



The dimensions of deep-layer soil desiccation and its impact on xylem hydraulic conductivity in dryland tree plantations

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Abstract. In water-limited areas, planted trees can extract substantial amounts of soil water from deep layers (> 200 cm) to meet their high water demand, resulting in deep soil desiccation, which influences not only regional water cycling but also the sustainability of trees per se in drylands. However, the specific dimensions of deep soil desiccation in relation to both the soil moisture limitation and the maximum root water uptake (RWU) depth are still not well determined. Whether the dimensions depend on tree species and how they will affect trees' xylem hydraulic conductivity are also unclear, restricting our ability to predict the fate of dryland tree plantations. Therefore, we studied the spatiotemporal distribution of deep soil moisture deficit (DSMD) for two typical planted trees, apple (*Malus pumila* Mill.) and black locust (*Robinia pseudoacacia* L.), based on published data and multiple field samplings on China's Loess Plateau. The results indicated that the lowest deep soil moisture (DSM; units of gravimetric percent, i.e., grav-%) occurred under the planted trees aged 24–28 years at all sites. The lowest DSMD varied around -0.6 , which was close to the DSMD at the permanent wilting point (PWP, grav-%), regardless of tree species and site, although shallow (< 200 cm) soil moisture was not reduced to the point of limitation. This suggests that PWP is a reliable indicator of the moisture limitation of deep-layer soil desiccation for

the tree species examined. The corresponding depth of soil moisture use reached 18.0–22.0 m for these old planted trees at different sites, while it was more than 25 m for *R. pseudoacacia* in the drier site of Mizhi. Furthermore, the mean values of native percentage loss of hydraulic conductivity of planted trees' branches xylem reached 74.9 %–96.5 % in the plantations sampled in this study, indicating that tree mortality may occur. The findings help predict the sustainability of planted trees in semi-arid regions with a thick vadose zone.

1 Introduction

Worldwide, tree planting is extensively advocated as a nature-based solution to facilitate the realization of Sustainable Development Goals (Griscom et al., 2020). China's Loess Plateau (LOP), with the greatest loess deposits in the world with a thickness of 30–200 m loess (Shao et al., 2018), has been one of the most active regions for tree planting during the last 4 decades, with the aim of controlling severe soil erosion and improving rural lives (Fu et al., 2016). Many trees of a variety of species (mostly exotic) have been planted during implementation of two large ecological projects: the Three-North Shelter Forest Program and the Grain For Green Program (Cao et al., 2021). So far, the area of tree planta-

tions on the LOP has reached 7.47×10^6 ha; the vegetation cover for this region increased from 31.6 % in 1999 to 65.0 % in 2017 (official data from the National Forestry and Grassland Administration, <http://www.forestry.gov.cn>, last access: 26 September 2023). As a result, a number of ecosystem services, including soil and water conservation, have been greatly improved on the LOP (Fu et al., 2016; Gao et al., 2021). In fact, the success of ecological restoration through tree planting on the LOP largely depends on the presence of an abundant “soil reservoir” deposited in deep loess deposits (200–2000 cm) (Zhu, 2006).

However, inappropriate tree planting in drylands can incur other ecological issues, such as soil desiccation due to excessive water consumption (Wang et al., 2010; Jia et al., 2017; Wu et al., 2021), preventing the sustainable development of revegetation, and may even result in tree mortality (Breshears et al., 2013; Choat et al., 2018; Stovall et al., 2019). Considerable studies have shown that planted trees extract substantial deep-layer soil moisture (DSM; units of gravimetric percent, hereafter “grav-%”) by developing deep roots to maintain their normal growth on the LOP (e.g., Gao et al., 2018; Su and Shangguan, 2019; Shi et al., 2021; Wang et al., 2023). However, continuous use of DSM over several years leads to severe soil desiccation (Wang et al., 2011; Gao et al., 2018; Tao et al., 2021) because DSM below 200 cm is hardly recharged by precipitation under tree plantations. In fact, deep-layer soil desiccation under tree plantations varies in degrees depending on tree age and local climate (Fang et al., 2016; Jia et al., 2017). Although soil desiccation has been widely reported in the above citations, the knowledge about variations in the lowest moisture of deep-layer soil desiccation that limits growth (hereafter “moisture limitation”) and/or the maximum depth from which trees can extract water (hereafter “maximum root water uptake (RWU) depth”) for different species and in different climatic regions remains limited. This is critical to estimate the amount of available soil water storage in deep soil and to predict the fate of planted trees based on soil water availability. Currently, the permanent wilting point (PWP; grav-%) is often used to represent the lowest limitation of soil moisture available to plants according to the definition of available water capacity (Tormena et al., 2017). However, the PWP determined in the lab and/or field is often related to annual crops rather than perennial mature trees (Lucia et al., 2020). In the literature, it is mainly estimated based on overall soil properties and uses soil moisture at -15 bar as the surrogate without considering differences between plant species (Torres et al., 2021). However, a recent synthesis indicated that the lowest soil moisture (moisture limitation) under tree plantations can be lower than the average PWP on the LOP (Li et al., 2021a). Therefore, it remains unclear whether the PWP is a reliable indicator of the moisture limitation of deep-layer soil desiccation for planted trees.

The occurrence of soil desiccation in deep layers can greatly affect the growth and eco-physiological traits of the

planted trees. It has been widely observed that the above-ground growth of planted trees is greatly constrained due to deep soil drying, resulting in so-called “dwarf aged trees” on the LOP (Hou et al., 1991; Zhu, 2000; Fang et al., 2016). Recently, Li et al. (2021) found that when deep soil below 200 cm is dry, the canopy transpiration of apple trees (*M. pumila*) with a rooting depth beyond 2000 cm is reduced by around 40 % relative to that without deep soil desiccation on the LOP. Furthermore, Yang et al. (2022) conducted a deep-soil–root partitioning experiment to quantify the effect of deep-layer (> 200 cm) root uptake on the transpiration and physiological features of *M. pumila*. They found that the canopy transpiration and the net photosynthetic rate of *M. pumila* without deep-layer root water uptake were reduced, respectively, by 36 % and 20 % on average, compared with the trees with deep roots at a semi-arid site on the LOP. Moreover, resistance of the xylem against embolism increases with a decrease in water availability, leading to a reduction in xylem hydraulic conductivity and even hydraulic failure of European beech (Schuldt et al., 2016; Stojnic et al., 2017). The effect of deep-layer soil desiccation on xylem hydraulic conductivity of planted trees, however, is still unclear, particularly in relation to the two dimensions.

To address the gaps, we built a dataset, including 11 980 observations of 258 soil profiles from 34 peer-reviewed publications and 4200 own observations of 72 soil profiles in-situ sampling across the LOP. The main objectives of this study were (i) to identify the moisture limitation and the maximum RWU depth for different tree species and the influencing factors and (ii) to explore their impact on xylem hydraulic conductivity of planted trees in different climatic regions on the LOP. Our hypotheses were that the moisture limitation and the maximum RWU depth under influence of deep-layer soil desiccation depend on tree species, as they differ in drought resistance, which is based on their water-use strategy under drought stress (Gessler et al., 2020). Because drought stress causes the formation of gas emboli and blockage of xylem conduits in woody plants, leading to a sharp decrease in xylem hydraulic conductivity (Gauthey et al., 2022), we also assume that the xylem hydraulic conductivity of planted trees varies among tree species and sites, as the sites represent gradients in aridity.

