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To monitor evaporation and isotopic fractionation under ambient conditions outside the ISCO autosampler, we prepared three additional ISCO bottles at the start of the monitoring period. For this purpose, we filled 400 ml of the reference water each into one open ISCO bottle (i.e., non-modified), one ISCO bottle that was retrofitted with evaporation protection, and one tightly sealed ISCO bottle on day 1 of the laboratory experiment. We placed these bottles <u>on the heater</u> inside the <u>climate controlledventilated</u> chamber, but outside the ISCO autosampler, for the duration of the experiment (24 days).

To mimic the field protocol (see Sect. 2.3), all sample bottles (i.e., inside and outside of the ISCO autosampler) were opened and sub-sampled at the end of day 24. For this, 1.5 ml of water from each sample bottle were immediately transferred into glass vials with screw caps and stored at 4°C until isotope analysis.

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Experiment 2

For Experiment 2, we prepared two ISCO autosamplers with alternating open and retrofitted bottles, analogously to Experiment 1. One sampler was stored indoors at approximately constant temperature and relative humidity, and the other sampler was stored outdoors at a sunny location where ambient conditions were more variable. Temperature and relative humidity were monitored inside and outside the ISCO autosamplers at both locations.

We filled all sample bottles on day 1 of the experiment to ensure that all samples underwent the same mixing and fractionation processes over the following 21 days. We alternatingly filled the bottle pairs (open and retrofitted) with two isotopically contrasting reference waters: one, which we will call RefA, was isotopically much heavier ($\delta^2 H \approx -40.5 \%$, $\delta^{18}O \approx -5.6 \%$) than the other, which we will call RefB ($\delta^2 H \approx -69.8 \%$, $\delta^{18}O \approx -9.7 \%$), with the isotopic difference

between the two reference waters being approximately 29.3 ‰ and 4.1 ‰ for δ²H and δ¹⁸O, respectively. To test whether smaller sample volumes were affected more substantially by vapor mixing and evaporation, we alternated the sample volumes between 200 and 400 ml. Thus, the carousel of each ISCO sampler contained three replicates of each possible combination of the two reference waters (RefA vs. RefB), the two sample volumes (200 vs. 400 ml), and two bottle types
(open vs. retrofitted with evaporation protection).

We placed four additional sample bottles into the center of each autosampler carousel on day 1 of the experiment. Two of these bottles contained 200 ml of RefA water and the other two bottles contained 200 ml of RefB water; all four bottles were tightly sealed.

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The bottom compartment of the autosampler (containing the sample bottles) and the middle compartment (containing the pump and control unit) remained attached for the entire duration of the experiment. Sample bottles were weighted at the start and end of the experiment to track potential changes in water volumes. After 21 days, the ISCO samplers were opened and all bottles were retrieved. We transferred 1.5 ml of the liquid sample water from each bottle into glass vials with screw caps and stored them at 4 °C until isotope analysis.

Experiment 3

To assess the effectiveness of the retrofitted bottles under central European climatic conditions, we monitored evaporative fractionation in two ISCO autosamplers during 62 two-to-three-week sampling periods between October 2015 and

195 June 2019. For this purpose, we installed the ISCO autosamplers at two different locations in the northern Swiss pre-Alps: at

the EIN site located near the city of Einsiedeln (8.75708°E, 47.13370°N, WGS84) at 910 m above sea level (m a.s.l) and at the ERL site located roughly 11 km southwest of Einsiedeln in the Erlenbach catchment (8.71502°E, 47.04249°N, WGS84) at 1228 m a.s.l.

At the beginning of each sampling period, we filled one tightly sealed, one open and one retrofitted sample bottle with 400 ml reference water each and placed them in the center of the ISCO carousel (the outer 24 bottles were reserved for conventional automatic precipitation sampling, not discussed here). The ISCO autosamplers remained at the field sites for roughly two to three weeks before all bottles were collected and replaced with new ones. After collecting the sample bottles, they were transported to the ETH Zurich laboratory and two-1.5 ml aliquots of sample water were transferred from each bottle into glass vials with screw caps; the vials were stored at 4 °C until isotope analysis.

To identify potential drivers of evaporative fractionation effects during these sampling periods, we used on-site air temperature and relative humidity measurements. These measurements were provided by the Swiss Federal Office of Meteorology and Climatology (MeteoSwiss) for the EIN site and by the Swiss Federal Institute for Forest, Snow and

210 Landscape Research (WSL) for the ERL site. In addition, we used daily maximum, minimum and average values of air temperature and relative humidity to calculate the daily vapor pressure deficit (VPD = $e_s - e_a$) following Allen et al. (1998):

$$e_T^0 = 0.6108 \cdot exp(\frac{17.27 \cdot T}{T + 237.3}) , \qquad (1)$$
$$e_s = \frac{e_{Tmax}^0 + e_{Tmin}^0}{2} , \qquad (2)$$

215 $e_a = \frac{e_{Tmin}^0 \cdot H_{max} + e_{Tmax}^0 \cdot H_{min}}{2} , \qquad (3)$

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where e_T^0 is the saturation vapor pressure at the air temperature T (kPa), e_s is the saturation vapor pressure (kPa), e_a is the actual vapor pressure (kPa), T is the air temperature (°C), H is the relative humidity (-), and the indices *min* and *max* indicate the minimum and maximum values of temperature and relative humidity observed during any day. To compare these potential drivers with the isotopice differences, we averaged the daily values of air temperature, humidity and VPD over the individual sampling periods.

