

Quantifying river water contributions to the transpiration of riparian trees along a losing river: lessons from stable isotopes and an iteration method

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Abstract. River water plays a critical role in riparian plant water use and riparian ecosystem restoration along losing rivers (i.e., river water recharging underlying groundwater). How to quantify the contributions of river water to the transpiration of riparian plants under different groundwater levels and the related responses of plant water use efficiency is a great challenge. In this study, observations of stable isotopes of water (δ^2 H and δ^{18} O), ²²²Rn, and leaf δ^{13} C were conducted for the deep-rooted riparian weeping willow (Salix babylonica L.) in 2019 (dry year) and 2021 (wet year) along the Chaobai River in Beijing, China. We proposed an iteration method in combination with the MixSIAR model to quantify the river water contribution to the transpiration of riparian S. babylonica and its correlations with the water table depth and leaf δ^{13} C. Our results demonstrated that riparian S. babylonica took up deep water (in the 80–170 cm soil layer and groundwater) by 56.5 % \pm 10.8 %. River water recharging riparian deep water was an indirect water source and contributed 20.3 % of water to the transpiration of riparian trees near the losing river. Significantly increasing river water uptake (by 7.0%) and decreasing leaf δ^{13} C (by -2.0%) of riparian trees were observed as the water table depth changed from 2.7 m in the dry year of 2019 to 1.7 m in the wet year of 2021 (p < 0.05). The higher water availability probably promoted stomatal opening and thus increased transpiration water loss, leading to the decreasing leaf δ^{13} C in the wet year compared to the dry year. The river water contribution to the

transpiration of riparian *S. babylonica* was found to be negatively linearly correlated with the water table depth and leaf δ^{13} C (p < 0.01). The rising groundwater level may increase the water extraction from the groundwater and/or river and produce a consumptive river-water-use pattern of riparian trees, which can have an adverse impact on the conservation of both river flow and riparian vegetation. This study provides new insights into understanding the mechanisms of the water cycle in a groundwater–soil–plant–atmosphere continuum and managing water resources and riparian afforestation along losing rivers.

1 Introduction

Ongoing climate warming and groundwater overexploitation have altered river runoff and bank storage globally, which have further resulted in widespread risks such as the flow of river water into underlying groundwater (i.e., "losing" river) and even drying up (Winter et al., 1998; Schindler and Donahue, 2006; Allen et al., 2015; Jasechko et al., 2021). Water replenishment for losing rivers and riparian revegetation has been applied worldwide to restore the river ecosystems (Smith et al., 2018; Long et al., 2020). The water replenishment to losing rivers contributed to bank storage and groundwater storage recovery by 40% (Long et al., 2020). However, large-scale riparian revegetation increased plant transpiration substantially, which in turn led to a great loss of riparian bank storage and even river runoff (Moore and Owens, 2012; Dzikiti et al., 2013; Missik et al., 2019; Mkunyana et al., 2019). Therefore, it is critical to determine what kind of water sources are used by riparian trees, how much river water is taken up by riparian trees, and what are the responses of tree water use characteristics to groundwater level variations (Wang et al., 2022). This can help us to regulate river runoff and vegetation water requirements in the revegetated riparian zones.

The potential water sources of riparian trees along a losing river are generally considered a mix of soil water at different depths, groundwater, and river water (Alstad et al., 1999; White and Smith, 2020). However, there is a debate on whether river water is a potential water source for riparian trees and how it becomes available to plants. Most previous studies considered river water as a direct water source to evaluate the river water contribution (RWC) to the transpiration of riparian trees (Alstad et al., 1999; Zhou et al., 2017; White and Smith, 2020). Based on the stable-isotope signatures of different water sources and plant stem water, these studies found that river water directly contributed up to 80 % to riparian plant transpiration (Dawson and Ehleringer, 1991; Busch et al., 1992; Alstad et al., 1999; Zhou et al., 2017; White and Smith, 2020). Nevertheless, some studies argued that river water was not a potential water source and rarely contributed to the transpiration of riparian trees (Dawson and Ehleringer, 1991; Bowling et al., 2017; P. Y. Wang et al., 2019). Dawson and Ehleringer (1991) first discovered that the mature streamside trees growing in or next to a perennial river did not use river water but depended on water from deeper strata. Similar findings have also been reported regarding riparian phreatophyte trees (Populus fremontii and Salix gooddingii) and riparian deep-rooted trees (Busch et al., 1992; Bowling et al., 2017; P. Y. Wang et al., 2019). Even under shallow groundwater with high salinity, no river water was directly taken up by riparian Eucalyptus coolabah alongside an ephemeral arid zone river in Australia (Costelloe et al., 2008). Growing evidence suggested that riparian trees rarely took up river water directly at a certain distance from the riverbank because their lateral roots could not reach the river (Mensforth et al., 1994; Thorburn and Walker, 1994). Nevertheless, riparian trees can indirectly utilize river water that recharges the deep zone (e.g., deep soil water and groundwater) when their roots tap into the groundwater level (Mensforth et al., 1994; P. Y. Wang et al., 2019). The RWC to the transpiration of riparian trees may be overestimated if the river water is considered a direct water source. How to separate and quantify the contributions of the indirect river water source to the transpiration of riparian trees near losing rivers is a great challenge.

Several approaches have been developed in recent years to determine plant root water uptake patterns. For example, the graphical inference and direct comparison of isotopic values between plant stem water and different water sources (Dawson and Ehleringer, 1991; Busch et al., 1992; Costelloe et al., 2008; Zhao et al., 2016), statistical two- or multi-source linear mixing models (Alstad et al., 1999; Zhou et al., 2017), and the MixSIAR Bayesian mixing model (P. Y. Wang et al., 2019; Wang et al., 2020; White and Smith, 2020; Li et al., 2021) that are integrated with stable isotopes of water $(\delta^2 H \text{ and } \delta^{18} O)$ have been extensively employed to identify the potential water sources taken up by riparian trees. The MixSIAR model has more advantages in quantifying water source contributions and accounting for uncertainties in the isotopic values (Stock and Semmens, 2013; Ma and Song, 2016). The indirect RWC to the transpiration of riparian trees can be estimated by quantifying both the direct water source contributions to the transpiration of riparian trees and the RWC to riparian deep water. A multi-iteration method (Marek et al., 1990; Zaid, 2010) is key to calculating the proportional contributions of total (old and current) river water to riparian deep water, which enhances the estimation accuracy of the indirect RWC to the transpiration of riparian trees. Radioactive radon (222Rn) has been broadly utilized for tracing groundwater origins and corresponding pathways in riparian zones (Close et al., 2014; Zhao et al., 2018). Based on ²²²Rn concentration, Stellato et al. (2013) estimated the river infiltration velocities into the riparian groundwater system in the Petrignano d'Assisi plain in central Italy, which varied from 1 to $39 \,\mathrm{m \, d^{-1}}$. It is helpful to estimate the residence time of recharged groundwater from river water and its effects on the RWC to the transpiration of riparian trees. A combination of these methods can give a more reliable quantification of the indirect RWC to the transpiration of riparian trees.

As far as we know, the trade-off between the RWC to the transpiration of riparian trees and plant ecophysiological characteristics is unclear, which is critical to understanding the relationships between river runoff and vegetation water requirements in the revegetated riparian zones. The RWC to the transpiration of riparian trees can substantially affect the leaf-level water-use efficiency (WUE) and the growth of riparian trees. Tree WUE is a key characteristic of plant water use, which can be defined as the ratio of photosynthetic rate to transpiration rate. Since leaf δ^{13} C values are positively related to tree WUE, leaf δ^{13} C has been widely employed as an indicator of tree WUE for C3 photosynthesis plants (Farquhar et al., 1989). For instance, based on leaf δ^{13} C measurements, Thorburn and Walker (1994) found that riparian Eucalyptus camaldulensis with more frequent access to river water had a higher tree WUE compared to those far away from the riverbank. Furthermore, the variations of the water table depth (WTD) at different distances from the riverbank in the riparian zones play a critical role in both the RWC to the transpiration of riparian trees and tree WUE (Horton and Clark, 2001; Liu et al., 2017; Xia et al., 2018). Li et al. (2022) elucidated that the water table decline led to an increase in deepwater contribution to riparian Salix babylonica

L. and tree WUE along the distance from the riverbank. Qian et al. (2017) reported a higher RWC to the transpiration of riparian *Ginkgo biloba* L. at the shallower WTD plot closer to the riverbank compared to the other two plots away from the riverbank along a losing river. However, little attention has been paid to quantifying the relationships between the RWC to the transpiration of riparian trees and tree WUE or WTD near a losing river.

