Seasonal variation in water uptake patterns of three plant species based on stable isotopes in the semi-arid Loess Plateau

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HIGHLIGHTS

• Seasonal variations of water uptake pattern were determined by dual stable isotopes (δD and δ¹⁸O) and MixSIAR model.
• Soil water in the 0–120 cm depth contributed 75–80% to the total water uptake in the growing season.
• Vitex negundo displayed larger degree of ecological plasticity to switch water between shallow and deep soil layers.
• Functionally dimorphic root systems were related to flexible water uptake pattern.

GRAPHICAL ABSTRACT

ABSTRACT

Water is a limiting factor and significant driving force for ecosystem processes in arid and semi-arid areas. Knowledge of plant water uptake pattern is indispensable for understanding soil–plant interactions and species coexistence. The ‘Grain for Green’ project that started in 1999 in the Loess Plateau of China has led to large scale vegetation change. However, little is known about the water uptake patterns of the main plant species that inhabit in this region. In this study, the seasonal variations in water uptake patterns of three representative plant species, Stipa bungeana, Artemisia gmelinii and Vitex negundo, that are widely distributed in the semi-arid area of the Loess Plateau, were identified by using dual stable isotopes of δD and δ¹⁸O in plant and soil water coupled with a Bayesian mixing model MixSIAR. The soil water at the 0–120 cm depth contributed 79.54 ± 6.05% and 79.94 ± 8.81% of the total water uptake of S. bungeana and A. gmelinii, respectively, in the growing season. The 0–40 cm soil contributed the most water in July (74.20 ± 15.20%), and the largest proportion of water (33.10 ± 15.20%) was derived from 120–300 cm soils in August for A. gmelinii. However, V. negundo obtained water predominantly from surface soil horizons (0–40 cm) and then switched to deep soil layers (120–300 cm) as the season progressed. This suggested that V. negundo has a greater degree of ecological plasticity as it could explore water sources from deeper soils as the water stress increased. This capacity can mainly be attributed to its functionally dimorphic root system. V. negundo may have a competitive advantage when encountering short-term drought. The ecological plasticity of plant water use needs to be considered in plant species selection and ecological management and restoration of the arid and semi-arid ecosystems in the Loess Plateau.

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1. Introduction

Water plays crucial roles in plant productivity and species diversity in both natural and anthropogenic ecosystems (Porporato et al., 2004), and determines the distribution and ecological functions of vegetation. Plants are able to obtain water from various sources over time, depending on species, growth forms and environment conditions (Dawson et al., 2002; Eggemeyer et al., 2009; Grossiord et al., 2017; Yang et al., 2015; Evaristo and McDonnell, 2017). The spatial and temporal variations in water sources absorbed by plants are referred to water uptake pattern. Plant water uptake plays an important role in understanding and modeling hydrological processes at the soil–vegetation–atmosphere interface (Chen et al., 2017; Sprenger et al., 2016; Vargas et al., 2017), and it provides critical insights into adaptive responses of plants to the changing environments (Dawson et al., 2002; Evaristo et al., 2016; Grossiord et al., 2017).

In arid and semi-arid ecosystems, soil moisture generated by precipitation is a limiting factor controlling vegetation structures and ecosystem processes (Gao et al., 2011; Moreno-de Las Heras et al., 2011). A healthy plant–soil ecosystem requires that water utilization by plants does not exceed the supply of soil water otherwise leads to water deficit (Chen et al., 2015). The capacity to adjust water use strategy along with the highly variable rainfall pattern and consequently soil water content is very critical for a plant to survive in these ecosystems (Wu et al., 2016; Grossiord et al., 2017). Plant derived the majority water from deep sources (e.g., deep soil water, groundwater) during the dry season while most water they used was derived from shallow sources supplied by upper soil layers during the wet season (Dawson and Pate, 1996; Asbjørnsen et al., 2008). Multiple species living in the same habitat may have distinct water use patterns, making it possible for the species to coexist (Eggemeyer et al., 2009; Wu et al., 2016). This phenomenon may be attributed to hydrological niche segregation (Silvertown et al., 2015). Several studies showed that grass and herbs tended to continuously rely on water in the shallow soil layer across the growing season (Asbjørnsen et al., 2008; Prechsl et al., 2015; Priyadarshini et al., 2016). In comparison, some studies showed that trees and shrubs tended to have the capacity to derive water sources from deeper soil layers (McCole and Stern, 2007; Wu et al., 2016). One of the reasons to explain the different water uptake patterns among varied plant functional types may relate to root morphology of plants. Some studies suggested that species with dimorphic roots have the capacity to switch water absorption zones between shallower and deeper soil layers (Dawson and Pate, 1996; Nie et al., 2010; Yang et al., 2015). The transposition capacity to absorb water sources from shallow to deep soil layers reflects plant ecological plasticity (Valladares et al., 2007). The greater ecological plasticity usually indicates a greater adaptation level to the environment. Therefore, a better understanding of plant water uptake patterns will improve our understanding of soil–plant interactions and guide ecosystem management practices in arid and semi-arid areas such as selection of plant species for ecological restoration (Jian et al., 2015; Lü et al., 2017; Sprenger et al., 2016).

