

Interactive comment on “Isotopic approaches to quantifying root water uptake and redistribution: a review and comparison of methods” by Yuri Rothfuss and Mathieu Javaux

M. Sprenger (Referee)

General comments

- 5 The manuscript by Rothfuss and Javaux on quantifying root water uptake by the means of isotopic approaches aims to provide both an overview of methods and a comparison of the methods with regard to their limitations for the interpretation. They further propose to include modeling approaches to better estimate the root water uptake patterns.

- 10 The manuscript is generally well prepared, with mostly sufficient references, in depth information, and proper visualizations. Due to its nature as a review, it is pretty long and I am not sure if the sections 2.1 and 2.2 are really necessary. I agree that it is necessary to understand the soil water isotopic composition in order to interpret the root water uptake with the means of stable isotopes. The authors focus on evaporation fractionation as one process to alter the soil water isotopic composition. From my point of view, also the precipitation input (and its variability in time) would then need to be considered. However, this has been reviewed recently and would blow up the manuscript.

I think that the manuscript is a good contribution to current issues in ecohydrology and will be of interest to a broad readership. Therefore, I suggest a publication after a minor revision.

Specific comments

- 20 On page 5, where you introduce into the theoretical backgrounds, I do not think that the isotope depth profiles are solely a result of fractionation effects. It seems that you miss the importance of the variability of the isotopic signal of the precipitation input and its consequences for the spatial variability of the soil water isotopes over depth.

Dear Matthias, we fully agree with this. A text was added (now P6 L4-6) on how soil upper-boundary isotopic condition (i.e., δ_{surf}) - one of the determinants of the soil water isotopic composition profile – is highly impacted both spatially and temporally by input precipitation isotopic composition.

- 25 As you write, the evaporation fractionation has been reviewed by Horita et al. (2008) and it was more recently reviewed by Soderberg et al. (2012). In order to streamline the manuscript and keep its focus on the root water uptake, I doubt that the section 2.1 and 2.2 are really necessary. However, I agree that it is necessary to account for the spatial and temporal variability of the soil water isotopes, but this is influenced by more than soil evaporation (see also my review Sprenger et al. (2016)).

- 30 *Sub-section 2.1 was removed from the manuscript, sub-sections 2.2 and 2.3 were shortened, merged together, and renamed “Flow of isotopologues in the soil-plant-system”. About the remark that “spatial and temporal variability is influenced by more than soil evaporation”, please refer to our answer to your specific comment below.*

- 35 I am wondering if the authors are aware of the work by Ogle et al. (2004) and Ogle et al. (2014), where they suggest to include biophysical conditions in a process-based mixing model (“root area profile and isotope deconvolution, RAPID”). I think it would be worth including this in the review of methods, since it goes in the same direction as the author’s proposal of including more physical basis of the root water uptake estimates.

- 40 *The authors are aware of the work of Ogle et al. (2004) where they could reconstruct “active root area” and RWU profiles from isotopic measurements (assuming, amongst other things, normal a priori distributions for the xylem water oxygen and hydrogen isotopic compositions and considering prior knowledge on x). Since we wanted to focus on the most used and cited methods for partitioning plant water sources, we choose not to incorporate that of Ogle in the model comparison (and for that same reason neither that of Romero-Saltos et al. (2005)). Nevertheless, we added their work in Section 5*

(Challenges and progresses), sub-section “Call for a coupled experiments-modelling approach for determination of plant water sources and redistribution on the basis of isotopic data” as it nicely fits there as you point out.

5 Please state that the recently published dependency on the carrier gas was found for a WS-CRDS (Picarro); I did not find this CO₂ dependency for Off-axis ICOS (Los Gatos) (currently in review).

Done. Thanks!

Technical corrections

P1 L12/13: I suggest using “studies” rather than “authors”

Done

10 P3 L11: Insert “and” for “in space,” and replace “but also on the root’s” with “and their”

Done

P3 L25: Not sure what you mean with “by reference”

We meant that the letter “S” makes reference to the “sink term”. We will use the correct formulation “in reference to” instead of the incorrect “by reference to”.

15 P3 L32: I suggest “distribution of S” instead of “S distribution”

Done

P3 L23: Introduce RLD here.

Done (now P3 L23).

P4 L21: Why not stating directly that Zarebanadkouki et al. (2012) used deuterated water?

20 *Done (now P4 L23)*

P4 L29: I suggest (or artificial enriched/depleted)

Since “artificial” stands for “isotopic abundance”, it cannot be followed by “enriched or depleted” but only “higher of lower” but then it reads a bit funny. We would like to keep the more general “artificial”

P5 L1: Please state here once again what “these methods” will be.

25 *Done*

P5 L8: Is this not a bit too simplified at this point? You would always also need some kind of info about vegetation isotopes. I don’t think the first sentence is necessary here.

The authors write that for reconstruction of S profiles, one needs both soil and plant isotopic information (“within the soil-plant system”).

30 P5 L9: Are you referring to S profiles or isotope depth profiles? I do not think that the isotope depth profiles are solely a result of fractionation effects.

We are referring to isotopic variations. It is implied here that these isotopic variations are solely due to the difference of physical properties of the different isotopologues (ultimately leading to isotopic fractionation) only in-between precipitation events, i.e., we rule out the partial to total reset of the isotopic profile due to each rain event.

35

We added the following sentence (now P6L4-6): “Finally, under natural conditions, the δ_S profile is not solely a result of isotopic fractionation but is as well highly impacted both spatially and temporally by input precipitation isotopic composition through modification of the upper boundary condition (δ_{surf}).”

P5 L 16: Consider splitting this long sentence.

5 *The entire sentence was removed from the current version of the manuscript*

P6 L 30: This line is not black anymore.

Thanks! Done

P7 L5: I think it would be worth including the definition by Barnes, Allison (1983) for the vapor region to be of total water potential about 15bar, while at lower potential there would be little connected water.

10 *Done (now P5 L22)*

P7 L8: This line is not grey anymore

Thanks! Done

P7 L15: Is this supported by data? I don't see it like that in Rothfuss et al. (2015), where you have shown that the slope for the depths above the EF (max -0.06 m) is still clearly below 8.

15 *We write that “As a result, an intermediate value for the slope is expected, depending on the mixing ratio of atmospheric water vapor to evaporated soil vapor at a given soil depth.” (now P6 L3-4), meaning that the value for slope should lie between 2 and 8 (which is in accordance with Rothfuss et al. (2015)).*

P7 L16: I have shown that for several studies in Sprenger et al. (2016).

20 *Indeed. The list of three references has been simply replaced with your review “Sprenger et al. (2016)” (now P6 L3)*

P10 L8: delete second “is”.

Thanks! Done

P10 L12: I summarized the uncertainty of the different methods to derive soil water isotope data in Sprenger et al. (2015b).

25 *The reference is cited now. Done (now P9 L2)*

P14 L25: I assume this is for $\delta^{18}\text{O}$? Please clarify.

Yes. We added “(δ_S)” after “soil water oxygen isotopic composition” (as well as “(T)” after “actual transpiration rate”) above in section 4.1.1 (now P12 L24)

30 P14 L24: It would be interesting which rooting depth and density profile was assumed for the modeling. Please provide.

This is explained (P12 L28-30):

“[All scenarios] relied on a common measured root length density vertical distribution of Festuca arundinacea”,

and in Appendix B2 (Running the model for the inter-comparison, P45 L15-17):

35 *“For this, H_S , δ_S , and RLD input data were interpolated at a 0.01 m vertical resolution...”*

P16 L2: cases

Done

P17 L28: What about the study by Dawson (1993) who provides volumes?

To the authors' understanding, Dawson (1993) provided a lateral gradient of the proportion of hydraulically-lifted water from maple trees used by neighboring plants, not HL water volumes.

5 P18 L27: Why do you not include the direct-equilibration method by Wassenaar et al. (2008)?

Done (now P17 L30)

P19 L7: I did not use in-situ in that study, but consider including a recent paper by Oerter et al. (2016).

Done (now P18 L12)

10 P19 L14: Please not that this is the case for a WS-CRDS (Picarro); I did not find this CO₂ dependency for Off-axis ICOS (Los Gatos) (currently in review)

Done (now P18 L19-20)

P19 L25: I suggest referring to Farquhar et al. (2007).

Done (now P18 L29)

15 Table 1: I believe there are more studies than the listed ones. I suggest considering the following: Meinzer et al. (1999); Kulmatiski et al. (2010); Kulmatiski, Beard (2013); Evaristo et al. (2016); Goldsmith et al. (2012); Liu et al. (2011); Bertrand et al. (2012); Meißner et al. (2012); Dawson (1996); Bijoor et al. (2012)

The authors agree! However we had to make a choice and focus on the non-exhaustive list of papers that were reviewed here... Note that, on the other hand, another reviewer says that Table 1 is too long in its current form.

20 Table 2: Why did you limit your analysis here to $\delta^{18}\text{O}$, while emphasizing that dual isotope approaches would be preferable on page 12 L4?

In the text (now P10 L24-28) we indeed underline the potential of the dual isotope approach:

25 *“As infrared laser-based spectrometry now enables simultaneous measurements of $\delta^{18}\text{O}$ and $\delta^2\text{H}$ at lower cost, we believe that this dual-isotope approach (referred as “D” in Table 1) will or should gain in importance in isotopic studies. This is especially useful when (i) under natural conditions the $\delta^{18}\text{O}$ - $\delta^2\text{H}$ slope is not constant over depth (Sprenger et al., 2016) or (ii) in the context of pulse labelling experiments, which can artificially change the value of the $\delta^{18}\text{O}$ - $\delta^2\text{H}$ slope at given locations in the soil profile. In these cases, two independent mixing equations are obtained, one for each isotopologue.”*

30 *As we wanted to concentrate on the different methods rather than on the consequences of the quality of the input data, we decided to go with a single isotope approach. But this would constitute, without a doubt, an interesting follow-up paper.*

A justification of the choice of single isotope approach was added (Now P12 L16-18):

“The inter-comparison of models was performed using a single isotope (^{18}O) approach as the focus here was the differences of outcomes rather than the impact of the input isotopic data on these results.”

35 Figure 1: Caption “negative towards the surface”

Done

Figure 2: Update the caption according to the color of the lines (blue).

Done

Figure 4: Caption “a detail is is presented for”

Done

Figure 5: Why standard deviation and standard error for the different approaches?

5 *Error bars for the RWU analytical model refer to the standard deviation associated with relative contributions to transpiration (x) across the 1000 model runs. For the TM approach, error bars are standard error of x as calculated with Equation (8b). This is now specified in the caption of the figure.*

Interactive comment on “Isotopic approaches to quantifying root water uptake and redistribution: a review and comparison of methods” by Youri Rothfuss and Mathieu Javaux

Anonymous Referee #2

5 General comments: The manuscript aims to compare different methods to locate root water uptake depth. I understand that it is a review paper, however, it is rather long, way too complex, and unfortunately hard to follow (it even comes with an appendix). This is mainly due to the many formulas that are presented and which disturb the text flow.

10 *The authors present only equations that are needed to understand all three methods (graphical inference, two to n end-member mixing models) plus the physically based approach of Couvreur et al. (2012). The appendix section was also meant in this way: not to disrupt the text flow.*

I agree that a method comparison is needed but the manuscript is blown up with a lot of “basic isotope knowledge” which for my feeling is not necessary in such extent. I would suggest to reduce the length of the whole manuscript and focus on what differentiates the three methods to be compared.

15 *Sub-section 2.1 was removed from the manuscript, sub-sections 2.2 and 2.3 were shortened, merged together, and renamed “Flow of isotopologues in the soil-plant-system”. About the remark that “spatial and temporal variability is influenced by more than soil evaporation”, please refer to our answer to your specific comment below.*

Further, the authors should only present equations which are really needed to understand the method comparison. I believe that this would increase the readability.

20 *Please see the answer to you first general comment.*

Concerning the presented figures, I would recommend to keep them simpler as they should generally be self-explaining and not as complex as they are now.

25 *Could you give some specifics? How the results were illustrated is directly inspired from the reviewed literature. Even though some figures might not be straightforward to all readers at first, the authors tried as much as possible to make them self-explaining by adding a thorough description in the caption. Note that the colors in Figure 1 were changed (but not the data).*

I would consider the manuscript ready for publication after major revision.

Specific comments:

Title: “quantify“ instead of “quantifying”

30 *Done*

p. 3ff: Introduction needs a better/clearer structure

The Introduction section is articulated thusly:

- 35
- 1- RWU definition;
 - 2- RWU controlling variables and factors (e.g., concept of compensation and extreme case of Hydraulic redistribution);
 - 3- Difficulty of measuring RWU;
 - 4- Water stable isotopic compositions as a way to measure RWU;
 - 5- Mention of objectives: review and comparison of the isotopic methods.

Where/what would you propose we change?

- p. 3 l. 2-6: References are missing
Done (Javaux et al., 2013) (now P3 L6)
- p.3 l. 7: “driven by transpiration taking place. . .”
Done
- 5 p. 3 l.11: “spatial distribution. . .is very variable in time and space”; spatial in space ! avoid duplication
Done
- p. 3 l. 13-14: Reference missing; permanent wilting point concept; what is a dry soil in this context?
Sentence was rewritten (now P3 L13-15):
“The flux of water depends also on soil water availability, i.e., the ability of the soil to provide water at the plant imposed rate (Couvreur et al., 2014): a highly conductive root segment will not be able to extract water from a dry soil.”
- 10
- p. 5 l. 1: “each other” without hyphen
Done
- p. 5 l. 3-6: Repetition from abstract
This sentences state two objectives of the paper addressed in the section “challenges and progresses”. This is why the authors mentioned it in both Abstract and Introduction sections. We do not feel this constitute repetition.
- 15
- p. 5 chp. 2.1: This chapter is too detailed; fundamentals of isotope hydrology do not have to be explained in such detail
- 20 *See answer to your second general comment*
- p. 6 l. 20: Reference missing
Text is no more part of the manuscript
- p. 7 l. 8: “grey” instead of “gray”
Thanks (color wasn’t actually grey but blue).
- 25
- p. 7 l. 12-14: Repetition
Sentence was erased.
- p. 7 l. 24: Mention this earlier in the manuscript
What exactly should we mention earlier?
If you are referring to “for plants growing in homogeneous external conditions, e.g., in hydroponic solution, root xylem sap water and external water have the same isotopic compositions”, the authors think this is the right place to mention this.
If you are referring to “In natural soils where the liquid phase is not homogeneous and a vertical gradient of isotopic composition due to evaporation exists...”, it is explained earlier (now P5ff).
- 30
- p. 9 l. 16: 21 studies: Based on which criteria have these studies been selected? Literature review using ISI web of knowledge? Please mention briefly.
We explain this briefly in the text (now P8 L5-7):
- 35

“This non-exhaustive list was drawn up according to either the number of citations and contribution importance (for studies published before 2015) or to the novelty of the publications (publication year \geq 2015).”

5 p. 9 l. 20: “unambiguously identified”! What about issues with regard to water extraction techniques which might be a cause for this?

At this point of the demonstration, there is, for sake of clarity and concision, no question of techniques-related issues but rather the concept behind the graphical method (i.e., \bar{z} is the depth where the soil water isotopic profile (δ_S) equals that of the tiller water (δ_{Ti})). Extraction techniques are evocated later (section 5).

10 p. 10 l. 7: grey ! correct throughout the manuscript

Done

p. 11 l. 26: Table 1 should rather go into the Introduction section, also it is too detailed

The authors agree that Table 1 was too detailed and is now simplified (e.g., less columns). However the authors feel it should remain in Section 3 as it illustrates the studies described in this very section.

15 p. 12 l. 22: Replace www. by an abbreviation for example EPA, 2015; same for p. 13 l.

The authors’ intention was to provide the readers with direct access links to the zip files for each model (as this is generally done in the literature with, e.g., R packages).

p. 13 l. 20ff: Why did the authors not intercompare the methods based on a dual isotope approach?

20 *As we wanted to concentrate on the different methods rather than on the consequences of the quality of the input data, we decided to go with a single isotope approach. But this would constitute, without a doubt, an interesting follow-up paper.*

A justification of the choice of single isotope approach was added (Now P12 L16-18):

“The inter-comparison of models was performed using a single isotope (^{18}O) approach as the focus here was the differences of outcomes rather than the impact of the input isotopic data on these results.”

25 How reliable/meaningful is a single isotope approach?

The point of the model inter-comparison is to determine if using the different approaches is meaningful in the context of a single isotopic approach. Note that the vast majority of published studies use(d) a single isotopic approach. Reliability depends on whether or not uncertainty is properly accounted for, which we address as well.

30 p. 14 l. 26: Is it necessary to mention the function?

Mention to the function was erased.

p. 18: Think about renaming the subsection e.g. method uncertainties and. . .

Subsection “5.2 High frequency isotopic data and sampling strategies” was split and renamed. We now have

35 *“5.2 Soil and plant water sampling strategies” (P16 L27)*

and

“5.3 Off-line destructive versus on-line non destructive isotopic measurements in plant and soil waters” (P17 L15)

p. 18 l. 1-20: This does not belong into the discussion section

Certainly. Please note that this section is not about discussing results but reviewing materials and isotopic techniques with focus on new developed developments.

p. 18 l. 21: Reference missing

5 *Reference was added (“e.g., Koeniger et al., 2011; West et al., 2006”) (now P17 L24-25)*

p. 18 l. 21-24: Is this water plant available? Does it make sense to extract at such conditions if plant available soil water pools are of interest? Please discuss briefly.

Although this adsorbed water might not be directly available to plants it is in equilibrium with the bulk water and needs to be accounted for as a potential source in the root zone for modeling purposes.

10 p. 18 l. 24ff: Methods are also not intercomparable and each method comes with a huge uncertainty (e.g. Sprenger et al., 2015; Orlowski et al., 2016). How reliable is such data in the end when utilized for RWU calculations? How would RWU depth vary if e.g. extraction method uncertainty is accounted for?

Uncertainty associated with extraction techniques are explicitly accounted for in each method and addressed in details in the manuscript:

- 15
- *through the uncertainty associated with measurement of δ_{Ti} (width of the vertical band) [graphical method];*
 - *through $\sigma_{\delta_{Ti}}$, σ_{δ_s} etc [two end-members method];*
 - *through the parametrization of the approach of Phillips and Gregg (2001) (tolerance parameter) and Parnell et al. (2013) (sources ($\delta_{S,j}$) and product (δ_{Ti}) uncertainties) [multi-sources mixing models]*
- 20
- *through the sensitivity analysis [the model of Couvreur et al. (2010) was run a 1000 times, see Appendix B]*

p. 19 l. 3ff: Again, does this represent plant available water?