2.3 Stable water isotope analysis and isotopic differences

For Experiment 3, all water samples collected between 6 October 2015 and 13 December 2017 were analyzed at the laboratory of the Swiss Federal Institute for Forest, Snow and Landscape Research (WSL) with an Isotopic Water Analyzer LGR IWA-45-EP (Los Gatos Research, ABB Los Gatos Research, San Jose, California, USA) with a measurement precision of 0.5 ‰ for δ¹⁸O and 1 ‰ for δ²H. All of Experiment 3's samples collected after 13th December 2017, and all water samples of Experiments 1 and 2, were analyzed with a Cavity Ring-down Spectrometer at the ETH Zurich laboratory

(L2140-*i* liquid isotope analyzer, Picarro Inc., Santa Clara, CA, USA) with a measurement precision of 0.2 ‰ for δ^{18} O and 1 ‰ for δ^{2} H. All isotope values in this study are reported in δ -notation relative to Vienna Standard Mean Ocean Water (V-SMOW) and the measurement uncertainty is provided as standard deviations calculated from 2 to3 repeated injections of each sample.

To quantify the isotopic change <u>effects</u> in the water samples, we calculated the isotopic difference (Δ^{18} O and Δ^{2} H, ‰) between the water sample at the end of the storage period, and the reference water at the beginning of the storage period:

 $\Delta^{i} E = \delta^{i} E_{\text{sample}} - \delta^{i} E_{\text{reference}} , \qquad (4)$

where $\delta^{i} E_{sample}$ and $\delta^{i} E_{reference}$ are the delta values of the isotope ⁱE in the sample water or the reference water, 235 respectively. For Experiment 1, we compared the isotope composition of the water samples from the open and retrofitted bottles ($\delta^2 H_{sample}$, $\delta^{18} O_{sample}$) to the isotope composition of the reference water from the storage tank ($\delta^2 H_{reference}$, $\delta^{18}O_{reference}$). Because each second day we collected one reference water sample from the tank and filled one open and one retrofitted bottle with reference water, the comparison of the isotopic changes differences ($\Delta^{i}E$) of samples from bottles 240 with and without evaporation protection assesses the effectiveness of the retrofitted sampler bottles in protecting against evaporative enrichment. For Experiment 2, we compared the isotopic composition of the RefA and RefB water samples from the various open and retrofitted bottles ($\delta^2 H_{sample}, \delta^{18} O_{sample}$) -to the isotopic composition of the closed sample bottles with the corresponding reference water RefA or RefB ($\delta^2 H_{reference}$, $\delta^{18} O_{reference}$) in order to quantify isotope effects due to vapor mixing and evaporation. For Experiment 3, we compared the isotopic composition of each sampling 245 period's water samples from the open and retrofitted bottles ($\delta^2 H_{\text{sample}}, \delta^{18} O_{\text{sample}}$) to the isotope values of the reference water in the tightly sealed bottles (i.e., $\delta^2 H_{reference}$, $\delta^{18} O_{reference}$) of the water in the tightly sealed bottles) for for each sampling period.

3 Results

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250 **3.1 Laboratory evaluation of the evaporation protection method: Experiment 1**

During Experiment 1, humidity outside the ISCO autosampler stayed relatively constant at approximately $11-\pm-3$ -% (mean±1standard deviation), while it continuously increased inside the <u>lower-ISCO bottom</u> compartment from 33 % to 100_-% between day 1 and 13, and then remained at 100 % until the end of the experiment (<u>Figure 2</u>Figure 2a). Air temperature outside the ISCO sampler was around 35-±-1_-°C with distinct diurnal variations (<u>a</u> 1.2_-°C temperature drop at the beginning of the 4th day was caused by moving the <u>humidity-</u>temperature logger from a position close to the heater to a

more higher representative position near the sampler's control unit to better represent the conditions inside the climate ventilated chamber). The air temperature inside the ISCO housing was 36-±-1 °C and did not exhibit strong diurnal patterns.

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The sample bottles stored outside the autosampler (i.e. at 11 % relative humidity and 36 °C temperature) experienced different degrees of evaporative fractionation between the start and end of the monitoring period (Table 1Table 1): while evaporative fractionation was insignificant in the closed bottle, we observed isotopic enrichment in both the retrofitted and the open bottle. Enrichment was substantially stronger for the sample in the open bottle (a change of roughly 100 ‰ in δ^2 H and 22 ‰ in δ^{18} O within the first 12_-days), compared to the sample in the retrofitted bottle (a change of 9 ‰ in δ^2 H and 2 ‰ in δ^{18} O over 24 days). We sampled tThe open bottle had been sampled already on day 12 of the experiment because we had observed substantial evaporated completely, while the loss of water volume was small in the retrofitted control-bottle (Figure 3Figure 3).

The δ^2 H values of the water samples from inside the ISCO <u>auto</u>sampler show that evaporative fractionation differed between samples from open and retrofitted bottles, and also varied with storage duration (<u>Figure 2Figure 2</u>b and c). The isotopic <u>changes differences</u> (eq. 4) were generally smaller for samples in retrofitted bottles compared to those in the open bottles. For δ^2 H, the isotopic <u>changes differences</u> (Δ^2 H) in the retrofitted bottles were mostly close to 0 ‰ independent of storage duration, while the isotopic <u>changes enrichment</u> in the open bottles ranged up to 5 ‰ (<u>Figure 2Figure 2</u>b). For δ^{18} O, we obtained less clear fractionation signals, and while the enrichment was always greater in δ^{18} O samples of open bottles compared to those in retrofitted bottles, we <u>also</u> observed isotopic depletion of up to <u>-</u>0.3 ‰ for samples <u>in retrofitted bottles</u> <u>that were</u> filled early on in the experiment (data for δ^{18} O are presented in Figure S1).

