Lawrence Berkeley National Laboratory

LBL Publications

Title

What Ecohydrologic Separation Is and Where We Can Go With It

Permalink https://escholarship.org/uc/item/999663x6

Journal Water Resources Research, 56(7)

ISSN 0043-1397

Authors Sprenger, Matthias Allen, Scott T

Publication Date 2020-07-01

DOI 10.1029/2020wr027238

Peer reviewed

eScholarship.org



Water Resources Research

COMMENTARY

10.1029/2020WR027238

Key Points:

- Isotope ratios of plant water should differ from water flowing in soils to streams, and so we need to move beyond confirming this difference
- To move beyond identifying ecohydrologic separation toward understanding it, we provide a framework for assessing soil-water flow processes
- By focusing on dynamics of how water infiltrates into the subsurface and becomes available to plants, we can better interpret past findings

Correspondence to:

M. Sprenger, mspreng@ncsu.edu

Citation:

Sprenger, M., & Allen, S. T. (2020). What ecohydrologic separation is and where we can go with it. *Water Resources Research*, 56, e2020WR027238. https://doi.org/ 10.1029/2020WR027238

Received 29 JAN 2020 Accepted 17 JUN 2020 Accepted article online 20 JUN 2020

©2020. American Geophysical Union. All Rights Reserved.

What Ecohydrologic Separation Is and Where We Can Go With It

Matthias Sprenger^{1,2} 🝺 and Scott T. Allen^{3,4} 🝺

¹Institute of Environmental Assessment and Water Research (IDAEA-CSIC), Barcelona, Spain, ²Department of Forestry and Environmental Resources, North Carolina State University, Raleigh, NC, USA, ³Department of Geology and Geophysics, University of Utah, Salt Lake City, UT, USA, ⁴Department of Environmental Systems Science, ETH Zürich, Zürich, Switzerland

Abstract The "ecohydrologic separation" hypothesis challenged assumptions of translatory flow through the rooting zone. However, studies aiming to test ecohydrologic separation have largely done so with insufficient discussion of infiltration and rooting zone recharge processes and instead have mostly focused on either isotopic differences between stream water and plant water or the presence of fractionated isotope ratios in plant water. Based on extensively observed heterogeneities in soils and watersheds, we posit that differences in isotopic compositions of water in plants, streams, and other subsurface pools are expected in most scenarios. Interpretation of those plant-and-stream water isotopic differences is important, but diagnosing the role of any specific process is typically confounded by the diversity of potential mechanisms contributing to those isotopic differences. Thus, we should progress from simply describing their occurrences and refocus the discussion of ecohydrologic separation on how heterogeneous infiltration and root uptake processes lead to such differences. Consequently, we outline areas where plant and soil-water stable isotope data may be useful for advancing our understanding and representation of soil-water transport and plant-water recharge.

1. From Ecohydrologic Separation to Two Water Worlds and Back Again

Ecohydrologic separation is a consequence of preferential flows through soils that result in soil-water heterogeneity. In 1864, Schumacher noted that "the permeability of a soil during infiltration is mainly controlled by big pores, in which the water is not held under the influence of capillary forces" (Schumacher, 1864 quoted in Beven & Germann, 1982). Over the century that followed, countless studies have described how water flows heterogeneously through soils: transport through soils is dominated by flow through macropores such that it may interact little with the water stored in smaller pores. Notable progress was made when Brooks et al. (2010) used stable isotopes of water (²H and ¹⁸O) to demonstrate a severe dichotomy between fast moving water and the slow moving storage that was presumably composed of smaller pore spaces (that consequently hold the water that is taken up by plants in the dry season). Roots took up water that fell as precipitation more than 6 months before the growing season, and there was little evidence that the tree-available water was displaced or diluted by the subsequent meters of precipitation that must have passed through those trees' root zones. Such temporal separation between the time of water input and its use by plants would not occur if water flow through the rooting zone is translatory (Figure 1a) or well mixed (Figure 1b). Instead, these data imply that preferential flows dominate: after plant-available pore spaces are refilled at the beginning of a wet season, precipitation mostly bypasses these storages (Figure 1c). In this commentary, we argue that evaluating the occurrence of this temporal ecohydrologic separation, as described by Brooks et al., is important because such patterns indicate which flow processes dominate infiltration (sensu Figure 1) and influence the interplay between root distribution and plant-water availability. Using stable isotope data to classify the relative expression of translatory, well-mixed, and preferential flow would constitute progress toward conceptualizing relationships between soil properties and soil-water transport; it would also support building upon simply considering plant uptake from shallow versus deep zones, toward also considering uptake across pores with faster and slower flow.