See answer to previous comment.

p. 19 l. 15: Gaj et al. (2015) is not a method comparison paper.

25 *The authors do not agree: Gaj et al. (2015) attempted to compare (i) on-line isotopic measurements, i.e., obtained non-destructively by sampling the soil atmosphere and analyzing with a WS-CRDS with (ii) off-line isotopic measurements, i.e., following destructive sampling and cryogenic vacuum extraction.*

Pratt et al. (2015) is wrong!

It is now Pratt et al. (2016). Thanks. Done

30 Orlowski et al. (2016) and please cite Sprenger et al. (2015) as review paper about extraction method comparisons.

Sprenger is now cited. What about Orlowski et al. (2016)? It is already cited.

p. 19 l. 27: “generalization of coupled approaches” ! What does that mean?

The sentence was reformulated as such (now P19 L2-6)

35 *“In order to fully benefit from the potential of water stable isotopic analysis as tools for partitioning transpiration flux, the authors call for the development of approaches making use of physically based models for RWU and isotopic fractionation to analyze experimental data, especially since several soil-vegetation-atmosphere transfer (SVAT) models are available that can simulate flow of isotopologues in the soil and the plant”*

Table 1: Too much information

Done: three columns were removed from the Table 1.

Table 3: Not sure if all these numbers are necessary to understand the method comparison/virtual experiments

Table 3 was simplified: we removed the columns where absolute differences between the outcome of both methods are reported (numbers are actually already mentioned in the text). Furthermore, results will be rounded to the next whole number for readability.

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- 35 Pratt, D. L., Lu, M., Barbour, S. L., and Hendry, M. J.: An evaluation of materials and methods for vapour measurement of the isotopic composition of pore water in deep, unsaturated zones, *Isotopes Environ. Health Stud.*, doi: 10.1080/10256016.2016.1151423, 2016. doi: 10.1080/10256016.2016.1151423, 2016.
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Isotopic approaches to quantifying root water uptake and redistribution: a review and comparison of methods

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Abstract. Plant root water uptake (RWU) and release (i.e., hydraulic redistribution – HR, and its particular case hydraulic lift – HL) have been documented for the past five decades from water stable isotopic analysis. By comparing the (hydrogen or oxygen) stable isotopic composition of plant xylem water to those of potential contributive water sources (e.g., water from different soil layers, groundwater, water from recent precipitation or from a nearby stream) [authors-studies](#) could determine the relative contributions of these water sources to RWU. Other [authors-studies](#) have confirmed the existence of HR and HL from the isotopic analysis of the plant xylem water following a labelling pulse.

15 In this paper, the different methods used for locating / quantifying relative contributions of water sources to RWU (i.e., graphical inference, statistical (e.g., Bayesian) multi-source linear mixing models) are reviewed with emphasis on their respective advantages and drawbacks. The graphical and statistical methods are tested against a physically based analytical RWU model during a series of virtual experiments differing in the depth of the groundwater table, the soil surface water status, and the plant transpiration rate value. The benchmarking of these methods illustrates the limitations of the graphical and statistical methods (e.g., their inability to locate or quantify HR) while it underlines the performance of one Bayesian mixing model, but only when the number of considered water sources in the soil is the highest to closely reflect the vertical distribution of the soil water isotopic composition. The simplest two end-member mixing model is also successfully tested when all possible sources in the soil can be identified to define the two end-members and compute their isotopic compositions. Finally, future challenges in studying RWU with stable isotopic analysis are evocated with focus on new
25 isotopic monitoring methods and sampling strategies, and on the implementation of isotope transport in physically based RWU models.

Keywords

Root water uptake; hydraulic redistribution; hydraulic lift; water stable isotopologues; isotope mixing model; physically
30 based root water uptake model

1 INTRODUCTION

Root water uptake (RWU) is defined as the amount of water abstracted by a root system from soil over a certain period of time. Understanding the relation between the distributions of soil water, roots, RWU location and magnitude, and root hydraulic properties is important for managing soil water and plant water status (e.g., by irrigation), developing new plant genotypes more tolerant to drought or tackling ecological questions in water-limited ecosystems, such as the competition for soil water by different plants (Javaux et al., 2013).

RWU is principally driven by ~~evaporative-transpiration~~ flux taking place in the leaves ~~(i.e., transpiration)~~ and its magnitude depends on the atmospheric evaporative demand and stomatal opening. The latter depends amongst others on leaf water status and stress hormonal signals from the roots transported to the leaves (e.g., Huber et al., 2015; Tardieu and Davies, 1993). Leaf water status and hormonal signals are related to the soil water potential distribution and to the plant hydraulic architecture (Huber et al., 2015). The ~~spatial~~ distribution of RWU is very variable in time ~~in-and~~ space, depends on the presence of roots ~~but also on the root's~~ and their ability to extract water. This ability is a function of radial conductivity but axial conductance may also limit water flow in younger roots or when cavitation occurs. The flux of water depends also on soil water availability, i.e., the ability of the soil to provide water at the plant imposed rate (Couvreur et al., 2014): a highly conductive root segment will not be able to extract water from a dry soil. Locally, this is the difference of water potential between the root and the soil which drives RWU, and its magnitude is controlled by the radial hydraulic resistances in the rhizosphere, at the soil root interface and in the root system (Steudle and Peterson, 1998). The actual RWU profile is thus a combination of different aspects: the root ability to extract water (characterized by the amount of roots and their hydraulic properties), the ability of the soil to fulfill the plant water demand, and the water potential difference between soil and root (Couvreur et al., 2014).

Plants have numerous mechanisms to cope with heterogeneous soil water distribution, and adapt their RWU rate distribution: adaptive root growth, adaptive root conductivity (Javaux et al., 2013) ~~(Javaux et al., 2013)~~, exudation (Carminati et al., 2016). A particular process, which has attracted the attention of plant breeders and ecologists is the ability of plants to extract water from non or less water limited soil areas with potentially low root lengths densities (RLD [$L L^{-3}$], usually expressed in $cm\ root\ per\ cm^3\ soil$), known as root water uptake compensation (Heinen, 2014). To describe the RWU rate in soils, we will use the root water uptake flow per volume of soil, defined as $S [L^3 L^{-3} T^{-1}]$ ~~by-in~~ reference to the sink term of the Richardson (1922) equation (Vereecken et al., 2016). According to Couvreur et al. (2012), root compensation is defined in the present article as the process that decreases or increases RWU at a certain location compared to the water uptake from that location when the soil water potential would be uniform in the root zone. Thus, the distribution of the $S(x,y,z)$ is a sum of two spatially distributed components:

$$S(x, y, z) = S_{uniH}(x, y, z) + S_{comp}(x, y, z) \quad (1)$$

where x , y and z are the 3-D spatial coordinates, S_{uniH} is a term proportional to the root distribution and S_{comp} the compensatory part of the RWU distribution. The first term on the right-hand side of Eq. (1) is always positive while the

second one can be either positive or negative. Figure 1 illustrates how this equation affects S distribution in a one-dimensional (1D) space. When there is no compensation ($S_{\text{comp}}(x,y,z) = 0$), the RWU distribution follows the root distribution (i.e., highest at the surface and lowest in the deepest layer, Fig. 1a). When $S_{\text{comp}}(x,y,z) < 0$ but its absolute value is lower than $S_{\text{uniH}}(x,y,z)$, then $S(x,y,z)$ is positive and different from the root vertical distribution. In case $S_{\text{uniH}}(x,y,z)$ is small, as in Fig. 1c, $S_{\text{comp}}(x,y,z)$ can locally be higher in absolute value and $S(x,y,z)$ can be locally negative which implies that there is a water efflux out of the root.

The water efflux at certain locations is called root hydraulic redistribution (HR, Burgess et al., 1998) or hydraulic lift (HL, Richards and Caldwell, 1987) as a specific case of HR in which fluxes in the root system are vertically upward. In their review, Neumann and Cardon (2012) discussed that the magnitude of HR observed in different studies varied from 0.03 mm d^{-1} (Brazilian Cerrado, Scholz et al., 2010) to 3.50 mm d^{-1} (*Artemisia Tridentata*, Ryel et al., 2003). Several authors have also raised the question of the “ecohydrological interest” for a plant to release water to the upper/dryer soil layers, therefore potentially providing water to shallow-rooted plants and enhancing competition for space and nutrients. Some studies suggested that HL could increase nutrient mobility and enhance biogeochemical processes by providing moisture to the dryer soil layers (Caldwell et al., 1998; Prieto et al., 2012; Snyder et al., 2008).

Despite its importance, there is a lack of measurements of RWU, related to the difficulty of measuring root and soil water fluxes. Often soil water content change is used as a proxy for RWU. Yet, as change of soil water content with time is not due to root extraction only (i.e., soil water redistribution can also occur), the assessment of RWU based on water content distribution alone is not possible in conductive soils (Musters and Bouten, 2000). Rather, the full soil water flow equation accounting for root uptake and soil water redistribution must be solved in an inverse mode, and, with an accurate knowledge of soil and root properties RWU distribution can be inferred (Guderle and Hildebrandt, 2015; Hupet et al., 2002; Musters and Bouten, 1999; Vandoorne et al., 2012). Nuclear magnetic resonance (NMR) imaging has been suggested as an adequate technique to measure water flow velocity in xylem vessels but no application exists yet on living roots in soils (Scheenen et al., 2000). More recently, Zarebanadkouki et al. (2012) could measure for the first time RWU in porous media by combining a tracer experiment (i.e., deuterated water) monitored by neutron tomography with inverse modelling of a transport equation. Yet, this was done under controlled conditions while there is no standard method to monitor three dimensional water uptake distribution of growing roots *in situ*. In woody plants, in which roots are thick enough, Nadezhkina et al. (2010; 2012; 2015) used sap flow measurements in roots to quantify hydraulic redistribution.

Since the seminal work of Zimmermann et al. (1967) which reported that RWU of *Tradescantia fluminensis* occurred in the absence of fractionation against water oxygen stable isotope, water stable isotopologues ($^1\text{H}^2\text{H}^{16}\text{O}$ and $^1\text{H}_2^{18}\text{O}$) have been frequently used to identify and quantify root water uptake and redistribution in soils through the measurements of their natural (and artificial) isotopic abundances. Methods include simple graphical inference to more sophisticated statistical methods, i.e., two-end members and multi-source linear mixing models. While the former attempts to locate the “mean root

water uptake” in the soil, the latter category of methods provides profiles of relative contributions to transpiration flux across a number of defined soil layers.

This present paper (i) aims at reviewing these methods ([i.e., graphical inference and statistical multi-source linear mixing models](#)) and (ii) proposes to compare them against each other during a series of virtual experiments differing in the water and isotopic statuses in the soil and the plant. Prior to the review and inter-comparison, the paper reports on the mechanisms at the origin of the spatiotemporal dynamics of natural isotopic abundances in soil and on the background knowledge of isotopic transfer of soil water to and from roots. Finally, we address future challenges to be undertaken such as the dynamic isotopic assessment of HR. We also evoke opportunities offered by novel isotopic monitoring tools which provide unprecedented high frequency isotopic measurements.

2 THEORETICAL BACKGROUNDSFLOW OF ISOTOPOLOGUES IN THE SOIL-PLANT SYSTEM

~~The temporal and spatial variations in natural isotopic abundances observed within the soil plant system allow for reconstruction of S profiles. These variations result from isotope specific fractionation between different phases at thermodynamic equilibrium and during non-equilibrium phase transition, i.e., when there is a net flux between different phases as for instance during evaporation. In this section, we briefly review process-based analytical models accounting for isotopic fractionation that were first proposed for (i) free water (section §2.1) and (ii) later for matric bound water in a bare soil (section §2.2). Finally, (iii) we report on the absence of isotopic fractionation during RWU for most of the documented plant species and on the simple mixing model which is at the basis of any isotopic study on RWU (section §2.3).~~

2.1 Isotopic effects during free water evaporation

~~In a closed liquid water water vapor isothermal system at water vapor saturation (relative humidity = 100 %) or at thermodynamic equilibrium, the difference between the liquid and vapor (hydrogen or oxygen) isotopic compositions (δ_l and δ_v [‰], expressed in ‰ relative to the Vienna Standard Median Ocean Water international isotope reference scale], Gonfiantini, 1978) is a function of the system temperature solely and is named “equilibrium isotopic fractionation factors” (α_{eq} [–]), for a complete list of symbols see Appendix A). Majoube (1971) and Horita and Wesolowski (1994), among other authors, gave empirical expressions (i.e., closed form temperature dependent equations) for these equilibrium fractionation factors.~~

~~When the system is no longer closed and a difference in water vapor partial pressure exists between the air layer in direct contact with the liquid surface and the atmosphere above (referred to as “free” atmosphere), water vapor is transferred from the liquid phase to the air layer, i.e., evaporation (E) occurs. In analogy to an electrical circuit (i.e., a Rideal Langmuir linear-resistance model, Brutsaert, 1982), E can be calculated from the vapor pressure difference and a transfer resistance (r) to vapor transport across the air layer between the evaporating surface and the free atmosphere. Following the same electrical~~

analogy, vapor transport of isotopologues (E_i) is a function of the difference of vapor isotopic composition between the air layer in contact with liquid water and the free atmosphere and a transfer resistance (r_i). In a simple yet comprehensive model Craig and Gordon (1965) divided the air layer into two consecutive layers with different aerodynamic conditions. In a first sub-layer (with transfer resistances r_{diff} and $r_{diff,i}$ for lighter and heavier isotopologues, respectively), vapor transfer is purely diffusive whereas in a second layer (with transfer resistances r_{ad} and $r_{ad,i}$ for lighter and heavier isotopologues, respectively) it is purely advective, i.e., controlled by turbulence. Since the diffusion coefficient of the heavier isotopologues are smaller than that of $^1\text{H}_2\ ^{16}\text{O}$, r_i is greater than r . Note that the difference between r_i and r originates from the difference between r_{diff} and $r_{diff,i}$ whereas $r_{ad} = r_{ad,i}$. In this model, thermodynamic equilibrium conditions still prevail in the air layer in contact with the evaporating surface, termed “liquid vapor interface” of isotopic composition $\delta_{L,v}$. The isotopic composition of the evaporated water vapor (δ_E), defined as the ratio E_i/E , depends on the ratio of the resistances r_i/r . The latter ratio, named “kinetic isotopic fractionation factor” (α_k , []) depends on the relative importance (or development) of each sub-layer and contributes producing an evaporated vapor depleted in heavy isotopologues with respect to the vapor at the liquid vapor interface (i.e., $\delta_E < \delta_{L,v}$). In turn, and depending on the turnover of the system (ratio E/V with V being the volume of the evaporating liquid), but also on the evaporation state (i.e., permanent or transient), the liquid phase enriches itself in the heavy isotopologues. Finally, when both E and V are constant over time, meaning that the loss of water is compensated by a source of constant isotopic composition δ_{source} , an “isotopic steady state” might be reached where, by mass balance, $\delta_E = \delta_{source}$. For a thorough review of the evaporation model of Craig and Gordon (1965), the reader is referred to Gat (1996) and to the more recent paper by Horita et al. (2008).

In a two dimensional ($\delta^{18}\text{O}$, $\delta^2\text{H}$) space, meteoric waters (e.g., precipitation, river water, groundwater) formed by equilibrium processes (i.e., condensation of water vapor) fall onto a line whose slope equals approximately eight and whose theoretical value is the ratio $(\alpha_{eq}^{2\text{H}} - 1)/(\alpha_{eq}^{18\text{O}} - 1)$ at the temperature of condensation. On the other hand, the water vapor produced during a non-equilibrium process, such as evaporation, fall onto a so-called “evaporation line” with a slope of generally lower than six and greater than 2 (Sprenger et al., 2016). This is explained by the fact that $\alpha_K^{18\text{O}} > \alpha_K^{2\text{H}}$ which leads to a greater depletion of $^1\text{H}_2\ ^{18}\text{O}$ with respect to $^1\text{H}^2\text{H}\ ^{16}\text{O}$ in the produced water vapor. Gat (1971) showed that the value of this slope was fairly approximated by the Craig and Gordon (1965) model, which was recently tested by Rothfuss et al. (2015).

2.2 Isotopic effects during bare soil evaporation and leaf transpiration

The Craig and Gordon (1965) model, originally developed for free evaporating water was later adapted to bound-to-matrix soil water. In a study that laid the basis for future work in isotopic ecohydrology, Zimmerman-Zimmermann et al. (1967) et al (1967) provided a steady-state analytical solution for soil water isotopic composition (δ_s), expressed in ‰ relative to the Vienna Standard Median Ocean Water international (VSMOW) isotope reference scale, Gonfiantini, 1978) in a water-saturated isothermal bare sand profile from which water evaporated at a constant rate. Under these steady-state and isothermal conditions, the upward (convective) liquid flux of isotopologues, triggered by evaporation (E) and rising from

deeper layers equals the downward (diffusive) isotopic flux from the evaporating surface which is enriched in the heavy stable isotopologues due to evaporation. Furthermore, by conservation of mass, the isotopic composition of evaporation equals that of its source (e.g., groundwater), i.e., $\delta_E = \delta_{\text{source}}$. A profile is obtained (Fig. 2a, ~~black-dark blue~~ line) whose exponential shape depends on boundary conditions, i.e., the source water (~~e.g., groundwater~~) and surface water isotopic compositions (δ_{source} and δ_{surf}), the diffusion coefficient of the isotopologues in water, and of a soil “tortuosity factor”, conceptually defined as the ratio of the geometrical to actual water transport distance. Barnes and Allison (1983) extended this formulation to a non-saturated sand column evaporating at isotopic steady state (~~$\delta_E = \delta_{\text{source}}$~~). In this case, the evaporating surface (i.e., the liquid-vapor interface) can be located below the soil surface and splits the profile into two regions where isotopic transport predominantly occurs either in the vapor phase above or in the liquid phase below it. In the “vapor region” relative humidity generally is still close to unity for sand total water potential below 15 bars. At isotopic steady state, the maximal isotopic enrichment is at the evaporation front (δ_{EF} at soil depth z_{EF}) and can be simulated with the Craig and Gordon (1965) model. The isotopic composition of the soil residual adsorbed water in the “vapor region” above the evaporation front can be obtained by assuming thermodynamic equilibrium conditions and by applying Fick’s law, and is shown to decrease linearly towards the value of the liquid water at the soil surface which is at thermodynamic equilibrium with the ambient atmospheric water vapor (Fig. 2a, ~~gray-light blue~~ line). Finally, note that at transient state ($\delta_E \neq \delta_{\text{source}}$), the maximal isotopic enrichment in the soil profile does not point to the location of the evaporation front as was demonstrated by Rothfuss et al. (2015). Instead, the depth where the steepest gradient in the isotopic profile is observed corresponds to the evaporation front.