Water samples filled <u>on and before day 14 of the Eexperiment 1</u>, i.e. samples with more than 10 <u>or more days of storage</u> time, showed substantially larger Δ²H values in the open bottles than in the retrofitted bottles; i.e. samples in open bottles
experienced stronger enrichment (Figure 2Figure 2b). Conversely, samples <u>introduced collected after on</u> day 16 <u>and later</u>, and thus stored for 8 days or less, experienced little or no evaporative fractionation, independent of the bottle type (i.e., Δ²H was not significantly different from zero). This decrease in evaporative fractionation in the later samples may have been due <u>caused by to the</u> increase in relative humidity to <u>approximately 90 % on day 10 and 100 % on day 12</u> inside the ISCO <u>autosampler compartment housing (Figure 2Figure 2a)</u> during the experiment, since the relative humidity inside the ISCO housing reached a value of approximately 90 % on day 10 and 100 % on day 12. Surprisingly, samples filled on days 10 and 12 showed stronger enrichment than those filled on adjacent days, both for open and retrofitted bottles. Because this isotope effect occurred in both bottle types-, it cannot be attributed to a specific process; -it may have been related to spillage during sample handling (thus resulting in smaller sample volumes which are more strongly affected by evaporative fractionation) or interferences in the isotope analyser.

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Figure 2Figure 2c compares the Δ^2 H values of samples from open and retrofitted bottles against the average relative humidity inside the ISCO sampler. While the change in relative humidity inside the ISCO <u>autosampler</u> did not seem to affect the samples in the retrofitted bottles, we obtained a nearly linear relationship for the open bottles indicating a 2.8_∞ enrichment per 10_∞ decrease in relative humidity (p < 0.005). Such a relationship is expected, because the vapor phase in the open bottle is in exchange with the vapor phase inside the ISCO housing, and evaporation from the liquid phase is generally faster when water vapor concentrations in the gas phase are lower (assuming constant temperature). Due to the much smaller contact area between the liquid and vapor phases in the retrofitted ISCO bottles, vapor exchange was reduced and evaporation from the liquid sample was much smaller (even when relative humilities were below 90 % inside the ISCO <u>autosampler</u> housing; Figure 2Figure 2c). Retrofitting made little difference at ≈100 % humidity, when vapor-pressure deficits, and thus evaporation rates, were minimal. Relative humidity outside the ISCO <u>autosampler</u> and temperatures inside and outside the ISCO <u>autosampler</u> remained nearly constant throughout the monitoring period, so no-their relationship with the observed isotopic composition could <u>not</u> be identified and their effect on evaporative fractionation in this laboratory experiment could not be assessed.

In spite of the high temperatures and low relative humidity <u>inside the ISCO autosampler</u>, the observed fractionation effects during Experiment 1 were not <u>excessively very</u> large. This may have been due to some limitations in the setup. For one, we did not use the built-in peristaltic pump of the ISCO system to fill our samples, which follows the protocol for collecting precipitation samples but is not suitable for streamwater grab sampling. In case of streamwater sampling, the pump is used and the tubing between <u>the</u> sampling location and pump is flushed with air before sampling. This process likely results in an intake of air into the ISCO and consequently enhanced vapor exchange with the surrounding atmosphere, which may enhance isotopic fractionation of the collected water sample. In addition, we could not measure the water volumes in the sample bottles at the time of filling and at the end of the experiment because we did not want to open the ISCO <u>autosampler</u> during the experiment period. While we took care to fill exactly 400 ml reference water into each sample bottle through the distributor arm, we cannot exclude that some spillage occurred <u>inside the ISCO</u> during the filling procedure <u>or that some</u> residual water remained in the sampling tube. It is therefore not possible to assess the exact amount of sample volume that was lost due to evaporation.

3.2 Assessing the effect of evaporative fractionation and mixing: Experiment 2

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Ambient conditions during Experiment 2 were colder and more humid compared to Experiment 1, and substantially more variable. Outside the ISCO autosamplers, air temperature (mean \pm standard deviation) was 13.3 \pm 6.2 °C in the outdoor setting and 18.6 \pm 4.7 °C indoors, while relative humidity was 73.5 \pm 23.0_% outdoors with distinct daily fluctuations and

44.8±8.9_% indoors (values of temperature and relative humidity measured outside the ISCO autosamplers are not shown but are provided in the Supplement). Temperature and relative humidity measured inside the autosampler housings exhibited similar but damped diurnal patterns (see Figure 4a, b). The temperature and relative humidity inside the outdoor ISCO were 16.7±6.7 °C and 86.0±13.6_%, respectively. For the indoor ISCO autosampler, the respective values were 18.2±2.9 °C and