2 Materials and methods

2.1 Data compilation

2.1.1 Data sources

Peer-reviewed research papers, published between 1999 and 2021, were extracted from the Web of Science (United States) and CNKI (China Knowledge Resource Integrated Database, China; 2000–2021) using the search terms (“Loess plateau” OR “Loess hilly region”) AND (“artificial forest”

OR “afforestation” OR “plantation”) AND (“soil moisture” OR “soil water content” OR “soil water deficit”). To avoid bias in the literature selection, the following criteria were set to filter 718 articles obtained:

1. Soil profiles for moisture extraction under deep-rooted artificial forests must exceed a depth of 500 cm at dense sampling intervals (≤ 100 cm). The reason for the criterion is that the 0–200 cm soil layer is greatly affected by rainfall, while the phenomenon of a 200–500 cm dry soil layer is common (Jia et al., 2020). Therefore, we considered data from a sampling depth greater than 500 cm in this study to explore the state of DSM.
2. Soil moisture was the gravimetric or volumetric moisture content measured by drying methods or moisture sensors. The volumetric moisture content was uniformly converted to gravimetric moisture content by means of the average soil bulk density in each study area.
3. Experiments with only shrub and grass communities or experiments with irrigation, mulching and additional pruning, in addition to regular canopy pruning and/or other measures, were excluded.
4. Experiments yielded paired data for DSM in *M. pumila* orchards or black locust (*R. pseudoacacia*) forests and for that in cropland or grassland (control).
5. Location, mean annual precipitation (MAP; mm), forest age, and vegetation types were clearly presented.

2.1.2 Data extraction and classification

A total of 34 peer-reviewed publications were selected for analysis based on the aforementioned criteria. The original data were either clearly obtained from tables in the selected papers or directly extracted from the intuitive figures in those papers using GetData Graph Digitizer (version 2.24, <https://getdata1.software.informer.com/>, last access: 11 October 2022). In total, we retrieved a total of 11 980 observations from 258 soil profiles, covering 83 sites across four provinces or municipalities (Fig. 1), and the specific information is shown in Table 1. Soil moisture data were collected at 200 cm intervals, with values averaged when multiple samples were obtained within a 200 cm interval (e.g., with an interval of 20 or 10 cm). Soil moisture data for the same stand age collected in different sampling years were analyzed independently to assess the influence of forest age on soil moisture. To avoid random errors, data extracted for the moisture limitation of deep-layer soil desiccation under *M. pumila* in Changwu and Luochuan were divided into tree age range for analysis. Simultaneously, we also collected basic data from each publication, including data source, geographic coordinates (longitude, latitude), MAP, tree species and soil texture. Given the limited published information on DSM in *R.*

pseudoacacia forests, we also conducted supplementary experiments to obtain relevant data, as described in Sect. 2.2.

2.2 Data measurement

2.2.1 Study area and sampling sites

The LOP covers a total area of 640 000 km², with an elevation range of 200–3000 m. It belongs to the continental monsoon region where annual precipitation ranges from 150 mm in the northwest to 800 mm in the southeast, 55 %–78 % of which falls from June to September (Wang et al., 2011). Figure 1 and Table 1 show the location of sites from which we collected field data across the entire LOP (34°43′–41°16′ N, 100°54′–114°33′ E), indicated by the small red crosses and the bold text, respectively. Geographic coordinates (altitude, longitude and latitude) for each site were obtained using a global positioning system (Zhuolin A8, with an accuracy of 1 m).

2.2.2 Soil moisture content

The sampling experiments for soil moisture were done from 9 April to 6 May 2022 (the early growing season of trees on the LOP) under the plantations of *M. pumila* and *R. pseudoacacia* aged 20–30 years as well as under the cropland/grassland as the control due to the intact DSM below 200 cm. Initially, the sampling depth was set as 0–10 m under different tree plantations; however, it was extended down to 25 m when the DSM reached the moisture limitation under one given aged plantation as well as under the paired control plots (sampling process is shown in the Fig. S1 in the Supplement). The information of the trees showing the lowest deep soil moisture deficit at each site sampled in the field is indicated in Table 2. The maximum sampling depth was set as 25 m because previous studies reported the maximum rooting depth less than 25 m on the LOP (Wang et al., 2009; Li et al., 2019). In order to minimize the spatial variability in soil moisture caused by soil properties, the straight-line distance between the planted trees and their controls was controlled at less than 500 m, and the average straight-line distance of the three replicates is shown in Table 2. All soil samples were collected by a hand auger ($\Phi = 6$ cm) at a depth interval of 20 cm and 1.5 m horizontal to the trunk of trees, and three trees with similar growth status were selected for each age as repeats. Collecting soil samples in the due north direction of the first tree, a clockwise rotation of 120° around the second tree from the due north direction was used as the second sampling point, and a clockwise rotation of 240° around the third tree from the due north direction was used as the third sampling point. Soil samples without roots were put into an aluminum specimen box and used to measure the gravimetric moisture content (grav-%), which was determined by a drying method at 105° for constant weight. A total of 4200 soil moisture data were collected at 72 soil profiles. Three repeats

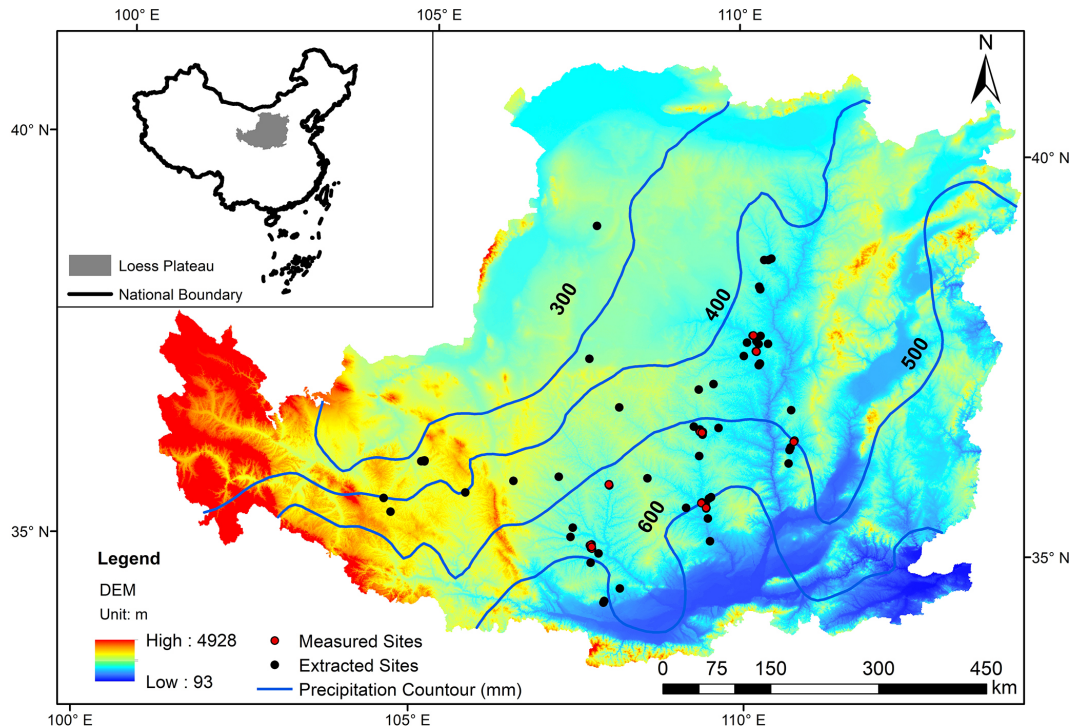


Figure 1. Distribution map of sampling sites on the Loess Plateau. The black dots represent the sampling sites extracted from the article, and the small red crosses represent the sampling sites supplemented in the field. Publisher's remark: please note that the above figure contains a disputed territory.

of soil moisture data under one given tree plantation were averaged when analyzing the moisture limitation and the maximum RWU depth under plantations to eliminate the effect of different sampling directions. Moreover, the soil sampling under all tree plantations was done once because the temporal variation of DSM below 200 cm under tree plantations was negligible in one growing season (Fig. S2; Gao et al., 2018; Li et al., 2019).