~~In a two dimensional ($\delta^{18}\text{O}$, $\delta^2\text{H}$) space, liquid soil water sampled below the evaporation front will plot on an “evaporation line” with a slope typically lower than six and greater than two, depending on atmospheric and isotopic forcing, as a result of kinetic processes during evaporation.~~ Above the evaporation front and at isotopic steady-state, soil liquid water is in equilibrium with a mixture of atmospheric water vapor ($\delta^{18}\text{O}$ - $\delta^2\text{H}$ slope ~ 8) and evaporated soil water vapor rising from the evaporation front ($2 < \delta^{18}\text{O}$ - $\delta^2\text{H}$ slope < 6) (~~Barnes and Allison, 1988; Brunel et al., 1995; DePaolo et al., 2004~~) (Sprengr et al., 2016). As a result, an intermediate value for the slope is expected, depending on the mixing ratio of atmospheric water vapor to evaporated soil vapor at a given soil depth. Finally, under natural conditions, the δ_s profile is not solely a result of isotopic fractionation but is as well highly impacted both spatially and temporally by input precipitation isotopic composition through modification of the upper boundary condition (δ_{surf}).

2.3 Isotopic transfer to and from roots

As opposed to the removal of water vapor by evaporation, RWU has been described in a number of studies and over a wide variety of plant species not to be associated with (kinetic) isotopic fractionation (Bariac et al., 1994; Dawson and Ehleringer, 1993; Thorburn et al., 1993; Walker and Richardson, 1991; Washburn and Smith, 1934; White et al., 1985; Zimmermann et al., 1967). Consequently, for plants growing in homogeneous external conditions, e.g., in hydroponic solution, root xylem sap water and external water have the same isotopic compositions. In natural soils where the liquid phase is not

homogeneous and a vertical gradient of isotopic composition due to evaporation exists, the root system takes up water at different depths having thus different isotopic compositions.

Assuming that water transport time in roots is negligible, the isotopic concentration of the xylem sap water at the root tiller (C_{Ti} [$M L^{-3}$]) can be modeled as the weighted average of the product of the soil water isotopic concentration (C_S [$M L^{-3}$]) and

5 $S(x,y,z)$:

$$C_{Ti} = \frac{\int_{x,y,z} C_S(x,y,z) \cdot S(x,y,z) \cdot dx \cdot dy \cdot dz}{\int_{x,y,z} S(x,y,z) \cdot dx \cdot dy \cdot dz} = \frac{\int_{x,y,z} C_S(x,y,z) \cdot S(x,y,z) \cdot dx \cdot dy \cdot dz}{J_{Ti}} \quad (2)$$

with J_{Ti} [$L^3 T^{-1}$] the xylem sap flux at the root tiller. Following Braud et al. (2005):

$$C = \rho \cdot R_{ref} \frac{M_i}{M_w} (\delta + 1) \quad (3)$$

with ρ [$M L^{-3}$] the volumetric mass of water, R_{ref} [-] the Vienna-Standard Mean Ocean Water (V-SMOW) hydrogen or oxygen isotopic ratio, M_w and M_i [$M L^{-3}$] the molar masses of $^1H_2^{16}O$ and isotopologue ($^1H_2H^{16}O$ or $^1H_2^{18}O$), respectively, the xylem sap water isotopic composition at the root tiller δ_{Ti} [-, expressed in ‰] can be expressed as:

$$\delta_{Ti} = \frac{\int_{x,y,z} \delta_S(x,y,z) \cdot S(x,y,z) \cdot dx \cdot dy \cdot dz}{J_{Ti}} \quad (4a)$$

with $\delta_S(x,y,z)$ [-, expressed in ‰] the isotopic compositions of soil water at coordinates (x,y,z) . Mostly, a one dimensional description of root water uptake is used assuming that δ_S and RWU do not vary in the horizontal direction and δ_S is obtained for discrete soil layers of depths z_j ($j \in [1,n]$) and thickness $\Delta z_j = z_{j+1} - z_j$. It is usually further hypothesized that J_{Ti} equals the transpiration flux T [$L^3 T^{-1}$] (low to no plant capacitance or phloem-xylem contact):

$$\delta_{Ti} = \frac{\sum_{j=1,n} \delta_S(z_j) \cdot S(z_j) \cdot \Delta z_j}{\sum_{j=1,n} S(z_j) \cdot \Delta z_j} = \frac{\sum_{j=1,n} \delta_S(z_j) \cdot S(z_j) \cdot \Delta z_j}{q_{Ti}} \quad (4b)$$

where $q_{Ti} = J_{Ti}/(\Delta x \Delta y) = T/(\Delta x \Delta y)$ represents the sap flow rate in the root tiller per unit surface area [$L T^{-1}$].

δ_{Ti} can be accessed at different locations in the plant depending on the species, but the sampling location should not be affected by evaporative enrichment in heavier isotopologues or back-diffusion of the isotopic excess accumulated at the sites of transpiration (stomatal chambers) in the leaf. For grasses and nonwoody plants, this is done by sampling the root crown (e.g., Leroux et al., 1995), the aerial nodal roots (e.g., Asbjornsen et al., 2007), the meristematic petiole, or else the collars (e.g., tillers) at the base of the plant (e.g., Dawson and Pate, 1996; Sánchez-Perez et al., 2008). In the case of ligneous plants the fully suberized stem (Asbjornsen et al., 2007) or sapwood (e.g., White et al., 1985) is sampled. On the other hand, δ_S is usually measured by sampling soil profiles destructively. Finally, water from plant and soil is predominantly extracted by

cryogenic vacuum distillation (Araguás-Araguás et al., 1995; Ingraham and Shadel, 1992; Koeniger et al., 2011; Orłowski et al., 2013; West et al., 2006).

Lin and Sternberg (1992) and Ellsworth and Williams (2007), amongst other authors, reported however that for some xerophyte (plants adapted to arid environments, e.g., *Prosopis velutina* Woot.) and halophytes species (plants adapted to saline environments, e.g., *Conocarpus erecta* L.), and mangrove species (e.g., *Laguncularia racemosa* Gaert.), RWU led to fractionation of water hydrogen isotopologues. For mangrove species, it was hypothesized that the highly developed Casparian strip of the root endodermis would force water moving symplastically (i.e., inside the cells) and therefore crossing cell membranes (Ellsworth and Williams, 2007). Water aggregates are then dissociated into single molecules to move across these membranes. This demands more energy for $^1\text{H}^2\text{H}^{16}\text{O}$ than for $^1\text{H}_2^{16}\text{O}$ and $^1\text{H}_2^{18}\text{O}$, thus preferentially affects $^1\text{H}^2\text{H}^{16}\text{O}$ transport and leads to a situation where xylem sap water is depleted in this isotopologue with respect to source water. Meanwhile, this affects to a much lesser extent $^1\text{H}_2^{18}\text{O}$ transport, so that no detectable isotopic fractionation of water oxygen isotopologues is observed. It can be concluded that, for the majority of the studied plant species, either RWU does not lead to isotopic fractionation or its magnitude is too low to be observable.

Hydraulic redistribution (e.g., hydraulic lift) can be conceptualized as a reverse RWU, defined as a negative S . In such case, Eq. (4b) should only account for the positive S . It can be done by assuming that in this equation δ_S is 0 when $S < 0$.

Finally, plant water samples will, similarly to soil water samples, also fall onto an “evaporation line” of a slope lower than eight in a two-dimensional ($\delta^{18}\text{O}$, $\delta^2\text{H}$) space (Javaux et al., 2016).

3 METHODS FOR CHARACTERIZING RWU FROM STABLE ISOTOPIC ANALYSIS

We distinguish two classes of methods: (i) the graphical method for inferring the “mean root water uptake depth” (\bar{z} [L]) (§3.1), and (ii) statistical methods based on end-member mixing analysis (EMMA) (Barthold et al., 2011; Christophersen and Hooper, 1992) for identifying x_j [-], the contribution to RWU of some plant water source j (e.g., water in some soil layer, groundwater, water from recent precipitation, or else from a nearby stream, §3.2). All methods have in common to use an inverse modeling approach: the RWU distribution is obtained by optimizing model input parameters until the simulated δ_{Ti} and/or the simulated soil isotopic profiles fit to the isotopic measurements.

Table 1 summarizes the 21 isotopic studies reviewed in this paper that use either one of the two classes of methods. [This non-exhaustive list was drawn up according to either the number of citations and contribution importance \(for studies published before 2015\) or to the novelty of the publications \(publication year \$\geq\$ 2015\).](#)

3.1 Graphical inference (GI)

This straightforward approach defines the “mean root water uptake depth” \bar{z} , as the depth where $\delta_S = \delta_{\text{Ti}}$. \bar{z} conceptually indicates to the soil depth where the plant root system, represented as one unique root, would extract water from.

There are cases where \bar{z} cannot be unambiguously identified (e.g., \bar{z}_1 and \bar{z}_2 of case 2, Fig. 2b) due to the non-monotonic character of the δ_s profile (shown in black dashed line, case 2 of Fig. 2b). In order to define a mean RWU depth for such a case one can derive a monotonously decreasing δ_s profile by smoothing the profile (shown as symbols in Fig. 2b), e.g., by averaging δ_s in a number of layers using the following mass balance:

$$5 \quad \delta_{s,J} = \frac{\sum_{j \in J} \delta_s(z_j) \cdot \theta(z_j) \cdot \Delta z_j}{\sum_{j \in J} \theta(z_j) \cdot \Delta z_j} \quad (5)$$

where J represents the set of depths that belong to the J^{th} soil layer, with θ [$\text{L}^3 \text{L}^{-3}$] and Δz_j [L] the soil volumetric water content and thickness of the soil layer centered around depth z_j . Due to this smoothing, the vertical resolution may be drastically reduced. In the example presented Fig. 2b where a uniform θ profile is assumed, the $\delta_{s,J}$ profile intersects with the vertical line of value δ_{Ti} deeper than for the initially non-monotonic δ_s profile, i.e., \bar{z} (case 2, integrated δ_s profile) $<$ $\bar{z}_2 <$ \bar{z}_1 .

10 Some authors rule out solutions in case of multiple mean root water uptake depths, e.g., by excluding the \bar{z} solutions where soil water content was low and/or soil water potential was high in absolute value (e.g., Li et al., 2007; see Table 1).

Note that while Eq. (5) provides a representative value for the isotopic composition that would be measured in soil layer J as a function of those of the water in the set of depths, $\delta_{s,J}$ is however equivalent to the isotopic composition “sensed by the plant” only if the root profile is homogeneous, i.e., when RLD is constant with depth in that particular soil layer J .

15 The method of graphical inference may not only provide \bar{z} but also its uncertainty caused by the uncertainty in measuring δ_{Ti} (e.g., based on the precision of the isotopic analysis and/or sampling natural variability, shown as gray stripe in Fig. 2b). The steeper the soil water isotopic profile, the larger ~~is~~ the uncertainty in determining \bar{z} is. Figure 2b illustrates this with estimated minimum and maximum \bar{z} for the monotonic δ_s profile and for the vertically averaged profile. In the latter case, the possible range of \bar{z} is the largest. These ranges give first quantitative indication of variance around \bar{z} . Finally, for a
 20 complete “graphical assessment” of the variance of \bar{z} , one should also consider the uncertainty associated with measurements of the δ_s profile (not shown [here; for a complete assessment of errors associated with determination of \$\delta_s\$, see Sprenger et al., 2015](#)).

3.2 Statistical approaches

3.2.1 Two end-member (TM) mixing model

25 The TM method is a particular case of end-member mixing analysis (EMMA) and is based on the concept that (i) a plant extracts water from two predominant water sources A and B (e.g., water in distinct upper and lower soil layers, or groundwater and recent precipitation water etc.) in given proportions, (ii) there is no isotopic fractionation during water uptake, and (iii) there is a complete mixing inside the plant of the contributing water sources A and B to RWU. The mass conservation for isotopologues gives:

$$J_{Ti}^i = J_i^A + J_i^B \quad (6a)$$

$$C_{Ti} \cdot J_{Ti} = C_A \cdot J_A + C_B \cdot J_B \quad (6b)$$

with J_A , J_B , and J_{Ti} [$L^3 T^{-1}$] (respectively J_i^A , J_i^B , and J_{Ti}^i [$M T^{-1}$]) the fluxes of water (respectively isotopologues) originating from water sources A and B, and at the plant tiller. C_A , C_B , and C_{Ti} [$M L^{-3}$] are the water sources A and B, and xylem sap water measured isotopic concentrations. By introducing $x = J_A / J_{Ti}$ and following Eq. (3), Eq. (6b) becomes:

$$\delta_{Ti} = x \cdot \delta_A + (1 - x) \cdot \delta_B \quad (7)$$

In this approach, δ_{Ti} is therefore defined as the mean value of the isotopic compositions of water sources A and B (δ_A and δ_B) weighted by the proportions to J_{Ti} of water volume extracted by the plant from water sources A and B, i.e., x and $(1 - x)$, respectively. The error associated with the estimation of x (σ_x [-, expressed in ‰]) can be calculated following Phillips and Gregg (2001):

$$\sigma_x^2 = \left(\frac{\partial x}{\partial (\delta_A)} \right)^2 \cdot \sigma_{\delta_A}^2 + \left(\frac{\partial x}{\partial (\delta_B)} \right)^2 \cdot \sigma_{\delta_B}^2 + \left(\frac{\partial x}{\partial (\delta_{xyl})} \right)^2 \cdot \sigma_{\delta_{Ti}}^2 \quad (8a)$$

$$\sigma_x = \sqrt{\sigma_x^2} = \frac{1}{(\delta_A - \delta_B)} \sqrt{(\sigma_{\delta_{Ti}}^2 + x^2 \cdot \sigma_{\delta_A}^2 + (1 - x)^2 \cdot \sigma_{\delta_B}^2)} \quad (8b)$$

with σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}}$ the standard errors associated with the measurements of δ_A , δ_B , and δ_{Ti} , respectively. The sensitivity of Eq. (8b) to different values of σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}}$ can be tested by considering either minimal possible errors, i.e., the analytical precision of the isotopic analyser (e.g., isotope ratio mass spectrometer, laser-based spectrometer), or by taking into account additional errors involved with sampling procedure and vacuum distillation technique (see e.g., Rothfuss et al., 2010). Equation (8b) also shows that, independently of the values considered for σ_{δ_A} , σ_{δ_B} , or $\sigma_{\delta_{Ti}}$, σ_x is inversely proportional to $1/(\delta_A - \delta_B)$, indicating that the two end-members should have as much as possible distinct isotopic compositions for a low standard error of x . Therefore, it is especially important, e.g., for partitioning between water from an upper and lower portion of the soil profile, to properly define the thickness of these layers, so that they have distinct isotopic compositions, and that the difference is considerably larger than the precision of the isotopic measurements. Figure 3 shows for example that when (i) x is evaluated at 10 % and (ii) σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}}$ are estimated being equal to 0.02 ‰ (dark blue solid line), $(\delta_A - \delta_B)$ should be greater than 0.75 ‰ (in absolute term) in order to reach a σ_x value lower than 5 %, i.e., more than 37 times the error made on δ_A , δ_B , and δ_{Ti} . To obtain the same standard error for x in case of a higher standard error on the estimation of δ_A , δ_B , and δ_{Ti} (e.g., σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}} = 0.1\%$), $(\delta_A - \delta_B)$ should be greater than 3.00 ‰ (in absolute term). This difference becomes much greater for σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}} = 1.00\%$ and reaches 42 ‰ (not shown in Figure 3). This certainly highlights the advantage of artificially labelling soil water with water enriched (or depleted) in heavy isotopologues for a more precise assessment of the relative contribution of soil water sources to RWU, as mentioned by Moreira et al.

(2000). In another study, Bachmann et al. (2015) labeled the upper and lower portion of the soil profile in a natural temperate grassland with ^{18}O -enriched and ^2H -enriched water, respectively. They defined two distinct (upper and lower) soil water sources, for which they calculated the corresponding $\delta^2\text{H}$ or $\delta^{18}\text{O}$ on the basis of measured soil water isotopic profiles and using Eq. (5). They could find evidence against the so-called hypothesis of “niche complementarity” regarding plant water use, which states that RWU of competitive plant species is spatially and temporally distinct, and that this distinction is stronger at high species richness. Figure 3 illustrates also that for given $(\delta_A - \delta_B)$, σ_{δ_A} , σ_{δ_B} , and $\sigma_{\delta_{Ti}}$ values, the “optimal x value” for a low σ_x is 50% (showed by the orange lines).

Table 1 displays a sample of studies that used the two end-member mixing approach. Authors could distinguish between uptake of irrigation and precipitation water (Goebel et al., 2015), precipitation and groundwater (White et al., 1985), soil water and groundwater (McCole and Stern, 2007), or else between stream water and soil water (Dawson and Ehleringer, 1991; McDonnell, 2014). Thorburn and Ehleringer (1995) could for instance locate the dominant source for RWU, i.e., groundwater for their mountain and floodplain test-site and water from the soil between 0.3 and 0.4 meters depth for their cold desert test-site. Other authors (e.g., Brunel et al., 1995) combined two mixing equations, i.e., one for each isotopologue, into a single one. As infrared laser-based spectrometry now enables simultaneous measurements of $\delta^{18}\text{O}$ and $\delta^2\text{H}$ at lower cost, we believe that this dual-isotope approach (referred as “D” in Table 1) will or should gain in importance in isotopic studies. This is especially useful when (i) under natural conditions the $\delta^{18}\text{O}$ - $\delta^2\text{H}$ slope is not constant over depth (Sprengrer et al., 2016) or (ii) in the context of pulse labelling experiments, which can artificially change the value of the $\delta^{18}\text{O}$ - $\delta^2\text{H}$ slope “disconnect” the strong correlation between soil water $\delta^{18}\text{O}$ and $\delta^2\text{H}$ at given locations in the soil profile, therefore in these cases, provide two independent mixing equations are obtained, one for each isotopologue.