The overall goal of this study was to clarify the impacts of river water on the water use of riparian trees along a gradient of WTD. Focusing on a losing river in Beijing, China, the specific objectives of this study were as follows: (1) proposing an iteration method in combination with the MixSIAR model and stable isotopes of water (δ^2 H and δ^{18} O) to quantify the RWC, (2) comparing the contributions of river water to the transpiration of riparian trees along a gradient of WTD at different distances from the riverbank in dry and wet years, (3) identifying the relationships between the RWC to the transpiration of riparian trees and tree WUE (indicated by leaf δ^{13} C values) as well as WTD. Our results will provide critical insights into plantation management, bank storage conservation, and ecosystem health maintenance for losing rivers.

2 Materials and methods

2.1 Study area

The study area was in the reach of the Chaobai River, located in Shunyi District, Beijing, China (40°07'30" N, 116°40'37" E) (Fig. 1). A temperate continental sub-humid monsoon climate prevails in this area, with the annual mean temperature and evaporation of 11.5 °C and 1175 mm, respectively. The average total precipitation from April to November between 1961 and 2021 was 532.8 mm, 84.5 % of which is occurring in the rainy season (from June to September) (Fig. 2a). Owing to continuous drought and groundwater overexploitation, the Chaobai River dried up from 1999 to 2007, and the riparian ecosystem seriously degraded. The "ecological water" (including reclaimed water, reservoir water, and diverted water by the South-North Water Transfer Project) has been supplied through a systematic water release by dams to restore this dry river since 2007. A total of 51.1×10^6 and 380×10^6 m³ of ecological water sources were released to the Chaobai River in 2019 and 2021, respectively. More than 33 km² of the riparian zone was revegetated until 2020. The deep-rooted riparian weeping willow (Salix babylonica L.) was one of the most widely planted species alongside the Chaobai River because the S. babylonica trees could adapt well to dramatic fluctuations in the WTD. Hence, this research selected S. babylonica trees as representative of riparian species. Three plots at different distances of 5 m (D05), 20 m (D20), and 45 m (D45) from the riverbank (one plot per distance) were also selected for field measurements and sample collection (Fig. 1).

2.2 Field measurements and data collection

The field measurements were carried out from April to November in both 2019 and 2021, with no field observations in 2020 due to COVID-19. The daily precipitation data from 1961 to 2021 and the daily mean temperature (T), relative air humidity (RH), solar radiation, and reference evapotranspiration (ET₀) data during the observation period in the Shunyi District were collected from the China Meteorological Data Service Centre (http://data.cma.cn/en, last access: 17 February 2023). Daily mean vapor pressure deficit (VPD) was calculated using the RH and *T* data (Wang et al., 2014; Schoppach et al., 2021).

The groundwater level in each plot was recorded monthly in 2019 and in 2021 via a pressure gauge (HOH-S-Y, King Water Co. Ltd., Beijing, China) installed in the groundwater monitoring well. The river water level was recorded using a water level gauge at the same time as the observed groundwater levels. The riparian ground surface level and riverbed level were 29.5 and 26 m, respectively (Fig. 3).

2.3 Sample collection and isotopic analyses

Twelve sampling campaigns on 5 May, 14 June, 26 July, 15 August, 26 September, and 5 November in 2019 and 24 April, 25 May, 26 June, 15 July, 1 September, and 5 November in 2021 were conducted to collect groundwater, river water, soil, stem, and leaf samples. Groundwater in each plot was sampled by a sucking pump from the monitoring well, and a plexiglass hydrophore water sample collector with a capacity of 1 L was utilized to collect the nearby river water. Precipitation was sampled after each precipitation event via a device consisting of a funnel, a polyethylene bottle, and a ping-pong ball. A total of 135 precipitation samples were collected throughout the whole years of 2019 (53 samples) and 2021 (82 samples). All precipitation, groundwater, and river water samples were stored in a refrigeration box with several ice bags to minimize evaporation in the field; then they were delivered to the laboratory and kept at 4 °C in the refrigerator until analysis of the stable isotopes $(\delta^2 H \text{ and } \delta^{18} O)$. The groundwater and river water were also collected with 100 mL brown glass vials to measure ²²²Rn concentrations in the field.

One riparian *S. babylonica* tree was selected in each plot (three trees in total) for δ^2 H and δ^{18} O measurements in xylem water as well as δ^{13} C analysis in plant leaves. The mean breast-height diameter of the three sampled trees at different distances of 5, 20, and 45 m from the riverbank was 28.6 ± 4.4 cm. Five mature and suberized stem samples were taken from the same riparian *S. babylonica* tree in each plot using an averruncator with a length of 5 m. We removed the bark and phloem of the sampled stems, and then



Figure 1. Schematic diagram of the study area and the three sampling plots (D05, D20, and D45). D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively.



Figure 2. Monthly average precipitation amount from 1961 to 2021 and monthly total precipitation amount for the observation years of 2019 and 2021 (a), daily precipitation amount and precipitation isotopes during 2019 (b), and 2021 (c).

we put the remaining xylem samples into three reduplicative 12 mL brown glass vials sealed with PARAFILM. These three reduplicative xylem samples were extracted, and stable isotopes of water were measured. Meanwhile, more than 50 mature leaves without petioles were sampled from the collected stems using pruning shears and mixed into one leaf sample for δ^{13} C analysis. The xylem and mature leaf samples were stored in a refrigeration box with several ice bags in the field. Then the xylem samples were transported to the laboratory and kept in a refrigerator at -10 °C before water extraction and isotope analysis. The mature leaves were oven-dried at 65 °C for 72 h on the day of sampling; then they were ground and passed through a 0.15 mm sieve to analyze leaf δ^{13} C (J. Wang et al., 2019; Cao et al., 2020).

Soils at depths of 0–5, 5–10, 10–20, 20–30, 40–60, 60– 80, 90–110, 150–170, 190–210, 250–270, and 280–300 cm in one soil profile near the selected *S. babylonica* trees were sampled by a power auger (CHPD78, Christie Engineering Company, Sydney, Australia). One portion of each soil sample was put into a 12 mL brown glass vial and stored at -10 °C before stable isotope analysis, and the other portion was packed into an aluminum box for gravimetric soil water content (SWC) measurement via the oven-drying method (J. Wang et al., 2019; Li et al., 2021).

The automatic cryogenic vacuum distillation system (LI-2100, LICA, Beijing, China) was employed to extract water from xylem and soil samples, which generally ran for at least 2.5 h. All the extracted water from the xylem and soil samples was filtered to remove impurities. We weighed all the xylem and soil samples before and after extraction as well as oven-dried samples. Subsequently, to ensure a water extraction efficiency above 99 % and to avoid isotopic fractionation during water extraction, the efficiency of water extraction was calculated as follows:

$$E_{\rm WE} = \frac{W_{\rm BE} - W_{\rm AE}}{W_{\rm BE} - W_{\rm OD}} \times 100\%,$$
 (1)

where E_{WE} represents the efficiency of water extraction; W_{BE} and W_{AE} represent the weights of xylem and soil samples before and after extraction, respectively; and W_{OD} represents the weights of oven-dried xylem or soil samples.

The δ^2 H and δ^{18} O values of soil water, river water, groundwater, and precipitation were analyzed using an isotopic ratio infrared spectroscopy (IRIS) system (DLT-100, Los Gatos Research, Mountain View, USA) (Li et al., 2021). The isotope ratio mass spectrometry (IRMS) system (MAT253, Thermo Fisher Scientific, Bremen, Germany), which could prevent organic pollution of plants, was used to measure δ^2 H and δ^{18} O values of xylem water as well as the leaf δ^{13} C value. There was the same measurement accuracy for both the IRIS and IRMS systems ($\pm 1 \%$ for δ^2 H and $\pm 0.1 \%$ for δ^{18} O). The Vienna Standard Mean Ocean Water (VSMOW) was utilized to calibrate and normalize the δ^2 H and δ^{18} O measurements in different waters, while Vienna Pee Dee Belemnite (V-PDB) was used for calibrating leaf δ^{13} C values.