Focusing on ecohydrologic separation, in the context of soil-water flow mechanisms, contrasts with the objectives of many recent ecohydrology studies that use soil-, plant-, and stream-water isotopic differences in dual-isotope space (δ^2 H vs. δ^{18} O) to confirm the *two water worlds* hypothesis. While often used



Soil-water isotope variations under different flow conditions

Figure 1. (a–c) Stable isotope ratios of soil water, as a function of different idealized transport conditions manifest where isotopic variations reflect water-age variations (left column) or evaporation effects on lc-excess as defined by Landwehr and Coplen (2006) (right column). Color codes represent the age of water (left; from white to black representing young to old), assuming that the age series reflects distinct isotopic variations, and values of lc-excess from undergoing evaporative fractionation (right; from white to red representing nonfractionated to kinetically fractionated). (d) Models often assume laterally homogeneous conditions, yielding assumed profiles that match the first two rows on the figure; however, observations demonstrate that laterally heterogeneous processes affect the isotope depth profiles and thus also which water is used by plants. The 51 modeling studies published since 2010 used for this diagram are listed in the supplementary material. Line-conditioned excess is used instead of deuterium excess because it is defined relative to local meteoric water lines, lc-excess = $\delta^2 H - a * \delta^{18}O - b$, with *a* and *b* being the slope and intercept of the local meteoric water line (Landwehr & Coplen, 2006).

interchangeably, we distinguish *ecohydrologic separation* (defined above) from the *two water worlds* which McDonnell (2014) introduced as a hypothesis of "vegetation and streams returning different pools of water to the hydrosphere." As stated by Pfister & Kirchner (2017), the two water worlds hypothesis is a test of a "homogeneity assumption" that would require the entire pool of waters contributing to streamflow, root water uptake, and evaporation being well mixed. Indeed, decades of theory and all recent isotope studies suggest that stream water and plant water should differ in δ^2 H, δ^{18} O, or lc-excess (Figure 1d); accordingly, we consider that assumption of homogeneity to be reasonably invalidated (regardless of whether or not corresponding representations are still useful). However, diverse processes cause stream water to differ isotopically from xylem water, and consequently, inferences drawn from stream-xylem differences are generally confounded.

We propose revisiting the original conceptualization of ecohydrologic separation (Brooks et al., 2010). This requires diagnosing the expression of different potential transport pathways, especially preferential flow,



through examining how soil-water isotope compositions respond to new inputs (as abstractly represented in Figures 1a– 1c). If we accept flow (and thus isotopic) heterogeneity as a given, we can move toward more precise interpretations of stable isotope data (and thus more precise understanding of how water infiltrates into soils and recharges plant-available water). Toward that objective, we offer four observation-driven statements that we believe should be considered when investigating ecohydrologic separation.

2. Soil Waters Seem to Almost Never Be Isotopically Homogeneous, But It Would Be Useful to Identify Under What Conditions They Are

In few scenarios do soil and plant waters have identical isotope ratios, let alone plant and stream waters. We have reviewed and synthesized dozens of soil- and plant-water isotope studies in previous works (Sprenger et al., 2016; Sprenger, Stumpp et al., 2019); none revealed isotopic uniformity across the critical zone, and no study found complete mixing of new precipitation with previously stored water, to contribute to both streams and transpiration. Even in the absence of the evaporation effects on isotope variations, stream water isotopes ratios should not equal those of soil water or plant water because water flows heterogeneously into and through the critical zone. Even in the improbable hypothetical scenario where isotopically varying precipitation percolates uniformly (i.e., translatory flow), we would still expect stream-isotope variations to differ from those of plants because most flow paths to roots are shorter than to the streams, resulting in higher variation in stem water than in stream water.