20 3.2.2 Multi-source (MS) mixing models

When there are more than two identified plant water sources contributing to RWU, e.g., water from different layers j ($j \in [1, N]$) in soil the profile, Eq. (7) becomes:

$$\delta_{Ti} = \sum_{j=1}^N x_j \cdot \delta_{S,j} \quad (9)$$

with N the number of plant water sources (e.g., soil layers) and $\sum_{j=1}^N x_j = 1$. As there are more water sources than (number of mixing equations + 1), there is not a unique solution but an infinite range of possible solutions. However, some of these solutions are not likely or possible based on background information or knowledge. A range of solutions that is most likely based on prior information can be obtained using Bayesian methods. In the method proposed by Phillips and Gregg (2003), the isotopic composition calculated for each considered x_j combination (δ_{Ti}) is compared with the measured value ($\delta_{Ti,m}$). The number of combinations depends on the value of contribution increment (i , %, typically 5 or 10 %) and the combinations for which δ_{Ti} meets the following requirement are selected:

$$\delta_{\text{T}} \leq |\delta_{\text{T},m} \pm \tau| \quad (10)$$

where τ [-, expressed in ‰], standing for “tolerance”, usually accounts for precision of the isotopic measurements or possible errors during sampling and vacuum distillation steps. This multi-source mixing model approach strongly depends on τ and i , which therefore should be carefully chosen by the user. A smaller i also refines the analysis. For this, the program “IsoSource” (https://www.epa.gov/sites/production/files/2015-11/isosourcev1_3_1.zip) is available (Phillips et al., 2005). Wang et al. (2010) compared the outcome of the GI and MS approaches and came to the conclusion that even though the latter did not solve the non-uniqueness problem and provided diffuse patterns of frequency that were difficult to interpret in some cases (e.g., in case of a non-monotonic isotopic profile), it had the advantage over the former method of providing a systematic and quantitative assessment of ranges of relative contributions.

10 Parnell et al. (2010) proposed to overcome two limitations of the approach of Phillips and Gregg (2003), i.e., its inability to (i) account for uncertainty in the estimations of δ_{T} and of the water sources isotopic compositions $\delta_{\text{s},j}$, and (ii) provide a optimal solution rather than ranges of feasible solutions. For this they use a Bayesian framework (for details see also Erhardt and Bedrick, 2013; Moore and Semmens, 2008; Parnell et al., 2013), which allows uncertainty in the x_j proportions and incorporates a residual error term ε_j (normally distributed with mean equal to zero and variance σ^2):

$$15 \quad \delta_{\text{T}} = \sum_{j=1}^N x_j \cdot \delta_{\text{s},j} + \varepsilon_j \quad (9')$$

Note that the terms of (i) trophic enrichment factor (TEF [-, expressed in ‰], see, e.g., meta-analysis of Vanderklift and Ponsard, 2003) and (ii) isotope concentration dependency (Koch and Phillips, 2002; Phillips and Koch, 2002) originally incorporated in the formulation of Parnell et al. (2010) for other applications are not present in Eq. (9') since (i) no isotopic fractionation during RWU is assumed and (ii) isotope concentration dependency applies only for situations where isotopic compositions of different elements are measured and available.

20 Parnell et al. (2010) developed the program “Stable Isotope Analysis in R” (SIAR, https://cran.r-project.org/src/contrib/siar_4.2.tar.gz) in which the initial (*a priori*) x_j distribution is by default the Dirichlet distribution, of which information can be partly specified by the user. A *posteriori* x_j distribution is obtained by fitting the linear model to data via a Metropolis-Hasting (Hastings, 1970; Metropolis et al., 1953) Markov Chain Monte Carlo algorithm.

25 Prechsl et al. (2015) apply both graphical and Bayesian approaches to evaluate the shift in \bar{z} and change of RWU profile following drought treatments (approx. 20 to 40 % precipitation reduction with transparent rainout shelters) in both extensively and intensively managed grasslands. From both approaches it appeared that a shift in \bar{z} was inexistent or not observable from isotopic analyses. Another recent application of the Bayesian approach was performed by Volkmann et al. (2016b), who took advantage of a newly developed soil isotopic monitoring method to confront high frequency δ_{s} profiles
30 time series to time series of δ_{T} (indirectly obtained from the isotopic measurement of the transpired water vapor and assuming isotopic steady state, i.e., $\delta_{\text{T}} = \delta_{\text{T}}$) following a labelling pulse (see Table 1 for details on the study).

4 INTER-COMPARISON OF METHODS

We tested and compared the different methods (GI, TM, MS) during a series of virtual experiments. Mean RWU depths (provided by the GI method) and x_j distribution (provided by the TM and MS methods) were determined from soil and xylem water oxygen isotopic composition distributions. While the former information was prescribed to the different methods, the latter was calculated with the physically based analytical RWU model (referred to as AM) of Couvreur et al. (2012). The inter-comparison of models was performed using a single isotope (^{18}O) approach as the focus here was the differences of outcomes rather than the impact of the input isotopic data on these results. The reader can refer to Appendix B1 for a description of the model of Couvreur et al. (2012) and to Appendix B2 on how it was run for the inter-comparison.

4.1 Methodology

4.1.1 Scenario definition

We developed eight virtual plausible scenarios of soil-plant systems under different environmental conditions. Each environmental condition was defined as a combination of different total soil water potential distributions (resulting from the location of the groundwater table and weather conditions), soil water oxygen isotopic composition (δ_s) profiles, and actual transpiration rate (T). The groundwater table was either shallow at -1.25 m depth (prefix “Sh”) or deep at -6 m depth (prefix “De”); the soil water potential was considered to be at static equilibrium below the groundwater level; the soil surface was either dry under evaporative conditions (suffix “Dr”), or wet, e.g., shortly after a rain event (suffix “We”); the transpiration rate was either low (e.g., relevant at night, $T = 0.01 \text{ mm h}^{-1}$, suffix “_IT”) or high ($T = 0.30 \text{ mm h}^{-1}$, suffix “_hT”). They all relied on a common measured root length density vertical distribution of *Festuca arundinacea*. Table 2 reports the input data. Note that, as hypothesized in Eq. (4b), transpiration and sap flow rates (i.e., per unit of surface area [L T^{-1}]) were considered as equal.

4.1.2 Setup of the models

The two end-member mixing approach (TM) was tested against the isotopic data for two different cases: (i) two conjoint soil layers spreading from 0 – 0.225 m and 0.225 – 2.00 m and (ii) two disjoint soil layers spreading from 0 – 0.225 m and 1.75 – 2.00 m. The latter case was designed to evaluate the impact of lacunar soil isotopic information on the calculation of x , i.e., when not all potential water sources are properly identified. Representative values of water oxygen isotopic compositions for these soil layers ($\delta_{s,J}$, $J \in [\text{I,II}]$) were obtained from the mass balance (Eq. (5)) after interpolation of the measured soil water content and δ_s profiles at a 0.01 m vertical resolution.

For the multi-source mixing approaches of Phillips and Gregg (2003) (MSPG) and Parnell et al. (2010) (MSPa), the number of potential water sources was initially fixed to three, i.e., water from the soil layers I (0.000-0.050 m), II (0.050-0.225 m), and III (0.225-2.000 m). Upper and lower boundaries of these layers were defined to reflect the exponentially shaped

- (monotonic) δ_s profiles (experiments ShDr and DeDr) or to smooth the non-monotonic δ_s profiles observed during experiments ShWe and DeWe. MSPG and MSPa were also tested for eight soil layers (i.e., as many layers as measurement points, I: 0.000-0.020, II: 0.020-0.050, III: 0.050-0.110, IV: 0.110-0.225, V: 0.225-0.400; VI: 0.400-0.750, VII: 0.750-1.500, and VIII: 1.500-2.000 m). Increment and tolerance of the MSPG method were fixed at 10 % and 0.25 %, respectively.
- 5 Similarly to the TM approach, profiles of $\delta_{s,J}$ ($J \in [I,III]$ or $[I,VIII]$) were obtained from the mass balance (Eq. (5)) after interpolation of the measured soil water content and δ_s profiles at a 0.01 m vertical resolution.
- Finally for the MSPa method, uncertainty associated with δ_s measurements was set to 0.2 ‰ and the number of iterations was fixed to function `siarmemedirichletv4` of the SIAR-R package was run 500000 and number of iterations to be discarded to times (of which 50000 runs were discarded).
- 10 For a detailed description of the inter-comparison methodology, refer to Appendix C.

4.2 Results and discussion

Figure 4 displays x_{AM} , the simulated $\frac{S_{AM}(z)dz}{T/(\Delta x \cdot \Delta y)}$ ratios (solid colored lines) simulated by the analytical model of (Couvreur et al., 2012) for the eight scenarios together with uncertainty (shaded areas) and the corresponding δ_{Ti_AM} ($\pm 1sd$) (for a description on how uncertainty was assessed, refer to Appendix C). In general, at high T the compensation was negligible and the S_{AM} profile was mainly proportional to the RLD profile (Fig. 4b, d, f, and h). The only exception was a soil with deep groundwater table and dry surface, where this dry layer limited root water uptake (DeDr_hT). At lower transpiration demand, the S profile predicted by the Couvreur et al. (2012) model generally differed from the RLD profile (Fig. 4a, c, e, and g) due to the fact that the second term of Eq. (1) (i.e., S_{comp} , see also Eq. (B4) and (B4') in Appendix B) was proportionally larger. Water uptake from the upper layer was always more than proportional to the RLD , when this layer was wetter, and vice versa. Water release to the soil (i.e., HR) was observed only for the soil with the deep groundwater table and dry upper layer (DeDr_hT, Fig. 4e). From the graphical method GI, either a single or two distinct solutions for \bar{z} (displayed as gray-shaded horizontal stripes) could be retrieved, depending on the monotonic/non-monotonic character of the δ_s profile, and ranged between -0.02 and -0.95 m.

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Figure 5a displays the relative contribution to T of the upmost layer 0 – 0.225 m in case of two conjoint soil layers as computed with the TM approach and a comparison with the results of the analytical model. Except for the very last two virtual experiments (i.e., DeWe_IT and DeWe_hT), there was a very good agreement between TM and AM results: absolute difference between x_{TM} and x_{AM} ranged between 1.5 % (ShDr_IT) and 6.3 % (ShDr_hT). During experiment DeDr_IT, The TM approach estimated that x was equal to 12.3 % while the analytical model simulated hydraulic redistribution, i.e., excluded the layer 0 – 0.225 m as potential source. The significant difference between TM and AM results during experiments DeWe_IT and DeWe_hT and the higher standard error associated with x_{TM} (σ_x , displayed in the form of error bars in Figure 5) were due to the small difference between the isotopic compositions of the defined soil water sources $\delta_{s,I}$ (–

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6.0 ‰) and $\delta_{s,II}$ (-5.3 ‰) as illustrated in section §3.2.1. Figure 5b gives the relative contribution to T of the layer 1.75 –2.00 m in case of two disjoint soil layers, i.e., when not all potential water sources are accounted for into the calculation of $\delta_{s,I}$ and $\delta_{s,II}$. In this case there were important disparities between x_{TM} and x_{AM} . The mean absolute difference between these two estimates was equal to 43,5 (± 17.8) %. Omitting some of the potential water sources contributing to T had in this second case the consequence of artificially overestimate the contribution of the lowest layer. We therefore suggest to always attempting to fully characterize the soil isotopic profile before aggregating the isotopic information when defining the two water sources.

Figure 6 gives the relative contributions from soil layers I, II, and III (upper, middle, and lower panel, respectively) to T following the method of Phillips and Gregg (2003) (x_{MSPG} , in %, displayed in the form of gray histograms) and the Bayesian method of Parnell et al. (2010) (x_{MSPa} , in %, gray probability density curves). The colored vertical lines are x_{I_AM} , x_{II_AM} , and x_{III_AM} , the simulated $\frac{S_{AM}(z)dz}{T/(\Delta x \cdot \Delta y)}$ ratios from layers I, II, and III. The color-shaded areas associated with x_{I_AM} , x_{II_AM} , and x_{III_AM} refer to their uncertainty by accounting for the uncertainty of the input data. As for Fig. 5, δ_{Ti_AM} is reported above each plot along with its standard deviation. x_{J_MSPG} probability distribution was observed to be either narrow (e.g., DeDr_IT / layer I, Fig. 5m) or broad (e.g., DeWe_hT / layer I), i.e., the range of the possible solutions for x_{J_MSPG} was relatively small or large (10 and 100 % respectively for these two examples). In general, both MSPG and MSPa statistical methods agreed well with each other: the x_{MSPa} most frequent value (MFV, at the peak of the density distribution curve) was in most cases either located near the median value of the x_{MSPG} probability range (e.g., ShWe_IT / layer I, Fig. 5g) or matched exactly the x_{MSPG} unique value (i.e., DeDr_IT / layer I, Fig. 5m). In contrast, the statistical methods succeeded best in providing x estimates similar to those of the model of Couvreur et al. (2012) in case of a shallow groundwater table and at low T only (Fig. 5a-c and g-i), thus when water availability was high and root compensation was low. In these cases, x_{I_AM} was included in the estimated x_{J_MSPG} range and the mean absolute difference (MD) between x_{J_AM} and x_{MSPa} MFV was equal to 8.6 %. This difference was the greatest (129.2 %) for experiment DeDr_IT when HR was simulated by the analytical model (Fig. 5m-o).

Considering eight soil layers instead of three added uncertainty in the assessment of their relative contribution to T as determined by the MSPG method: the estimated probability ranges increased in most of the cases (results not shown). However it considerably improved the results of the MSPa method: the mean absolute difference between x_{J_AM} and the most frequent x_{MSPa} value was equal to 4.7 % for the scenarios with a shallow groundwater table and low transpiration rate and equal to 52.1 % in case of HR (Table 3).

Independent of the number of defined soil layers, lowering the value of increment to 5 % in the MSPG method refined the analysis where the probability distribution was already narrow (i.e., in the case of a well identified x_{MSPG} value, e.g., Fig. 5m) while it produced distributions that were flatter and contained less gaps when no clear solutions had emerged before (results not shown). Artificially increasing the value of tolerance had the consequence that more solutions to Eq. (10) were found for

each experiment / transpiration value / layer combinations and *vice versa* (results not shown). An increase or decrease of a factor 2 of the number of runs as well as the number of runs to be discarded from the analysis had only a marginal impact on the density distribution curves obtained with the MSPa method in the case of three or four soil layers.

The modelling exercise illustrated the disparities of outcome between the graphical method on the one hand and the statistical and mechanistic methods on the other: there simply cannot be a single or multiple “root water uptake depths” but rather a continuous RWU profile (AM) or statistical solutions of contribution to transpiration (MSPG and MSPa). Significant changes of δ_{Ti} do not necessarily mean important changes in the depth of RWU but rather slight (nevertheless significant) modification of the RWU profile. The authors believe that the relatively novel statistical tools MSPG and MSPa presented in this review should be therefore preferred over the GI method, especially since the two former are available as user-friendly programs and packages and do not require significant computing time, therefore can be run locally on a personal computer. As highlighted in this series of virtual experiments, the Bayesian method showed much more convincing results than the method of Phillips and Gregg (2003), especially in the case of eight soil layers, illustrating the interest of reaching the best vertical resolution and maximizing the number of identified potential sources.

One can also show from this inter-comparison of methods that labelling of soil water in either ^{18}O or ^2H has potentials for improving the different methods presented here theoretically if water is taken up by the roots from the labeled region predominantly. However this was never the case looking at the results of the analytical model. A dual isotope (^{18}O or ^2H) labelling pulse experiment that would artificially disconnect the strong link between $\delta^{18}\text{O}$ and $\delta^2\text{H}$ would on the other hand much more constrain the inverse problem and provide accurate estimate of contribution of S to transpiration flux.

5 CHALLENGES AND PROGRESSES

20 5.1 Isotopic assessment of hydraulic redistribution

HR (e.g., HL) has been observed using isotopic measurements in a number of studies (e.g., Caldwell and Richards, 1989; Dawson, 1993; Kurz-Besson et al., 2006). However, in contrast to nondestructive “traditional” methods allowing for direct monitoring of redistribution dynamics (i.e., psychrometry, time domain reflectometry, and frequency domain capacitance, Brooks et al., 2002; Dawson, 1996; Richards and Caldwell, 1987; Wan et al., 2000), isotopic methods provide a destructive and indirect assessment. These methods are based on (i) labelling of soil or roots of deep-rooted plants at a given depth in the soil or at a certain location in the experimental field and (ii) measuring the δ_{Ti} of plants not having access to labeled water (i.e., of which the roots do not reach the isotopic labeled depth or location). When HR occurs, the xylem sap water (of measured isotopic composition $\delta_{Ti,m}$) of these plants can be conceptualized as a mixture of antecedent soil water (at natural isotopic abundance) and isotopically enriched water released to the soil by the deep-rooted vegetation. From simple mass balance at the release location, δ_s at a given depth z in the soil and at time $(t+\Delta t)$ deviates from that at time t as a function of the (negative) S (i.e., HR or HL) at time t and change of soil volumetric water content (θ):

$$\delta_s(z, t + \Delta t) \cdot \theta(z, t + \Delta t) - \delta_s(z, t) \cdot \theta(z, t) = \delta_{Ti,m} \cdot V_{HR}(t)/V(z) = \delta_{Ti,m} \cdot |S(z, t)| \cdot \Delta t \quad (11)$$

If θ and δ_s at times t and $t+\Delta t$, and δ_{Ti} are measured, the water volume transported by the roots (V_{HR} , L^3) can be calculated knowing the volume of soil representative of the hydric and isotopic measurements (V , L^3). Note that HR is observable at a certain soil depth if and only if uptake and release locations in the soil have distinct water isotopic compositions. Finally, the obtained volume can be compared with the water volume transpired by the vegetation on the following day.

To the authors' knowledge, no precise observation (other than the study of Zegada-Lizarazu and Iijima, 2004) of change of soil water isotopic composition has been attributed with certainty to hydraulic redistribution and simultaneously provided amount of water involved in the process. Such observations however should be feasible under controlled experimental conditions where (i) the initial soil water isotopic profile before labelling is known and (ii) natural isotopic changes (due to, e.g., soil redistribution and moisture input from a precipitation event) can be avoided, and (iii) the lateral heterogeneity of soil water and isotopic composition profiles can be minimized (see for instance the setups of Armas et al., 2012; Querejeta et al., 2012). As highlighted in section 2.3, HR can be conceptualized as a negative S (Eq. (4b)) and should therefore be exempt of isotopic fractionation. However, to the authors' knowledge this point has not yet been proven experimentally.