Water sources utilized by plants can be determined by many methods, such as root system excavation (Xu and Li, 2006), sap flow techniques (Delzon and Loustau, 2005), electrical resistivity (Mares et al., 2016), GIS tools (Howard and Merrifield, 2010), radioactive tracer tritium (Zhang et al., 2017). Although these methods could determine plant water sources in some extent, stable isotope technique provides an effective, powerful and nondestructive approach for identifying and partitioning the different potential water sources used by plants (Ma and Song, 2016; Geris et al., 2017; Hardanto et al., 2017; Rothfuss and Javaux, 2017). Previous studies have proven that no isotopic fractionation of water occurs during water absorption by roots and transportation along the root–shoot conduit before transpiration starts in terrestrial plants (Brunel et al., 1995; Dawson et al., 2002; Ehleringer and Dawson, 1992), except for some coastal wetland species (Lin and Sternber, 1993) and woody xerophytes (Ellsworth and Williams, 2007), for which isotope fractionation occurs for hydrogen but not for oxygen during plant water uptake. Evaristo et al. (2017) reported that plant water sources quantification methods should consider the possible effects of hydrogen isotope fractionation. Nevertheless, the stable isotopes of δD and δ18O in water still can be used to determine water uptake patterns (Ehleringer et al., 1991; Li et al., 2007; Moreno-Gutiérrez et al., 2015; Swaffer et al., 2014; Yang and Fu, 2017). Although previous studies on water uptake patterns have been conducted in various ecosystems (Eggemeyer et al., 2009; Liu et al., 2010; Schwendenmann et al., 2015), there are still uncertainties regarding plant water uptake patterns (Eggemeyer et al., 2009; Yang et al., 2015). Only soil samples of 0–100 cm depths were collected and extracted as the main soil water sources in most studies (Meißen et al., 2012; Prechsl et al., 2015; Yang et al., 2015). This may underestimate the proportional contribution of soil water to plant water sources and correspondingly overestimate that of other water sources (e.g., spring, ground water, fog), particularly for shrubs and trees with deeper roots. Therefore, including deep soil sampling is necessary to improve the accuracy of water uptake quantification from varied sources and understanding of plant–water interactions (Querejeta et al., 2007; Wu et al., 2014).

The ‘Grain for Green’ project that was implemented in the Chinese Loess Plateau in 1999 is the largest vegetation restoration project in the world (Chen et al., 2015). Trade-offs between vegetation recovery and water depletion has attracted extensive attention (Chen et al., 2015; Jian et al., 2015; Wang et al., 2010). Unsuitable species selection for vegetation restoration leads to soil desiccation, vegetation degradation and difficulties in renewal and reforestation (Chen et al., 2008; Jian et al., 2015; Wang et al., 2010). Although some studies explored soil water migration (Yang and Fu, 2017) and plant water use strategy for natural and planted shrub (Lü et al., 2017), the difference between species and seasonal variations of water uptake pattern remain poorly understood in this region.

In this study, we examined the ecological plasticity of three plant species, Stipa bungeana (herb), Artemisia gmelinii (subshrub) and Vitex negundo (shrub) in terms of the water uptake patterns across the vertical soil profile during the growing seasons by using the isotopic technique. We hypothesized that the three species have different ecological plasticity in plant water uptake. The objectives of this study were to: (i) investigate the isotopic compositions of soil water and their vertical gradients along the soil profile, (ii) quantify the seasonal variations in water uptake patterns and identify their differences among the three species.

2. Materials and methods

2.1. Study area

This study was conducted in the Yangjuangou catchment in the central region of the Loss Plateau in Shanxi Province, China (36°42′45″ N, 109°31′45″ E). This catchment is a typical Loess hilly and gully region with a total area of 2.02 km². The elevation is between 1050 and 1298 m, and the slope gradients range from 10 to 30° (Liu et al., 2012). This area is influenced by the semi-arid continental climate. The annual average precipitation and air temperature from 1960 to 2016 were 537 mm and 10 °C, respectively (Fig. 1). Approximately 80% of the precipitation is concentrated between May and September, with large inter-annual variability. The growing season for most plant species starts in late April and ends by early October. Droughts frequently occur in spring and early summer because there is little precipitation during this period (Liu et al., 2012; Wang et al., 2012). The soil in the study area was loessial soil with the texture of silty loam, consisting of 25% clay (<0.002 mm), 55% silt (0.002–0.05 mm) and 20% sand (>0.05 mm) (Li et al., 2003). The soil is vulnerable to erosion because of weak resistance. The soil in the study area was derived from loss, and the soil depth is usually 50–200 m depending on the topography
Renewal grass and usually occurs in middle or late successional stage during natural recovery. It usually occurs in middle or late successional stage during natural recovery. The Loess Plateau. Five plots of 2 m × 2 m dominated by hilly and gully region of the Loess Plateau. It usually occurs in middle successional stage during natural recovery.

2.2. Experimental design and sample collection

Soil, plant and rainwater samples were collected once a month from May to September 2016. *S. bungeana* is a widely distributed native perennial grass and usually occurs in middle or late successional stage during natural recovery from abandoned cropland. *A. gmelinii* is a perennial subshrub (e.g., a low-growing perennial plant having a woody lower stem) and a primary constructive species of natural grasslands on the hilly and gully region of the Loess Plateau. It usually occurs in middle successional stage during natural recovery. *V. negundo* is a perennial deciduous shrub that has been widely planted in ecological restoration on the Loess Plateau. Five plots of 2 m × 2 m dominated by *S. bungeana* and *A. gmelinii*, respectively, and three plots of 5 m × 5 m dominated by *V. negundo* were selected for study (Table 1). These plots for each species were close to each other to ensure that microenvironment of each plot was consistent, including soil condition, micro-geomorphology. The average basal diameter of *V. negundo* was 13.00 mm. Three individuals of *V. negundo* were randomly chosen for plant sampling from three plots on each sampling date. Each plant sample was collected from four canopy directions at a height of approximately 100 cm above the ground. The phloem tissue of *V. negundo* was removed to avoid isotopic fractionation of xylem water and contamination by isotopic enriched water (Dawson, 1996; Querejeta et al., 2007). For *S. bungeana* and *A. gmelinii*, 8–10 individuals were selected as a sample, and three samples were collected from one plot on each sampling date. The root crown of each plant was collected from the surface soil because it correlates with the source water (Barnard et al., 2006; Wu et al., 2016). All of the plant samples were cut into 3–4 cm segments, immediately placed into glass vials with screw caps, sealed with polyethylene parafilm and kept frozen in a freezer (−20 °C) for isotopic analysis. A total of 45 plant samples for three species were collected during the study periods.