Thus, we argue that isotopic homogenization is a very interesting case, because it indicates efficient mixing and mobilization of subsurface water by newly infiltrating water. Sprenger, Llorens et al. (2019) showed that even rising water tables briefly saturate soils but do not flush older waters and isotopically homogenize the soil-water pools. We can hypothesize that there are some conditions where stream, soil, and plant isotope ratios would be similar, for example, with plant uptake from saturated zones with near-complete mixing of waters supplying streams and plants in riparian zones, if soils are coarse enough that near-stream groundwaters are indistinguishable from stream waters. Another cause could be a lack of isotopic variation in the input; however, precipitation isotope ratios vary systematically in most places on earth (Allen, Jasechko et al., 2019). Regardless, for quantifying and conceptualizing the transport of carbon, nitrogen, or other solutes or pollutants, it is imperative to identify instances when inputs (either vertical or lateral) do seem to completely replace, displace, or mix with the previously stored soil waters. Thus, we should not consider a lack of ecohydrologic separation to be an unimportant null finding (unless it is due to a lack of input signal variation and lack of ability to detect separation) but actually one that contrasts with expectations.

3. Trees Do Not Prefer Water Under High Tensions, But They Might Root in Less Conductive Pores

The finding that plant water is isotopically similar to bulk soil water (from cryogenic vacuum distillation) but not to suction-lysimeter water often spurs comments on how mysterious (or impossible) it is that plants "preferentially" take up so-called bound waters that are under high tensions (Berry et al., 2017). We believe that confusion underlies this discussion. Indeed, cryogenic vacuum distillation of soil water can extract waters under higher tensions than can suction lysimeters, but this does not mean that bulk soil waters are always reflecting a higher tension pool. A fully saturated soil sample will experience high water tensions during vacuum extraction, and only small amounts of water remain in the soil; however, this does not mean that it was under high tension in the field when plants were potentially accessing it. Perhaps more significant is that suction lysimeters pull water; larger pores are more conductive and thus will transport water to lysimeters more quickly (Weihermüller et al., 2005). Thus, we posit that the difference between lysimeter and bulk soil-water isotope ratios may often reflect the pore sizes sampled more so than their relative tensions.

We can see that tension does not always explain isotopic variations because they occur when tension is less relevant (e.g., in saturated conditions): in addition to the aforementioned Sprenger, Llorens et al. (2019) study that showed lysimeter water and bulk soil water isotopically differed even after saturation, soils in the Brooks et al. (2010) study were extremely wet throughout the winter when bypass flow occurred. For another example, Morales (2019) submerged intact soil monoliths into buckets of water and found that the water in these clayey soils did not homogenize even after 1 month. These examples highlight that mixing and equilibration across pores can take a very long time even without the influence of tension (due to

saturation). Thus, isotopic heterogeneities in soils are not only necessarily a product of tension variations but also a product of contrasts in how water flows through different pores.

Accordingly, when plant water matches bulk soil water but not suction-lysimeter water, this does not imply a preference for higher tension water. Instead, such differences imply that roots are located in less conductive pores where water tends to travel more slowly and can reside for longer times, potentially across seasons. These variations in conductivity are not only reflections of roots occupying smaller pores, which occurs partially because roots grow toward water due to the so-called hydropatterning mechanism (Bao et al., 2014), but also that plants reduce conductivity around roots through compaction and mucilage production (York et al., 2016). It has been long observed that soil surrounding roots has lower porosity (Dexter, 1987; Young, 1998). Furthermore, given that fungal hyphae can have much smaller diameters than roots (Smith et al., 2010), hypothetically, mycorrhiza associations may further facilitate access to slower moving pools. Importantly, these pools are not dead storage as these waters can move at various velocities, inconsistent with the common problematic description of these subsurface waters as "immobile" or "bound." We therefore advocate for referring to this as bulk soil water, which can include water in finer pores and is less likely to include water in the coarsest pores (that already drained).

We note that hydrologists often ignore the role of nutrients in determining root distributions (see recent exception by Muñoz-Villers et al., 2020), despite extensive research showing root uptake patterns track nutrients (e.g., Caldwell et al., 1991; de Kroon & Mommer, 2006; Kulmatiski et al., 2017), even at short timescales (Jackson et al., 1990). Given that pore-scale variations in water retention affect nitrogen dynamics as well (Hall et al., 2016), questions on how roots differentially access pores of different filling/draining dynamics may also be central to investigating how factors other than water availability further control rooting habits.