- 96.7 \pm 2.7<u>%</u>. In contrast to Experiment 1, the relative humidity inside the <u>ISCOautosampler housing</u> did not increase gradually to 100 % over several days but remained high throughout the experiment, probably because all sample bottles were filled from the start, instead of successively as in Experiment 1.
- 330 The changes in isotopic composition, $\Delta^2 H_{RefA}$ and $\Delta^2 H_{RefB}$, were calculated following eq. (4) with sample being RefA or RefB water in the open or retrofitted bottles and *reference* being RefA or RefB water in the closed bottles. We observed no significant change in the isotopic composition of samples in retrofitted bottles (both $\Delta^2 H_{RefA}$ and $\Delta^2 H_{RefA} \approx 0$ %, red filled markers in Figure 4c-f; results for δ^{18} O were similar, see Figure S2), whereas the isotopic composition of samples in open bottles changed over the course of the experiment by up to 10 % (blue open markers in Figure 4c-f). The observed change 335 in isotopic composition differences of samples in open bottles was more pronounced for smaller sample volumes (comparison of circles and diamonds in Figure 4c-f). It was also larger in the outdoor setting compared to indoor conditions (comparison of Figure 4c, e with Figure 4d, f), even though the average temperature was lower in the outdoor setting. This may indicate that the average temperature is less important for causing isotope effects than the magnitude of the temperature fluctuations that may trigger gas volume exchanges. In contrast to Experiment 1 (ventilated chamber), Experiment 2 was 340 characterized by significant daily temperature fluctuations (20 °C and more inside the autosampler and; that was situated outdoors outdoor setting; Figure 4a). During the daytime when the air temperature increased, the air inside the autosampler ISCO expanded and some of the (moist) air was pushed out of the device; when temperatures dropped (at night), the opposite happened, i.e. the air inside the autosampler ISCO-contracted and sucked in fresh air from outside. This "sampler breathing" probably happened on a daily basis, resulting in a greater air exchange with the outside, which in turn may have 345 reduced the humidity inside the autosampler ISCO and resulted in more evaporation (Figure 4b). In addition, the greater potential for wind-driven ventilation of the outdoor-autosampler located outdoors ISCO-may also have enhanced evaporative fractionation effects in the open sample bottles, or it may reflect the greater potential for wind-driven ventilation of the outdoor ISCO. Samples which underwent a stronger change in isotopic composition also experienced a greater loss of sample volume between the start and end of the experiment. We observed a larger decrease in sample volumes in open 350 bottles compared to the retrofitted sample bottles, a larger decrease in sample volumes in the outdoor setting compared to the indoor setting, and a larger relative decrease in the 200 ml samples compared to the 400 ml samples (see Figure S3 in the

Supplement).

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In the outdoor setting, the samples in the open bottles became isotopically heavier, with larger changes observed in the 200 ml samples than in the 400 ml samples (e.g., δ^2 H in RefB water increased by 9.6 ‰ and 3.9 ‰ in 200 ml and 400 ml

samples, respectively; Figure 4e), likely due to evaporative fractionation. For the open bottles in the indoor setting, however, RefA samples became isotopically lighter by about 1 % in δ^2 H (-1.2 % and -0.9 % for 200 ml and 400 ml, respectively; Figure 4e) while RefB samples became roughly 2 % heavier (2.6 % and 1.6 % for 200 ml and 400 ml, respectively; Figure 4f). The isotopic lightening of RefA samples may be explained by mixing in the vapor phase of isotopically heavier RefA water with isotopically lighter RefB water, and subsequent condensation in both samples. This isotopic exchange should make RefA samples isotopically lighter and RefB samples isotopically heavier, in addition to any isotopic fractionation due to net evaporative losses from both samples. Thus, a large part of the observed enrichment in RefB water in the indoor setting may have been due to isotopic exchange with the heavier RefB-RefA water, in addition to any evaporative fractionation. A likely reason why the mixing effect was only visible in the indoor setting may be that evaporation was smaller compared with the outdoor setting; the greater evaporative losses (and thus evaporative fractionation) in the outdoor setting may have overprinted the vapor mixing effect. In either case, mixing and/or evaporative

- fractionation only affected the isotopic composition in the open sample bottles, while no measurable effect was observed in samples from the retrofitted bottles.
- 370 We can quantify the isotopeie changes effects due to mixing and evaporative fractionation in the different settings under the assumption that a) evaporative fractionation and mixing have additive effects, b) the per mil change due to evaporative fractionation is the same for RefA and RefB waters, and c) mixing has an exactly inverse effect on the two waters (i.e., it results in the same degree of isotopic depletion in the heavier RefA water and enrichment in the lighter RefB water):

 $m_{RefA} + f = \Delta^2 H_{RefA}$ 375 $m_{RefB} + f = \Delta^2 H_{RefB}$ (6) $m_{RefA} = -m_{RefB}$ (7),

(5)

where mixing-induced isotopic change is denoted by m for RefA and RefB waters, and the change in isotopic composition due to evaporative fractionation is denoted by f.

380 The results of this analysis are illustrated in Figure 5 for $\delta^2 H$, confirming that isotope effects due to mixing and evaporation are small in samples from the retrofitted bottles (red filled markers). In the open bottles (blue open markers), the isotope effects due to evaporative fractionation were between 1.5 to 2 times larger than the isotopic change due to mixing in the outdoor setting, but fractionation was less important than mixing in the indoor setting. Both the mixing- and fractionationinduced isotope effects were roughly twice as large in the 200-ml samples volumes compared tothan in the 400-ml samples volumes (diamonds vs. circles, respectively). Applying eqs. (5)-(7) to δ^{18} O yielded similar results (see Figure S4 in the 385 Supplement).

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In summary, the results of Experiment 2 confirmed the findings from Experiment 1 that the retrofitted ISCO bottles efficiently protected the collected water samples from undergoing isotopic changes due to both evaporative fractionation and vapor mixing.