Soil samples were collected simultaneously with plant tissue sampling. Three soil cores were taken from three plots for *V. negundo* and one plot for *S. bungeana* and *A. gmelinii* on each sampling date at depths of 0–5 cm, 5–10 cm, 10–20 cm, every 20 cm in the 20–120 cm range, every 40 cm in the 120–200 cm range, and at 50 cm intervals in the 200–300 cm layers. A power auger with the petrol engine-driven post driver (CHPD78, Christie Engineering Company, Sydney, Australia) was used to collect soil samples. Compared with the traditional hand auger, the post driver auger can obtain less compacted soil samples, improving the reliability of soil sample data. Soil samples were divided into two: one portion was stored in a freezer for isotopic analysis and the other was used to obtain gravimetric soil water content (SWC, %) as determined by drying at 105 °C for 24 h. In addition, we investigated the root distributions of the three species by excavating soil layers per 10 cm until no root was found in the soil at the end of the growing season. The excavating depth of *S. bungeana*, *A. gmelinii* and *V. negundo* was 60 cm, 100 cm, 200 cm, respectively. Root surface area (m²/m³) was analyzed by WinRHIZO system (Regent Instruments, Quebec, Canada).

A polyethylene bottle and funnel were used to collect rainwater. A plastic ball was placed on the funnel to prevent evaporation (Yang et al., 2015). The rainwater samples were collected immediately after the rainfall terminated once a month from May to September and a total of 46 rainwater samples were taken. Similarly, rainwater samples were put into polyethylene bottles, wrapped with parafilm and stored in a freezer until subsequent isotopic analysis.

### Table 1

Basic information of sampled plant species.

| Species        | Family     | Life form     | Height (m) (mean ± SD) | Leaf phenology | Vegetation coverage (%) |
|----------------|------------|---------------|------------------------|----------------|-------------------------|
| *S. bungeana*  | Poaceae    | Tussok grass  | 0.2 ± 0.05             | Perennial herb | 59–65                   |
| *A. gmelinii*  | Compositae | Subshrub      | 0.6 ± 0.08             | Deciduous      | 61–68                   |
| *V. negundo*   | Verbenaceae| Large shrub   | 1.5 ± 0.10             | Deciduous      | 62–70                   |

![Fig. 1](image-url) The mean rainfall, temperature pattern and isotopic composition in rainwater: (a) the monthly variation of multi-year precipitation (1960–2016), temperature and precipitation in 2016; (b) the temporal variation of isotopic compositions in rainwater.
water from plant tissues using cryogenic vacuum distillation can co-distill organic materials (e.g., methanol and ethanol) that may affect the spectroscopy and lead to erroneous stable isotope values when analyzing with IRIS. However, trace amounts of organic matter did not affect the measurement results of IRMS method (West et al., 2010). The δ2H and δ18O for water extracted from soil samples (n = 18) and rainfall water (n = 5) were measured with both IRIS and IRIS. The results showed that the mean discrepancies between IRMS and IRIS were 0.34‰ for δ2H and 0.06‰ for δ18O in water from soil samples, and 0.34‰ for δ2H and 0.08‰ for δ18O in rainfall water. There was no significant difference between IRMS and IRIS for δ2H (p = 0.93 > 0.05) and δ18O (p = 0.93 > 0.05) in soil water and rainfall water. No contamination of organic matter was found in soil water and rainfall water after quantifying using the spectral contamination identifier post-processing software (Los Gatos Research Inc., Mountain View, CA, USA) (Schultz et al., 2011; Zhao et al., 2011). Schultz et al. (2011) also found there was no contamination observed in soil water measurements analyzed with IRIS. In addition, previous studies reported that no discrepancies between the isotopic compositions of soil water measured using both IRIS and IRMS methods (Schultz et al., 2011; Zhao et al., 2011). Thus, we supposed that isotopic ratios of the two measurement methods would not affect our subsequent analyses. The calculation of isotopic ratios was expressed as follows:

\[ \delta X = \frac{R_{\text{sample}}}{R_{\text{standard}}} - 1 \]  

where \( X \) represented δ2H or δ18O, \( R_{\text{sample}} \) and \( R_{\text{standard}} \) represented the molar abundance ratios \( \delta(2H/1H) \) and \( \delta(18O/16O) \), respectively, of the sample and standard (Standard Mean Ocean Water), respectively.

The weighted averages of \( \Delta \delta2H \) and \( \Delta \delta18O \) for precipitation per month (\( \delta_{p,\text{mean}} \)) during the sampling dates were calculated as:

\[ \delta_{p,\text{mean}} = \frac{\sum_{i=1}^{n} \delta_i \times PPT_i}{\sum_{i=1}^{n} PPT_i} \]  

where \( \delta_i \) and \( PPT_i \) represented the isotopic composition and amount of each precipitation event, respectively.

2.4. MixSIAR model

Groundwater was not used as a potential water resources because the table depth of groundwater in the Loess Plateau is buried deeply (40–100 m below soil surface), precluding any upward capillary flow into the root zone (Chen et al., 2008; Chen et al., 2007; Huang et al., 2013). The roots of plants had difficulty reaching the groundwater. Moreover, there was no surface runoff because of the high infiltration and no irrigation in the study area. Thus, soil water from different layers was regarded as the main potential resources for plant water uptake.

To identify the depths of water used by plants, stable isotopes of xylem water were compared with those of potential water sources, and the intersections between xylem water and potential water sources were detected. The intersection of the xylem water isotopic vertical line and the soil water isotopic profile was used to determine the main depth of the plant water uptake (Brunel et al., 1995; Yang et al., 2015). This graphical inference method was used as a preliminary judgment on water uptake depths.