4. Concepts of Plant-Water Availability Should Account for Lateral Heterogeneities

For decades, root-water uptake studies have primarily focused on identifying the depth of water uptake, by matching plant-water isotope ratios to isotope depth profiles. However, such practices are challenged by the findings of Brooks et al. (2010) that demonstrate the additional need to also focus on the difference between faster and slower flowing waters in soils (e.g., Figures 1c). Isotope variations reflect only depth (and thus support inferring depth of root-water uptake) in conditions controlled by evaporation or by translatory flow (Figure 1a). In reality, soil structure is three-dimensional, and not only do vertical flow rates vary laterally (as conceptualized in Figure 1c), but water also flows laterally at different rates. Both may yield lateral isotopic variations that can be as large as the vertical variations (Goldsmith et al., 2019; Penna & van Meerveld, 2019; Yang et al., 2016), which further complicate identifying depths of root-water uptake. Theoretically, where preferential flow and heterogeneous transport dominate, depth of uptake may not be inferable from isotope variations unless a strong evaporation signal is present (Figure 1c). Instead, if preferential flows dominate, stable isotopes are better suited for identification of different pore sizes containing water of different ages, because these pores will be filled and drained at different time scales (e.g., Gerke & van Genuchten, 1993). For example, Allen, Kirchner, et al., 2019 showed that broadleaf trees used isotopically different water than collocated conifers despite having similarly shallow root distributions, suggesting uptake from different pore domains. Hypothetically, this may result from the broadleaf trees-which isotope ratios matches winter precipitation-having smaller diameter fine roots and thus occupying smaller pores than conifers, as has been described of mixed broadleaf-conifer systems elsewhere (Bauhus & Messier, 1999; Comas & Eissenstat, 2004). Such findings challenge common assumptions that isotopic differences among tree xylem waters exclusively reflect rooting depth differences (vertical variability), rather than access to different soil pores that contain waters percolating at different velocities (horizontal variability).

Considering lateral variations in plant-water sources, rather than solely focusing on vertical variations (root depth) as in most concepts and model representations, supports testing of new hypotheses. For example, we posit that tree roots often do not use the most recently infiltrated water (e.g., Allen, Kirchner et al., 2019; Brooks et al., 2010; Evaristo et al., 2019; Hervé-Fernández et al., 2016; Knighton et al., 2019; McCutcheon et al., 2017) because they root in fine pores (with slower moving waters) where water is more reliably held, consistent with the needs of a long-lived organism to endure varying conditions (including droughts). Alternatively, we hypothesize that plants with more ephemeral life histories (e.g., opportunistic or weedy

species) dramatically reduce transpiration when water is unavailable and may use more quickly moving waters when they are transiently availably. This dichotomy has been evaluated in hot desert summers, where cacti and herbaceous plants tend to use recent precipitation whereas co-occurring woody plants rely more on precipitation from past seasons that is stored in deep soil layers (Ehleringer et al., 1991). Isotope measurements have previously been used to test ecological niche partitioning theories with respect to depth (see review by Silvertown et al., 2015), but the evidence of plants selecting older water prompts questions on how niches separate by pore size. We are not the first to ask how roots occupy different sized pores (e.g., Stewart et al., 1999). Nonetheless, using stable isotope data to ask how plants associate with different pore or soil structures could lead to better understanding which and how much infiltrating water becomes available to plants, ultimately allowing for framing niche partitioning findings in an ecohydrologic separation context.

5. Preferential Flows Should Be Represented to Accurately Model Soil-Water Transport and Plant Uptake

Although soil-water isotope data are laterally heterogeneous (Figure 1d), indicating preferential flows (Figure 1c), most hydrological modeling studies use soil storage conceptualizations that neglect the processes that can create this heterogeneity; this affects our ability to model how infiltrating water reaches roots and enters the transpiration stream (Figure 1d). Recently, many studies have shown improved model representation of soil-water stable isotope dynamics through parameterizing preferential flow fractions (e.g., Knighton et al., 2019; Sprenger et al., 2018; Stumpp & Maloszewski, 2010), regardless of how those flow paths are oriented within soils' three-dimensional structure. At catchment scales, modeling studies show the need to represent heterogeneous incomplete mixing to match stream-water isotope observations (Fenicia et al., 2010; Knighton et al., 2017; McMillan et al., 2012; van der Velde et al., 2015). However, such practices remain an exception in tracer-aided models representing soil-water transport: about two thirds of their applications over the last 10 years assumed that soil evaporation and root water uptake are supplied by a well-mixed soil-water pool (Figure 1d). Thus, if isotope-enabled models do not consider subsurface heterogeneous flow, they may be misrepresenting soil-water transport processes given that they are calibrated to tracer data that mostly show evidence of heterogeneous, preferential flow through soils (see, e.g., model comparisons in Mueller et al., 2014; Sprenger et al., 2018; Stumpp & Maloszewski, 2010). While various decades-old soil physical models account for heterogeneous solute/tracer transport (e.g., Gerke & van Genuchten, 1993; Hutson & Wagenet, 1995), dual-porosity or multiregion simulation of subsurface isotope tracer dynamics remains under used (considering the current state of observations). Although the role of vapor-phase exchange across the pore space in dual-porosity modeling approaches may diminish the heterogeneities (Sprenger et al., 2018), we admittedly only have limited evidence to guide representation of those processes.