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3.3 Evaluation of the evaporation protection in the field

During the field experiment (Experiment 3, October 2015 to June 2019), we observed distinct seasonality in air temperature, but no seasonal pattern in relative humidity at both field sites, and slightly higher temperatures and humidity at the EIN site (Figure 6Figure 6). At both field sites, the vapor pressure deficit (VPD), which is strongly correlated with air temperature, peaked around June and July and was lowest in December and January. Because these climatic variables exhibited very similar behavior at both field sites, we decided to pool the isotope data sets from both sites for analysis. The isotope data from 8 March 2016 were excluded from this analysis because the water samples in the retrofitted bottles were isotopically lighter than the reference water (e.g., Δ^2 H ranged from -4.7 to -3.5 ‰) for unknown reasons. In addition, we removed the data points from 14 February 2017 from our analysis because of an anomalous δ^{18} O measurement of the water sample from the retrofitted bottle at the ERL site.

The field experiment with the open and retrofitted bottles in two ISCO samplers Experiment 3 resulted in 244 usable samples (i.e., 61 samples per site and bottle type) for which the storage duration varied between 12 and 23 days. The isotopic differences of the samples in open bottles relative to the reference water exhibited substantial scatter, substantially with values between --2‰ and +3.5‰, but the average isotopic differences (mean±1 standard error) were 1.45±0.22 ‰ for Δ^2 H 405 and 0.27 \pm 0.05 ‰ for Δ^{18} O, and thus deviated statistically significantly from zero. Conversely, when the retrofitted bottles were used, the isotopic differences were statistically not significantly larger than zero, i.e., 0.10 ± 0.11 % for Δ^2 H and 0.05 ± 0.03 % for $\Delta^{18}0$. Figure 7 Figures 7 and Figure S5 show that the isotopic differences of the samples in open bottles relative to the reference water wereas positively correlated with average air temperature, and thus with VPD. Pearson 410 correlation coefficients between air temperature (average, minimum and maximum) and $\Delta^2 H$ were r>0.70 (p<0.001), and r > 0.60 (p < 0.001) for the same relationships with Δ^{18} O. For VPD, the correlation coefficients were r > 0.56 (p < 0.001) for both isotopes. No statistically significant relationships were evident for the samples from the retrofitted bottles (r < 0.17, p>0.1 for both isotopes; see also the red data points in Figure 7Figures 7 and S5). Similarly, higher climatic variability at our field sites (represented by the maximum changes in air temperate and relative humidity (as an indicator for climatic 415 variability at our field sites) were positively associated with correlated with the larger isotopic differences only for the open bottles (the relationship was statistically significant only for the change in air temperature, with r > 0.40 and p < 0.0001 for both isotopes; Figure 8Figure 8 and Figure S6).

Overall, our results indicate that the retrofitted sample bottles significantly reduced isotopic fractionation compared to the
 open sample bottles when deployed over two- to three-week periods under the ambient climatic conditions at our two field sites.

4 Practical implications

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In the three experiments presented above, we assessed how storage duration, temperature and humidity fluctuations, a<u>s well</u> asnd sample volume influenced isotopic shifts due to evaporative fractionation and vapor mixing in samples stored inside the ISCO autosampler. In all three experiments we found that the observed change in isotopic composition was substantially smaller in sample<u>s stored in</u> bottles <u>that were</u> retrofitted for evaporation protection.

We can use the relationship between isotopic fractionation and air temperature from Experiment 3 to estimate the expected 430 isotopic change in the water samples collected in the laboratory during Experiment 1. If we apply the linear regression slopes shown in Figure 7 Figure 7 b to estimate calculate the expected isotopic change difference in the samples in open bottles at the average air temperature of 35 °C maintained during Experiment 1, we obtain an expected change in isotopic composition of Δ^2 H Δ^2 H=9.4±1.1 ‰ (± 1 standard error) and Δ^{18} O Δ^{18} O=1.6±0.3 ‰. These values estimates are substantially larger than the measured differences obtained after 24 days of Experiment 1 (i.e., $\Delta^2 H=5 \% \text{ in } \Delta^2 H$ and $\Delta^{18} O=1$ 435 $\% \frac{1}{10} \frac{1}{10}$ Sect. 3.1). The larger isotopic change observed during the field deployment of Experiment 3 may be attributed to more variable climatic conditions (e.g., due to diurnal temperate variations) causing "sampler breathing", and possibly also to variable ventilation by wind, whereas during Experiment 1 the sampler was placed in a windless chamber with constant less variable temperature and relative humidity during Experiment 1. This hypothesis is supported by the results from Experiment 2, where isotopic differences were larger in the open bottles in the outdoor setting; here, the average 440 temperature was lower in both the indoor and outdoor settings compared to Experiment 1, but the change in isotopic composition was comparable (in Experiment 1 we observed an increase in δ^2 H of approx. 3 ‰ in the open sample bottle with a storage duration of 21 days; in Experiment 2, compared to the indoor setting (e.g., Δ^2 H was the increase was 1.7-3.9 ‰ and vs. 1.64 - <u>min the indoor and outdoor settings</u>, respectively, for the 400--ml RefB samples of nopen bottles-RefB). Larger temperature and humidity contrasts due to diurnal fluctuations in outdoor conditions may have resulted in repeated 445 evaporation and condensation inside the ISCO-autosampler housing and "sampler breathing", and in enhanced vapor exchange between the sample bottles and the outside atmosphere ("sampler breathing"), in contrast to the very stable conditions during Experiment 1.