2.2.3 Native percentage loss of hydraulic conductivity

To explore the influence of deep-layer soil desiccation on the native percentage loss of hydraulic conductivity (NPLC; %) of planted trees across different sites, while collecting soil moisture, six current-year branches that were 18–20 cm in length and 2–5 mm in diameter were cut off in the morning from the southern crown of *M. pumila* and *R. pseudoacacia* trees at the tree age when the plantations reached the moisture limitation, and three trees with similar growth status were selected as repeats for each site. Sampled branches were sprayed with water and enclosed in humidified black plastic bags before excising to minimize water loss. After bringing these branches to the laboratory, we quickly cut 3–5 cm from the middle of the branch under water and immediately measured xylem hydraulic conductivity of it using a xylem embolism measurement system (Xylem embolism meter, Bronkhorst, Montigny-lès-Cormeilles, France). To

prevent water splitting from the branch, the initial xylem hydraulic conductivity (K_{ini} , $\text{kg s}^{-1} \text{MPa}^{-1}$) was measured with 1 mmol L^{-1} CaCl_2 solution ($\text{pH} = 6$) through a filter with a filter aperture of $0.22 \mu\text{m}$ at a low pressure of 0.5–1.0 kPa. And at a high pressure of 0.2 MPa, the branch was flushed with 1 mmol L^{-1} CaCl_2 solution until there were no air bubbles, and then the xylem hydraulic conductivity was determined again at a low pressure of 0.5–1.0 kPa, and the flushing was repeated until a maximum conductivity was attained (K_{max} , $\text{kg s}^{-1} \text{MPa}^{-1}$). The NPLC was calculated by

$$\text{NPLC} = \left(1 - \frac{K_{ini}}{K_{max}}\right) \times 100. \quad (1)$$

2.2.4 Basic parameters of plantation

To accurately target the sample plantation, tree age was determined using the plant growth cones (Haglof, CO300-52, Sweden) before sampling, and the crown diameter, basal diameter and diameter at breast height were measured with a tape (Table 2). The sampling of trees' roots was carried out simultaneously with soil moisture sampling, and soil samples with roots were stored in a sealed sample bag to minimize the loss of fine roots ($\leq 2 \text{ mm}$) and then gently rinsed in a 100-mesh sieve, removing residual roots from the sieve with tweezers. Each fine-root sample was dried at 65° to a constant weight and weighed using an electronic balance with a precision of 0.0001 g to determine root dry weight. Finally,

Table 1. Sample information used in this study.

Sites	Species	NEP (pair)	NSP (pair)	MAP (mm)	2021-P (mm)	Climate type zone
Luochuan	<i>M. pumila</i>	24	10	608	957	Semi-humid
Fufeng	<i>M. pumila</i>	1	–	606	–	Semi-humid
	<i>R. pseudoacacia</i>	4	–		–	
Yongshou	<i>R. pseudoacacia</i>	4	–	600	–	Semi-humid
Changwu	<i>M. pumila</i>	44	8	578	910	Semi-humid
	<i>R. pseudoacacia</i>	6	10			
Baishui	<i>M. pumila</i>	2	–	577	–	Semi-humid
Jingchuan	<i>M. pumila</i>	1	–	571	–	Semi-humid
Qingcheng	<i>M. pumila</i>	6	–	561	–	Semi-humid
Yan'an	<i>M. pumila</i>	7	–	530	–	Semi-humid
	<i>R. pseudoacacia</i>	15	10		704	
Daning	<i>M. pumila</i>	2	–	528	–	Semi-humid
Zichang	<i>M. pumila</i>	1	–	480	–	Semi-arid
	<i>R. pseudoacacia</i>	1	–		–	
Zizhou	<i>M. pumila</i>	1	–	424	–	Semi-arid
Mizhi	<i>M. pumila</i>	9	10	421	441	Semi-arid
	<i>R. pseudoacacia</i>	1	10		458	
Total	–	129	58	–	–	–

Note that NEP and NSP indicate the number of data pairs from the literature and field sampling, respectively. A data pair refers to the moisture content of a soil profile from plantation and that of a soil profile from cropland/grassland. MAP is the mean annual precipitation (mm); based on MAP, these sites are classified as semi-arid (250 mm < MAP < 500 mm) and semi-humid (500 mm < MAP < 800 mm) climate zones, respectively (Wang et al., 2011). The bold text indicates the supplementary sampling in this study with reference to the existing data, and these are representative plots for the different climatic regions upon which we focused.

the fine-root dry-weight density (g m^{-3}) was calculated using the formula given by Zhao et al. (2022).

2.2.5 Soil texture and permanent wilting point

The sampling of soil properties was done simultaneously as soil moisture was sampled, and the data are shown in Table 3. Soil samples without roots were air-dried, ground, and then passed through a 16-mesh sieve for measuring soil particle composition (i.e., clay, silt and sand contents) and soil pH and through a 100-mesh sieve for determining soil organic carbon (SOC), respectively. Soil particle composition was determined by laser diffraction using a Mastersizer-2000 laser granularity analyzer (Malvern Instruments, Malvern, England) (Wang et al., 2008). Soil pH was determined by the DELTA 320 pH meter. SOC was measured using the potassium dichromate volumetric method (Fu et al., 2010), and soil organic matter was converted from SOC (Eq. 4). Due to the challenges of collecting undisturbed soil cores within the 10 m profile, soil bulk density and the PWP of each site were estimated using the pedo-transfer functions developed

by Wang et al. (2012) and Balland et al. (2008) based on the above indicators, as follows:

$$\text{PWP}_i = \text{FC}_i \left(1 - \exp \left(\frac{-0.511 \times \text{clay}_i - 0.865 \times \text{SOM}_i}{\text{FC}_i} \right) \right) \quad (2)$$

$$\begin{aligned} \text{FC}_i = & 46.481 - 4.757 \times \text{SOC}_i - 14.028 \times \log_{10}(\text{clay}_i) \\ & - 13.991 \times \log_{10}(\text{sand}_i) + 42.261 \times \log_{10}(\text{SOC}_i) \\ & - 11.763 \times \text{sand}_i^{-1} + 19.198 \times \text{SOC}_i^{-1} - 5.448 \\ & \times \text{BD}_i + 0.044 \times \text{SOC}_i^2 + 1.975 \times \text{BD}_i \times \text{SOC}_i \end{aligned} \quad (3)$$

$$\text{SOM}_i = \text{SOC}_i \times 1.724 \quad (4)$$

$$\begin{aligned} \text{BD}_i = & 1.8284 + 0.0429 \times \log_{10}(\text{clay}_i) + 0.0205 \\ & \times \text{clay}_i^{0.5} - 0.0125 \times \cos(\text{clay}_i) - 0.0061 \times \text{silt}_i \\ & + 0.0001 \times \text{silt}_i \times \text{SG} - 0.0098 \times \text{SG} - 0.0071 \\ & \times \text{SOC}_i - 0.0505 \times \text{SOC}_i^{0.5} + 0.0002 \times \text{SOC}_i^2, \end{aligned} \quad (5)$$

where FC is field capacity (grav-%); SOC and SOM are soil carbon content and soil organic matter (g kg^{-1}); BD is the

Table 2. Basic information for the trees showing the lowest deep soil moisture deficit at each site sampled in the field.