A consequence of neglecting preferential flows includes underestimating the tail of soil-water residence time distributions and overestimating the amount of young water in soils. If all else are equal, greater contrasts between the ages of waters stored in soils versus those percolating through the profile occur when water flows more preferentially (Berghuijs & Allen, 2019). If preferential flows are not considered in simulations of soil-water systems, an inaccurately large fraction of precipitation contributes toward the displacement of previously stored soil water. Thus, if a model that does not represent preferential flow is fitted to data reflecting preferential flow, the turnover time of water in the matric would be underestimated, and the rate of flushing of stored solutes would be overestimated.

Another consequence of neglecting preferential flows is poorly estimated evaporation- or transpiration-flux isotope ratios, which may diminish the value of constraining earth system models with isotope values. The parameterization of soil and canopy evaporation and transpiration in tracer-aided hydrology models is mostly through optimizing the tracer signal in catchment runoff and not explicitly calibrated to the soil-transport processes that are explicitly represented. Improving our ability to simulate soil and plant isotope variations is key to interpreting isotope ratios in the atmosphere (Aemisegger et al., 2014; Dee et al., 2015; Good et al., 2015; Risi et al., 2016) or in plant waters (e.g., Knighton et al., 2019; Kuppel et al., 2018) to estimate the relative importance of evaporation and transpiration fluxes. Moving toward these



larger scale applications in a nonspeculative way requires an accurate representation of processes at the soil-profile scale.

6. Conclusion

Many recent studies demonstrate isotopic differences between soil or saprolite, plant, and stream waters, but it is often unclear how these observations differ from expectations. Underlying these differences is more than the unsurprising observation that plants use evaporatively fractionated soil water. While those observations are often invoked as evidence of ecohydrologic separation, ecohydrologic separation should be seen as a manifestation of diverse flow heterogeneities in the critical zone. We need to move beyond simply identifying "ecohydrologic separation" or "two water worlds" behavior and dig deeper into the questions that originally prompted those hypotheses. Regardless, future work should be grounded in seeking to understand the physical and biological processes that underlie the tracer observations caused by the interplay of roots and the heterogeneous subsurface. Accordingly, we discuss several of such opportunities to advance the topic of ecohydrologic separation in isotope ecohydrology.