5.2 High frequency isotopic data and sSoil and plant water sampling strategies

For determination of δ_s , soil profiles are usually destructively sampled, typically with an auger down to a depth of a few centimeters (Rothfuss et al., 2010) to a few meters (Moreira et al., 2000) (see Table 1), depending on the depths of the root system and of the water table. The sampling depth interval should, when possible, match the exponential decrease of isotopic composition (Wang et al., 2010) due to fractionating evaporation and it should capture sudden variations with time at the soil surface due to precipitation, i.e., be minimal at the surface and maximal deeper in the soil profile where isotopic dynamics are less pronounced. A minimal sampling interval at the surface is also crucial as it provides the isotopic composition of the layer contributing the most to transpiration in the case of a low T flux (e.g., morning transpiration) under non-limiting water availability. Not measuring this maximum soil isotopic composition (between precipitation event) can lead to a situation where source partitioning is not feasible from isotopic measurements. Under field conditions (i.e., ~95 % of the studies reviewed in this work, summarized in Table 1) soil material is generally not a limiting factor, thus can be sampled twice or thrice to average out or characterize lateral heterogeneity without significant disturbance of the soil (Leroux et al., 1995).

For grasses and nonwoody plants the root crown (e.g., Leroux et al., 1995), the aerial nodal roots (e.g., Asbjornsen et al., 2007), the meristematic petiole, or else the collars (e.g., tillers) at the base of the plant (e.g., Dawson and Pate, 1996; Sánchez-Perez et al., 2008) can be sampled for determination of δ_{Ti} . In the case of ligneous plants the fully suberized stem (Asbjornsen et al., 2007) or sapwood (e.g., White et al., 1985) is sampled. Special attention is to be given to remove any organ potentially transpiring that would have a water isotopic composition not representative of δ_{Ti} , e.g., dead material or the sheath of the oldest living leaf around tillers (Barnard et al., 2006; Durand et al., 2007). Non-conducting heartwood should be discarded from conducting sapwood (White et al., 1985).

5.3 Off-line destructive versus on-line non destructive isotopic measurements in plant and soil waters

Water from plant and soil materials is predominantly extracted by cryogenic vacuum distillation (Araguás-Araguás et al., 1995; Ingraham and Shadel, 1992; Koeniger et al., 2011; Orlowski et al., 2013; West et al., 2006). This consists in (i) introducing the plant or soil sample into an extraction flask attached to one end of the extraction line, while at the other end a collection tube is connected, (ii) freezing the sample by immersing the collection flask into liquid nitrogen (temperature ~ – 200°C), (iii) pumping the extraction line down to a pressure of $\sim 10^{-3}$ mbar, (iv) heating the sample to a certain temperature ($\sim 60 < T < \sim 100^\circ\text{C}$) depending on its nature while immersing the trap into liquid nitrogen. The water vapor produced condenses in the trap following a stepwise procedure (~lasting one to a few hours), in order to avoid condensation elsewhere on the water vapor path between sample and collection trap. Accuracy of this extraction method was shown to be maximal at higher water content and for sandy soils and lower for soils with high clay content (e.g., Koeniger et al., 2011; West et al., 2006). In the latter case, extraction times should be longer and temperatures higher to mobilize water strongly bound to clay particles, which has a distinct isotopic composition from that of pore “bulk” water (Araguás-Araguás et al., 1995; Ingraham and Shadel, 1992; Oerter et al., 2014; Sofer and Gat, 1972). In other studies, plant and soil waters are extracted following azeotropic distillation with kerosene as solvent (e.g., Brunel et al., 1995; Thorburn and Ehleringer, 1995), or direct equilibration with CO₂ (Asbjornsen et al., 2007) following the method of Scrimgeour (1995), or liquid water – water vapor equilibration (Wassenaar et al., 2008), or else the mild vacuum method (Dawson and Pate, 1996; Jeschke and Pate, 1995).

Certainly one of the main limitations of all isotopic approaches for quantifying RWU and HR is the destructive character of isotopic sampling (see section 3.1) and associated offline analyses (sections 2.2 and 2.3). This usually leads to poor spatial (maximum a few cm²) as well as temporal (minimum hourly) resolution of the inferred results, when comparing with measuring frequency of other soil and plant state variables, e.g., soil water content and potential, and leaf water potential (section 3.2.2). In addition, one may question the representativeness of plant samples, in which tissues (and thus water) with very different water residence time is mixed. Similarly, given the expected high lateral and temporal variability of the HR process, the representativeness of δ_s should be questioned for soils, in particular when combined with 1D models.

Recently developed methods take advantage of laser-based spectroscopy which allows on-line and continuous isotopic measurements in the gas phase. These methods rely on coupling a laser spectrometer (e.g., Wavelength-Scanned Cavity Ring-Down Spectroscopy – WS-CRDS, Picarro Inc., Santa Clara, CA, USA; Cavity-Ringdown Laser Absorption Spectroscopy – CRLAS and Off-Axis Integrated Cavity Output Spectroscopy - ICOS, Los Gatos Research, Los Gatos, USA) with specific soil gas sampling probes consisting of gas-permeable microporous polypropylene membranes or tubing. These membranes or tubing exhibit strong hydrophobic properties, while their microporous structures allow the intrusion and collection of soil water vapor. Several authors (Gaj et al., 2015; Gangi et al., 2015; Herbstritt et al., 2012; Oerter et al., 2016; Rothfuss et al., 2013; Sprenger et al., 2015a; Volkman and Weiler, 2014) could determine the soil liquid water isotopic composition in a nondestructive (yet invasive) manner from that measured in the collected soil water vapor considering thermodynamic equilibrium between vapor and liquid phase in the soil. In contrast to “traditional” isotopic methods, these

novel isotopic monitoring methods have also the distinct advantage of determining soil liquid water isotopic composition at very low water content, since water vapor, in contrast to soil liquid water, is not limiting for analysis. These novel methods allow a vertical resolution down to 1 cm and an approximately hourly time-resolution. However, they do not allow horizontal resolution along the tube and ~~the laser spectrometers could are be~~, as pointed out by Gralher et al. (2016) ~~for the specific case of a Picarro WS-CRDS greatly sensitive to the carrier gas used for a Picarro CRDS, greatly sensitive to the carrier gas used~~. In their opinion papers, McDonnell (2014) and Orłowski et al. (2016) also urged for a comparison between methods, which was addressed by ~~Gaj et al. (2015) and~~ Pratt et al. (2016).

Leaf and plant gas chamber systems provide indirect means for a nondestructive determination of δ_{T_i} , i.e., by either assuming full steady-state conditions at the evaporative sites of the leaves ($\delta_{T_i} = \delta_T$) (e.g., Dubbert et al., 2014; Volkmann et al., 2016b). In the coming years, effort should be made towards developing novel methods for a direct and nondestructive determination of δ_{T_i} based on the use of gas-permeable membranes, which was recently initiated for trees (Volkmann et al., 2016a). This should be further investigated to test applicability to other (non-woody) plant species. This will imply the major challenge of not disrupting the water columns in the active xylem vessels when installing such a membrane-based system. Another potential issue to be investigated is the species-specific extent of water exchange between xylem and phloem conductive tissues which might lead to isotopic “contamination” of the xylem sap water (Farquhar et al., 2007).

5.3.4 Call for a coupled experiments-modelling approach for determination of plant water sources and redistribution on the basis of isotopic data

In order to fully benefit from the potential of water stable ~~isotopologue~~ isotopic analysis as tools for partitioning transpiration flux, the authors call ~~for the development of approaches making use of physically based models for RWU and isotopic fractionation to analyze experimental data, especially since several soil-vegetation-atmosphere transfer (SVAT) models are available that can simulate flow of isotopologues in the soil and the plant (i.e., SiSPAT-Isotope, Braud et al., 2005; Soil-litter-iso, Haverd and Cuntz, 2010; TOUGHREACT, Singleton et al., 2004; HYDRUS, Sutanto et al., 2012)~~ ~~for a generalization of coupled approaches based on the confrontation of experimental data with a physically based understanding of RWU processes.~~

~~To the authors' knowledge, there are only a few studies which attempted to do so. An example is the work of~~ Campbell (1991); Ogle et al. (2004) ~~Rothfuss et al. (2012) who ran an experiment under controlled laboratory conditions where they measured on four dates (corresponding to four different stages of vegetation and therefore root development) soil water potential and isotopic composition profiles, and root length density distribution profiles. In their experiment, the isotopic composition of transpiration was also known. The authors used a global optimization algorithm to obtain the set of parameter of SiSPAT-Isotope (Braud et al., 2005) that best reflected the experimental dataset. Distributions of RWU could be determined on these four dates. Also, in the study of Mazzacavallo and Kulmatiski (2015), the RWU model of HYDRUS could also be parameterized during a labelling (heavy water $^2\text{H}_2\text{O}$) pulse experiment on the basis of measurements of xylem~~

water hydrogen and oxygen isotopic compositions. This provided insights into the existence of niche complementarity between tree (mopane) and grasses species. Note however that this HYDRUS version did not incorporate isotopic transport through the soil and the roots.

5 Another example is the work of Ogle et al. (2004) who could reconstruct active root area and RWU profiles from isotopic measurements using the 1D analytical macroscopic model of Campbell (1991) in a Bayesian framework (root area profile and deconvolution algorithm – RAPID). By assuming normal a priori distributions for the xylem water oxygen and hydrogen isotopic compositions and considering prior knowledge on RWU distribution (i.e., synthetic information based on measurements of other studies), Ogle et al. (2004) obtained a posteriori distributions of x of a desert shrub (*Larrea tridentate*).

10 Simple analytical models, such as the formulation of Couvreur et al. (2012), can be applied and confronted with isotopic data. In comparison with statistical tools, such physical models provide profiles with high spatial resolution and lower uncertainty, on the condition that all required (isotopic) data is available. We recognize that in comparison with the statistical and conceptual methodologies presented in this review, using a physical (analytical or numerical) model implies the measurements of additional state variables to be fed as input to the model, and of one parameter (K_{plant}) (when considering

15 the assumption $K_{\text{plant}} = K_{\text{comp}}$ valid, see Appendix B). Some of these variables are laborious to obtain (e.g., *RLD*) or not straightforward to measure (H_s , H_L , and T) – especially in the field – but are mandatory to be able to determine contributions to T across a set of identified water sources. In addition, they are necessary to gain insights into soil-plant interactions, e.g., dynamics of root function (active versus non-active roots in the soil profile) in water uptake and thus quantify the disconnection between measured *RLD* and the prognostic variable *SSF* (see Appendix B1). For doing this, controlled

20 conditions in state-of-the-art climatic chambers are ideal, as they allow reducing the inherent spatial heterogeneity present under natural conditions and, thus, the deconvolution of environmental effects on RWU. Experimental facilities that not only control atmospheric forcing (soil upper boundary conditions for latent and heat flow), but impose lower boundaries for the soil compartment (e.g., drainage and capillary rise dynamics) and provide means to close the hydrological balance are required. Moreover, macrocosm experiments ($\sim\text{m}^3$ scale) should be favored over mesocosm ($\sim\text{dm}^3$ scale) experiments to

25 avoid or reduce inherent side effects that would ultimately hamper mimicking natural conditions.

6 CONCLUSION

Root water uptake is a key process in the global water cycle. More than 50% of total terrestrial evapotranspiration crosses plant roots to go back to the atmosphere (Jasechko et al., 2013). Despite its importance, quantification of root water uptake remains difficult due to the opaque nature of the soil and the spatial and temporal variability of the uptake process.

- 5 Water stable isotopic analysis is a powerful and valuable tool for the assessment of plant water sources and for the identification of hydraulic redistribution. In an inverse modelling framework, isotopic analysis of plant tissues and soil also allow for obtaining species-specific parametrization of physically-based analytical and numerical RWU models. They provide at the plant scale a unique way to tackle the difficulty of disentangling actual RWU profiles with root traits and characteristics.
- 10 In this review we tried to highlight the importance of systematically reporting uncertainties along with estimates of contribution to T of given plant water sources. The inter-comparison exercise could quantify the impact of the definition of the plant water sources (i.e., whether they are spatially disjoint or not and whether their isotopic compositions values are significantly different or not) on the outcome of the two end-member mixing model. The inter-comparison also illustrated the limitations of the graphical inference method and the multi-source mixing model of Phillips and Gregg (2003), whereas it
- 15 underlined the performance of the Bayesian approach of (Parnell et al., 2010), which uses a more rigorous statistical framework, if the number of considered water sources matches the number of isotopic measurements in the soil profile. However, contrary to the analytical model none of the graphical and statistical methods could locate and quantify hydraulic redistribution of water.
- 20 Finally, the authors call for (i) the further development of nondestructive and on-line isotopic measurement methodologies to circumvent the necessity of sampling soil material and plant organs destructively, and (ii) a generalization of coupled approaches relying on the confrontation between labelling experiments under controlled conditions and three dimensional RWU numerical modelling.

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AUTHOR CONTRIBUTION

Y. Rothfuss reviewed the published literature. Y. Rothfuss and M. Javaux designed the virtual experiments, analysed, and discussed the obtained results.

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TABLES

Table 1

Table 1: Summary of the reviewed studies that use one of either the three methods (graphical inference (GI), two-end members mixing model (TM), and “multi-source mixing models (MS)) for plant water sources partitioning.

Experimental conditions (field: F / laboratory: L)	Authors
F (tropical)	Graphical inference (GI) Leroux et al., 1995
F (semi-arid; cold desert, subhumid)	Thorburn and Ehleringer, 1995
F (temperate semi-arid savanna)	Weltzin and McPherson, 1997
F (tropical forest / savanna)	Jackson et al., 2009
F (eastern Amazon)	Moreira et al., 2000
F (desert)	Chimner and Cooper, 2004
F (shrub-steppe)	Kulmatiski et al., 2006
F (cold continental -semiarid)	Li et al., 2007
F (warm temperate / monsoon climate)	Wang et al., 2010
F (tropical)	Stahl et al., 2013
F (temperate)	Two-end members mixing model (TM) White et al., 1985
F (riparian zone)	Dawson and Ehleringer, 1991
F (aeolian sand dune)	Brunel et al., 1995
F (mediterranean)	Dawson and Pate, 1996
Field conditions (subtropical)	McCole and Stern, 2007
F (semi-arid)	Goebel et al., 2015
F (cornfield, prairie, oak savanna, and woodland)	Multi-source mixing model (MS) Asbjornsen et al., 2007
F (warm temperate / monsoon climate)	Wang et al., 2010
F (desert)	Huang and Zhang, 2015
F (temperate)	Prechsl et al., 2015
F (temperate)	Volkman et al., 2016

Plant species

Hyparrhenia diplandra, *Andropogon schirensis*, *Imperata cylindrica* (grasses), *Cussonia barteri*, *Crossopteryx febrifuga*, *Bridelia ferruginea* (shrubs)

Eucalyptus largiflorens, *camaldulensis*, *Acer negundo* and *grandidentatum*, *Atriplex canescens*, *Chrysothamnus nauseosus*, *Vancleavea stylosa*

Quercus emoryi Torr., *Trachypogon montufari* (H.B.K.) Nees.
(deciduous) *Qualea grandiflora* Mart., *Q. parviflora* Mart., *Kielmeyera coriacea* (Spr.) Mart., *Pterodon pubescens* Benth., *Dalbergia myscolobium* Benth. / (evergreen) *Didymopanax macrocarpum*, *Sclerolobium paniculatum*, *Miconia ferruginata*, *Roupala montana*

(invasive) *Solanum crinitum* Lamb. (native) *Panicum maximum* Jacq.

Sarcobatus vermiculatus, *Chrysothamnus nauseosus*, and *Chrysothamnus Greenei*

(invasive) *Centaurea diffusa*, (native) *Pseudoroegneria spicata*; *Bromus tectorum* L.

Larix sibirica

Summer corn and cotton (species not specified)

tropical rainforest trees

Taxodium distichum / *Pinus strobus*

Acer grandidentatum Nutt., *A. negundo* L., *Quercus gambelii* Nutt.

"Mallee Tree" (*Eucalyptus* sp.)

Banksia prionotes, *Dryandra sessilis*, *Grevillea* (species unknown)

Juniperus ashei

Gossypium hirsutum L.