Sites	Plant species	Longitude	Latitude	Elevation (m)	SD (m)	Stand age (yr)	PH (m)	BD (cm)	DBH (cm)	CD (m)
Mizhi	<i>M. pumila</i>	110°10'22"	37°52'13"	1135.1	430	27	3.5 ± 0.25	5.8 ± 0.53	4.9 ± 0.21	5.73 ± 0.35
	Cropland	110°11'09"	37°52'21"	1139.6	–	–	–	–	–	–
	<i>R. pseudoacacia</i>	110°13'58"	37°40'08"	1086.2	433	24	9.3 ± 0.40	8.9 ± 0.43	7.3 ± 0.17	5.65 ± 0.75
	Grassland	110°13'46"	37°40'16"	1076.4	–	–	–	–	–	–
Luochuan	<i>M. pumila</i>	109°21'59"	35°46'44"	1086.9	182	28	2.85 ± 0.35	8.8 ± 0.82	8.1 ± 0.52	4.51 ± 16.7
	Cropland	109°21'58"	35°46'38"	1083.9	–	–	–	–	–	–
Yan'an	<i>R. pseudoacacia</i>	109°23'22"	36°40'02"	1325.1	412	25	14.35 ± 0.25	9.3 ± 0.19	7.9 ± 0.86	5.81 ± 0.77
	Grassland	109°22'29"	36°40'31"	1333.2	–	–	–	–	–	–
Changwu	<i>M. pumila</i>	107°41'01"	35°14'12"	1231.7	390	24	4.42 ± 0.25	6.1 ± 0.29	5.4 ± 0.73	4.77 ± 0.13
	Cropland	107°40'43"	35°14'20"	1232.6	–	–	–	–	–	–
	<i>R. pseudoacacia</i>	107°41'45"	35°12'50"	1215.2	212	28	13.35 ± 0.25	9.4 ± 0.66	8.1 ± 0.51	5.96 ± 0.01
	Grassland	107°42'12"	35°12'36"	1219.4	–	–	–	–	–	–

Note that $A \pm B$, where A represents the average of the three replicates of each parameter, and B represents the standard deviation. SD – average of straight-line distance between three trees and their controls (cropland or grassland). PH – average plant height. BD – average basal diameter. DBH – average diameter at breast height. CD – average crown diameter.

soil bulk density (g cm^{-3}); clay, silt and sand are clay, silt and sand content (%), respectively; SG is the slope gradient at each sample location ($^{\circ}$); and i represents the i th soil depth.

2.2.6 Meteorological parameters

The precipitation, air temperature, and relative humidity of different sites were from the European Centre for Medium-Range Weather Forecasts ERA5-Land reanalysis (hereafter ERA5-Land) data, the accuracy of which has been verified by Araújo et al. (2022). Based on the above meteorological data, the vapor pressure deficit (VPD; kPa) was calculated using Eq. (6).

$$\text{VPD} = 0.61078 \times e^{\frac{17.27 \times T_a}{T_a + 237.3}} \times (1 - \text{RH}), \quad (6)$$

where T_a is air temperature ($^{\circ}$), and RH is relative humidity (%).

2.3 Data analysis

2.3.1 Meta-analysis

In a traditional meta-analysis, the mean values of the treatment and control with their standard deviations or standard errors are extracted from the publications. However, most published papers selected for this study reported their mean values without standard deviations. To maximize the number of observations, the following unweighted meta-analysis approach was adopted (Li et al., 2021b):

$$\text{RR} = \text{Ln}(X_e/X_c), \quad (7)$$

where response ratio (RR) is a unit-free index with positive and negative values indicating either increasing or decreasing soil moisture in response to planted trees' growth. X_e and X_c are values of soil moisture under trees and controls for each study, respectively. The soil depth interval analyzed was 200 cm, as mentioned in Sect. 2.1.2. We use a 95 % confidence interval (95 % CI) to test the statistical difference between the RR of soil moisture and zero (Deng et al., 2016). The calculation method of 95 % CI is shown in Eqs. (8) and (9):

$$\text{SE}_{\text{total}} = \sqrt{\frac{V_s}{N}} \quad (8)$$

$$95\% \text{ CI} = 1.96 \times \text{SE}_{\text{total}}, \quad (9)$$

where SE_{total} is the standard error of RR, and V_s and N are the variances of RR and the number of observations, respectively. In this study, the statistical difference between the RR of soil moisture and zero was considered significant if the 95 % CI did not include zero, and the differences were not considered significant if the 95 % CI included zero.

2.3.2 Quantitative index for moisture limitation of soil desiccation

The effect of different deep-rooted planted trees' RWU on soil moisture was expressed as soil moisture deficit, which is the relative difference in soil moisture for given trees and adjacent cropland/grassland. Because root distribution of the latter is too shallow (< 2 m) to influence the change of DSM (Gao et al., 2014; Zhu et al., 2016; Gao et al., 2022), the soil moisture of grassland/farmland measured simultane-

Table 3. Basic soil properties of field sampling sites.

Sites	Plant species	Soil type	Clay	Silt	Sand	Soil organic matter (g kg ⁻¹)	Bulk density (g cm ⁻³)	pH
Mizhi	<i>M. pumila</i>	Cultivated loessial soils	15.5	24.3	60.2	2.94	1.28	7.64
	Cropland	Cultivated loessial soils	14.9	25.1	60.0	2.33	1.29	8.44
	<i>R. pseudoacacia</i>	Cultivated loessial soils	13.6	25.0	61.4	3.09	1.28	8.03
	Grassland	Cultivated loessial soils	13.8	25.4	60.8	2.35	1.29	8.36
Luochuan	<i>M. pumila</i>	Dark loessial soils	21.5	37.1	41.4	4.62	1.31	7.54
	Cropland	Dark loessial soils	22.2	38.5	39.3	4.38	1.33	8.17
Yan'an	<i>R. pseudoacacia</i>	Cultivated loessial soils	18.1	33.1	48.8	4.87	1.29	8.07
	Grassland	Cultivated loessial soils	19.6	31.5	48.9	4.43	1.30	8.53
Changwu	<i>M. pumila</i>	Dark loessial soils	24.6	38.7	36.7	4.76	1.31	7.35
	Cropland	Dark loessial soils	25.8	38.7	35.5	4.35	1.32	8.21
	<i>R. pseudoacacia</i>	Dark loessial soils	22.9	38.2	38.9	5.01	1.29	7.96
	Grassland	Dark loessial soils	22.7	38.4	38.9	4.26	1.31	8.23

Note that the sampling depth of basic soil properties, including soil particles (which was divided into clay (< 0.002 mm, %) , silt (0.002–0.02 mm, %) and sand (0.02–2 mm, %) according to the international soil particle classification standard), soil organic carbon (g kg⁻¹) and pH, was 10 m for different plant species, and the average values of these parameters in different soil layers are presented here. The soil bulk density presents the average value of the 1 m soil profile.

ously with that of plantation can be used as a background value to eliminate the influence of climate difference. The soil moisture deficit in the *k*th layer of the *j*th plantations' calculation method is shown in Eq. (10). And the soil moisture deficit of the soil layer below 2 m is the deep soil moisture deficit (DSMD). In order to compare the difference between the DSMD of plantations and their PWP, the soil moisture deficit corresponding to PWP in the *k*th layer of the *j*th plantations along the soil profile was calculated by Eq. (11).

$$SMD_{j,k} = \frac{SMC_{j,k} - SMC_{0,k}}{SMC_{0,k}} \tag{10}$$

$$SMD_PWP_{j,k} = \frac{PWP_{j,k} - SMC_{0,k}}{SMC_{0,k}}, \tag{11}$$

where $SMC_{j,k}$ and $PWP_{j,k}$ represent soil moisture content and permanent wilting point in the *k*th layer of the *j*th plantations (grav-%), respectively; $SMC_{0,k}$ indicates soil moisture content in the *k*th layer of the control (cropland or grassland) (grav-%); and $SMD_{j,k}$ and $SMD_PWP_{j,k}$ represent the soil moisture deficit corresponding to $SMC_{j,k}$ and $PWP_{j,k}$, respectively.

2.3.3 Quantitative index for the maximum RWU depth

The paired-sample *t* test in SPSS was used to analyze the significant difference of soil moisture between plantations and their controls. Three duplicate values were taken for each plantation and its control per site, and the straight-line distance between them was less than 500 m. If the difference

of soil moisture content in the plantation and its' control becomes non-statistically significant ($p > 0.05$) at a given layer, it is defined that this depth corresponds to the maximum RWU depth for the planted trees sampled in this study. On the other hand, if it is always statistically significant ($p < 0.05$) along the whole measured profile, this indicates the maximum RWU depth exceeds the measurement depth. A list of all acronyms, the corresponding definitions and units is given in Table 4.