The fractions of water sources used by plants was estimated by the Bayesian isotope mixing model MixSIAR (version 3.1.7), which incorporates uncertainties associated with multiple sources, discrimination factors (Stock and Semmens, 2013). The raw xylem isotope values \( \delta2H \) and \( \delta18O \) of three plant species were used as mixture data inputs into MixSIAR. The source data had no concentration dependence. Individual effects as a random occurrence were included in all analyses. The discrimination data were set to zero for both \( \delta2H \) and \( \delta18O \) because isotopic fractionation did not occur during plant water uptake (Brunel et al., 1995; Dawson and Ehleringer, 1991). The run length of the Markov chain Monte Carlo (MCMC) was set to ‘long’ (chain length = 300,000; burn = 200,000; thin = 100; chains = 3). MCMC was used to converge on the posterior distributions of all the variables in the model. It was essential to determine that the model had converged before accepting the output of MixSIAR. Gelman–Rubin and Geweke diagnostic tests were used to determine whether the model was close to convergence (Stock and Semmens, 2013). ‘Residual only’ was specified as the error structure in the model. The uninformative prior was set in the model. Water uptake patterns did not necessarily match the root distribution (Thorburn and Ehleringer, 1995) unless sufficient water in shallow soil layers generated by rainfall met vegetation requirements (Knight, 1999; Liu et al., 2011). Previous study found that the maximum water utilization depth of S. bungeana and A. gmelinii could reach 300 cm in the study area (Fan et al., 2016; Wang et al., 2002). V. negundo had larger lateral root and only considered the vertical water use patterns. Moreover, loessial soil is characterized with porous and homogeneous soil water retention (Huang et al., 2003). Thus, soil water sources in each soil layer were considered equal likelihood to plant water sources. The median values (50% quartiles) were presented as the predictions of MixSIAR. The water sources from different soil layers were combined into three larger layers (0–40, 40–120 and 120–300 cm) to facilitate the subsequent analysis and comparison (Eggemeyer et al., 2009). Three layers were identified by the following (Wu et al., 2016):

1. Shallow soil layer (0–40 cm): the isotopic ratios in soil water and SWC were larger variability and were vulnerable to rainfall pulse and evaporation with season.
2. Middle soil layer (40–120 cm): the isotopic ratios in soil water and SWC were lower variations and milder monthly changes than that in the shallow soil layer.
3. Deep soil layer (120–300 cm): the isotopic ratios in soil water and SWC showed relatively stable variations.

2.5. Data analyses

In order to make stable isotope analysis more reasonable, two assumptions were made in this study (Brunel et al., 1995; Liu et al., 2011): (1) the isotopic composition in soil water was laterally homogeneous within the rooting area; and (2) the time delay between sampling and ascent of water transport in plant was not significant.

The Kolmogorov–Smirnov (K–S) test showed that the isotopic ratios of plant tissues and soil water met normal distribution. A two-way analysis of variance was conducted to detect the differences in the isotopic ratios of plant tissues between sampling dates and species. A one-way analysis of variance followed by post-hoc Tukey’s least significant difference test (p < 0.05) was used to detect the differences in the isotopic compositions of soil water among the different depths for each species. All of the statistical analyses were conducted in SPSS 20.0 (SPSS Inc., Chicago, IL, USA).

3. Results

3.1. Precipitation distribution and isotopic composition

The total precipitation was 478 mm in the year of study (2016) and 91.3% occurred during the growing season. In addition, the highest monthly precipitation was 184.40 mm in July, accounting for 38.94% of the year’s total precipitation. The rainfall amount in 2016 was 24.40% greater than that of 2015 and 11.79% less than that of the multi-year mean in this region. The monthly variation was consistent with that of the multi-year mean (1960–2016) (Fig. 1).
The isotopic composition of the rainwater presented a large fluctuation across the sampling dates. The δ¹⁸O ranged from −1.37 to −12.03‰, with an average of −5.40‰, and δ²H ranged from −13.06 to −90.25‰, with a mean value of −36.92‰ (Fig. 1). The average monthly precipitation-weighted values of δ²H and δ¹⁸O were −40.32‰ and −6.29‰, respectively. The distribution of isotopic values of the monthly weighted precipitation amounts during the sampling periods was bimodal with two peaks appearing in June and August (Fig. 2). For the greatest amount of precipitation, which occurred in July, the monthly weighted rainwater isotopic composition was −62.03‰ for δ²H and −8.83‰ for δ¹⁸O.

The local meteoric water line (LMWL) was fitted based on the precipitation data (Fig. 3). The slope and intercept of the LMWL were less than those of the global meteoric water line (GMWL) (Craig, 1961), indicating that the rainfall enriched isotopic composition was caused by evaporation during the rainfall. Almost all of the isotopic values of the soil water from the three species' plots were plotted to the right of the LMWL, implying that soil water was derived from the precipitation and underwent enrichment associated with evaporation in this area. The δ²H and δ¹⁸O of soil water from V. negundo plots were steeper compared with those of the two other species. The co-variation between δ²H and δ¹⁸O in the soil waters from S. bungeana and A. gmelinii plots were similar during the sampling periods (Fig. 3).

3.2. Isotopic composition and variation in xylem water

The isotopic values in xylem water varied among plant species and sampling periods. S. bungeana had the highest isotopic values, which ranged from −34.22 to −66.52‰ for δ²H, with an average of −51.47‰, and δ¹⁸O ranged from −3.23 to −8.08‰, with an average of −5.84‰ during the sampling periods. The average isotopic compositions for A. gmelinii and V. negundo were −56.22 and −58.95‰ for δ²H, respectively, and −7.15 and −6.22‰ for δ¹⁸O, respectively. Averaging xylem water isotopic values over all sampling dates showed no significantly different (p > 0.05) among the three species, implying that all of the species absorbed water from similar soil layers. The isotopic compositions of xylem water of the three species were significantly different (p < 0.01) among sampling dates, suggesting that plant water uptake had significant temporal variability. Both δ²H and δ¹⁸O in the xylem water from V. negundo gradually decreased from May to August and increased in September. The trend of change in δ¹⁸O was consistent with that of δ²H in S. bungeana, in which the greatest values were in June and the lowest values were in August (Fig. 2). The isotopic compositions of S. bungeana and A. gmelinii were distributed close to those of soil water (Fig. 3), indicating that the two plants primarily obtained water from various soil horizons. However, the isotopic values of V. negundo were outside the soil mixing space, implying that hydrogen isotope fractionation may occur during water uptake. The isotopic values of xylem water were within the range of isotopic ratios in soil water from 0 to 120 cm soil layers (Fig. 3).