Quercus macrocarpa, *Umus americana* L. (trees), *Zea mays* L. (crop), and *Andropogon gerardii* (grass)

Summer corn and cotton (species not specified)

Caragana korshinskii and *Artemisia ordosica*

Phleum pratense, *Lolium multiflorum*, *Poa pratensis*, *Taraxacum officinale*, *Trifolium repens*, *Rumex obtusifolius*, *Trisetum flavescens*, *Phleum rhaeticum*, *Carum carvi*, and *Achillea millefolium*, *Rumex alpestris*, *Taraxacum officinale* and *Trifolium pratense*.
Quercus petraea and *Fagus sylvatica*

<p>SAMPLES FOR ISOTOPIC MEASUREMENTS</p> <p>Soil profile (soil depth: SD [m] / increment: I [m] / number of profiles: Nps / Replicates)</p>	<p>Roots profile (soil depth: SD [m] / increment: I [m] / number of profiles: Npr)</p>
<p>0.10 < SD < 2.00 / 0.01 < I < 0.10 / R = 3</p> <p>No profiles, but soil directly surrounding roots are sampled SD = 1.50 / I = 0.05 / 3 < R < 4</p> <p>SD = 5.00 (depending on the site) R = 2 / 0.05 < I < 0.20 4 < SD < 6 / 0.05 < I < 2.00</p> <p>(1st campaign) SD = 0.6 / I = 0.10 / R = 2 at each site / 4 < Nps (per site) < 6 / (2nd campaign) SD = 2.1 / 0.2 < I < 0.30 / Nps = 10 1.05 < SD < 2.20 / 0.10 < I < 0.60</p> <p>SD = 1.00 / 0.05 < I < 0.30</p> <p>SD = 1.50 / 0.05 < I < 0.30 / Nps = 7</p> <p>SD = 2.00 / 0.20 < I < 0.30 / R = 6</p> <p>none</p> <p>SD = 0.50</p> <p>50 < SD < 4.2 / 0.10 < I < 0.25</p> <p>none (sand around the root system)</p> <p>SD = 0.30 / 0.05 < I < 0.10 / 4 < Nps (per site) < 5</p> <p>SD = 0.3 / I ~ 0.02 / Nps = 4 per irrigation treatment</p> <p>1.40 < SD < 2.00 / 0.05 < I < 0.20 / R=2 per site</p> <p>SD = 1.50 / 0.05 < I < 0.30 / Nps = 7</p> <p>SD = 2.00 / 0.05 < I < 0.50</p> <p>0.30 < SD < 0.40 / 0.04 < I < 0.10 / R = 3</p> <p>SD = 0.60 / 0.05 < I < 0.10 / Nps > 17</p>	<p>SD = 1.80 / I = 0.10 / R = 16</p> <p>No profiles, but single roots sampled after excavation none</p> <p>none</p> <p>SD = 4 / 0.25 < I < 0.50 / R = 3</p> <p>none (reference made to Cooper and Chimmer, unpublished data)</p> <p>1.20 < SD < 2.20 / I = 0.15</p> <p>none</p> <p>none</p> <p>none (reference made to other literature)</p> <p>none</p> <p>none</p> <p>none</p> <p>none (qualitative observation)</p> <p>none (reference made to other literature) none</p> <p>none</p> <p>none</p> <p>none</p> <p>SD = 0.30 / 0.075 < I < 0.125 / 6 < Npr < 7</p> <p>SD = 0.60 / 0.05 < I < 0.10 / R = 4</p>

<p>Additional isotopic meas. (groundwater: G / spring: S / river: R / precipitation: P)</p>	<p>Plant (organ: O / number of samples: Ns / replicates: R / temporal resolution: TR [hrs])</p>
<p>none</p>	<p>O: sapwood trunks (shrub), Crown (grasses) / 8 < Ns < 24 / 0.5 < TR < 1</p>
<p>none</p>	<p>O: non-green, suberized stems / Ns = 3</p>
<p>P</p>	<p>O: stem with phloem tissue / R = 4 (tree and sapling) / O: stem without green tissue / R = 4 (seedling) / O: culm base with sheaths removed / 3 < R < 4</p>
<p>none</p>	<p>O: wood or suberized, mature, stem segments (outer bark and phloem are removed) / R = 2</p>
<p>none</p>	<p>O: well suberized stems (trees) / thick fleshy culms covered with dry leaves (grass) / 3 < R < 5</p>
<p>GW / P / R</p>	<p>O: fully suberized stem sections from the base of plants / (1st campaign, S. Vermiculatus) 4 < Ns (per site) < 8 / R = 3 / (1st campaign, C. nauseosus) 5 < Ns (per site) < 6 / R = 3 / (2nd campaign) 3 < Ns < 5 (depending on the species)</p>
<p>P / Snow</p>	<p>O: stem</p>
<p>P / R</p>	<p>O: stem / Ns (per sampling date) = 5</p>
<p>P / GW / Irrigation water</p>	<p>O: stems (epidermis contacted with air was removed) / Ns = 7 / TR = 1 per vegetation stage</p>
<p>none</p>	<p>O: branch (length = 0.07 m; diameter = 0.1 – 0.3 m). Bark tissue is immediately removed</p>
<p>P / GW / R</p>	<p>O: wood samples taken at breast height</p>
<p>P / Stream / S</p>	<p>O: mature suberized stems</p>
<p>none</p>	<p>O: twigs (bark is removed) / R = 2</p>
<p>GW / P</p>	<p>O: roots, trunks, stem base</p>
<p>P / S</p>	<p>O: stem</p>
<p>P</p>	<p>O: meristematic petiole reduced in size to 5mm. / 14 < Ns < 17 (depending on the irrigation treatment) / R = 2 / TR = 1</p>
<p>none</p>	<p>O: stem (trees); aerial nodal roots just above the soil surface (Zea mays L.); stem (grass) / Ns = 2 per specie</p>
<p>P / GW / Irrigation water</p>	<p>O: stems (epidermis contacted with air was removed) / Ns = 7 / TR = 1 per vegetation stage</p>
<p>P</p>	<p>O: twigs (1–2 cm of stem with bark immediately removed) / R = 3</p>
<p>none</p>	<p>O: root crown / 2 < Ns < 10</p>
<p>none</p>	<p>none (measurement of transpiration isotopic composition)</p>

Natural Abundance: NAb / labelling Experiment: LE	Identified plant-water sources	Water extraction method (cryogenic vacuum distillation: Cr / azeotropic distillation Az / direct equilibration: GI / mild vacuum distillation: MI)	NON-ISOTOPIC MEASUREMENTS soil water Content: SWC / soil water potential: SWP / soil temperature: ST / leaf water potential: LWP / stem water potential: StWP / sap flow (heat pulse technique): SF / transpiration flux T / precipitation amount: Pa / leaf area index: LAI / depth of water table: DWT
Nab	Soil water across depths	Cr	SWC / LWP (not shown)
Nab	Soil water across depths	Cr / Az	SWP / StWP / SF
Nab	Soil water across depths	Cr	SWC / Pa
Nab	Soil water across depths	Cr	SWP / ST / StWP / SF
LE	Soil water at isotopic labeled depth	Cr	none
Nab	Soil water across depths	Cr	Pa / LAI
Nab	Soil water across depths	Cr	SWC / Pa / DWT
Nab	Soil water across depths	Cr	Pa
Nab	Soil water across depths	Cr	SWP
LE	Soil water across depths	Cr	SWC / Predawn LWP
Nab	GW / P	none (Sap flow water)	none
Nab	GW / Stream water	Cr	none
Nab	Upper / lower soil profile	Az	DWT
Nab	Upper / lower soil profile	Mi	none
Nab	Upper / lower soil profile	Cr	Pa
Nab	Precipitation / Irrigation	Cr	Pa
Nab	Soil water across depths	Di	none
Nab	Soil water across depths	Cr	SWP
Nab	Soil water across depths	Cr	Pa
NAb	Soil water across depths	Cr	SWC / Pa / Aboveground biomass dry weight
LE	Soil water across depths	none	SWC / T

Single: S / Dual: D /
"Double Single": DS"
isotope approach

S ($\delta^{18}\text{O}$)

S ($\delta^2\text{H}$)

D

S ($\delta^2\text{H}$)

S ($\delta^2\text{H}$)

S ($\delta^{18}\text{O}$)

DS

DS

DS

D

S ($\delta^2\text{H}$)

S ($\delta^2\text{H}$)

D

S ($\delta^2\text{H}$)

S ($\delta^{18}\text{O}$)

S ($\delta^{18}\text{O}$)

S ($\delta^{18}\text{O}$)

DS

DS

DS

S ($\delta^2\text{H}$)

Main results (RWU depth: z_{RWU} [m] / soil depth z [m] / fraction of transpiration: x / Source: So)

$0.00 < z_{RWU} < 0.05$ (grasses, early morning) / $0.05 < z_{RWU} < 0.10$ (grasses, midday) / $z_{RWU} = 0.30$ or $z_{RWU} > 1.50$ (shrubs, no unique solution)

Dominant source: groundwater (Mountain and floodplain) / $0.3 < z_{RWU} < 0.4$ (cold desert)

$z_{RWU} > 0.50$ (trees and sapling) / $z_{RWU} < 0.15$ (2-months-old seedling) / $0.20 < z_{RWU} < 0.35$ (1- and 2-year-old seedlings and grasses)

$z_{RWU} < 2.00$ (four evergreen and one deciduous species) / $z_{RWU} > 2.00$ (three deciduous and one evergreen species)

Fraction of root water uptake from the labeled region $x = 0.20$ (Solanum crinitum Lamb.) $z_{RWU} < 1.00$ (P. maximum Jacq.)

$0.00 < z_{RWU} < 0.50$ (C. nauseosus, pre-monsoon and monsoon periods) / (S. vermiculatus and C. nauseosus) dominant source: groundwater (pre-monsoon period) and switched to precipitation recharged water ($0.30 < z_{RWU} < 0.40$, during monsoon).

$z_{RWU} = 0.15$ (Bromus tectorum L., early season) / $z_{RWU} = 1.20$ (Centaurea diffusa Lam, late season)

$0.10 < z_{RWU} < 0.40$ from $\delta^{18}O$ meas / $0 < z_{RWU} < 0.80$ from δ^2H meas
 $z_{RWU} = 0.10$ (Corn, jointing stage) / $z_{RWU} = 0.50$ (Corn, flowering stage) /

$z_{RWU} = 0.10$ (Corn, full ripe stage) / $z_{RWU} = 0.40$ (Cotton, seedling stage) / $z_{RWU} = 0.50$ (Cotton, bud stage) / $z_{RWU} = 1.10$ (Cotton, bolls open stage)

$z_{RWU} > 1.00$ (dry periods, tall trees) / More diffuse z_{RWU} for shorter trees

Groundwater: $0.46 < x < 0.64$ (wet site) / $0.16 < x < 0.25$ (intermediate site)

So: Groundwater (streamside mature trees) / So: stream water (younger streamside trees) / So: Precipitation (younger non-streamside trees)

$0 < z_{RWU} < 0.4$ with $0.7 < x < 0.9$ (1st site) / $0 < z_{RWU} < 1.5$ with $x = 0.07$ (2nd site)

So: deeper soil layers (dry season) / So: shallow soil layers (wet season)

$z_{RWU} > 0.30$ m (hot, dry summer) / $0.10 < z_{RWU} < 0.30$

Evidence for shifting to rainwater predominantly

$0.0 < z_{RWU} < 0.20$ m with $x < 45\%$ and $x < 36\%$ (crop and grass, resp.) / $0.00 < z_{RWU} < 0.20$ m with $x < 40\%$ and $x < 20\%$ and $z_{RWU} > 0.60$ m with $x < 60\%$ and $x < 80\%$ (Q. macrocarpa and U. americana L., resp.)

$0 < z_{RWU} < 0.20$ with $96 < x < 99$ (Corn, jointing stage) / $0.20 < z_{RWU} < 0.50$ cm with $58 < x < 85$ (Corn, flowering stage), $0.00 < z_{RWU} < 0.20$ with $69 < x < 76$ (Corn, full ripe stage) // $0.00 < z_{RWU} < 0.20$ with $27 < x < 49$ (Cotton, seedling stage), $0.20 < z_{RWU} < 0.50$ cm with $79 < x < 84$ (Cotton, bud stage) / $0.50 < z_{RWU} < 0.90$ with $30 < x < 92$ (Cotton, blooming stage) / $z_{RWU} > 0.90$ cm with $69 < x < 92$ (Cotton, boll open stage)

$0.1 < z_{RWU} < 1.0$ (wet seasons) / z_{RWU} not affected by small rainfall events / $z_{RWU} > 1.00$ with x observed to increase from z (± 0.7) to 10 (± 1.4) % for both plants after large rainfall event.

$0 < z_{RWU} < 0.10$ with $0.43 < x < 0.68$ (Drought treatment) / $0 < z_{RWU} < 0.10$ with $0.04 < x < 0.37$ and $0.20 < z_{RWU} < 0.35$ with $0.29 < x < 0.48$ (control treatment)

constant RWU depth profile with $0.15 < x < 0.18$ (beech) / $x < 0.15$ for $z < 0.20$ and $0.15 < x < 0.25$ for $z > 0.30$ (beech/oak mixture and oak monoculture)

1 **Table 2**

SOIL DATA z (m)	Shallow groundwater table (Sh)						Deep groundwater table (De)						RLD (cm cm ⁻³)
	Dry surface conditions (ShDr)			Wet surface conditions (ShWe)			Dry surface conditions (DeDr)			Wet surface conditions (DeWe)			
	θ (cm ³ cm ⁻³)	H_s (cm)	δ_s (‰)	θ (cm ³ cm ⁻³)	H_s (cm)	δ_s (‰)	θ (cm ³ cm ⁻³)	H_s (cm)	δ_s (‰)	θ (cm ³ cm ⁻³)	H_s (cm)	δ_s (‰)	
-0.01	0.235	-454	5	0.372	-2	-7	0.044	-9875	11	0.372	-51	-5	6.0
-0.03	0.325	-267	3	0.372	-8	-6	0.055	-3581	7	0.371	-77	-5,5	3.0
-0.07	0.347	-215	1	0.372	-11	-5	0.081	-1661	1	0.372	-14	-7	2.0
-0.15	0.360	-179	-4	0.372	-70	-6	0.105	-1165	-3,5	0.135	-869	-3,5	0.8
-0.30	0.367	-155	-6	0.370	-125	-6,5	0.122	-989	-4	0.134	-889	-4	0.5
-0.50	0.371	-135	-7	0.371	-135	-7	0.165	-730	-5	0.165	-730	-5	0.4
-1.00	0.372	-125	-7	0.372	-125	-7	0.210	-620	-6	0.210	-620	-6	0.3
-2.00	0.372	-125	-7	0.372	-15	-7	0.259	-600	-7	0.259	-600	-7	0.2
PLANT DATA	T (mm h ⁻¹)	H_L (cm)		T (mm h ⁻¹)	H_L (cm)		T (mm h ⁻¹)	H_L (cm)		T (mm h ⁻¹)	H_L (cm)		
IT	0.01	-587		0.01	-491		0.01	-2347		0.01	-918		
hT	0.30	-12330		0.30	-12234		0.30	-14090		0.30	-12661		

2
3 **Table 2: Soil, plant, and isotopic input data for the different modelling approaches (depth (z) profiles of soil water**
4 **content θ , total soil water potential H_s , soil water oxygen isotopic composition δ_s , root length density RLD ,**
5 **transpiration rate T , and leaf water potential H_L) “collected” during eight virtual experiments differing in the depth**
6 **of the groundwater table (Shallow –Sh / Deep – De) and the water status at the soil surface (Dry – Dr / Wet – We).**

1 Table 3

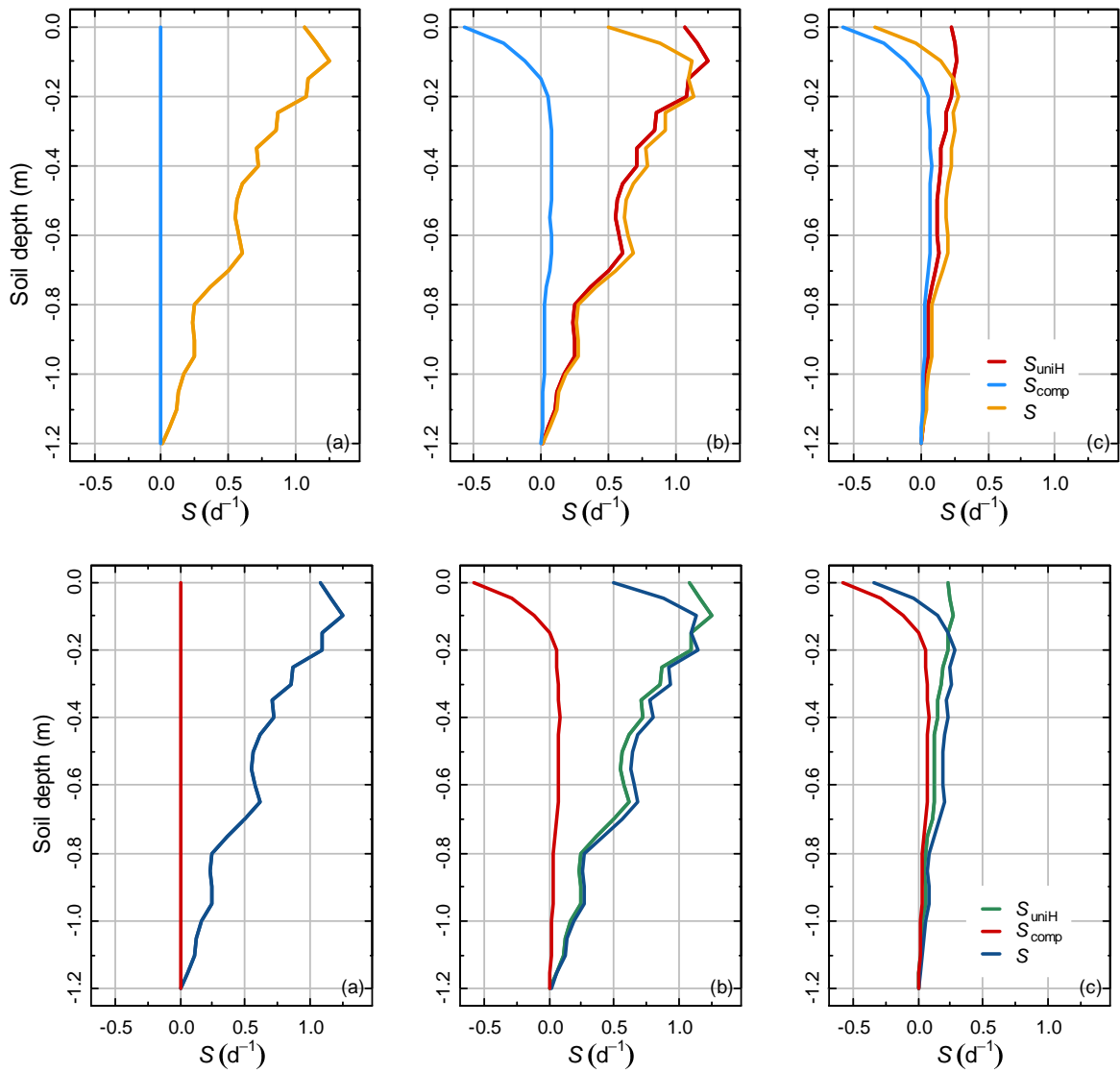
Soil layer r (m)	Shallow groundwater table (Sh)							
	Dry surface conditions (ShDr)				Wet surface conditions (ShWe)			
	Low T (ShDr_IT)		High T (ShDr_hT)		Low T (ShWe_IT)		High T (ShWe_hT)	
x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	
0-0.02	1,4(0-34,65)	4,85(9,71)	6(0-37)	10,91(14,31)	17,58(0-48)	13,4(1,4)	5,86(0-52,93)	11,2(1,3)
0.02-0.05	1,4(0-35,4)	7,3(0,91)	5(0-37,8)	8,79(10,9)	13,4(0-41,52)	10,4(1)	7,4(0-42,83)	8,89(10,9)
0.05-0.11	2,53(0-40,81)	10,71(1,2)	9,810(0-48,4)	11,3(1,1)	0,91(0-40,51)	13,2(1,1)	7,4(0-41,2)	11,4(1,1)
0.11-0.225	4,55(0-56,67)	9,910(0,71)	13,94(0-47,2)	9,710(0,51)	11,3(0-45,66)	10(0,51)	3,4(0-42,93)	9,710(10,5)
0.225-0.4	19,4(0-56,57)	10,61(0,3)	16(0-54,65)	9,810(0,4)	5,86(0-52,83)	9(0)	16,4(0-48,59)	9,810(7,4)
0.4-0.75	16,2(0-54,95)	16,3(0,2)	16,7(0-47,98)	14,4(0,4)	17,58(0-44)	12,73(0,51)	15,2(0-48,4)	14
0.75-1.5	17,4(0-52,4)	26,77(1,82)	18,4(0-46,4)	23,2(1,92)	5,86(0-47,98)	20,81(1,92)	16,2(0-52,83)	14
1.5-2	16,67(0-59,2)	13,74(1,82)	17,4(0-47,2)	14,92(1,72)	15,3(0-52,2)	10,71(1,62)	16(0-51,2)	14
		3,8		4,9		4,4		4,7
		6,2		3,7		2,7		10,3
	4)	8,3	2)	1,5	3)	2,2	4)	4,4
		4,6		4,2		1,4		11,6,3
		8,8		6,1		6,9		6,3
		0,0		2,2		4,8		0,8
		9,6		4,9		4,9		12,6,8
		2,8		4,6		3,9		4,4
	MD	5,5		4,1		3,9		13
Soil layer r (m)	Deep groundwater table (De)							
	Dry surface conditions (DeDr)				Wet surface conditions (DeWe)			
	Low T (DeDr_IT)		High T (DeDr_hT)		Low T (DeWe_IT)		High T (DeWe_hT)	
x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	x_{MSPa} mfv(range) (%)	x_{AM} (1sd) (%) Abs.-diff. (%)	
0-0.02	1(0-42,2)	-170(16)	1(0-40,91)	5,4(0,71)	4,52(0-48,9)	23,64(2,3)	10,4(0-52,3)	14,51(9,3)
0.02-0.05	1(0-41,72)	-17(1)	2,2(0-45,2)	7,98(0,91)	6,67(0-54,85)	18,4(1,62)	13,2(0-54)	2,8(1)
0.05-0.11	1,2(0-44)	19(6)	5,3(0-47,2)	14,62(1,2)	16,4(0-)	21(1,52)	16(0-50,51)	11,72(1,1)
0.11-0.225	2,63(0-55)	28(5)	11,3(0-51,2)	10,3(0,71)	5758,6)	3(0,2)	12(0-43)	9,2(1,51)
0.225-0.4	6,97(0-75)	33(4)	17(0-50,1)	10,61(0)	4,21(0-38,99)	1,3(0)	8,79(0-38)	9,510(0)
0.4-0.75	15(0-68)	57(3)	17(0-56)	16(0)	0,8(0-38)	7(1)	15(0-53)	14(0)
0.75-1.5	16(0-74)	98(1)	16(0-54)	26(2)	5(0-46)	17(2)	16(0-45)	23(2)
1.5-2	17(0-76)	51(4)	18(0-53)	13(2)	16(0-51)	9(2)	16(0-46)	24(2)
								25
0.4-0.75	5)	-169,5(15,8)	5)	2)	8)	4)	4)	26,4)
0.75-1.5	14,5(0-67,7)	-16,9(1,4)	17,1(0-55,9)	15,8(0,3)	4,5(0-46,2)	6,9(0,8)	15,4(0-52,5)	14,2(7,4)
1.5-2	16,4(0-73,8)	18,9(6)	16,2(0-54,4)	25,6(1,8)	16,1(0-51,2)	16,6(2,3)	16(0-45,4)	22,8(1,9)
	16,7(0-76,4)	27,7(6)	17,5(0-53,3)	43,2(1,7)	17,9(0-53,2)	9,2(1,8)	15,9(0-46,2)	11,5(1,7)
		33,3(3,8)		4,4		22,2		29,1,4
		57,3(3,3)		5,6		1,9		30,4,4
		98(0,6)		6,3		4,5		4,3
		51,1(3,5)		1,1		1,8		31,2,6
		170,4		6,3		0,4		32,0,9
		17,8		1,3		2,4		33,1,3
		17,7		9,3		0,5		6,9
		25,1		4,3		8,8		34,1,4
		26,4						35
		42,9						36
		81,6						37
		34,4						38
	MD	52,4		4,8		5,3		