2.3.4 Statistical analysis

Redundancy analysis (RDA) was performed to evaluate the contributions of environmental factors to soil moisture deficit and soil moisture using Canoco 5 (Microcomputer Power Ithaca, New York, USA). Least-significant-difference post hoc tests in the SPSS 22.0 software package (SPSS 22.0, SPSS Institute Ltd., USA) were used to analyze the significant differences in NPLC of current-year branches of planted trees in different sites (takes the average of six branches as the NPLC per tree, and three duplicate trees were treated as independent in statistical analysis); $\alpha = 0.05$ was considered statistically significant. Origin 2021 software (Origin-Lab Corporation, USA) was used to draw the figures.

Table 4. List of acronyms included in the study, with definition and units.

Acronym	Definition	Units
Soil properties		
DSMD	Deep-soil moisture deficit	–
DSM	Deep-soil moisture	grav-%
PWP	Permanent wilting point	grav-%
SOC	Soil organic content	g kg^{-1}
Research objects		
LOP	China's Loess Plateau	–
<i>M. pumila</i>	Apple trees (<i>Malus pumila</i> Mill.)	–
<i>R. pseudoacacia</i>	Black locust (<i>Robinia pseudoacacia</i> L.)	–
SD	Average of straight-line distance between three trees and their controls	m
Plant properties		
PH	Average plant height	m
BD	Average basal diameter	cm
DBH	Average diameter at breast height	cm
CD	Average crown diameter	m
NPLC	Native percentage loss of hydraulic conductivity	%
K_{ini}	The initial branch xylem hydraulic conductivity	$\text{kg s}^{-1} \text{MPa}^{-1}$
K_{max}	The maximum branch xylem hydraulic conductivity	$\text{kg s}^{-1} \text{MPa}^{-1}$
RWU	Root water uptake	mm
Environmental conditions		
VPD	Vapor pressure deficit	kPa
MAP	mean annual precipitation	mm
Data analysis		
RDA	Redundancy analysis	–
95 % CI	95 % confidence interval	–

3 Results

3.1 Soil moisture under planted trees of the different ages

The soil moisture under *M. pumila* orchards and *R. pseudoacacia* forests varied greatly with tree age across the entire soil layer (Fig. 2). As tree age increased, the negative effect of *M. pumila* (0–21 m soil layer) and *R. pseudoacacia* (0–10 m soil layer) growth on DSM initially increased and then stabilized. Specifically, the negative effect value of DSM (4–10 m) under *R. pseudoacacia* forests older 30 years decreased by 17.4 % compared with those aged 20–30 years. Therefore, the two plantations had the greatest effect on DSM at the age range of 20–30 years. Consequently, plantations within this age range were considered the main subjects when analyzing the moisture limitation and the maximum RWU depth under the influence of deep-layer soil desiccation for planted trees in Sect. 3.2 and 3.3.

3.2 The dimensions of deep-layer soil desiccation for planted trees

The DSM under a given tree species can vary substantially among different years, even at the same age, due to varying annual precipitation. For instance, the DSM of 15-year-old *M. pumila* in Changwu varied greatly across different sampling years (Fig. 3a). To account for climate differences between sampling years, we used soil moisture deficit, which was defined as the relative difference in DSM between the given trees and nearby controls. The DSM of control adjacent to the plantation was regarded as the background soil moisture to reflect local annual precipitation because it was not influenced by root water extraction from annual crops with the shallow roots (< 2 m). We found that the variation coefficient of DSMD (Fig. 3b) was lower than that of DSM; thus, DSMD was used for the following in-depth analyses.

3.2.1 Moisture limitation of deep-layer soil desiccation

We combined the extracted (Fig. 4a and b) and measured (Fig. 4c–f) data to explore the moisture limitation of deep-

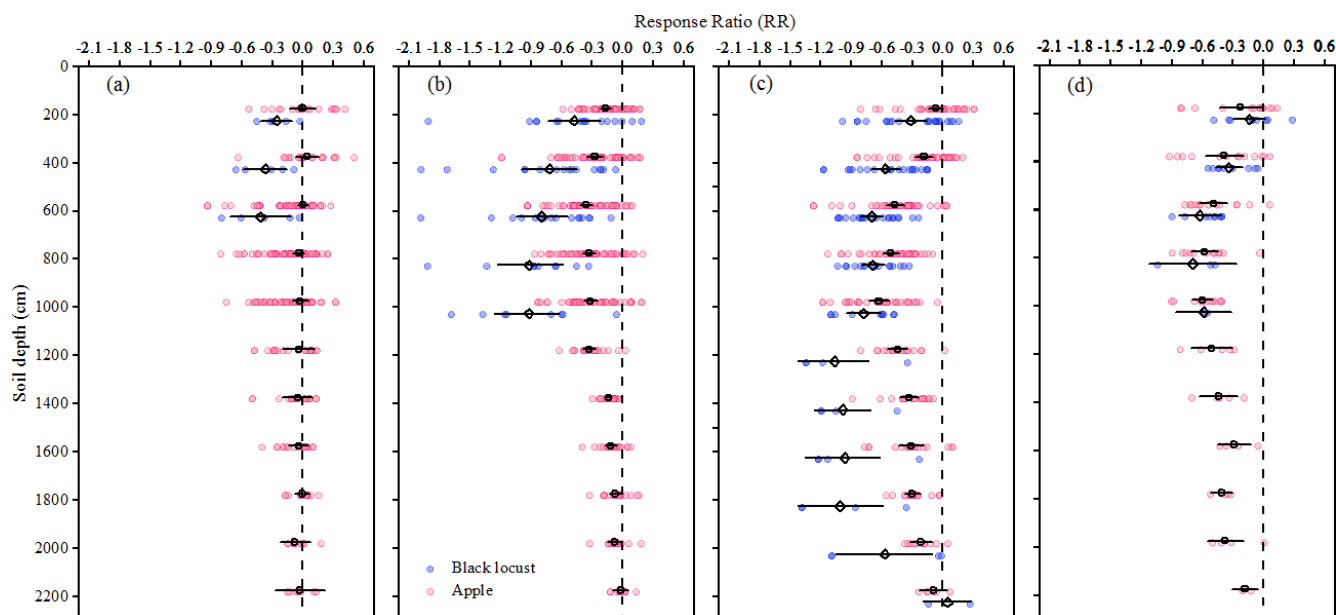


Figure 2. Mean effect sizes indicating the mean value of response ratio (RR; a unit-free index with positive or negative value means soil moisture increases or decreases with the older of plantations) of soil moisture with the age of plantation on the Loess Plateau are divided into (a) tree age < 10 years, (b) 10 years ≤ tree age < 20 years, (c) 20 years ≤ tree age < 30 years and (d) tree age ≥ 30 years. The vertical black dotted lines indicate RR = 0. Blue and red dots, which includes 11 980 literature extraction data and 4200 field sampling data, represent black locust (*R. pseudoacacia*) forests and apple orchard RRs, respectively; diamonds and circles represent the mean RR of *R. pseudoacacia* and apple (*M. pumila*) orchard, respectively; and the error bars represent 95 % CI.

layer soil desiccation. The DSMD fluctuation in *M. pumila* over 19 years was relatively stable in Changwu, Luochuan and Mizhi (Fig. 4a–c). Interestingly, with the increase in tree age, the lowest DSMD of *M. pumila* occurred at different sites and tree age between 24 and 28 years was associated with the lowest DSMD at different sites. After reaching the lowest value, *M. pumila* could no longer extract water from the deep soil. Notably, although there were differences in soil moisture among different sites and tree species (Fig. S3), the lowest value of DSMD at three sites varied around -0.6 , which was very close to the DSMD value corresponding to the local PWP. In addition, DSMD of the *R. pseudoacacia* in Changwu, Yan’an and Mizhi was similar to that of *M. pumila* (Fig. 4d–f). Namely, the lowest values of DSM consumption were the same as those of *M. pumila* at different sites. This suggests that PWP is a reliable indicator of the moisture limitation of deep-layer soil desiccation, regardless of tree species and site. In addition, compared with *M. pumila*, the variation range in DSMD for *R. pseudoacacia* decreased greatly in Changwu and Yan’an.