The ²²²Rn concentration in the groundwater and river water samples (C_{Water} , BqL⁻¹) was determined based on the air ²²²Rn concentration values (C_{Air} , Bq m⁻³) measured by a ²²²Rn monitor (Alpha GUARD PQ2000 PRO, Bertin Instruments, Germany). A volume of 100 mL of the water sample was slowly poured into the airtight glass bottles and then purged with air in a closed gas-cycling system. The C_{Air} in the ²²²Rn monitor was recorded at 10 min intervals. The air inside the measurement setup maintained a certain ²²²Rn concentration right before the water sample injection (C_{System} , Bq m⁻³). It is generally assumed that when C_{System} is around or lower than 80 Bq m⁻³, the existing C_{System} can be ignored accordingly (Saphymo, 2017). We conducted more than four intervals to ensure that C_{System} was lower than 80 Bq m⁻³. The measurement range of C_{Air} was 2–2 000 000 Bq m⁻³ with a measurement precision of 3 % (Saphymo, 2017). The C_{water} was calculated as follows:

$$C_{\text{Water}} = \frac{C_{\text{Air}} \times \left(\frac{V_{\text{System}} - V_{\text{Sample}}}{V_{\text{Sample}}} + k\right) - C_{\text{System}}}{1000},$$
 (2)

where V_{System} stands for the interior volume of the measuring setup (mL), which is 1122 mL in this study; V_{Sample} symbolizes the volume of the water sample (mL); *k* denotes the ²²²Rn distribution coefficient of water/air (–), which can be set as 0.26 within the specified temperature range around the mean room temperature of 20 °C (Clever, 1985).

We identified the average residence time (T_{res} , d) of recharged groundwater from river water based on the ²²²Rn isotopes (Hoehn and Von Gunten, 1989), which was described as follows:

$$T_{\rm res} = \frac{1}{\lambda} \times \ln\left(\frac{C_{\rm e} - C_{\rm r}}{C_{\rm e} - C_{\rm g}}\right),\tag{3}$$

where λ represents the decay coefficient (0.181 d⁻¹) (Hoehn and Von Gunten, 1989); C_e signifies the ²²²Rn concentration of background groundwater when the equilibrium between radon production and decay is reached (The measuring of ²²²Rn concentration of groundwater in aquifers more than 100 m away from the riverbank remained constant in this study (with an average value of 7400.0 ± 35.4 Bq m⁻³), suggesting that C_e can be defined as 7400.0 Bq m⁻³.); C_r indicates the ²²²Rn concentration of river water (Bq m⁻³); and C_g stands for the ²²²Rn concentration of riparian groundwater (Bq m⁻³).

2.4 Determination of RWC to the transpiration of riparian trees

In this study, stable isotopes of water (δ^2 H and δ^{18} O) were integrated within the MixSIAR model, and an iteration method was proposed to identify the contributions of the indirect river water that recharged riparian deep water to the transpiration of riparian *S. babylonica* trees (Figs. 4–5). First, the

direct water source (including soil water in different layers and groundwater) contributions to the transpiration of riparian trees were determined via δ^2 H and δ^{18} O values of different waters and the MixSIAR model. Second, the proportional contributions of river water to riparian deep water (i.e., riparian groundwater and deep soil water in the 80–170 cm layer) were determined by the MixSIAR model and stable isotopes of water. Finally, the proposed iteration method was applied to quantify the proportions of the indirect river water source taken up by riparian trees (Figs. 4–5).

The MixSIAR model is a Bayesian mixing model which can be integrated with stable isotopes of water to quantify the proportions of source contributions to a mixture (Stock and Semmens, 2013). The input data of the MixSIAR model include mixture data, source data, and discrimination data. In this study, the mean and standard deviation (SD) of the isotopic values of each water source for riparian trees or riparian deep water were inputted as source data into MixSIAR, while the measured isotopic values of xylem water or riparian deep water were input as raw mixture data into MixSIAR. The discrimination data for both δ^2 H and δ^{18} O were set to zero because the input δ^2 H and δ^{18} O values in the MixSIAR were non-fractionated or δ^2 H corrected. The Markov chain Monte Carlo parameter was set to the run length of "very long". The trace plots and three diagnostic tests (i.e., Gelman-Rubin, Heidelberger-Welch, and Geweke) were adopted to determine whether the MixSIAR model converged (Stock and Semmens, 2013). Then, the mean and SD values of different water source contributions could be estimated using the MixSIAR model.

2.4.1 Quantifying proportional contributions of direct water sources to riparian trees

Soil water was an important direct water source for the transpiration of riparian S. babylonica trees. We measured soil water isotopes at 11 depths in each plot at a distance of 5, 20, and 45 m from the riverbank. To reduce errors in the analytical procedure, four soil layers (0-30, 30-80, 80-170, and 170-300 cm) were determined to identify the main root water uptake depth of riparian trees according to seasonal variations in the SWC, water isotopes, and WTD. The average soil water isotope values for the 0-30 cm soil layer were determined as the average of the soil water isotope values of 0-5, 5-10, 10-20, and 20-30 cm soil layers because the water isotopes underwent strong evaporation and SWC changed considerably seasonally. We determined the average soil water isotope values for the 30-80 cm (average of 40-60 and 60-80 cm soil layers) and 80-170 cm (average of 90-110 and 150-170 cm soil layers) soil layers because the water isotopes and SWC were almost stable. The average soil water isotope values for the 170-300 cm soil layer were determined as the average of the soil water isotope values of 190-210, 250-270, and 280-300 cm soil layers, which varied with the fluctuations of groundwater levels. Groundwater could also

be considered a direct water source for phreatophyte riparian trees (Dawson and Ehleringer, 1991; Busch et al., 1992). As the isotopic composition of soil water in the 170-300 cm layer $(-57.6\% \pm 2.0\%)$ for δ^2 H and $-7.3\% \pm 0.1\%$ for δ^{18} O) was similar to that of groundwater (-57.7 % ± 1.4 %) for δ^2 H and $-7.4\% \pm 0.1\%$ for δ^{18} O), they were considered to be one water source (groundwater). Mensforth et al. (1994) and Thorburn and Walker (1994) characterized the projected edge of the canopy as the extension range of lateral roots. In this way, it is possible to determine whether or not riparian trees take up river water directly. The projected edge of the canopy in our study was less than 5 m for the riparian S. babylonica trees which were closest to the river (5 m away from the riverbank). This indicated that the lateral roots of S. babylonica trees could not tap into the river. Therefore, river water was not regarded as a direct potential water source for tree water uptake, while groundwater and soil water in the 0-30, 30-80, and 80-170 cm layers were used as direct potential water sources for riparian S. babylonica.

The δ^2 H offsets between the xylem water in riparian trees and its corresponding potential source waters were observed in this study, which possibly resulted from $\delta^2 H$ fractionation in the plant water use processes (Li et al., 2021; Cernusak et al., 2022). These δ^2 H offsets could lead to large errors in estimating the water source contributions using the MixSIAR model. To eliminate the δ^2 H offsets of xylem water from its potential water sources, the measured xylem water δ^2 H values were corrected by the potential water source line (PWL) proposed by Li et al. (2021). The PW-excess (PW-excess = $\delta^2 H - a_p \delta^{18} O - b_p$, where a_p and b_p are slope and intercept of the PWL, respectively) was calculated to determine the δ^2 H deviation from the PWL, which was subsequently subtracted from the measured xylem water $\delta^2 H$ values. To quantify the contributions of direct water sources to the transpiration of riparian S. babylonica, the corrected $\delta^2 H$ and raw δ^{18} O in xylem water were set as the mixture data in the MixSIAR model.

2.4.2 Quantifying water source contributions to deep soil water and groundwater

The MixSIAR model in conjunction with stable isotopes of water (δ^2 H and δ^{18} O) was applied to quantify the proportional contributions of current (between previous sampling time t - 1 and current sampling time t) river water to riparian deep water (i.e., deep soil water in the 80–170 cm layer or groundwater). The potential water sources of riparian deep soil water in the 80–170 cm layer at t included the in situ (i.e., water that was already in the deep soil layer or groundwater) soil water in this layer at t - 1, soil water in the 0–80 cm layer at t - 1 and t, and groundwater between t - 1 and t (Fig. 4a). We considered the in situ groundwater at t - 1, soil water in the 0–170 cm layer at t - 1, river water between t - 1 and t - 1.