Table 3: Most frequent value (mfv) and range of the density distribution curve of the relative contribution to transpiration across eight defined soil layers as determined by the Bayesian method of Parnell et al. (2010) (x_{MSPa} , %) and mean relative contribution (with standard deviation) provided by the analytical model of Couvreur et al. (2012) (x_{AM} , %). Profiles of relative contribution were computed for eight soil-plant virtual experiments differing in the depth of the

39 groundwater table (shallow – Sh / deep –De), the soil surface water status (dry – Dr / wet – We), and the plant
 40 transpiration rate (low – IT / high – hT). The absolute difference between the x_{MSPa} -mfv and x_{AM} for each soil layer
 41 (Abs.-diff., %) and the mean absolute difference (MD) for each soil-plant experiment are also reported.

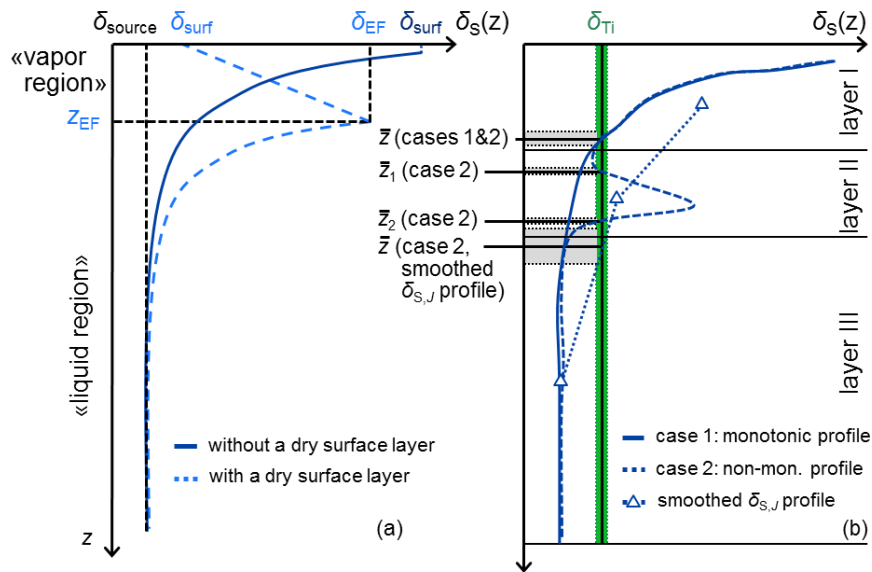
1 FIGURES

2 Figure 1



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5 Figure 1: Some examples of root water uptake sink term (S , in d^{-1}) profiles (orange-blue lines) conceptualized as the
6 sum of two vertically distributed components, the root water uptake term proportional to root distribution (S_{uniH} , blue
7 green lines) and the compensatory root water uptake (S_{comp} , red lines) accounting for heterogeneous soil water
8 potential distribution. (a) $S_{comp} = 0$ (no root compensation, i.e., soil water potential profile is homogeneous) leading to
9 a situation where $S = S_{uniH}$. (b) S_{comp} is (i) different than zero and (ii) becomes negative **at towards** the surface but
10 remains smaller (in absolute term) than S_{uniH} . (c) S_{comp} is (i) different than zero and (ii) becomes negative at the
11 surface while becomes greater (in absolute term) than S_{uniH} at the surface ($z > -0.08$ m). In the last case, S is negative
12 at the surface, meaning hydraulic redistribution – more specifically hydraulic lift – is observed.

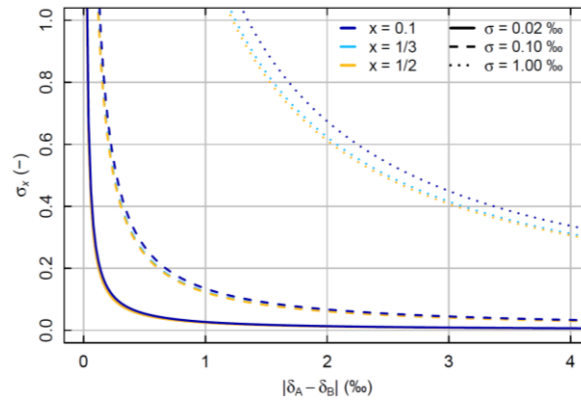
1 **Figure 2**



2

3 **Figure 2: (a) Simulated soil water isotopic composition (δ_s) profiles under isothermal conditions for a water saturated**
 4 **(~~black-dark blue~~ line) and unsaturated (~~gray-light blue dotted~~ line) soil following Zimmermann et al. (1967) and**
 5 **Barnes and Allison (1983). Indices “surf” and “EF” refer to soil surface and Evaporation Front. “vapor” and “liquid”**
 6 **regions refer to soil regions where water flow occurs predominantly in the liquid and vapor phase, respectively. (b)**
 7 **Illustration of the “graphical inference” (GI) method for determining the “mean root water uptake depth” (\bar{z}) as the**
 8 **soil depth where $\delta_s = \delta_{Ti}$ with “Ti” standing for the sap xylem water at the plant tiller. Case 1 represents the condition**
 9 **for which a unique solution is found and case 2 the condition with more than one solution due to a heterogeneous δ_s**
 10 **profile. In the latter case, a possibility is to smooth the δ_s profile (using Eq. (5)) until it is monotonous (smoothed**
 11 **profile is designated by the symbols). In this example, the δ_s profile is smoothed over three layers and a uniform soil**
 12 **water content profile is implied. \bar{z} range (gray horizontal stripes) is determined by taking into account the uncertainty**
 13 **associated with measurement of δ_{Ti} (i.e., precision of the isotopic analysis; green vertical stripe).**

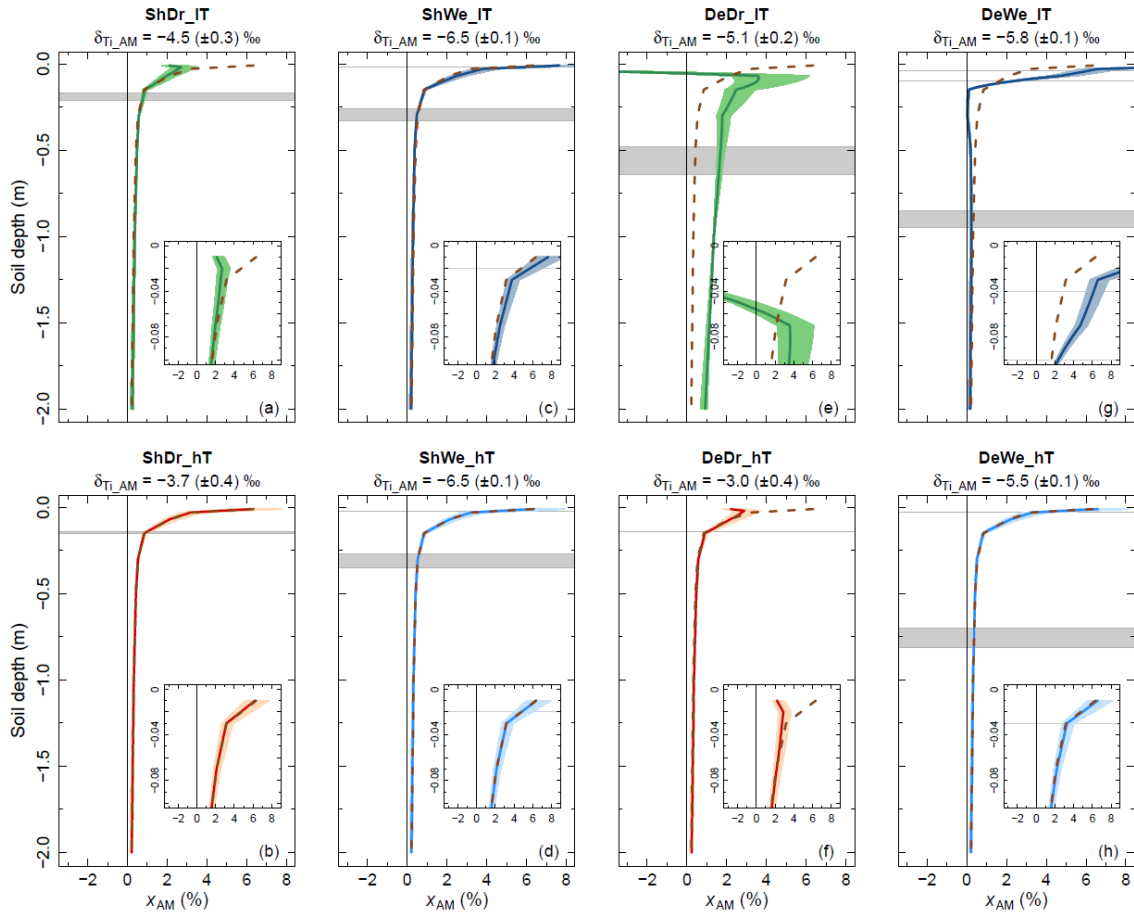
1 **Figure 3**



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Figure 3: Standard error (σ_x) associated with the estimation of the relative contribution (x) of source A water to root water uptake in case of two distinct sources (A and B of isotopic compositions δ_A and δ_B). Following Eqs. (8a) and (8b) (Phillips and Gregg, 2001), σ_x is a function of the difference ($\delta_A - \delta_B$) (represented in absolute term in this Figure), of the value of x (three values are tested here: 0.1, 1/3, and 1/2, corresponding to dark blue, light blue and orange lines), and of the standard errors associated with sampling and measurement of δ_A , δ_B , and of the isotopic composition of the tiller sap water (δ_{Ti}) (three values are tested, i.e., 0.02, 0.10, and 1.00, corresponding to solid and dashed lines).

1 **Figure 4**

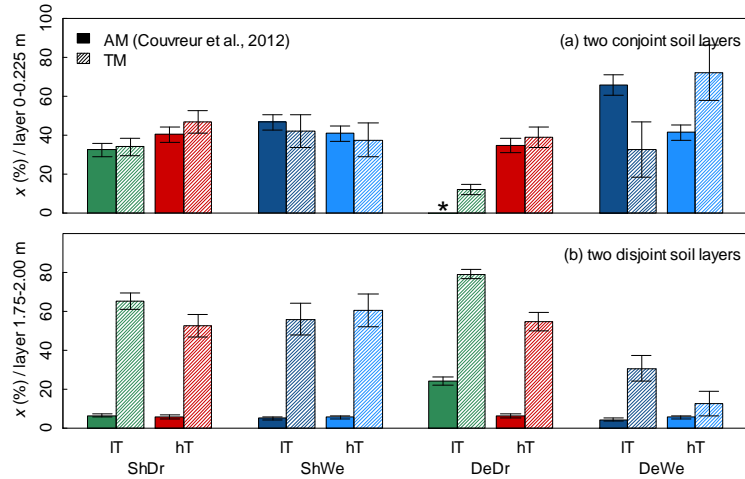


2

3 **Figure 4: Simulated depth (z , in m) profiles of x_{AM} (%) (solid colored lines), the simulated ratio $\frac{S_{AM}(z)dz}{T/(\Delta x \cdot \Delta y)}$**

4 provided by the model of Couvreur et al. (2012) (Eq. (B4'), see Appendix B) on the basis of input data reported in
 5 Table 2 for experiments “ShDr” (soil with shallow groundwater table and relatively dry soil surface), “ShWe” (soil
 6 with shallow groundwater table after a rainfall event), “DeDr” (soil with deep groundwater table with relatively dry
 7 soil surface), “DeWe” (soil with deep groundwater table after a rainfall event). Suffices “IT” and “hT” refer to “low”
 8 and “high” transpiration rate simulations. Note that negative x_{AM} means hydraulic redistribution (HR) by the roots.
 9 The color-shaded areas depict the results of 1000 model runs where for each input data variable (soil water potential,
 10 $\delta^{18}O$, and root length density – RLD) a single offset randomly selected between -5 and $+5$ cm, -0.2 and $+0.2$ ‰, and $-$
 11 0.1 and $+0.1$ cm cm^{-3} respectively for each variable was added to the initial values reported in Table 2. This should
 12 represent the uncertainty of the model estimates on the account of the precision of the measurements. The horizontal
 13 gray-shaded areas delimit the soil layers that contribute to RWU as obtained by the “graphical inference” (GI)
 14 method, i.e., by locating the depth of the intersection between a vertical line of value $\delta_{TL,AM}$ and the soil water isotopic
 15 profile. Layers’ lower and upper boundaries are inferred from the uncertainty of the $\delta_{TL,AM}$ estimates for each
 16 scenario ($\delta_{TL,AM}$ is given above each plot along with its standard deviation). At the bottom right corner of each plot is
 17 ~~represented~~ a detail ~~presented~~ for $z \geq -0.10$ m. Finally, results from the first term of the model of Couvreur et al.
 18 (2012) which considers uptake proportional to RLD (S_{uniH} , Eq. (B4)) is plotted as a dashed brown line for comparison.

Figure 5



5 | Figure 5: Comparison between relative contributions to transpiration (x , in %) from one soil layer simulated by the analytical RWU model of Couvreur et al. (2012) and the two end-member mixing model (TM, see section §3.2.1) in case of two defined soil layers. Figure 5a displays x from the topmost soil layer (0 – 0.225 m) in case of a two conjoint soil layers (0 – 0.225 m and 0.225 – 2.00 m) whereas Figure 5b displays x from the lowest soil layer (1.75 – 2.00 m) in case of a two disjoint soil layers (0 – 0.225 m and 0.225 – 2.00 m), i.e., information on soil water isotopic composition is lacking between 0.225 and 1.75 m. “Sh” (“De”) stands for the virtual experiments where the soil has a shallow (deep) groundwater table while “Dr” and “We” stand for when the soil is dry or wet at the surface (e.g., shortly after a rain event). Suffices “IT” and “hT” refer to “low” and “high” transpiration rate simulations. “*” refers to when hydraulic redistribution is simulated by the analytical model, leading to a negative x . Error bars refer to either one standard deviation (for the RWU analytical model) or one standard error (for the TM approach, [calculated using Equation \(8b\)](#)).