3.2.2 The maximum RWU depth under the influence of deep-layer soil desiccation

Based on the measured and literature data, we analyzed the maximum RWU depth of plantations that reached the moisture limitation of deep-layer soil desiccation in different regions (Fig. 5). With an increase in depth, the soil moisture

of plantations was close to or even higher than that of the control in the top 200 cm, except for the *M. pumila* orchards and *R. pseudoacacia* forests in Mizhi. Therefore, we calculated the difference significance of soil moisture between the plantations and control using the paired-sample *t* test, treating the points with no significant ($p > 0.05$) soil moisture difference as the maximum RWU depth for the planted trees sampled in this study. In this way, the maximum RWU depth reached 18.0–22.0 m for *M. pumila* orchards at different sites and 18.4–18.8 m for *R. pseudoacacia* forests in Changwu and Yan’an, whereas it was deeper than 25 m in the drier site of Mizhi.

3.3 Effect of factors on the dimensions of deep-layer soil desiccation

Based on all extracted and measured data, including soil moisture deficit, soil moisture, soil texture (clay, silt) and fine-root dry-weight density, along with ERA5-Land meteorological data (MAP, mean annual temperature, relative humidity and VPD), we used RDA to analyze the key factors that determined the moisture limitation and the maximum RWU depth under the influence of deep-layer soil desiccation (Fig. 6). The result indicated that the explanatory variables (soil properties, meteorological variables and plant characteristic) accounted for 67.7 % of the total variation of soil moisture and soil moisture deficit. Soil moisture deficit was significantly ($p < 0.01$) positively correlated

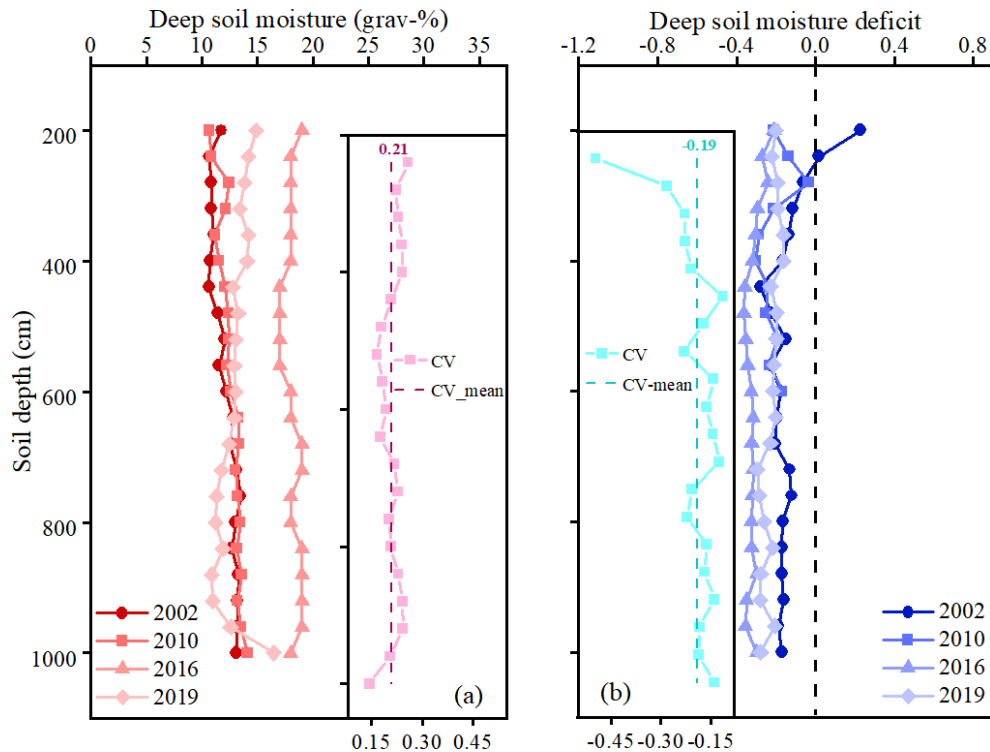


Figure 3. Distribution of deep soil moisture (grav-%, **a**) and deep soil moisture deficit (**b**) of 15-year-old apple (*M. pumila*) orchards in Changwu in 2002, 2010, 2016 and 2019, which is all based on literature extraction data. The variation coefficient (CV) of deep soil moisture and deep soil moisture deficit below 2 m are shown in the small figures, the dashed lines of which represent the mean values of the CV for deep soil moisture and deep soil moisture deficit, respectively.

with VPD and negatively correlated with clay, silt, relative humidity, mean annual temperature and MAP (Fig. 6a). The clay, VPD, silt, relative humidity and fine-root dry-weight density provided statistically significant ($p < 0.01$) explanations for the distribution of soil moisture deficit and soil moisture (Fig. 6b). The clay content determined the lower limit of water that plants could uptake through its adsorption of water molecules. Therefore, soil texture and climatic conditions were the main factors influencing the dimensions of deep-layer soil desiccation in different sites. However, the relationship between MAP and soil moisture deficit was not significant, which might be because the MAP of each site diminished the differences of precipitation in each sampling year.

3.4 Effect of the dimensions of deep-layer soil desiccation on hydraulic conductivity

To further elucidate how the moisture limitation and the maximum RWU depth under the influence of deep-layer soil desiccation affect the hydraulic conductivity in plants, we explored the variations of hydraulic conductance of branches for different tree species and sites (Fig. 7). Please note that the data used for analysis here were from only the plantations where the moisture limitation and the maximum RWU depth

were determined in this study (Figs. 4 and 5). The NPLC was higher than 50 % for all tree species and sites and exhibited distinct variations between tree species and sites. The NPLC ranged from 74.9 %–96.5 % for *R. pseudoacacia* trees and 83.0 %–92.7 % for *M. pumila* trees. Particularly, the NPLC of *R. pseudoacacia* trees in Mizhi was significantly higher than that in other sites ($p < 0.05$), indicating that the *R. pseudoacacia* trees in Mizhi had greatest mortality risk in relation to the severe deep-layer soil desiccation.

4 Discussion

4.1 The mechanism underlying for the dimensions of deep-layer soil desiccation

Although DSM (> 200 cm) declined as the planted trees aged and even reached the lowest limitation, shallow soil moisture (< 200 cm) remained stable (Figs. 2 and 4). This is attributed to the strong effect of rainfall and evapotranspiration on shallow soil moisture. However, DSM is often in a negative balance, being depleted by deep-rooted trees and unable to recover for a long time (Fang et al., 2016; Gao et al., 2020). As the critical water resource for plants to cope with extreme droughts, DSM of various sites was exhausted to the lowest