Figure 3. Seasonal variations of the river water level, the water table depth (WTD), and groundwater level (GWL) at distances of 5, 20, and 45 m from the riverbank during the observation period in 2019 (**a**) and 2021 (**b**). The red arrow indicates the riparian ground surface level (29.5 m). The riverbed level is 26 m.



Figure 4. Schematic diagram of potential water sources: riparian deep soil water in the 80–170 cm layer (**a**) and groundwater (**b**). The red box represents riparian deep soil water in the 80–170 cm layer in panel (**a**) and groundwater in panel (**b**). The dark blue arrow indicates different potential water sources of riparian deep water.

and t, and precipitation between t - 1 and t as the potential water sources for riparian groundwater at t (Fig. 4b). The isotopic changes from t-1 to t (such as fractionation during this period) were negligible when calculating the contribution of upper soil water (i.e., in the 0–80 or 0–170 cm layers) at t-1 to deep moisture (i.e., soil water in the 80–170 cm layer or groundwater). The δ^2 H and δ^{18} O values of riparian deep wa-

ter at *t* were set as the mixture data in the MixSIAR model, while the water isotopes of their potential water sources were regarded as the source data.



Figure 5. Flowchart for quantifying the proportional contributions of river water to the transpiration of riparian trees. P_s and P_g denote the contributions of riparian deep soil water in the 80–170 cm layer and groundwater to the transpiration of riparian trees, respectively. s_r^{t-1} and g_r^{t-1} represent the proportional contributions of the old river water (before t - 1) to riparian deep soil water in the 80–170 cm layer and groundwater, respectively. s_s^{t-1} , s_r^t , and s_g^t signify the proportional contributions of in situ soil water in the 80–170 cm layer at t - 1, river water during t - 1 to t, and groundwater during t - 1 to t for riparian deep soil water in the 80–170 cm layer at t, respectively. g_g^{t-1} and g_r^r stand for proportional contributions of in situ groundwater at t - 1 and river water from t - 1 to t to riparian groundwater at t, respectively.

2.4.3 An iteration method to determine the RWC to the transpiration of riparian trees

After determining both riparian deepwater contributions to the transpiration of trees and the RWC to riparian deep water, the proportional contributions of the river water between t-1and t to the transpiration of riparian trees were quantified. It is worth noting that riparian deep soil water (80-170 cm) and groundwater could be recharged by river water continuously when the groundwater levels lay below the riverbeds (i.e., losing rivers). Therefore, the proportional contribution of the old river water (before t - 1) to riparian deep water should not be ignored. The total RWC to riparian deep water should be quantified explicitly during the entire period of the river flow into the riparian deep zone since 2007. We suppose that the contributions of old river water to riparian in situ deep water are identical to those of current river water (between t-1 and t) to riparian in situ deep water. We proposed an iteration method using the following expression to quantify the total RWC to the transpiration of riparian S. babylonica trees near the losing rivers:

$$\begin{aligned} \text{RWC} &= P_{\text{s}} \cdot S_{\text{r}} + P_{\text{g}} \cdot G_{\text{r}} \\ &= P_{\text{s}} \cdot \left(s_{\text{r}}^{t} + s_{\text{r}}^{t-1}\right) + P_{\text{g}} \cdot \left(g_{\text{r}}^{t} + g_{\text{r}}^{t-1}\right) \\ &= P_{\text{s}} \cdot \left(s_{\text{r}}^{t} + s_{\text{r}}^{t} \cdot s_{\text{s}}^{t-1} + s_{\text{r}}^{t} \cdot \left(s_{\text{s}}^{t-1}\right)^{2} + s_{\text{g}}^{t} \cdot g_{\text{r}}^{t} \\ &+ s_{\text{g}} \cdot g_{\text{r}}^{t} \cdot g_{\text{g}}^{t-1} + s_{\text{g}}^{t} \cdot g_{\text{r}}^{t-1} + g_{\text{r}}^{t} \cdot \left(g_{\text{g}}^{t-1}\right)^{2}\right) \\ &+ P_{\text{g}} \cdot \left(g_{\text{r}}^{t} + g_{\text{r}}^{t} \cdot g_{\text{g}}^{t-1} + g_{\text{r}}^{t} \cdot \left(g_{\text{g}}^{t-1}\right)^{2}\right) \\ &= \left(P_{\text{s}} \cdot s_{\text{r}}^{t} + P_{\text{g}} \cdot g_{\text{r}}^{t} + P_{\text{s}} \cdot s_{\text{g}}^{t} \cdot g_{\text{r}}^{t}\right) \\ &+ \left(P_{\text{s}} \cdot s_{\text{r}}^{t} \cdot s_{\text{s}}^{t-1} + P_{\text{g}} \cdot g_{\text{r}}^{t} \cdot g_{\text{g}}^{t-1} + P_{\text{s}} \cdot g_{\text{r}}^{t} \cdot s_{\text{g}}^{t} \cdot g_{\text{g}}^{t-1}\right) \\ &+ \left(P_{\text{s}} \cdot s_{\text{r}}^{t} \cdot \left(s_{\text{s}}^{t-1}\right)^{2} + P_{\text{g}} \cdot g_{\text{r}}^{t} \cdot \left(g_{\text{g}}^{t-1}\right)^{2} \right) \\ &+ P_{\text{s}} \cdot s_{\text{g}}^{t} \cdot g_{\text{r}}^{t} \cdot \left(g_{\text{g}}^{t-1}\right)^{2}\right), \end{aligned} \tag{4}$$

where S_r and G_r represent total RWC to riparian deep soil water in the 80–170 cm layer and groundwater, respectively; P_s and P_g represent the contributions of riparian deep soil water in the 80–170 cm layer and groundwater to the transpiration of riparian trees, respectively; s_r^{t-1} and g_r^{t-1} denote the proportional contributions of the old river water (before t-1) to riparian deep soil water in the 80–170 cm layer and groundwater, respectively; s_s^{t-1} , s_r^t , and s_g^t signify the proportional contributions of in situ soil water in the 80–170 cm layer at t-1, river water during t-1 to t, and groundwater during t-1 to t to riparian deep soil water in the 80–170 cm layer at t, respectively; and g_g^{t-1} and g_r^t symbolize the proportional contributions of in situ groundwater at t-1 and river water from t-1 to t to riparian groundwater at t, respectively.

The expression " $P_s \cdot s_r^t + P_g \cdot g_r^t + P_s \cdot s_g^t \cdot g_r^t$ " in Eq. (4) was proposed to determine the current river water (between t - 1and t) contributions to the transpiration of riparian trees. The second iteration ($P_s \cdot s_r^t \cdot s_s^{t-1} + P_g \cdot g_r^t \cdot g_g^{t-1} + P_s \cdot g_r^t \cdot s_g^t \cdot g_g^{t-1}$) and the third iteration ($P_s \cdot s_r^t \cdot (s_s^{t-1})^2 + P_g \cdot g_r^t \cdot (g_g^{t-1})^2 + P_s \cdot s_g^t \cdot g_g^t \cdot (g_g^{t-1})^2$) were applied to quantify the proportional contributions of old river water that recharged riparian in situ deep water to trees (Fig. 5). We only applied three iterations because the differences between the RWCs in the third iteration and the next iteration were smaller than 0.1%. Using this proposed iteration method, we accurately estimated the total proportions of old and current river waters to the transpiration of riparian trees.

2.5 Statistical analysis

For each variable, we tested the homogeneity of variance between the two studied years and between the three plots using Levene's test. The one-way analysis of variance (ANOVA) was applied to examine differences in each variable among three plots in 2019 and 2021 (p < 0.05). The variables included the WTD, SWC, δ^2 H values, and δ^{18} O values of different water sources and xylem water, ²²²Rn concentration of river water and groundwater, contributions of different water sources to riparian deep water or trees, and leaf δ^{13} C values. The linear regression model was fitted to the whole dataset in both years to obtain the general relationships between the WTD, leaf δ^{13} C values, and the RWC to the transpiration of riparian trees. The statistical analysis was carried out in Microsoft Excel (v2016) and SPSS (24.0, Inc., Chicago, IL, USA).