3.3. Soil moisture and isotopic composition

The SWC values in the plots of S. bungeana and A. gmelinii displayed similar temporal variations in each of the soil depths (Fig. 4). The SWC increased with soil depths above 40 cm soil horizons in May and August. However, the SWC exhibited the opposite trend in June, July and September. The SWC decreased from depths over 40 cm in the V. negundo plot, except for in August. The SWC from shallow soil layers (0–40 cm) was highly variable compared with those of the other layers during the sampling periods. In contrast, the SWC from the deeper soil layers (120–300 cm) was relatively stable and showed no distinct differences among species and sampling dates.

The isotopic composition of the soil water changed with the soil depths and plant species (Figs. 5, 6). The δ¹⁸O of soil water in the three species plots were −8.53 ± 2.18‰ for S. bungeana, −8.52 ± 1.80‰ for A. gmelinii, and −8.86 ± 1.85‰ for V. negundo. The δ²H of the soil water in the S. bungeana, A. gmelinii and V. negundo plots were −64.78 ± 14.49‰, −63.40 ± 11.81‰ and −66.38 ± 13.50‰, respectively. The isotopic ratios in soil water of the three species plots showed no significant differences (p > 0.05). The isotopic compositions in the soil water became more negative as the soil depth increased over the sampling period. During the sampling period, except for August, the shallow soil (0–40 cm) water had higher isotopic values and variances than those of middle (40–120 cm) and deep (120–300 cm) soil water. For A. gmelinii and V. negundo plots, both δ²H and δ¹⁸O in soil water were significantly different in shallow (p < 0.001), middle (p < 0.001) and deep (p < 0.001) soil layers during the sampling period. However, the isotopic compositions of the soil water in S. bungeana plots had significant differences, except for between shallow and middle soil water isotopes (p > 0.05).

3.4. Variations in the proportion of plant water uptake

According to the graphical inference method, the three plant species derived most of their water from the 0–40 cm soil layers during the sampling period. The isotopic ratios of xylem water and soil water overlapped at several soil depths during July and August (Figs. 5, 6), implying that plants might derive water from several depths. Nevertheless, this graphical inference method only provided the main uptake depth and may lead to erroneous interpretation when plan water sources derived simultaneously from several soil layers.

The MixSIAR model predicted that the water uptake fraction of S. bungeana was 49.32 ± 5.65% for shallow soil, 30.22 ± 4.81% for middle soil and 20.56 ± 5.10% for deep soil during the sampling period. Although S. bungeana mainly obtained water from shallow soil during the growing season, the proportional contribution of soil water varied over the sampling times. The fraction of water uptake from shallow soil was the greatest in May (55.20 ± 16.23%) and the lowest in August (40.00 ± 8.50%) (Fig. 7). The proportions of water that S. bungeana derived from shallow and middle soils over the sampling periods showed no significant differences (p > 0.05). The proportions of water that A. gmelinii extracted from shallow, middle, deep soils during the sampling periods were 57.08 ± 13.20%, 22.86 ± 5.68% and 20.04 ± 0.08%, respectively. In July, the shallow soil contributed the most water (74.20 ± 15.20%), and the largest proportion of water (33.10 ± 7.05%) was absorbed from deep soils in August (Fig. 7). Compared with A. gmelinii, V. negundo obtained a relative lower proportion of...
water from shallow soils (49.24 ± 14.14%) during the sampling dates. However, *V. negundo* absorbed the largest proportion of soil water (24.62 ± 5.56%) from 120 to 300 cm soil layer during the sampling periods. The percentage of water from deep soil increased from 14.50% in May to 42.40% in August (Fig. 7). For both *A. gmelinii* and *V. negundo*, the proportional contribution of soil water had significant difference from shallow (p < 0.05), middle (p < 0.05) and deep soil (p < 0.001) over the sampling periods.

### 4. Discussion

#### 4.1. Vertical gradient of isotopic composition in soil water

The isotopic vertical gradient of soil water was primary influenced by two processes, evaporation and infiltration, simultaneously, with the mixing of old and new water from rainwater. The δ²H and δ¹⁸O in soil water became more depleted as the soil depth increased. The
isotopic compositions in surface soil water were enriched as compared to deeper soil profiles in May, June and September (Figs. 5 and 6). This was attributed to less rainfall and more evaporation from the surface soil horizons (Gazis and Feng, 2004; Geris et al., 2017). Rainwater mixed with heavy isotope water in the surface soil and infiltrated through the soil matrix (Mathieu and Bariac, 1996; Brooks et al., 2009). However, this trend was reversed in July in surface soil because of the influence of recent rainfall having negative isotopic values, suggesting that precipitation was also an important factor controlling the isotopic compositions of soil water. The isotopic ratios of surface soil displayed larger variances than those of deep soil as the combined influences of evaporation and precipitation. Our findings were agreement with previous studies in arid and semi-arid ecosystems (Gazis and Feng, 2004; Wu et al., 2016).

Nevertheless, the δ2H and δ18O in water from 60 to 300 cm soil layers had less variation with both depths and sampling dates (Figs. 5 and 6). This pattern was consistent for the three species. The effect of evaporation was limited in deep soil and the groundwater was deep at the study site. Thus, the rainwater moved through soil pores that bypassed superficial soil in the soil profile (Eggemeyer et al., 2009; Gazis and Feng, 2004) and subsequently mixed with the stationary soil water and resulted in the vertical variation. A similar result was identified in a previous study (Brooks et al., 2009) in which the isotopes of the deep soil water were affected by infiltration mixed new and old water through preferential flow paths. In our study, there was no significant difference in soil water isotopes among three species, indicating that vegetation types had no influence on the isotopic ratios of soil water, because both plant water uptake and transpiration do not cause the isotopic fractionation of water (Brunel et al., 1995; Ehleringer and Dawson, 1992; Allison et al., 1983).

4.2. Differences in seasonal water uptake patterns among species

The isotopic compositions of xylem water in three species approached that of soil water from 0 to 120 cm soil (Fig. 3), suggesting that the potential water sources absorbed by plants derived mostly from 0 to 120 cm soil during the sampling periods. The MixSIAR outcomes indicated that S. bungeana derived 80% of its water from 0 to 120 cm soil during the growing season, and utilized the largest proportion (85%) from 0 to 120 cm soil in July. In the semi-arid grasslands of Nebraska (USA), Eggemeyer et al. (2009) found that grasses predominantly utilized water from the upper soil layers (5–50 cm). The water uptake pattern of S. bungeana displayed the smallest fluctuation during the growing season, indicating a low level of plasticity in water source utilization.