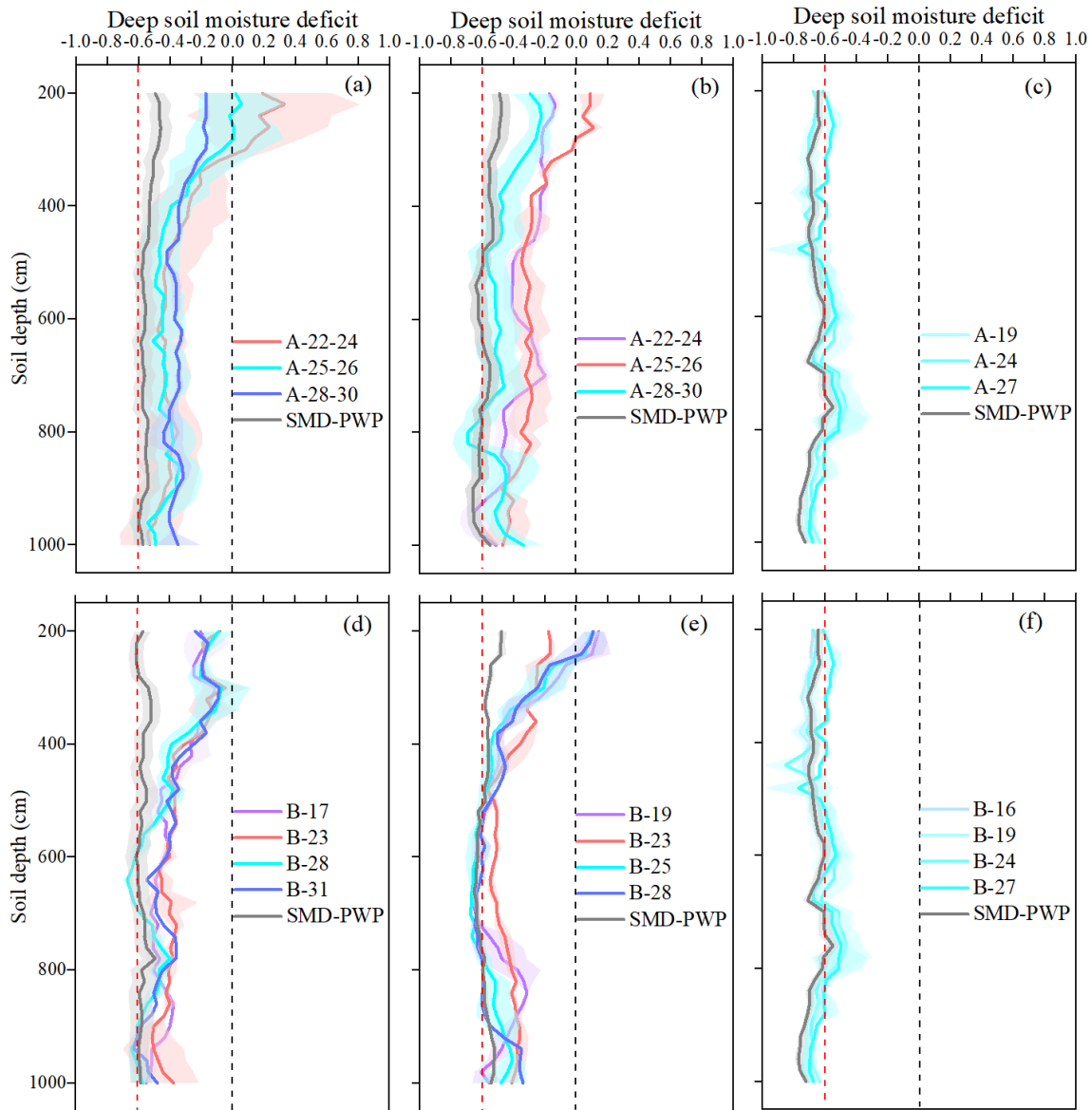


Figure 4. Deep-soil moisture deficit (DSMD) distribution under apple (*M. pumila*) orchards of different ages in Changwu (a), Luochuan (b) and Mizhi (c) and deep soil moisture deficit under black locust (*R. pseudoacacia*) forests of different ages in Changwu (d), Yan’an (e) and Mizhi (f). The capital letters A and B in the legend represent *M. pumila* and *R. pseudoacacia*, respectively, and the numbers after the letters represent tree age; for example, A-22 represents 22-year-old *M. pumila*. Panels (a) and (b) show data extracted from literature, while panels (c–f) show field-measured data. In order to reduce the randomness caused by different sampling years and locations between the extracted data, the data for *M. pumila* trees with similar ages in Changwu (a) and Luochuan (b) were combined. The gray line indicates soil moisture deficit at the permanent wilting point (SMD_PWP) in each sampling site, and the dashed red line indicates the soil moisture deficit equal to -0.6 . The shaded portion corresponding to each line represents the positive and negative values of the standard deviation.

limitation, posing a serious threat to the sustainability of tree growth under future climate conditions.

In our study, the lowest DSMD varied around -0.6 regardless of tree species and site (Fig. 4). This challenges the assumption that moisture limitation of deep-layer soil desiccation is dependent on tree species. Based on the results of this study and opinions from the literature, Fig. 8 shows the mechanism by which plants regulate the moisture limita-

tion of deep-layer soil desiccation. Soil water potential, when the soil hydraulic conductivity (K_s) drops to lower values than the root hydraulic conductivity (K_r), can be regarded as the critical soil matric potential, at which the soil begins to limit the water flux in the soil–plant continuum (Fig. 8; Cai et al., 2022). Thus, the soil with coarser texture has higher K_s when soil starts limiting trees’ RWU. Correspondingly, the K_r value is higher in areas with less rainfall, which may