3 Results

3.1 Hydro-meteorological conditions

The observation period (from April to November) in 2021 was wet with total precipitation of 802.5 mm, which was 1.8 times greater than for the drier year 2019 (445.6 mm) (Fig. 2a). The precipitation amount during the rainy season accounted for 75.4 % and 97.0 % of the whole precipitation in 2019 and 2021, respectively. The annual mean temperature during the observation period in 2019 and 2021 was 22.4 and 21.8 °C, respectively. The average daily VPD during the observation period was significantly greater in the dry year of 2019 (1.1 kPa) than in the wet year of 2021 (0.9 kPa)

(p < 0.05) (Fig. S1a and b in the Supplement). There was a significant difference in the average daily ET₀ from June to September between the dry year of 2019 (5.0 mm d⁻¹) and the wet year of 2021 (4.3 mm d⁻¹) (p < 0.05), but no significant difference was observed during the rest of the observation period (i.e., April, May, October, and November) between the 2 years (p > 0.05) (Fig. S1c and d). No significant difference was found in the daily mean net radiation during the observation period between the dry year of 2019 and the wet year of 2021 (p > 0.05) (Fig. S1c and d).

The river water level fluctuated between 27.9 and 28.9 m in 2019 and between 27.7 and 29.3 m in 2021 (Fig. 3). The mean WTD across the three plots was significantly (p < 0.05) deeper in 2019 $(2.7 \pm 0.3 \text{ m})$ than in 2021 $(1.7 \pm 0.5 \text{ m})$. The WTD increased with increasing distances from the riverbank in both 2019 and 2021 (Fig. 3). The river water continuously recharged the groundwater system ("losing" river) during the observation periods in 2019 and 2021, which was indicated by a lower groundwater level than the river water level (Fig. 3). Significantly higher SWC was observed in 2021 compared to 2019 (p < 0.05) (Fig. S2). The SWC of each soil layer at D45 was significantly lower than that at D05 and D20 in 2021 (p < 0.05), while no pronounced difference was observed in the SWC in the 0–30 cm layer among the three plots in 2019 (p > 0.05) (Fig. S2).

3.2 Direct water source contributions to the transpiration of riparian trees

Precipitation was significantly more depleted in $\delta^2 H$ and δ^{18} O in 2021 (-52.9% ± 30.2% for δ^{2} H and $-8.1\% \pm 3.8\%$ for δ^{18} O) than in 2019 (-29.2\% \pm 18.8\%) for δ^2 H and $-4.1\% \pm 3.0\%$ for δ^{18} O) (p < 0.05) (Fig. 6). The slope of the local meteoric water line in 2021 (7.8) was significantly higher than in 2019 (5.5) (p < 0.05), suggesting that the falling raindrops underwent stronger sub-cloud evaporation in 2019 (Zhao et al., 2019). The δ^2 H and δ^{18} O values of the surface soil water (above 30 cm depth) were significantly lower and more variable in 2021 than in 2019 (p < 0.05) (Fig. 6). In contrast, there were slightly higher water isotopic compositions in the 30-170 cm soil layer in 2021 compared to 2019. No significant difference was observed in the isotopic compositions of the soil water below 170 cm depth or groundwater between 2019 and 2021 (p > 0.05). The δ^2 H and δ^{18} O values of soil water in the 80– 170 cm layer were significantly lower than those of groundwater in 2019 (p < 0.05), while no significant difference was observed between soil water isotopes in the 80-170 cm layer and groundwater isotopes in 2021 (p > 0.05). Groundwater was significantly more depleted in $\delta^2 H$ and $\delta^{18} O$ compared to river water in both years (p < 0.05) (Fig. 6). The δ^2 H and δ^{18} O values of xylem water during the observation periods in 2019 and 2021 were not significantly different (p > 0.05), but they were gradually lower with the increasing distance from the riverbank.



Figure 6. Dual-isotope (δ^2 H and δ^{18} O) biplots of different water bodies in the three plots (D05, D20, and D45) for the observation years of 2019 and 2021. The local meteoric water line (LMWL) was determined for each year from the precipitation samples taken over each year. The soil water line (SWL) was determined for each plot using the soil water samples taken over each year. D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. The error bars indicate standard deviations.

The contributions of the surface soil water to the transpiration of riparian trees in 2019 (20.1 $\% \pm 9.7 \%$) were similar to those of 2021 (19.0 $\% \pm 10.5 \%$). No significant difference was observed in the soil water contributions to the transpiration of riparian S. babylonica in the 30-80 cm layer between both years (p > 0.05) (Fig. 7). The S. babylonica principally relied on riparian deep water below the 80 cm depth in both 2019 (55.9%) and 2021 (57.1%). There was no significant difference in the riparian deepwater contributions to the transpiration of S. babylonica trees between the three distances from the riverbank (p > 0.05) (Fig. 7). Nevertheless, the soil water contributions in the 80-170 cm layer to the transpiration of riparian trees decreased with increasing distance from the riverbank in both years, whereas the proportions of groundwater taken up by riparian trees increased from D05 to D45 in both 2019 (from 27.6 % to 32.1 %) and 2021 (from 17.0% to 32.2%) (Fig. 7). It was found that groundwater contributions to the transpiration of riparian S. babylonica trees increased significantly (p < 0.05) from April to July

in both years. They plummeted significantly (p < 0.05) and reached a minimum in September 2021.

3.3 Water source contributions to riparian deep soil water and groundwater

The primary water sources of riparian deep soil water in the 80–170 cm layer were the in situ soil water in this layer (with a mean value of 33.1 %) and groundwater capillary rise (with a mean value of 25.3 %) in 2019 (Fig. 8). However, the in situ soil water in the 80–170 cm layer (with a mean value of 23.9 %), groundwater capillary rise (with a mean value of 24.6 %), and river water (with a mean value of 24.4 %) contributed almost equally to riparian deep soil water in 2021. The in situ soil water contribution to riparian deep soil water was significantly higher in 2019 than in 2021 (p < 0.05). However, the river water contributed less to riparian deep soil water in 2019 (with a mean value of 15.7 %) compared to 2021 (p < 0.05). The RWC to riparian deep soil water was the lowest in August 2019 (11.3 % ± 4.5 %) and in June 2021



Figure 7. Seasonal variations in the proportional contributions of soil water and groundwater to the transpiration of riparian trees in the three plots (D05, D20, and D45) for the observation years of 2019 (**a**–**c**) and 2021 (**d**–**f**). D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. The error bars indicate standard deviations.

 $(13.6\% \pm 3.8\%)$, respectively. The in situ soil water contributions to riparian deep soil water showed a significant increase with increasing distance from the riverbank, while the RWC to riparian deep soil water decreased from D05 to D45 in both years (p < 0.05) (Fig. 8).

The in situ groundwater contribution was significantly higher in 2019 (with a mean value of $56.0\% \pm 11.2\%$) than in 2021 ($37.1\% \pm 16.7\%$) (p < 0.05) (Fig. 9). The average contribution of the river water to riparian groundwater was $28.1\% \pm 12.1\%$ during the observation period. There was a significantly higher RWC to riparian groundwater in 2021 (with a mean value of $35.1\% \pm 11.9\%$) than in 2019 (with a mean value of $21.1\% \pm 7.2\%$) (p < 0.05). The lowest RWC ($13.0\% \pm 1.2\%$) occurred in August with the lowest groundwater level of 3.1 m in 2019, whereas the contribution of river water to riparian groundwater ($47.1\% \pm 13.2\%$) was the highest in July with a higher groundwater level of 1.8 m in 2021 (Figs. 3 and 9). The proportional contribution of the in situ groundwater to riparian groundwater increased with increasing distance from the riverbank during the observation periods, while the RWC to riparian groundwater decreased significantly from D05 to D45 (p < 0.05) (Fig. 9). There was a significant increase of ²²²Rn activity in groundwater from D05 (494.5 ± 107.5 Bqm⁻³) to D45 (787.4 ± 153.2 Bqm⁻³) (p < 0.05) (Table 1). The $T_{\rm res}$ of recharged groundwater from river water increased from D05 (0 d) to D45 (0.15 ± 0.13 d) (Table 1). This also indicated that the river recharged riparian deep strata rapidly and frequently, particularly more significant in the plots closer to the riverbank.