Compared with S. bungeana, A. gmelinii extracted more water (~60%) from shallow soil during the sampling period. The proportion of water absorbed by A. gmelinii from 0 to 120 cm soils showed no significant differences (p = 0.25 > 0.05) during the sampling periods. However, the proportions of water uptake from deep soils over the sampling dates presented significant differences (p < 0.001). A. gmelinii absorbed the largest proportion of water from shallow soil in July (74.2%) (Fig. 7), indicating that this may be related to precipitation because of precipitation peak in July (Fig. 1), which resulted in ample surface soil water at this time. This result was similar to that of the previous study (Wu et al., 2016) in which subshrubs primarily relied on water from 0 to
Fig. 6. Seasonal variations of $\delta^{18}O$ in soil horizons (0–300 cm) and xylem water (vertical dash) from *S. bungeana*, *A. gmelinii*, and *V. negundo*. Error bars represent standard deviation (N = 3).

Fig. 7. Seasonal variations in proportion of water uptake from different soil layers based on MixSIAR for *S. bungeana*, *A. gmelinii*, and *V. negundo*. Error bars represent standard deviation (N = 3).
30 cm soils throughout the growing season in a water limited ecosystem owing to the extensive horizontal root area. 

Asbjornsen et al. (2008) observed that two herbaceous species predominantly utilized water from 0 to 20 cm soil horizons and exhibited a relatively low range of ecological plasticity in water uptake. However, Lü et al. (2017) found that A. gmelinii absorbed water from 0 to 40 cm soil when SWC was sufficient, but from 40 to 80 cm when the shallow soil was dry. In contrast, V. negundo derived the largest proportion of soil water from deep soil among the three species and progressively increased the contributory proportion of deep soil water during the growing season (Fig. 7). It displayed the greatest degree of plasticity in water uptake. This pattern may be attributed to the gradual increase in transpiration during the continuous growing season. Similar results were identified in other ecosystems (Asbjornsen et al., 2008) in which woody shrubs progressively increased their water uptake depths over the growing season.

Many plants species in arid and semi-arid ecosystems possess a functionally dimorphic root system, which has a zone of lateral roots that derive water from deep soil layers when surface water is unavailable, and another zone, which predominantly acquires water from surface soil water. Dawson and Pate (1996; Nie et al., 2010) reported that the maximum water uptake depths of S. bungeana on abandoned land in an erodable small watershed on the Loess Plateau was 300 cm. In contrast, V. negundo possessed a deeper root system (Fig 9) and was able to extract water from deep soil layers. Deep soil water contributed 22% of the total plant water sources for three species over the sampling periods. Deeper roots increased the tolerance to drought by increasing the probability to absorb deeper water sources when shallow soil water was low (Ellsworth and Sternberg, 2015; Markesteijn and Poorter, 2009). Moreover, some ecological characters might enable plant species exhibited specific water uptake patterns in specific environment (Nie et al., 2010; Swemmer et al., 2006), such as more rapid recovery of photosynthesis after wilting, more rapid leaf turnover, greater allocation to roots, reduced allocation to flowering.

4.3. Implications

Although there is relatively low precipitation in arid and semi-arid regions, individual rainfall events cause water source enrichments in the short-term, which play important roles in shaping plant adaptation in water use strategies (Schwinning and Ehleringer, 2001). Plant species that switch water uptake patterns can avoid interspecific competition with shallow rooted species at the same site and have a greater probability of survival owing to their improved competitiveness by adapting to droughts in the semi-arid regions (Ehleringer and Dawson, 1992).

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**Fig. 8.** Picture of field investigation for V. negundo roots. The view of the entire root system (fine roots were invisible) (a) a zoom-in view for lateral and tap roots (b).

**Fig. 9.** Vertical distribution of root surface area (m²/m³) for S. bungeana, A. gmelinii, and V. negundo.
Some deciduous species adapted to droughts by different strategies. One was to avoid drought by losing their leaves and remaining dormant in the dry season (Ellsworth and Sternberg, 2015), and another was to switch water sources absorbed by plants depending on water available (Wu et al., 2016; Liu et al., 2010). V. negundo was deciduous woody species and progressively switched its water source during the growing season. This flexible water use strategy will be more advantageous under the predicted increasingly frequent drought events of the future and will promote population growth and the plant’s survival probability. S. bungeana and A. gmelinii mainly derived water from surface soil layers and had lower variation levels during the whole growing season. This demonstrated that shallow-rooted plant species, especially A. gmelinii, were sensitive to precipitation. The roots were beneficial for the plant to make better use of precipitation. Extracting shallow water in semi-arid ecosystems might exhibit a high tolerance for limited water source environments, which is dependent on the ability of fine roots to quickly switch absorption patterns in response to precipitation pulses and water shortages (Asbjørnsen et al., 2008). The combinations of different water uptake patterns of plant species might be more conducive to ecological restoration. In this manner, V. negundo was conducive to being combined with S. bungeana or A. gmelinii. In addition, the variation in water uptake patterns could lead to ecological niches separation complementary uses of resources and, in turn, facilitate species coexistence and ecosystem functions (Asbjørnsen et al., 2008). Stable isotope technology provided insights into plant species selection and management during ecological restoration in the arid and semi-arid ecosystems. More physiological and ecological characteristics coupled with the continuous monitoring of stable isotopic ratios may provide further insights into water uptake patterns.

5. Conclusions
The MixSIAR model based on dual stable isotopes δ18H and δ18O was applied to determine the seasonal variations of water uptake patterns of a grass (S. bungeana), a subshrub (A. gmelinii) and a shrub (V. negundo) on the Chinese Loess Plateau. The isotopic compositions of the shallow soil water were more enriched and had more variation than that of the deep soil water. The MixSIAR model predicted that S. bungeana and A. gmelinii predominately derived 80% of water from 0 to 120 cm soil layers and displayed a relatively low level of variation during the growing season. However, V. negundo exhibited a distinct water uptake pattern that progressively switched its water source from shallow to deep soil horizons as the season progressed, indicating a greater degree of ecological plasticity. This was mainly attributed to the functionally dimorphic root system. The flexible water uptake patterns of plants might facilitate plant species making full use of water and nutritional sources, adapting to the changeable environment. This study provides a useful method that identified partitioning and proportional contributions of water resources, and the research results will be significant for vegetation restoration in the arid and semi-arid ecosystems.