References

- Aemisegger, F., Pfahl, S., Sodemann, H., Lehner, I., Seneviratne, S. I., & Wernli, H. (2014). Deuterium excess as a proxy for continental moisture recycling and plant transpiration. *Atmospheric Chemistry and Physics*, 14(8), 4029–4054. https://doi.org/10.5194/acp-14-4029-2014
- Allen, S. T., Jasechko, S., Berghuijs, W. R., Welker, J. M., Goldsmith, G. R., & Kirchner, J. W. (2019). Global sinusoidal seasonality in precipitation isotopes. *Hydrology and Earth System Sciences*, 23(8), 3423–3436. https://doi.org/10.5194/hess-23-3423-2019
- Allen, S. T., Kirchner, J. W., Braun, S., Siegwolf, R. T. W., & Goldsmith, G. R. (2019). Seasonal origins of soil water used by trees. Hydrology and Earth System Sciences, 23(2), 1199–1210. https://doi.org/10.5194/hess-23-1199-2019
- Bao, Y., Aggarwal, P., Robbins, N. E., Sturrock, C. J., Thompson, M. C., Tan, H. Q., et al. (2014). Plant roots use a patterning mechanism to position lateral root branches toward available water. Proceedings of the National Academy of Sciences of the United States of America, 111(25), 9319–9324. https://doi.org/10.1073/pnas.1400966111
- Bauhus, J., & Messier, C. (1999). Soil exploitation strategies of fine roots in different tree species of the southern boreal forest of eastern Canada. Canadian Journal of Forest Research, 29(2), 260–273. https://doi.org/10.1139/x98-206
- Berghuijs, W. R., & Allen, S. T. (2019). Waters flowing out of systems are younger than the waters stored in those same systems. *Hydrological Processes*, 33(25), 3251–3254. https://doi.org/10.1002/hyp.13569
- Berry, Z. C., Evaristo, J., Moore, G., Poca, M., Steppe, K., Verrot, L., et al. (2017). The two water worlds hypothesis: Addressing multiple working hypotheses and proposing a way forward. *Ecohydrology*, 11(3), e1843. https://doi.org/10.1002/eco.1843
- Beven, K., & Germann, P. (1982). Macropores and water flow in soils. Water Resources Research, 18(5), 1311–1325. https://doi.org/10.1029/ WR018i005p01311
- Brooks, J. R., Barnard, H. R., Coulombe, R., & McDonnell, J. J. (2010). Ecohydrologic separation of water between trees and streams in a Mediterranean climate. *Nature Geoscience*, 3(2), 100–104. https://doi.org/10.1038/NGEO722
- Caldwell, M. M., Manwaring, J. H., & Jackson, R. B. (1991). Exploitation of phosphate from fertile soil microsites by three Great Basin perennials when in competition. *Functional Ecology*, 5(6), 757. https://doi.org/10.2307/2389538
- Comas, L. H., & Eissenstat, D. M. (2004). Linking fine root traits to maximum potential growth rate among 11 mature temperate tree species. *Functional Ecology*, 18(3), 388–397. https://doi.org/10.1111/j.0269-8463.2004.00835.x
- de Kroon, H., & Mommer, L. (2006). Root foraging theory put to the test. Trends in Ecology & Evolution, 21(3), 113–116. https://doi.org/ 10.1016/j.tree.2005.11.021
- Dee, S., Noone, D., Buenning, N., Emile-Geay, J., & Zhou, Y. (2015). SPEEDY-IER: A fast atmospheric GCM with water isotope physics. Journal of Geophysical Research: Atmospheres, 120(1), 73–91. https://doi.org/10.1002/2014JD022194
- Dexter, A. (1987). Compression of soil around roots. Plant and Soil, 97(3), 401-406. https://doi.org/10.1007/BF02383230
- Ehleringer, J. R., Phillips, S. L., Schuster, W. S. F., & Sandquist, D. R. (1991). Differential utilization of summer rains by desert plants. Oecologia, 88(3), 430–434. https://doi.org/10.1007/BF00317589
- Evaristo, J., Kim, M., Haren, J., Pangle, L. A., Harman, C. J., Troch, P. A., & McDonnell, J. J. (2019). Characterizing the fluxes and age distribution of soil water, plant water, and deep percolation in a model tropical ecosystem. Water Resources Research, 55(4), 3307–3327. https://doi.org/10.1029/2018WR023265
- Fenicia, F., Wrede, S., Kavetski, D., Pfister, L., Hoffmann, L., Savenije, H. H. G., & McDonnell, J. J. (2010). Assessing the impact of mixing assumptions on the estimation of streamwater mean residence time. *Hydrological Processes*, 24(12), 1730–1741. https://doi.org/10.1002/ Hyp.7595
- Gerke, H. H., & van Genuchten, M. T. (1993). A dual-porosity model for simulating the preferential movement of water and solutes in structured porous media. *Water Resources Research*, 29(2), 305–319. https://doi.org/10.1029/92WR02339
- Goldsmith, G. R., Allen, S. T., Braun, S., Engbersen, N., Romero González-Quijano, C., Kirchner, J. W., & Siegwolf, R. T. W. (2019). Spatial variation in throughfall, soil, and plant water isotopes in a temperate forest. *Ecohydrology*, 12(2), e2059. https://doi.org/ 10.1002/eco.2059
- Good, S. P., Noone, D., & Bowen, G. (2015). Hydrologic connectivity constrains partitioning of global terrestrial water fluxes. Science, 349(6244), 175–177. https://doi.org/10.1126/science.aaa5931
- Hall, S. J., Weintraub, S. R., & Bowling, D. R. (2016). Scale-dependent linkages between nitrate isotopes and denitrification in surface soils: Implications for isotope measurements and models. *Oecologia*, 181(4), 1221–1231. https://doi.org/10.1007/s00442-016-3626-1
- Hervé-Fernández, P., Oyarzún, C., Brumbt, C., Huygens, D., Bodé, S., Verhoest, N. E. C., & Boeckx, P. (2016). Assessing the "two water worlds" hypothesis and water sources for native and exotic evergreen species in south-central Chile. *Hydrological Processes*, 30(23), 4227–4241. https://doi.org/10.1002/hyp.10984

Acknowledgments

M.S. was supported by the Deutsche Forschungsgemeinschaft (project no. 397306994). This commentary was improved by comments from two anonymous reviewers, Editors Holly Barnard and Ilja van Meerveld, Todd Dawson, and Wouter Berghuijs, and has been influenced by conversations with Renee Brooks and many others.