3.4 Seasonal variations in RWC to the transpiration of riparian trees

The proportional contributions of river water to the transpiration of riparian *S. babylonica* trees were significantly higher in 2021 (with a mean value of 23.8 % \pm 7.8 %) than in 2019 (with a mean value of 16.8 % \pm 4.7 %) (p < 0.05). Specifically, the most significant monthly difference in the RWC to the transpiration of riparian *S. babylonica* trees between the



Figure 8. Seasonal variations in the contributions of different water sources to riparian deep soil water in the 80–170 cm layer in the three plots (D05, D20, and D45) for the observation years of 2019 (**a–c**) and 2021 (**d–f**). D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. The error bars represent standard deviations.

Table 1. The ²²²Rn values in river water, background groundwater and riparian groundwater in three plots (D05, D20, and D45), and the average residence time of recharged groundwater from river water (T_{res} , d) in 2021. The background groundwater indicates groundwater in aquifers more than 100 m away from the riverbank. The "negative T_{res} values" were set to "0".

	River water	Background groundwater	Riparian groundwater		
			D05	D20	D45
$\frac{222}{T_{\rm res}} \frac{({\rm Bq}{\rm m}^{-3})}{({\rm d})}$	610.1 ± 212.3 0	7400 ± 35.4 Null	$\begin{array}{c} 494.5\pm107.5\\ 0\end{array}$	763.3 ± 118.3 0.13 ± 0.1	$787.4 \pm 153.2 \\ 0.15 \pm 0.13$

Note: D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. Null represents no data for the residence time of background groundwater.

dry year of 2019 and the wet year of 2021 was up to 19.8 % (p < 0.01). The monthly maximum RWC to the transpiration of *S. babylonica* trees was significantly higher in the wet year of 2021 ($35.2 \% \pm 7.0 \%$) compared to the dry year of 2019 ($24.2 \% \pm 3.0 \%$) (p < 0.05).

The riparian S. babylonica took up the most river water in July 2021 $(35.2\% \pm 7.0\%)$, whereas the highest

RWC to the transpiration of riparian trees occurred in June 2019 (24.2 $\% \pm 1.6 \%$). The minimum river water up-take for riparian *S. babylonica* in 2021 was in September (17.7 $\% \pm 2.7 \%$), while trees took up the least river water in August 2019 (13.2 $\% \pm 1.9 \%$). Although the precipitation amount in the rainy season was much higher than in the drought season (p < 0.01), no significant difference in the



Figure 9. Seasonal variations in the contributions of different water sources to riparian groundwater in the three plots (D05, D20, and D45) for the observation years of 2019 (**a–c**) and 2021 (**d–f**). D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. The error bars indicate standard deviations.

RWC to the transpiration of riparian *S. babylonica* trees was observed between the rainy and drought seasons in the same year (p > 0.05) (Figs. 2 and 9). The difference values of the RWC to the transpiration of riparian trees between the rainy and dry seasons were not significantly different (p > 0.05) in both 2019 (-4.0%) and 2021 (-4.4%) (Fig. 9). This suggested that there were no significant seasonal variations in the RWC to the transpiration of riparian trees within a year (p > 0.05).

The water uptake of river water by riparian *S. babylonica* was significantly different between the three plots in 2019 (p < 0.05), while no difference was observed between the three plots in 2021 (p > 0.05) (Fig. 10). In particular, the RWC to the transpiration of riparian trees decreased significantly by 6.9 % from D05 (20.0 %) to D45 (13.1 %) in 2019 (p < 0.05), whereas there was no significant difference in 2021 (p > 0.05) (Fig. 10).

3.5 Relationships between leaf δ^{13} C, RWC to the transpiration, and WTD

The leaf δ^{13} C of riparian *S. babylonica* trees was significantly higher in 2019 (-27.7% ± 1.0%) than in 2021 (-29.7% ± 0.7%) (p < 0.05) (Table 2). There was a significant increase of the leaf δ^{13} C from D05 (-28.8%) to D45 (-27.0%) in 2019 (p < 0.05), while no significant difference in the leaf δ^{13} C was observed between the three plots in 2021 (p > 0.05). The lowest leaf δ^{13} C value of riparian trees occurred on 15 August 2019 and 14 July 2021. These minimum values of leaf δ^{13} C occurred when intense rainfall had not occurred in both years.

There was a significantly negative relationship between the RWC to the transpiration of riparian trees and WTD $(R^2 = 0.57; p < 0.01)$ (Fig. 11a). The leaf δ^{13} C of riparian *S. babylonica* was found to be negatively correlated with the RWC to the transpiration of riparian trees ($R^2 = 0.61$; p < 0.01) but positively linearly related to WTD ($R^2 = 0.37$; p < 0.01) (Fig. 11b and c). This demonstrated that deeper



Figure 10. Contributions of river water to the transpiration of riparian trees in the three plots (D05, D20, and D45) for each sampling campaign for the observation years of 2019 (a) and 2021 (b). Different letters show a significant difference in the river water contribution to the transpiration of riparian trees between three plots for each sampling campaign (p < 0.05). D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively.

Table 2. Leaf δ^{1} .	³ C values of ripa	rian <i>S. babylonica</i> in th	e three plots (D05, D	D20, and D45) during	g the observation	period in 2019 and 2021
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	Leaf δ^{13} C value (% <i>o</i>)							
	2019							
	5 May	14 Jun	26 Jul	15 Aug	26 Sep	5 Nov	Mean	SD
D05	-28.8	-29.2	-29.7	-30.4	-28.1	-27.4	-28.8	1.0
D20	-27.1	-26.7	-27.1	-27.5	-27.4	-27.2	-27.1	0.2
D45	Null	-27.2	-26.9	-27.4	-26.9	-26.5	-27.0	0.3
				2021				
	24 Apr	25 May	26 Jun	14 Jul	1 Sep	5 Nov	Mean	SD
D05	-29.7	-29.5	-29.5	-31.0	-29.5	-29.1	-29.7	0.6
D20	-28.8	-29.1	-29.4	-30.4	-30.1	-30.3	-29.7	0.7
D45	-29.0	-29.0	-29.4	-30.8	-30.1	-30.0	-29.7	0.9

Note: D05, D20, and D45 are the plots at distances of 5, 20, and 45 m from the riverbank, respectively. SD represents standard deviation.

WTD $(2.7 \pm 0.3 \text{ m})$ resulted in lower RWC to the transpiration of riparian *S. babylonica* and higher leaf-level WUE in the drier year of 2019. In contrast, the riparian *S. babylonica* under shallower WTD $(1.7 \pm 0.5 \text{ m})$ gave rise to higher RWC but lower leaf-level WUE in the wetter year of 2021.

4 Discussion

4.1 RWC to the transpiration and effects of distance from the river on RWC

We identified deep-rooted riparian trees near the losing river to use a small proportion of river water (less than 25%) for transpiration (Fig. 10). The small RWC to the transpiration of riparian trees may originate from three non-exclusive pro-



Figure 11. Relationships between the contributions of river water to the transpiration of riparian trees and the water table depth (**a**), between the leaf δ^{13} C values and the water table depth (**b**), and between the leaf δ^{13} C values and proportions of river water contributions to riparian trees (**c**). The red line represents the linear relationship fitted by the monthly data in three plots in 2019, while the blue line represents the linear relationship fitted by the monthly data in three plots in 2019, while the blue line represents the linear relationship fitted by the monthly data in three plots in both years. The WTD, leaf δ^{13} C values, and river water contributions to the transpiration of riparian *S. babylonica* are monthly data at each plot at a distance of 5, 20, and 45 m from the riverbank during the observation period in both years.