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References
Allison, G., Barnes, C., Hughes, M., Leahey, F., 1983. Effect of climate and vegetation on oxygen-18 and deuterium profiles in soils. Isot. Hydrox. 105–123.
Ashjornsen, H., Shepherd, G., Helmers, M., Mora, G., 2008. Seasonal patterns in depth of water uptake under contrasting annual and perennial systems in the Corn Belt Region of the Midwestern U.S. Plant Soil 308, 69–92.
Barnard, R.L., de Bello, F., Gilgen, A.K., Buchmann, N., 2006. The δ18O of root crown water best reflects source water δ18O in different types of herbaceous species. Rapid Commun. Mass Spectrom. 20, 3790–3802.
Brand, W.A., Geilmann, H., Crosson, E.R., Relia, C.W., 2009. Cavity ring-down spectroscopy versus high-temperature conversion isotope ratio mass spectrometry: a case study on δ18O and δ13C of pure water samples and alcohol/water mixtures. Rapid Commun. Mass Spectrom. 23, 1879–1884.
Brooks, J.R., Barnard, H.R., Coulombe, R., McDonnell, J.J., 2009. Ecohydrologic separation of water between trees and streams in a Mediterranean climate. Nat. Geosci. 3, 103–104.
Brunel, J.-P., Walker, G.R., Kenneth-Smith, A.K., 1995. Field validation of isotopic procedures for determining sources of water used by plants in a semi-arid environment. J. Arid Environ. 32, 351–368.
Chen, L., Huang, Z., Gong, J., Fu, B., Huang, Y., 2007. The effect of land cover/vegetation on soil water dynamic in the hilly area of the loess plateau, China. Catena 70, 205–222.
Dawson, T.E., 1996. Determining water use by trees and forests from isotopic energy balance and transpiration analyses: the roles of tree size and hydraulic lift. Tree Physiol. 16.
Dawson, T.E., Ehleringer, J.R., 1991. Streamside trees that do not use stream water. Nature 351, 335–337.
Dawson, T.E., Pate, J.S., 1996. Seasonal water uptake and movement in root systems of Australian phreatophytic plants of dimorphic root morphology: a stable isotope investigation. Oecologia 107, 13–20.
Delzon, S., Loustau, D., 2005. Age-related decline in stand water use: sap flow and transpiration in a pine forest chronosequence. Agric. For. Meteorol. 129, 105–119.
Eggenberger, K.D., Awada, T., Harvey, F.E., Wedin, D.A., Zhou, X., Zanner, C.W., 2009. Seasonal changes in depth of water uptake for encroaching trees Juniperus virginiana and Pinus ponderosa and two dominant C4 grasses in a semiarid grassland. Tree Physiol. 29, 157–169.
Ehleringer, J., Dawson, T., 1992. Water uptake by plants: perspectives from stable isotope composition. Plant Cell Environ. 15, 1073–1082.
Ehleringer, J.R., Phillips, S.L., Schuster, W.S., Sandquist, D.R., 1991. Differential utilization of summer rains by desert plants. Oecologia 88, 430–434.
Ellsworth, P.Z., Sternberg, L.S., 2015. Seasonal water use by deciduous and evergreen woody species in a scrub community is based on water availability and root distribution. Ecosystems 8, 538–551.
Ellsworth, P.Z., Williams, D.G., 2007. Hydrogen isotope fractionation during water uptake by woody xerophytes. Plant Soil 291, 93–107.
Evaristo, J., McDonnell, J.J., 2017. Prevalence and magnitude of groundwater use by vegetation: a global stable isotope meta-analysis. Sci Rep 7, 4110.
Evaristo, J., McDonnell, J.J., Scholl, M.A., Bruijnzeel, L.A., Chian, K.P., 2016. Insights into plant water uptake from xylem-water isotope measurements in two tropical catchments with contrasting moisture conditions. Hydrox. Process. 30, 3210–3227.
Evaristo, J., McDonnell, J.J., Clemens, J., 2017. Plant source water apportionment using stable isotopes: a comparison of simple linear, two-compartment mixing model approaches. Hydrox. Process. http://dx.doi.org/10.1002/hyp.11233.
Fan, J., Wang, Q., Jones, S.B., Shao, M., 2016. Soil water depletion and recharge under different land cover in China’s Loess Plateau. Ecol. Hydrol. 119, 4068–4086.
Gao, X., Wu, P., Zhao, X., Shi, Y., Wang, J., Zhang, B., 2011. Soil moisture variability along transects over a well-developed gulley in the Loess Plateau, China. Catena 87, 39–47.
Gazi, C., Feng, X., 2004. A stable isotope study of soil water: evidence for mixing and preferring flow paths. Geoderma 119, 97–111.
Geris, J., Tetzlaff, D., McDonnell, J.J., Soulsby, C., 2017. Spatial and temporal patterns of soil water storage and vegetation water use in humid northern catchments. Sci. Total Environ. 595, 486–493.
Grossiord, C., Sevanto, S., Dawson, T.E., Adams, H.D., Collins, A.D., Dickman, L.T., et al., 2017. Warming combined with more extreme precipitation regimes modifies the water sources used by trees. New Phytol. 213, 584–596.
Hardanto, A., Röll, A., Hölscher, D., 2017. Tree soil water uptake and transpiration in mono-cultural and jungle rubber stands of Sumatra. For. Ecol. Manag. 397, 67–77.
Howard, J., Merrifield, M., 2018. Mapping groundwater dependent ecosystems in California. PLoS One 5, e11249.
Huang, M., Dang, T., Gallichand, J., Goulet, M., 2003. Effect of increased fertilizer application to wheat crop on soil-water depletion in the Loess Plateau, China. Agric. Water Manag. 58, 267–278.
Huang, T., Pang, Z., Edmunds, W.M., 2013. Soil profile evolution following land-use change implications for groundwater quantity and quality. Hydrox. Process. 27, 1238–1252.
Jian, S., Zhao, C., Fang, S., Yu, K., 2015. Effects of different vegetation restoration on soil water storage and water balance in the Chinese Loess Plateau. Agric. For. Meteorol. 206, 83–96.
Knight, J.H., 1999. Root distributions and water uptake patterns in Eucalypts and other species. In: Landsberg, J. (Ed.), The Ways Trees Use Water. RRDC, AU, pp. 55–85.
Li, Y., Xoosae, J., Yang, J., Fu, R., Zhang, J., 2003. Evaluating gully erosion using 127Cs and 210Pb/137Cs ratio in a reservoir catchment. Soil Tillage Res. 69, 107–115.
Li, S.G., Romero-Salton, H., Tsijimura, M., Sugimoto, A., Sasaki, L., Davaa, G., et al., 2007. Plant water sources in the cold semiarid ecosystem of the upper Kherlen River catchment in Mongolia: a stable isotope approach. J. Hydrox. 33, 109–127.
Lin, G.H., Sterner, L.S., 1993. Hydrogen isotope fractionation by plant roots during water uptake in coastal wetland plants. Stable Isotopes and Plant Carbon/ Water Relations. Academic Press, New York, pp. 497–510.
Liu, W., Liu, W., Li, P., Duan, W., Li, H., 2010. Dry season water uptake by two dominant canopy tree species in a tropical seasonal forest of Xishuangbanna, SW China. Agric. For. Meteorol. 150, 380–388.