Our evaporation protection reduced the contact area between the water surface in the sample bottle and the atmosphere inside the ISCO autosampler by a factor of approximately 5500 (comparing the cross-sectional area of the bottle to that of

the silicone tube attached to the syringe), and also the area for diffusion of vapor through the bottle opening by a factor of approximately 1300 (comparing the area of the bottle opening to the cross-sectional area of the silicone tube). Consequently, isotopic fractionation and mixing should be substantially reduced in samples in retrofitted bottles compared to those in open sample bottles. However, because the syringe housing does not entirely seal the ISCO sample bottle (because air needs to be released when water samples are introduced into the bottle), some vapor exchange may still occur between the sample bottle and the <u>atmosphere</u> inside of the <u>ISCO-autosampler compartmenthousing</u>. This vapor exchange will likely be stronger if air temperature is high and relative humidity inside the <u>ISCO-autosampler</u> housing is low (Experiment 1). Experiment 2 also suggested that strong diurnal variations or windy conditions may occur during extremely dry and warm summer days so that automatically collected water samples should be retrieved sooner than 24 days if possible. However, Experiment 1 showed that in the absence of wind, the relative humidity inside the <u>ISCO-autosampler</u> can build up over time, even if the relative humidity outside is very low.

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We furthermore showed that in open bottles, 400-ml samples exhibited smaller isotope effects than 200-ml samples, simply
because the ratio between the water volume affected by mixing and fractionation (i.e., the uppermost water layer that is in
exchange with the atmosphere) and the total sample volume is two times smaller for the 400-ml sample than for the 200-ml sample (Experiment 2). We therefore recommend that streamwater samples collected with our evaporation protection
method should comprise at least 400 ml (but note that due to the narrow silicone tube, care has to be taken to not exceed a filling rate of approximately 100 ml min⁻¹). When collecting precipitation samples, larger sample volumes can be achieved
by using larger funnels; e.g., 1 mm of rain collected with a 45-cm diameter funnel results in approximately 160 ml sample volume, while a 20-cm diameter funnel would only yield around 30 ml. Control samples with known isotopic composition in open, retrofitted and closed bottles, placed in the autosampler for the entire storage duration, should be used to monitor composite isotope effects and to allow for a retrospective quality assessment of the automatically collected samples.

While we have discussed the performance of the retrofitted ISCO 1-litre sample bottles with respect to stable water isotopes, the new bottle design may also be useful for water quality studies. Experiment 2 showed that evaporation from open sample bottles result<u>eds</u> in reduced water sample volumes, implying evapo-concentration of solutes in the samples. The importance of this effect likely depends on the storage duration and the sample volume, and will therefore be greater for small samples and for samples collected early on in the sampling period. Consequently As a result, water quality data from water samples automatically collected in open bottles and stored over periods of days and weeks may not be directly comparable. The results from Experiment 2 suggest that our retrofitted sample bottle may reduce evapo-concentration effects in the water samples. To further adapt the presented evaporation protection for water quality studies, our design modification could also be combined with a gravitational filtration system similar to the ones presented by(e.g., Kim et al., -(2012) added

between the syringe outlet and the silicone tubing₁₇ but further studies would be needed to assess this <u>filtration approach</u> in a more systematic manner.

5 Conclusions

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We tested whether retrofitting the <u>1-litre</u> sample bottles of the conventional <u>ISCO-6712</u> Full-size Portable Sampler (Teledyne ISCO., Lincoln, US) with a modified syringe housing and silicone tube reduces evaporative fractionation and vapor mixing in water samples collected for subsequent analysis of-stable water isotopes <u>analysis(δ^2 H, δ^{18} O)</u>. Laboratory and field tests under different temperature and humidity conditions showed that water samples in <u>retrofitted</u> bottles with evaporation protection-were far less altered by evaporative fractionation and vapor mixing than samples <u>stored</u> in conventional open bottles.

We showed that retrofitting sample bottles with a modified syringe housing and silicone tube is a cost-efficient approach to
 upgrade autosampler bottles to protect water samples from isotopic fractionation during storage in the field. The setup
 described here can likely be adapted without difficulty (e.g., by using a different syringe size) to be compatible with bottles
 in other autosamplers, such as the Maxx P6L-Vacuum System (Maxx GmbH, Rangendingen, Germany), or the smaller ISCO
 6712C and 3700C Compact Portable Samplers (Teledyne ISCO., Lincoln, US) that use 500_-ml sample bottles. These
 adapted evaporation protections will require further testing, because the observed results partly depend on the size of the air
 space and thus the buildup of humidity inside the automatic-sampler compartment, which influences the buildup of humidity
 inside the autosampler. _Different systems autosampler designs may also be more or less tightly sealed from the surrounding atmosphere, likely resulting in stronger or weaker-differing rates of vapor exchange.

ISCO-Conventional automatic water samplers are generally available in many laboratories, but researchers may be reluctant to use them for isotope studies due to a fear of evaporative fractionation and vapor mixing occurring in the water samples, particularly if sample volumes are small, and/or-weather conditions are dry and warm, and/or samples are stored for multiple weeks. We showed that retrofitting 1-litre ISCO sample bottles with a modified syringe housing and silicone tube iscan be a cost-efficient approach to upgrade the 6712 Full-size Portable Samplerautosampler so that bottles to protect-water samples are protected from isotopic fractionation during storage in the field-during storage in the field. Our This inexpensive and robust method may thus provide a new possibility for water sample collection at remote locations can be used to

510 robust method <u>may thus provide a new possibility for water sample collection at remote locations can be used to</u> substantially reduce evaporative fractionation in water samples collected at daily or sub-daily frequency frequencies over periods of up to 24 days.