cesses. First, the lateral roots of riparian trees further than 5 m away from the riverbank rarely took up river water directly when their projected edges of the canopy (less than 5 m in our study) were out of reach of the river (Busch et al., 1992; Thorburn and Walker, 1994). Instead, they took up riparian deep soil water and/or groundwater recharged by river water, which likely restricted the RWC to the transpiration of riparian trees. Second, the ecohydrological separation (Brooks et al., 2010; Evaristo et al., 2015; Allen et al., 2019; Sprenger et al., 2019) possibly resulted in large isotopic discrepancies between fast-moving water flow and immobile water for plant water uptake. Although the residence time of recharged groundwater from river water was extremely short (less than 0.28 d) (Table 1), only one-third of riparian groundwater was replaced by the lateral seepage of river water (Fig. 9). Our finding probably indicates that river water recharged mobile groundwater quickly but could not completely replace water held tightly in the soil pores (Brooks et al., 2010; Evaristo et al., 2015; Allen et al., 2019). This is consistent with Sprenger et al. (2019), who found that the lateral seepage of river water or rising groundwater level could briefly saturate riparian soils but could not entirely replace/flush immobile waters or homogenize different water pools isotopically. Third, several recent studies showed that phreatophytic or deep-rooted trees predominantly extended roots into fine pores to take up immobile soil water (Evaristo et al., 2015, 2019; Maxwell and Condon, 2016). As mentioned above, the immobile water could not be completely replaced by infiltrating river water, which eventually resulted in a small contribution of river water to deep-rooted riparian trees. This ecohydrological separation perspective that plantaccessible water pools were separated from the fast-moving water can also be supported by our findings that no significant difference in RWCs to the transpiration of riparian trees was observed between rainy and drought seasons in both dry and wet years (Fig. 9). This is because riparian *S. babylonica* trees preferred to rely on immobile water in fine soil pores, and they would not change the river water uptake patterns when the fast-moving precipitation input increased (Brooks et al., 2010; Sprenger et al., 2019).

Compared to the small RWC to the transpiration of riparian S. babylonica trees (less than 25%) in our study, Alstad et al. (1999) found that riparian Salix monticola trees near a losing river on the northeast side of the Rocky Mountain National Park, Colorado, relied on rivers for approximately 80% of their transpiration. The significant difference in the RWC to the transpiration of riparian trees between the two studies can be attributed to the potential water source determination as well as the calculation method. First, Alstad et al. (1999) only considered river water and precipitation as potential water sources for riparian S. monticola, which resulted in an overestimation of the RWC to the transpiration of riparian S. monticola. This is because the RWC estimation in Alstad et al. (1999) also included the proportions of the indirect river water, in situ soil water, and in situ groundwater contributions to the transpiration of riparian S. monticola. Second, river water can seep into the saturated and/or vadose zone across the riparian riverbank and be further taken up by riparian trees indirectly in the form of river-recharged deep soil water and/or groundwater. In our study, we separated and determined the contributions of indirect river water sources (i.e., the river-recharged deep soil water in the 80-170 cm layer and groundwater also contained river water) to the transpiration of riparian trees. Accurately quantifying the indirect RWC to deep-rooted riparian trees could assist us to determine the effect of riparian plant water use on river runoff along the losing river.

We observed substantial variations in the RWC to the transpiration of riparian trees at interannual (between 2 years) and spatial (between three distances from the riverbank) scales (Fig. 10). The RWC to the transpiration of riparian S. babylonica trees in the wet year of 2021 was 1.4 times greater on average than in the dry year of 2019 (Fig. 10). Nevertheless, riparian S. babylonica trees presented similar root architecture (i.e., phreatophyte) associated with similar water source proportions between dry and wet years (Fig. 7). This suggested that the higher groundwater level in the wet year induced higher RWC to riparian deep water compared to the dry year, which further resulted in a higher indirect RWC to the transpiration of riparian phreatophyte trees in the wet year than in the dry year. Although there was no significant difference in the deep water (below the 80 cm layer) contributions to the transpiration of riparian trees between the three plots, we observed a substantial impact of the declining groundwater level with increasing distance from the riverbank on the decreased indirect RWC to the transpiration of riparian trees in the dry year of 2019 (Fig. 10). The declining water table and increasing residence time of recharged groundwater from D05 to D45 could consequently lead to the decreasing RWC to riparian deep water along the distance away from the riverbank. Thus, the interannual and spatial variabilities of the RWC to the transpiration of riparian S. babylonica trees were generally attributed to the various RWCs to riparian deep water rather than the water uptake patterns of riparian trees. This result is in contrast to that of a previous study conducted by Qian et al. (2017), who reported a significant increase in the RWC to the transpiration of G. biloba trees in response to the groundwater level decline. This discrepancy was ascribed to the fact that riparian G. biloba had a dimorphic root system and shifted their main water sources from the shallow soil layer to the deeper soil layer. However, the potential root growth rate of riparian phreatophyte S. babylonica trees can reach $1-13 \text{ mm d}^{-1}$, which allows the riparian S. babylonica trees to remain in contact with a rising or declining groundwater level and to maintain constant water uptake proportions from deep strata below the 80 cm depth (Naumburg et al., 2005).

4.2 Link between RWC, WUE, and WTD as well as the implications

The water uptake patterns of riparian *S. babylonica* trees generally followed the characteristics of a phreatophyte. We observed that leaf WUE of all *S. babylonica* trees across three plots in both dry and wet years was negatively correlated with the indirect RWC to the transpiration of riparian trees and positively related to WTD (Figs. 10 and 11b and c). These relationships are consistent with previous studies (Cao et al., 2020; Ding et al., 2020; Behzad et al., 2022). Higher leaf

WUE associated with lower RWC to the transpiration of riparian trees and lower groundwater levels are likely because water stress restricts the stomatal conductance and further reduces the transpiration rate of riparian trees. Specifically, the dry year of 2019 was characterized by higher water demand (indicated by higher VPD) and lower water availability compared to the wet year of 2021, but the energy resource (indicated by net radiation) for riparian trees was similar between the 2 years (Figs. S1 and S2 in the Supplement). Hence, we suppose that water limitation rather than energy limitation regulates the leaf-level stomatal conductance of riparian S. babylonica trees. The high water demands but low river water availability in the dry year likely resulted in the stomatal closure of riparian trees to minimize water loss, which eventually led to a decrease in transpiration rate and even photosynthetic rate (Fabiani et al., 2022; Behzad et al., 2022). Aguilos et al. (2019) further found that water stress would enhance radiation-normalized WUE, because the lack of water availability induced a stronger reduction in transpiration than photosynthesis. With no difference in the average net radiation between dry and wet years, the lower river water availability in a dry year probably increased leaf WUE. It can be inferred that riparian S. babylonica trees took up more river water and possibly exhibited a consumptive river-water-use pattern in the wet year compared to the dry year. This agreed well with previous investigations during which the woody plants showed lower leaf WUE and consumptive water use patterns in the rainy season, while they showed higher leaf WUE and conservative water use patterns with lower soil water availability in the dry season (Horton and Clark, 2001; Cao et al., 2020; Behzad et al., 2022). However, consumptive river water taken up by riparian trees could result in a great loss of river water, which should be avoided in the riparian zone of a losing river that is under restoration by "ecological water".

The WTD played a critical role in the river water uptake of riparian trees near a losing river (Mensforth et al., 1994; Horton and Clark, 2001; Qian et al., 2017; Zhou et al., 2017). We observed that the proportional contributions of the river water to the transpiration of riparian trees decreased linearly in response to groundwater level decline, leading to a proportional increase in leaf WUE (Fig. 11a and b). Our finding was consistent with Horton and Clark (2001), who reported an exponential increase in the leaf WUE of riparian Salix gooddingii with increasing WTD. As mentioned above, we emphasized the key role of reduced water availability in decreasing transpiration rate and thus in enhancing leaf WUE in this study. Nevertheless, there were some controversial views that the leaf WUE of plant species increased initially and then decreased with increasing WTD (Antunes et al., 2018; Xia et al., 2018). This can be justified by the fact that riparian trees can tolerate reduced water availability only within a speciesspecific threshold, beyond which xylem cavitation and even crown mortality occur (Naumburg et al., 2005). This indicates that optimal WTD for plant species is related to the

highest leaf WUE, under which plant species can consume less water for transpiration to maximize CO_2 assimilation (Antunes et al., 2018; Xia et al., 2018). The breakpoint of WTD was not observed in this study (Fig. 11a and b). Further investigations would need to be conducted under deeper groundwater levels (WTD > 4 m) to optimize the WTD and riparian plant-water relations.