10 |

Figure 6

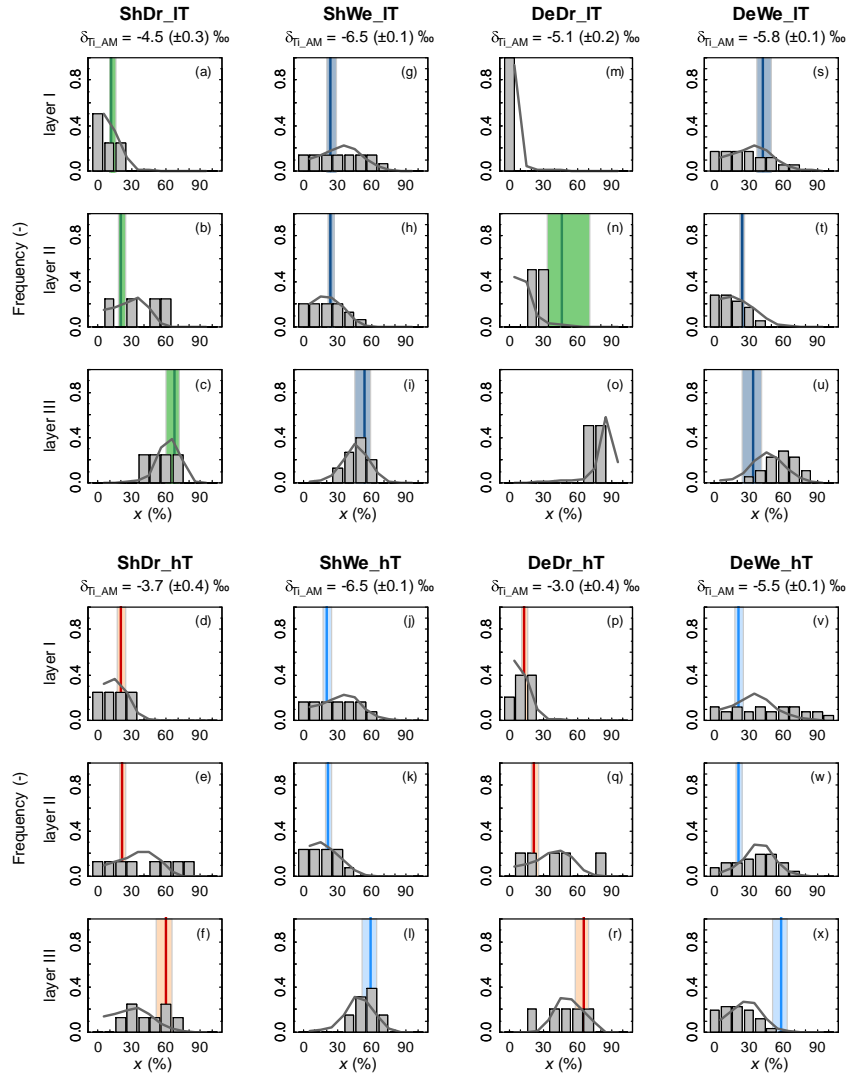


Figure 6: Simulated ranges of possible relative contributions to transpiration from three defined soil layers (I: 0.00 – 0.05 m, II: 0.050 – 0.225 m, and III: 0.225 – 2.000) following the method of Phillips and Gregg (2003) (x_{MSPG} , in %, displayed in the form of gray histograms). Density distributions following the Bayesian approach of Parnell et al. (2010) (x_{MSPa} , gray lines). x_{MSPG} and x_{MSPa} were obtained from the confrontation of soil water $\delta^{18}\text{O}$ profiles (Table 1) and the δ_{TI} simulated by the model of Couvreur et al. (2012) (i.e., $\delta_{\text{TL,AM}}$, given above each plot along with its standard deviation). Tolerance of the MSPG was set equal to the standard deviation of $\delta_{\text{TL,AM}}$. “Sh” (“De”) stands for the virtual experiments where the soil has a shallow (deep) groundwater table while “Dr” and “We” stand for when the soil is dry or wet at the surface (e.g., shortly after a rain event). Suffices “IT” and “hT” refer to “low” and “high” transpiration rate simulations. The colored vertical lines give $x_{\text{I,AM}}$, $x_{\text{II,AM}}$, and $x_{\text{III,AM}}$, the ratios $\frac{S_{\text{AM}}(z)dz}{T/(\Delta x \cdot \Delta y)}$ from layers I, II, and III to transpiration rate as simulated by the analytical model of Couvreur et

al. (2012). The color-shaded areas associated with $x_{\text{I,AM}}$, $x_{\text{II,AM}}$, and $x_{\text{III,AM}}$ vertical lines refer to their uncertainty associated with input data uncertainty (i.e., ± 5 cm for soil water potential, ± 0.2 ‰ for soil water $\delta^{18}\text{O}$, and ± 0.1 cm $^{-3}$ for root length density).

APPENDIX

Appendix A: List of symbols

Symbol	Description	Dimension	Equation number	Measured (m) / simulated (s) / prescribed (p)
C, C_S, C_A, C_B, C_{Ti}	Water stable isotopic concentration, soil water stable isotopic concentration, sources A and B water stable isotopic concentrations, xylem sap water isotopic concentration, root water uptake isotopic concentration	$M L^{-3}$	2, 3, 6a, 6b	m s
E, E_i	Evaporation rate for $^1H_2^{16}O$ isotopologue, Evaporation rate for $^1H_2^{16}O$ or $^1H_2^{18}O$ isotopologue	$L^3 T^{-1}$	B1-B4	m/s
h	Matric head	L		m
H_{eq}, H_L, H_S	Soil water equivalent and leaf water potentials, total soil water potential	P		m
J_A, J_B, J_{Ti}	Fluxes of water originating from water sources A and B, and at the plan tiller	$L^3 T^{-1}$	6b	m
$J_i^A, J_i^B, \text{ and } J_{Ti}^i$	Fluxes of isotopologues originating from water sources A and B, and at the plan tiller	$M T^{-1}$	6a	m
K_{plant}, K_{comp}	Plant and compensatory conductances to water flow	$L^3 P^{-1} T^{-1}$	B1-B4	m/p
M_w, M_i	Molar masses of water and isotopologue ($^1H_2^{16}O$ or $^1H_2^{18}O$)	$M L^{-3}$	3	m
RLD RLD_{1D}	Root length density Root length density per unit of surface area	$L L^{-3}$ $L L^{-1}$	B3	m/p m/p
R_{ref}	Vienna-Standard Mean Ocean Water (V-SMOW) hydrogen or oxygen stable isotopic ratio	-	3	m
$S, S_{uniH}, S_{comp}, S_{AM}$	Root water uptake sink term, Root water uptake sink term under uniform soil water potential distribution, compensatory root water uptake sink term, total root water uptake sink term as simulated by the analytical model of Couvreur et al. (2012)	$L^3 L^{-3} T^{-1}$	1-4, 6b, B4,B5, 6b	s
SSF	Standard sink fraction	-	B2, B4, B4'	m/p
$t, \Delta t$	Time, time step	T	11	m
T	Transpiration flux	$L^3 T^{-1}$	2, 4a, 4b, B1, B3, B4	m
x, x_j $x_{AM}, x_{J_AM}, x_{J_MSPG},$ x_{J_MSPa}	Contributive proportion to transpiration, source j contributive proportion to transpiration, continuous and integrated (layer J) contributive proportions to transpiration as simulated by the analytical model of Couvreur et al. (2012), integrated (layer J) contributive proportions to transpiration as determined by the statistical approaches of Phillips and Gregg (2003) and Parnell et al. (2010). Contributive proportion to transpiration under conditions of uniform soil water potential	-	7, 8b, 9, 9'	s
$z, z_j, z_{j+1}, \Delta z_j, z_{max},$ \bar{z}_{RWU}	Soil depth, soil depth of layers j and $j+1$, thickness of soil layer j , depth of the root system, "mean root water uptake depth"	L	4b, 5, B2-B4'	m/p

α_{eq}, α_K $\alpha_{eq}^{2H}, \alpha_{eq}^{18O}, \alpha_K^{2H}, \alpha_K^{18O}$	Equilibrium and kinetic isotopic fractionation factors, hydrogen and oxygen equilibrium isotopic fractionation factors, hydrogen and oxygen kinetic isotopic fractionation factors	-		m/s
$\delta, \delta^2H, \delta^{18O}, \delta_{li}, \delta_{v}, \delta_{int}, \delta_{source}, \delta_{surf}, \delta_{sim}, \delta_{S_i}, \delta_{S_j}, \delta_{S_j}, \delta_A, \delta_B, \delta_{Ti}, \delta_{Ti,m}, \delta_{Ti,AM}, \delta_{E_i}, \delta_{F_i}$	Water stable isotopic composition, water hydrogen and oxygen stable isotopic compositions, liquid, vapor, liquid-vapor interface, source, soil surface, and simulated water, isotopic compositions, soil water isotopic composition, soil layer <i>j</i> and <i>J</i> water isotopic composition, sources A and B water stable isotopic compositions, isotopic composition of xylem sap water at the plant tiller, isotopic composition of xylem sap water measured at the plant tiller, isotopic composition of xylem sap water at the plant tiller as simulated by the model of Couvreur et al. (2012), isotopic composition of transpiration	- (expressed in ‰)	3-5, 7-9, 11	m/s
ϵ_j	Residual error term	- (expressed in ‰)	9'	s
θ	Soil volumetric water content	$L^3 L^{-3}$	5, 11	m
ρ	Volumetric mass of water	$M L^{-3}$	3	m
$\sigma_x, \sigma_{\delta_A}, \sigma_{\delta_B}, \sigma_{\delta_{Ti}}, \sigma_{\delta_{Ti,AM}}, \sigma_x$	Standard errors associated with the measurements of $x, \delta_A, \delta_B, \delta_{Ti}$ and estimated uncertainty of $\delta_{Ti,AM}$ as simulated by the analytical model of Couvreur et al. (2012), error associated with the estimation of the contributive proportion to <i>T</i> of water source A in the case of two distinct sources	- (expressed in ‰)	8a, 8b	s
τ	Isotopic tolerance	- (expressed in ‰)	10	p

Appendix B: The macroscopic RWU model of Couvreur et al. (2012)

B1: Presentation of the model

In the approach of Couvreur et al. (2012), RWU is based on physical equations describing the water flow processes but without the need of the full knowledge of the root system architecture and local hydraulic parameters. Instead, three
 5 macroscopic parameters are needed. The first equation defines plant transpiration:

$$J_{Ti} = K_{\text{plant}} \cdot (H_{\text{eq}} - H_L) \quad (\text{B1})$$

where J_{Ti} [$\text{L}^3 \text{T}^{-1}$] is the sap flow rate in the root tiller and considered to be equal to the transpiration rate, K_{plant} [$\text{L}^3 \text{P}^{-1} \text{T}^{-1}$] is the plant conductance to water flow (the first macroscopic parameter of Couvreur et al., 2012's model). H_L [P] is the leaf water potential and H_{eq} [P] the “plant averaged soil water potential” defined as the mean soil water potential “sensed” by the
 10 plant root system in the one dimensional (vertical) space:

$$H_{\text{eq}} = \int_z SSF(z) \cdot H_S(z) \quad (\text{B2})$$

where z is the soil depth, H_S [P] is the total soil water potential, and SSF [-] the standard sink term fraction (the second macroscopic parameter of the model of Couvreur et al., 2012). SSF is defined as the RWU fraction under the condition of totally uniform soil water potential (i.e., when $H_S(x,y,z) = H_S = \text{cst}$). Under such conditions, if all the root segments had the
 15 same radial conductivity (and the xylem conductance would not be limiting), the RWU distribution in a uniform soil water potential profile would be exactly the same as the root length density per unit of surface area (RLD_{1D} of dimension [L L^{-1}]) profile. SSF could be defined as:

$$SSF(z) = \frac{S_{\text{uniH}}(z) dz}{q_{Ti}} \approx \frac{RLD_{1D}(z) \cdot dz}{\int_z RLD_{1D}(z) \cdot dz} \quad (\text{B3})$$

where $q_{Ti} = J_{Ti}/(\Delta x \Delta y)$ represents the sap flow rate in the root tiller per unit surface area [L T^{-1}], S_{uniH} [T^{-1}] is the RWU sink term under uniform soil water potential profile. The RWU under conditions of heterogeneous soil water potential is described with the following equation:

$$S(z) = S_{\text{uniH}}(z) + S_{\text{comp}}(z) = q_{Ti} \cdot SSF(z) + K_{\text{comp}} \cdot \frac{(H_s(z) - H_{\text{eq}}) \cdot SSF(z)}{V(z)} \quad (\text{B4})$$

where K_{comp} [$\text{L}^3 \text{P}^{-1} \text{T}^{-1}$] is the compensatory conductance and S_{comp} [$\text{L}^3 \text{T}^{-1}$] the compensatory RWU accounting for the non-uniform distribution of the soil water potential and $V(z)$ is the volume of soil considered. If the soil water potential is
 25 uniform, this term vanishes from the equation, as $H_S = H_{\text{eq}}$ for any z , and water is extracted from the soil proportionally to RLD . When the water potential at a certain location is smaller (more negative, which means drier) than H_{eq} , less water is extracted from this location. On the other hand, when the soil is wetter (H_S less negative), a larger amount of water can be taken up from the same location as compared. Note that if $H_S < H_{\text{eq}}$ and if the compensatory term is higher than the first one,

S can become positive, and water is released to the soil (i.e., hydraulic redistribution – HR occurs). From Eq. (B4), it can be concluded that HR will preferably occur when q_{Ti} is small and when large soil water potential gradients exist. Plant root hydraulic characteristics will control compensation through the K_{comp} term. The importance of the compensatory RWU term has been discussed in the literature for a long time (e.g., Jarvis, 1989). Except if plants activate specific mechanisms to avoid it, compensation always takes place under natural conditions due to the spatially heterogeneous distribution of soil water potential (Javaux et al., 2013).

A simplifying hypothesis that can be made (Couvreur et al., 2014; Couvreur et al., 2012) is to consider that K_{plant} and K_{comp} are equal, which substituted in Eq. (B4) leads to:

$$S(z) = SSF(z) \cdot K_{plant} \cdot (H_s(z) - H_L) / V(z) \quad (B4')$$

10 Finally, the uptake of water stable isotopologues, i.e., the “isotopic sink term” (S_i [$M T^{-1}$]) is defined as:

$$S_i(z) = S(z) \cdot C(z) \quad (B5)$$

where C [$M L^{-3}$] is the water isotopic concentration.

B2: Running the model for the inter-comparison

The root water uptake (S_{AM}) depth profiles and corresponding δ_{Ti_AM} were simulated using the model of Couvreur et al. (2012) (Eq. (B4')) for all eight scenarios. For this, H_s , δ_s , and RLD input data were interpolated at a 0.01 m vertical resolution and the resistance of the xylem vessels was assumed to be negligible so that $H_{Ti} = H_L$. A K_{plant} value of $2.47 \cdot 10^{-6} h^{-1}$ was taken and was determined based on concomitant T , H_{eq} and H_L data measured for *Festuca arundinacea*. δ_{Ti_AM} was then calculated from Eq. (4b) (section §2.3). From these simulations, the depth profiles of x_{AM} (%), the ratio $\frac{S_{AM}(z)dz}{T/(\Delta x \cdot \Delta y)}$ at each

interpolated depth z was determined, and x_{J_AM} , the ratio $\frac{S_{AM}(z)dz}{T/(\Delta x \cdot \Delta y)}$ from each of the integrated soil layers J ($J \leq III$ or $J \leq$

20 VIII) were calculated. In order to account for uncertainty of the input data (i.e., total soil water potential and oxygen isotopic composition H_s and δ_s , and root length density RLD), the model was run a 1000 times where a single offset randomly selected between -5 and $+5$ cm, -0.2 and $+0.2$ ‰, and -0.1 and $+0.1$ $cm\ cm^{-3}$ was added to the initial values (reported Table 2) of H_s , δ_s , and RLD , respectively. By doing this we obtained *a posteriori* distributions of S_{AM} and corresponding δ_{Ti_AM} standard deviations ($\sigma_{\delta_{Ti_AM}}$);

25

Appendix C: Inter-comparison methodology

The graphical inference method (GI), the two end-member mixing model (TM), and the statistical methods of Phillips and Gregg (2003) (MSPG) and Parnell et al. (2010) (MSPa) were compared to each-other in the following manner for each of the eight virtual experiments:

- 5 (i) Single (or multiple) mean RWU depth(s) (\bar{z}) were graphically identified following the GI method as the depth(s) of the intersection between a vertical line of value δ_{Ti_AM} with the δ_s profile. The uncertainty of method GI was determined on the basis of the δ_{Ti_AM} *a posteriori* distribution: by taking into account $\sigma_{\delta_{Ti_AM}}$, \bar{z} results were translated into “RWU layers”;
- 10 (ii) relative contribution of RWU to transpiration (x_{TM} , %) to two defined soil layers (either conjoint: 0 – 0.225 m and 0.225 – 2.00 m or disjoint: 0 – 0.225 m and 1.75 – 2.00 m) were determined using the TM approach. For this, representative values for the water oxygen isotopic compositions of these soil layers were computed using Eq. (5) which uses soil volumetric water content (θ , in $m^3 m^{-3}$) as input data. θ distribution was obtained from H_s distribution and the van Genuchten (1980) closed-form equation. Values for its different parameters, i.e., the soil residual and saturated water contents (θ_{res} and θ_{sat}), and the shape parameters related to air entry potential and pore-size distribution
- 15 (α and n) were equal to 0.040 and $0.372 m^3 m^{-3}$, $0.003 cm^{-1}$, and 3.3, respectively;
- (iii) Possible range of x_{J_MSPG} , the relative contribution of RWU to transpiration for each of the integrated soil layers following the MSPG method was computed based on smoothed $\delta_{s,J}$ profile and δ_{Ti_AM} by solving the following equation:
- $$\sum_J x_{J_MSPG} \cdot \delta_{s,J} \leq |\delta_{Ti_AM} \pm t| \quad (C1)$$
- 20 with $\tau = \sigma_{\delta_{Ti_AM}}$.
- $\delta_{s,J}$ was computed similarly to for the TM method;
- (iv) Density distribution of x_{J_MSPa} , the relative contribution of RWU to transpiration for each of the three (or eight) soil layers following the MSPG method was determined based on smoothed $\delta_{s,J}$ profile and δ_{Ti_AM} data as well. To compare with the MSPG method (i) the number of δ_{Ti} replicates was fixed to three and equal to $\delta_{Ti_AM} - \sigma_{\delta_{Ti_AM}}$, δ_{Ti_AM} , and
- 25 $\delta_{Ti_AM} + \sigma_{\delta_{RWU_AM}}$, and (ii) x_{J_MSPa} was computed at a 10 % increment (*i*);
- (v) Results obtained at steps (i)-(iv) were compared to each other;
- (vi) Sensitivity of the MSPG method to the values of *i* and τ , and of the MSPa method to number of δ_{Ti} replicates, and to values of arguments *iterations* and *burnin* were finally briefly tested.