Data availability

The stable water isotope measurements from the three experiments are provided as supplementaryal material information.

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Author contributions

JF, JLAK and JWK <u>conceptualized_designed</u> the study. JF, JLAK, AR and BS performed the experiments and analyzed the isotope data, JF, JLAK, and AR analyzed the datasets, JF and JLAK prepared the manuscript with contributions from all coauthors.

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Competing interests

The authors declare that they have no conflict of interest.

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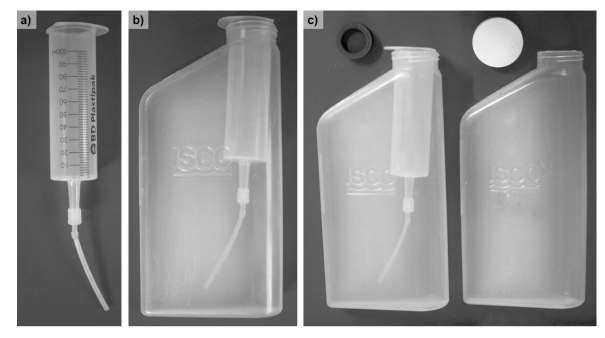
600 Tables

Table 1: Isotopic changes differences between a reference water and in the water samples in a closed, open and retrofitted relative to the references water in three different bottles that were stored outside the ISCO <u>auto</u>sampler during Experiment 1. Isotopic changes differences are expressed as mean±1 standard deviation.

| Bottle type | Storage duration [days] | Δ ² Η [‰] | Δ ¹⁸ Ο [‰] |
|-------------|-------------------------|----------------------|-----------------------|
| Closed | 24 | -0.42 ± 0.27 | -0.08 ± 0.12 |
| Retrofitted | 24 | 9.05 ± 0.54 | 1.63 ± 0.15 |
| Open | 12 ^a | 100.46 ± 1.04 | 21.97 ± 0.20 |

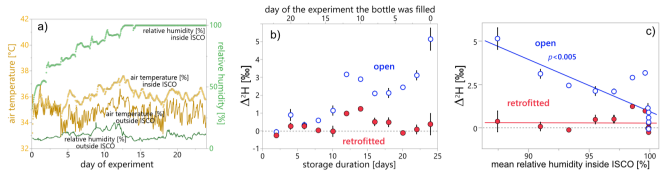
^a For the open bottle, the change in isotopic composition between day 0 and day 12 is provided, because the water sample was fully evaporated from the open bottle by day 24.

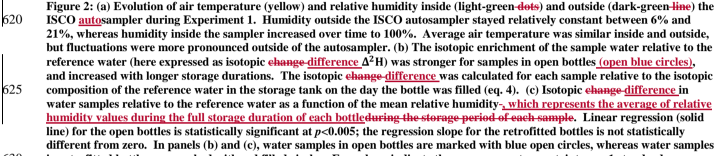
Figures



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Figure 1: (a) Modified 100 ml syringe housing with Luer-tip adapter and fitted silicone tubing for extending the syringe outlet towards the bottom of the bottle; the barrel flange at the syringe housing was trimmed on the outer side to ensure the retrofitted bottles fit into the autosampler. (b) Retrofitted sample bottles with evaporation protection using the modified syringe shown in (a). (c) The sample bottles with evaporation protection can be sealed for transport with black rubber piston stoppers (left); bottles 5 without evaporation protection can be sealed with a screw lid (right).





630 in retrofitted bottles are marked with red filled circles. Error bars indicate the measurement uncertainty as ±1 standard deviation.

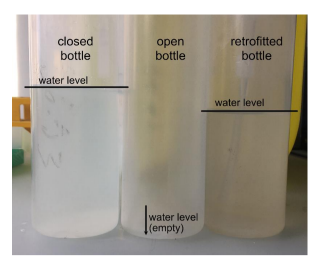


Figure 3: Water levels in the <u>control three</u> bottles <u>stored outside of the autosampler after on the 24-day laboratory experiment</u> (<u>day 24 of</u> Experiment 1). The water level in the closed bottle is identical to the water level in all three bottles at the start of the experiment. By the end of the experiment the water level in the retrofitted bottle had only decreased slightly, while the water from the open bottle was completely evaporated after approximately 12 days.

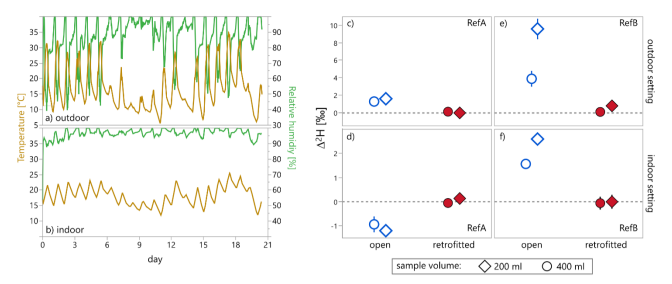




Figure 4: (a,b): Temperature (yellow) and relative humidity (green) measured inside the ISCO <u>auto</u>samplers that were located outdoors (a) and indoors (b) over the 21 days of Experiment 2. (c-f): Mean change in isotopic composition of samples relative to the reference waters. Each data point is calculated from the three replicates of each combination of the two reference waters (RefA vs. RefB), the two sample volumes (200 vs. 400 ml), and <u>the</u> two bottle types (open vs. retrofitted with evaporation protection). Please note the different y-axis scales between panels c), e) and d), f). Error bars denote standard errors of the three replicates of each condition, and account for measurement uncertainty and the standard error of the sample means.