- Hutson, J. L., & Wagenet, R. J. (1995). A multiregion model describing water flow and solute transport in heterogeneous soils. Soil Science Society of America Journal, 59(3), 743–751. https://doi.org/10.2136/sssaj1995.03615995005900030016x
- Jackson, R. B., Manwaring, J. H., & Caldwell, M. M. (1990). Rapid physiological adjustment of roots to localized soil enrichment. Nature, 344(6261), 58–60. https://doi.org/10.1038/344058a0
- Knighton, J., Saia, S. M., Morris, C. K., Archiblad, J. A., & Walter, M. T. (2017). Ecohydrologic considerations for modeling of stable water isotopes in a small intermittent watershed. *Hydrological Processes*, 31(13), 2438–2452. https://doi.org/10.1002/hyp.11194
- Knighton, J., Souter-Kline, V., Volkmann, T. H. M., Troch, P. A., Kim, M., Harman, C., et al. (2019). Seasonal and topographic variations in ecohydrological separation within a small, temperate, snow-influenced catchment. *Water Resources Research*, 55(8), 6417–6435. https:// doi.org/10.1029/2019WR025174
- Kulmatiski, A., Adler, P. B., Stark, J. M., & Tredennick, A. T. (2017). Water and nitrogen uptake are better associated with resource availability than root biomass. *Ecosphere*, 8(3), e01738. https://doi.org/10.1002/ecs2.1738
- Kuppel, S., Tetzlaff, D., Maneta, M. P., & Soulsby, C. (2018). EcH2O-iso 1.0: Water isotopes and age tracking in a process-based, distributed ecohydrological model. *Geoscientific Model Development*, *11*(7), 3045–3069. https://doi.org/10.5194/gmd-11-3045-2018
- Landwehr, J. M., & Coplen, T. B. (2006). Line-conditioned excess: A new method for characterizing stable hydrogen and oxygen isotope ratios in hydrologic systems (Eds.), International Conference on Isotopes in Environmental Studies (pp. 132–135). Vienna: IAEA.
- McCutcheon, R. J., McNamara, J. P., Kohn, M. J., & Evans, S. L. (2017). An evaluation of the ecohydrological separation hypothesis in a semiarid catchment. *Hydrological Processes*, 31(4), 783–799. https://doi.org/10.1002/hyp.11052
- McDonnell, J. J. (2014). The two water worlds hypothesis: Ecohydrological separation of water between streams and trees? Wiley Interdisciplinary Reviews Water, 1(4), 323–329. https://doi.org/10.1002/wat2.1027
- McMillan, H., Tetzlaff, D., Clark, M., & Soulsby, C. (2012). Do time-variable tracers aid the evaluation of hydrological model structure? A multimodel approach. Water Resources Research, 48, W05501. https://doi.org/10.1029/2011WR011688
- Morales, S. R. (2019). Matrix recharge in a shrink-swell floodplain forest soil. Master of Science: Agricultural and Mechanical College, Louisiana State University.
- Mueller, M. H., Alaoui, A., Kuells, C., Leistert, H., Meusburger, K., Stumpp, C., et al. (2014). Tracking water pathways in steep hillslopes by δ¹⁸O depth profiles of soil water. *Journal of Hydrology*, *519*(A), 340–352. https://doi.org/10.1016/j.jhydrol.2014.07.031
- Muñoz-Villers, L. E., Geris, J., Alvarado-Barrientos, M. S., Holwerda, F., & Dawson, T. (2020). Coffee and shade trees show complementary use of soil water in a traditional agroforestry ecosystem. *Hydrology and Earth System Sciences*, 24(4), 1649–1668. https://doi.org/10.5194/ hess-24-1649-2020
- Penna, D., & van Meerveld, H. J. (2019). Spatial variability in the isotopic composition of water in small catchments and its effect on hydrograph separation. *Wiley Interdisciplinary Reviews Water*, 6(5), e1367. https://doi.org/10.1002/wat2.1367