Liu, Y., Xu, Z., Duffy, R., Chen, W., An, S., Liu, S., et al., 2011. Analyzing relationships among water uptake patterns, rootlet biomass distribution and soil water content profile in a subalpine shrubland using water isotopes. European Journal of Soil Biology 47, 380–386.

Liu, Y., Fu, B., Lü, Y., Wang, Z., Gao, G., 2012. Hydrological responses and soil erosion potential of abandoned cropland in the Loess Plateau, China. Geomorphology 138, 39–48.

Lü, T., Zhao, X., Gao, X., Pan, Y., 2017. Soil water use strategy of dominant species in typical natural and planted shrubs in loess hilly region. Chin. J. Plant Ecol. 41, 175–185.

Ma, Y., Song, X., 2016. Using stable isotopes to determine seasonal variations in water uptake of summer maize under different fertilization treatments. Sci. Total Environ. 550, 471–483.

Mares, R., Barnard, H.R., Mao, D., Revil, A., Singka, K., 2016. Examining diel patterns of soil and xylem moisture using electrical resistivity imaging. J. Hydrol. 536, 327–338.

Markesteijn, L., Poorter, L., 2009. Seedling root morphology and biomass allocation of 62 tropical tree species in relation to drought- and shade-tolerance. J. Ecol. 97, 311–325.

Mathieu, R., Bariac, T., 1996. An isotopic study (2H and 18O) of water movements in clayey soils under a semiarid climate. Water Resour. Res. 32, 779–789.

McCole, A.A., Starn, L.A., 2007. Seasonal water use patterns of Juniperus ashei on the Edwards Plateau, Texas, based on stable isotopes in water. J. Hydrol. 342, 238–248.

Meißner, M., Kühl, M., Schwendemann, L., Hilscher, D., 2012. Partitioning of soil water among canopy trees during a soil desiccation period in a temperate mixed forest. Biogeosciences 9, 3465–3474.

Moreno-de Las Heras, M., Espigares, T., Merino-Martín, L., Nicolau, J., 2011. Water-related ecological impacts of rill erosion processes in Mediterranean-dry reclaimed slopes. Catena 84, 114–124.

Moreno-de Las Heras, M., Battaglia, G., Cherubini, P., Delgado Huertas, A., Querejota, J.J., Thompson, K., 2015. Pine afforestation decreases the long-term performance of understory shrubs in a semi-arid Mediterranean ecosystem: a stable isotope approach. Ecol. Evol. 5, 15–25.

Nie, Y.P., Chen, H.S., Wang, K.L., Tan, W., Deng, P.Y., Yang, J., 2010. Seasonality of hydraulic redistribution by trees to grasses and changes in water uptake depth in response to summer drought of two lowland and sub-alpine thin soils in subtropical China. Plant Soil 341, 399–412.

Porporato, A., Daly, E., Rodriguez-Iiturbe, I., 2004. Soil water balance and ecosystem response to climate change. American Naturalist 164, 625–632.

Prechil, U.E., Burri, S., Gällen, A.K., Kahmen, A., Buchmann, N., 2015. No shift to a deeper water uptake depth in response to summer drought of two lowland and sub-alpine C3(+) grassland species in Switzerland. Oecologia 177, 97–111.

Priyadarshini, K.V.R., Prins, H.H.T., de Bie, S., Heitkötter, I.M.A., Woodborne, S., Gort, G., et al., 2016. Seasonality of hydraulic redistribution by trees to grasses and changes in their water-source use that change tree-grass interactions. Ecology Hydrology 9, 218–228.

Querejota, J.J., Estrada-Medina, H., Allen, M.F., Jimenez-Osornio, J.J., 2007. Water source partitioning among trees growing on shallow karst soils in a seasonally dry tropical climate. Oecologia 152, 26–36.

Rothfuss, Y., Javava, M., 2017. Reviews and syntheses: isotopic approaches to quantify root water uptake: a review and comparison of methods. Biogeosciences 14, 2199.

Schenk, H.J., Jackson, R.B., 2005. Mapping the global distribution of deep roots in relation to climate and soil characteristics. Geoderma 125, 129–140.

Schulz, N.M., Griffith, T.J., Lee, X., Baker, J.M., 2011. Identification and correction of spectral contamination in 1H/2H and 18O/16O measured in leaf, stem, and soil water. Rapid Commun. Mass Spectrom. 25, 3360–3368.