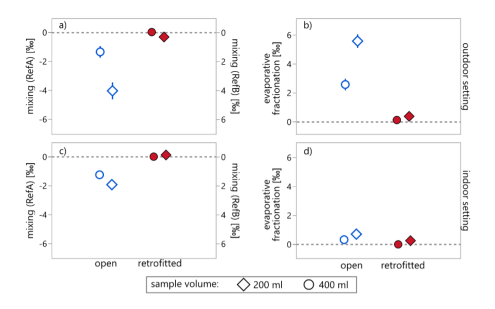


Figure 5: A comparison of the isotope effects due to mixing (panels a and c) and evaporative fractionation (panels b and d) in 650 water samples stored outdoors (panels a and b) and indoors (c and d) during Experiment 2. Both mixing and evaporative fractionation effects are small in samples from the retrofitted bottles (filled red markers), and larger in samples from the open bottles (open blue markers). In addition, the isotope effects were larger for the 200 ml samples (diamonds) than for the 400 ml samples (circles). Error bars indicate standard errors determined from the three replicates of each combination of the two reference waters (RefA vs. RefB), the two sample volumes (200 vs. 400 ml), and two bottle types (open vs. retrofitted with 655 evaporation protection), and they account for measurement uncertainty and the standard error of the sample means.

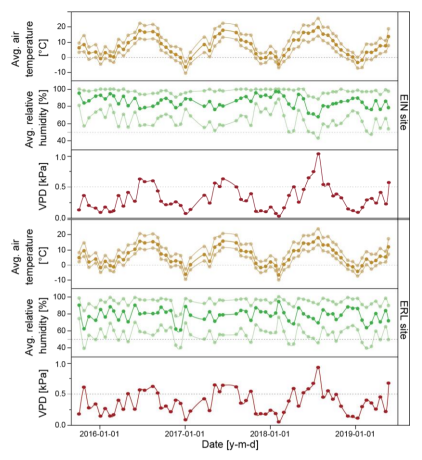


Figure 6: Mean, minimum, and maximum values of air temperature and relative humidity, as well as vapor pressure deficit (VPD), averaged over two- to three-week storage periods during Experiment 3 at the two field sites EIN and ERL. Dashed horizontal lines in each panel indicate 0 °C air temperature, 50 % relative humidity and 0.5 kPa VPD for easier comparison between sites.

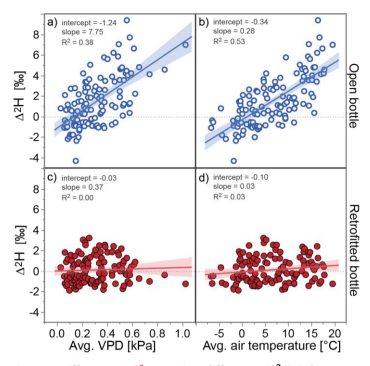


Figure 7: Differences in δ^2 H-euterium differences (Δ^2 H) in between samples stored in open and retrofitted in the ISCO bottles and their accompanying completely closed relative to a reference water samples over storage periods of 12-23 days during Experiment 3 at the EIN and ERL sites, and their relationships with the average vapor pressure deficits (VPD) and the average air temperatures during the respective storage periods at the EIN and ERL sites. Samples in oOpen bottles (open blue circles) show a substantial isotopic enrichment with higher VPD and air temperature, whereas samples in retrofitted bottles (filled red circles) do not indicate a systematic fractionation effect. No relationship with relative humidity was found. The uncertainties of the individual Δ^2 H values were on average 0.52 ‰; linear regression fits are indicated by solid lines, as well as and slope, intercept and R^2 values; the shaded areas represent the 95 % confidence intervals of the fitted lines.

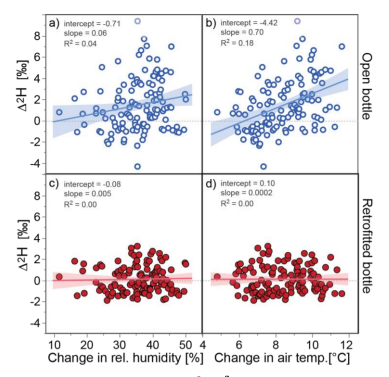


Figure 8: Deuterium differences in δ^2 H-(Δ^2 H) between in samples stored in open and retrofitted in the ISCO bottles relative to aand their accompanying completely closed reference water samples over storage periods of 12-23 days during Experiment 3 at the EIN and ERL sites, and their relationships with the maximum changes in relative humidity and air temperature within the 675 respective storage periods at the EIN and ERL sites. Large changes in relative humidity resulted in some isotopic enrichment in the samples stored in open bottles (open blue circles), but not in the retrofitted bottles (filled red circles). Samples in oOpen bottles showed the strongest isotopic enrichment when temperature contrasts were large (>10°C), whereas samples in retrofitted bottles seemed to be unaffected by temperature changes. The relationship between the change in air temperature and $\Delta^2 H$ of samples in open bottles was statistically significant (r=0.42, p<0.0001). The uncertainties of the individual Δ^2 H values were on average 680 0.52 %; linear regression fits are indicated by solid lines, as well as and slope, intercept and R² values; the shaded areas represent the 95 % confidence intervals of the fitted lines.