Pfister, L., & Kirchner, J. W. (2017). Debates—Hypothesis testing in hydrology: Theory and practice. Water Resources Research, 53(1792), 1792–1798. https://doi.org/10.1002/2016WR020116

- Risi, C., Ogee, J., Bony, S., Bariac, T., Raz Yaseef, N., Wingate, L., et al. (2016). The water isotopic version of the land-surface model ORCHIDEE: Implementation, evaluation, sensitivity to hydrological parameters. *Hydrology: Current Research*, 07(04), 258. https://doi. org/10.4172/2157-7587.1000258
- Schumacher, W. (1864). Die Physik des Bodens. Berlin: Wiegandt & Hempel.
- Silvertown, J., Araya, Y., & Gowing, D. (2015). Hydrological niches in terrestrial plant communities: A review. *Journal of Ecology*, 103(1), 93–108. https://doi.org/10.1111/1365-2745.12332
- Smith, S., Facelli, E., Pope, S., & Smith, F. A. (2010). Plant performance in stressful environments: Interpreting new and established knowledge of the roles of arbuscular mycorrhizas. *Plant and Soil*, 326(1-2), 3–20. https://doi.org/10.1007/s11104-009-9981-5
- Sprenger, M., Leistert, H., Gimbel, K., & Weiler, M. (2016). Illuminating hydrological processes at the soil-vegetation-atmosphere interface with water stable isotopes. *Reviews of Geophysics*, 54, 674–704. https://doi.org/10.1002/2015RG000515
- Sprenger, M., Llorens, P., Cayuela, C., Gallart, F., & Latron, J. (2019). Mechanisms of consistently disjunct soil water pools over (pore) space and time. *Hydrology and Earth System Sciences*, 23(6), 2751–2762. https://doi.org/10.5194/hess-23-2751-2019
- Sprenger, M., Stumpp, C., Weiler, M., Aeschbach, W., Allen, S. T., Benettin, P., et al. (2019). The demographics of water: A review of water ages in the critical zone. *Reviews of Geophysics*, 57(3), 800–834. https://doi.org/10.1029/2018RG000633
- Sprenger, M., Tetzlaff, D., Buttle, J. M., Laudon, H., Leistert, H., Mitchell, C. P. J., et al. (2018). Measuring and modelling stable isotopes of mobile and bulk soil water. Vadose Zone Journal, 17(1), 170,149. https://doi.org/10.2136/VZJ2017.08.0149
- Stewart, J. B., Moran, C. J., & Wood, J. T. (1999). Macropore sheath: Quantification of plant root and soil macropore association. Plant and Soil, 211(1), 59–67. https://doi.org/10.1023/A:1004405422847
- Stumpp, C., & Maloszewski, P. (2010). Quantification of preferential flow and flow heterogeneities in an unsaturated soil planted with different crops using the environmental isotope 8¹⁸O. *Journal of Hydrology*, 394(3–4), 407–415. https://doi.org/10.1016/j. jhydrol.2010.09.014
- van der Velde, Y., Heidbüchel, I., Lyon, S. W., Nyberg, L., Rodhe, A., Bishop, K., & Troch, P. A. (2015). Consequences of mixing assumptions for time-variable travel time distributions. *Hydrological Processes*, *29*(16), 3460–3474. https://doi.org/10.1002/hyp.10372
- Weihermüller, L., Kasteel, R., Vanderborght, J., Pütz, T., & Vereecken, H. (2005). Soil water extraction with a suction cup. Vadose Zone Journal, 4(4), 899–907. https://doi.org/10.2136/vzj2004.0156
- Yang, J., Chen, H., Nie, Y., Zhang, W., & Wang, K. (2016). Spatial variability of shallow soil moisture and its stable isotope values on a karst hillslope. *Geoderma*, 264, 61–70. https://doi.org/10.1016/j.geoderma.2015.10.003
- York, L. M., Carminati, A., Mooney, S. J., Ritz, K., & Bennett, M. J. (2016). The holistic rhizosphere: Integrating zones, processes, and semantics in the soil influenced by roots. *Journal of Experimental Botany*, 67(12), 3629–3643. https://doi.org/10.1093/jxb/erw108
- Young, I. M. (1998). Biophysical interactions at the root-soil interface: A review. *The Journal of Agricultural Science*, 130(1), 1–7. https://doi.org/10.1017/S002185969700498X