Y. Li et al.: Lessons from stable isotopes and iteration method

Our results have important implications for untangling the trade-offs between riparian tree water use and river runoff management. The proportion of the RWC to the transpiration of riparian trees was compared between dry and wet years to investigate the impacts of river water availability on the water use characteristics of riparian trees. The riparian S. babylonica trees showed the highest leaf WUE and the lowest river water uptake proportion under the lowest groundwater level condition (with a WTD of 4 m). The rising groundwater level may encourage riparian trees to exhibit a consumptive river-water-use pattern, which should not be recommended in revegetated riparian zones beside an ecological water-recharged losing river. Thus, the relationships between the RWC to the transpiration of riparian trees, leaf-level physiological characteristics (e.g., leaf WUE), and hydro-meteorological conditions are critical to protecting the revegetated riparian zones and maintaining river runoff sustainability.

4.3 Advantages and limitations of the MixSIAR model and the iteration method

The iteration method in combination with the MixSIAR model and stable isotopes of water is particularly useful for separating and quantifying the proportional contributions of river water to the transpiration of riparian trees near a losing river. This integration of methods is more accurate than previous studies (Alstad et al., 1999; Zhou et al., 2017; White and Smith, 2020), which only considered river water as a direct water source of riparian trees without considering their distances from the riverbank and the extents of the lateral roots. The primary advantage of the combined method is that

it explicitly identifies the direct and indirect water sources of riparian trees based on the distance from the riverbank, the extent of lateral roots, and the process of riparian deepwater recharging by the river. To ensure the convergence of the MixSIAR model, both the trace plots and three diagnostic tests (i.e., Gelman–Rubin, Heidelberger–Welch, and Geweke) were adopted (Stock and Semmens, 2013). Besides, the MixSIAR model explicitly considers the uncertainties in the isotopic values and the estimates of source contributions compared to the simpler linear mixing models (Stock and Semmens, 2013; Ma et al., 2016). The strength of the newly proposed multi-iteration method is that it can determine the total contributions of the indirect river water source to the transpiration of riparian trees. The multi-iteration will not stop until there is no significant difference between the results of the last two iterations. This reduces the calculation errors of the RWC to the transpiration of riparian trees.

However, there are still some limitations that should be further investigated in future studies. First, the riparian deepwater sources were identified using the water isotopic data collected in campaigns taking place at an interval of about 1 month. The riparian soil water movement was complex, and the stable isotopes of water might not be uniform between the two campaigns along the losing river. Nevertheless, the isotopic changes from t - 1 to t (such as fractionation during this period) were negligible when calculating the contribution of upper soil water (i.e., in the 0-80 or 0-170 cm layers) at t - 1 to deep moisture (i.e., soil water in the 80-170 cm layer or groundwater). Assuming the isotopic uniformity over such a time interval may cause uncertainties in estimating the RWC to the transpiration of riparian deep water. Second, we supposed that the contributions of old river water (before initial time (t - 1)) to riparian in situ deep water were identical to the contributions of current river water (during the observation period between t - 1 and t) to riparian in situ deep water. This can induce some uncertainties in the estimations of the RWC to riparian deep water and the RWC to the transpiration of riparian trees. To minimize this issue, water samples would need to be collected more frequently to quantify the contributions of river water to riparian deep water and tree transpiration. Third, we inferred the approximate lateral root extent based on the projected edge of the canopy of S. babylonica, which indicated that S. babylonica trees could not tap into the river or take up river water directly. However, the lateral roots of S. babylonica trees should be directly investigated in further research to confirm our inference. Fourth, the riparian WTD along the studied reach of the Chaobai River (from Dam 5 to Dam 4) ranged from 0.2 to 4.3 m in two studied years (these data have not been published yet). The selected site in this study was the most representative site since there was a significant water table variation (ranging from 0.3 to 4.0 m) in the 2 studied years. However, the implications of quantifying the effects of river water on the water use of riparian trees in this study are only applicable to relatively shallow water table conditions (with the WTD less than 4 m). Further investigations should be conducted at deep-WTD sites to better understand and regulate river runoff and tree water needs.

5 Conclusion

We presented a new iteration method in combination with the MixSIAR model and stable isotopes of water (δ^2 H and δ^{18} O) to separate and quantify the proportional contributions of river water to the transpiration of riparian *S. babylonica* in the dry year of 2019 and the wet year of 2021 along a losing river in Beijing, China. We found that the infiltrating river water was exchanged with riparian mobile water quickly but was not completely mixed with waters held tightly in the fine

pores. Riparian trees near the losing river generally extended roots into fine pores to access immobile water sources. The isotopic discrepancies between the fast-moving water flow and the immobile water taken up by the roots led to a small RWC (20.3 %) to the transpiration of riparian trees. The water deficit in the dry year probably induced stomatal closure and a larger reduction in transpiration compared to the photosynthesis of riparian trees, thus leading to an evident increase of leaf WUE compared to the wet year. The leaf WUE exhibited a negative correlation with the RWC to the transpiration of riparian trees but was positively linearly related to WTD (p < 0.01). Riparian S. babylonica trees maintained the highest leaf WUE and the lowest river water uptake proportion under deep-groundwater conditions (with a WTD of 4 m). This suggests that rising groundwater levels may encourage riparian trees to increase the river water uptake and show a consumptive river-water-use pattern, which cannot be beneficial to the water resource management of a losing river that is under restoration by ecological water. This study provides valuable insights into riparian afforestation that is related to water use and ecosystem health.

Appendix A: Acronym dictionary

MixSIAR	A Bayesian mixing model that uses stable-
	isotope data to estimate the proportions of
	source contributions to a mixture
RWC	River water contribution
WUE	Leaf-level water use efficiency
WTD	Water table depth
Т	Temperature
RH	Relative air humidity
ET ₀	Reference evapotranspiration
VPD	Vapor pressure deficit
SWC	Soil water content
IRIS	Isotopic ratio infrared spectroscopy
IRMS	Isotope Ratio Mass Spectrometry
VSMOW	Vienna Standard Mean Ocean Water
C_{Water}	²²² Rn concentration of the water samples
$C_{\rm Air}$	Air ²²² Rn concentration of the water samples
C_{System}	Air ²²² Rn concentration of the measurement
2	system
V _{System}	The interior volume of the measuring setup
V _{Sample}	The volume of water sample
$T_{\rm res}$	The average residence time of recharged
	groundwater from river water
k	The ²²² Rn distribution coefficient of
	water/air
λ	The decay coefficient
Ce	The ²²² Rn concentration of background
	groundwater when the equilibrium between
	radon production and decay is reached

$C_{\rm r}$	The ²²² Rn concentration of river water
C_{g}	The ²²² Rn concentration of riparian ground-
	water
PWL	The potential water source line
ap	The slope of the PWL
<i>b</i> _p	The intercept of the PWL
PW-excess	The δ^2 H deviation of riparian tree xylem
G	water from the PWL
Sr	The total RWC to riparian deep soil water
	in the $80-170 \text{ cm}$ layer (throughout the river
G	losing-flow period since 2007)
$G_{\rm r}$	The total RWC to riparian groundwater
	(throughout the river losing-flow period since 2007)
P_{s}	The contribution of riparian deep soil water
5	in the 80–170 cm layer to riparian trees
P_{g}	The contribution of riparian groundwater to
8	riparian trees
s_r^{t-1}	The proportional contribution of the old river
1	water (before $t-1$) to riparian deep soil water
	in the 80–170 cm layer
g_r^{t-1}	The proportional contribution of the old river
	water (before $t - 1$) to riparian groundwater
s_{s}^{t-1}	The proportional contribution of in situ soil
-	water in the 80–170 cm layer at $t - 1$ to
	riparian deep soil water in the 80-170 cm
	layer at t
s_{r}^{t}	The proportional contributions of river water
	from $t - 1$ to t to riparian deep soil water in
	the $80-170$ cm layer at t
s_{g}^{t}	The proportional contribution of ground-
	water from $t - 1$ to t to riparian deep soil
	water in the $80-170$ cm layer at t
g_{g}^{t-1}	The proportional contribution of in situ
	groundwater at $t - 1$ to riparian groundwater
<i>t</i>	at t
$g_{ m r}^{\prime}$	The proportional contribution of river water
	from $t - 1$ to t to riparian groundwater at t
ANOVA	One-way analysis of variance
LMWL	Local meteoric water line

Data availability. The data that support the findings of this study are available from the corresponding author upon request.

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Author contributions. YL: investigation, methodology, formal analysis, writing (original draft and review and editing); YM: methodology, formal analysis, conceptualization, writing (review and editing); XS: supervision, writing (review and editing), project

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