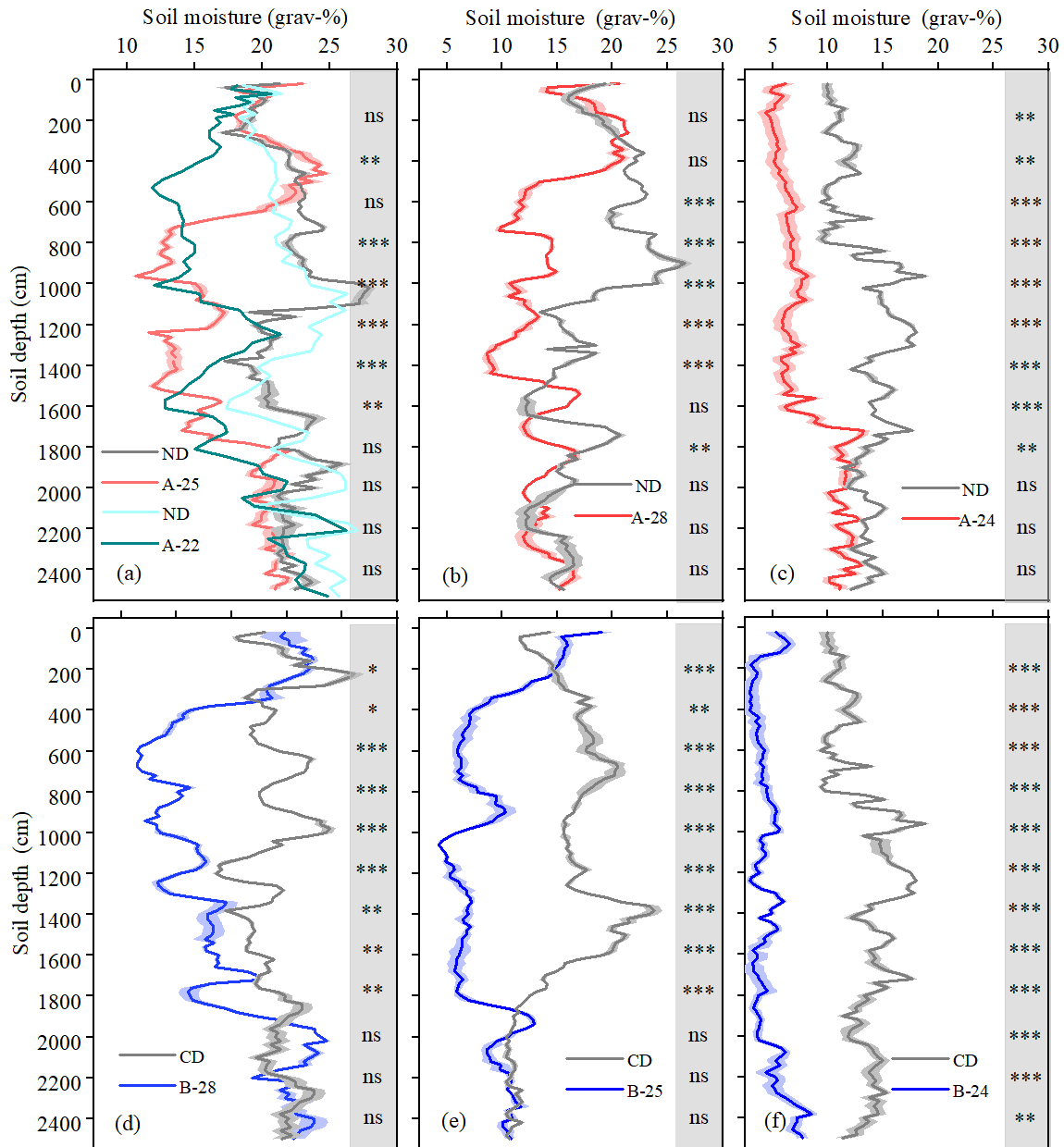


Figure 5. Dynamic distribution of maximum water absorption depth after reaching the moisture limitation of deep soil desiccation. The 22-year-old apple (*M. pumila*) and cropland (dark green) data in (a) have been extracted from the literature, and other data are field-measured data. The capital letters A and B in the legend represent *M. pumila* and black locust (*R. pseudoacacia*), respectively, and the numbers after the letters represent tree age; for example, A-22 represents 22-year-old *M. pumila*. Panels (a) and (d) indicate Changwu; panels (b) and (e) indicate Luochuan and Yan'an, respectively; and panels (c) and (f) indicate Mizhi. The gray line represents soil moisture (grav-%) of grassland (CD) or farmland (ND). ***, **, * and ns in the gray bands indicate the significance level of measured data obtained by the paired-sample *t* test for $p < 0.001$, $p < 0.01$, $p < 0.05$ and no significant differences, respectively. The shaded portion corresponding to each line represents the positive and negative values of the standard deviation.

be maintained by allocating more biomass to fine roots than in areas with more rainfall (Fig. S5). In addition, VPD in the areas with less rainfall is relatively high, and the evaporative demand increases (Will et al., 2013; Hayat et al., 2019), causing a larger water potential gradient between roots and leaves under the same soil moisture and thus enabling roots to ab-

sorb more moisture. Therefore, although the physiological and growth characteristics of *M. pumila* and *R. pseudoacacia* across different sites were markedly different (Table 2; Figs. 7, S5), the lowest DSMD of all tree species at each site fluctuated around -0.6 . This indicates that the amount of soil moisture absorbed by plants is determined by VPD and soil

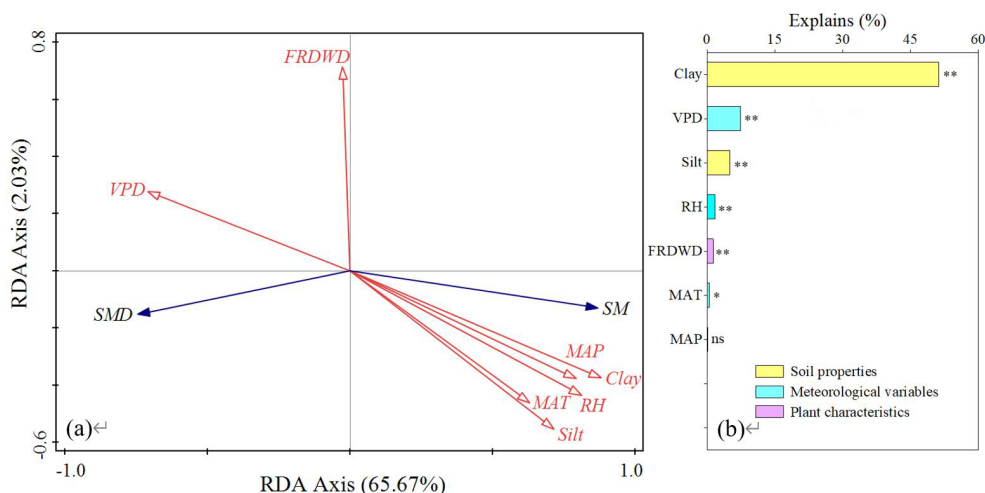


Figure 6. (a) Redundancy analysis (RDA) of “response variables (soil moisture deficit (SMD), soil moisture (SM; grav-%)) and environmental factors” is based on the literature and field-measured data. Descriptors (red arrows) are soil properties (clay and silt), meteorological variables (vapor pressure deficit (VPD; kPa), mean annual precipitation (MAP; mm), relative humidity (RH), mean annual temperature (MAT; °)) and plant characteristic (fine-root dry-weight density (FRDWD; g m^{-3})). Axis 1 (65.67 %, $p < 0.01$) and Axis 2 (2.03 %, $p < 0.01$) are shown. Panel (b) shows the explanation provided by each of the environmental variables for the response variables. **, * and ns following the bar chart indicate $p < 0.01$, $p < 0.05$ and not significant, respectively.

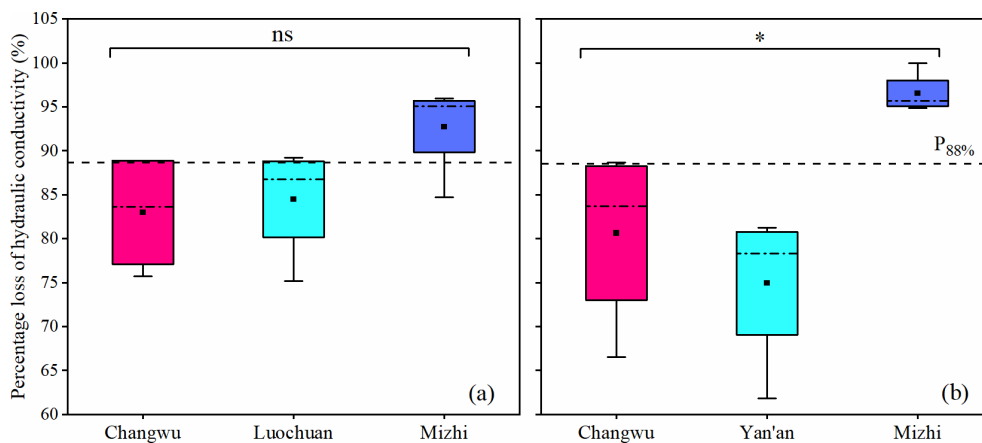


Figure 7. Differences in percentage loss of hydraulic conductivity of annual branches of apple (a, *M. pumila*) and black locust (b, *R. pseudoacacia*) measured in different sites (Changwu, Luochuan, Yan’an and Mizhi). * and ns indicate that the significance level is $p < 0.05$ and there is no statistically significant difference, respectively. The n_{sample} in (a) and (b) is 54, respectively.

water availability, which is mainly affected by the precipitation and soil structure (Wu et al., 2015). Notably, the lowest DSMD was very close to the DSMD value corresponding to the local PWP, regardless of tree species (Fig. 4). This might be attributed to different plant species wilted at similar water contents in soils with low clay content (Torres et al., 2021). Overall, our study confirms that PWP is a reliable indicator of the moisture limitation of deep-layer soil desiccation for the tree species examined. These findings provide important information for the sustainable development of plantations on the LOP.

When soil water in the profile becomes limited, plants extend their roots into the deep soil to expand their water-hunting range during the dry season (Fig. 8; Ivanov et al., 2012). Our results showed that for the sampled trees, the maximum RWU depth reached 18.0–22.0 m for *M. pumila* orchards across different sites and 18.4–18.8 m for *R. pseudoacacia* forests in Changwu and Yan’an, while in Mizhi it was more than 25 m (Fig. 5). These depth ranges are similar to the maximum RWU depth of the artificial *Caragana microphylla* forest reported by Wang et al. (2009). Divergence in the maximum RWU depth of the same tree species examined at different sites was mainly due to trees adopting differ-

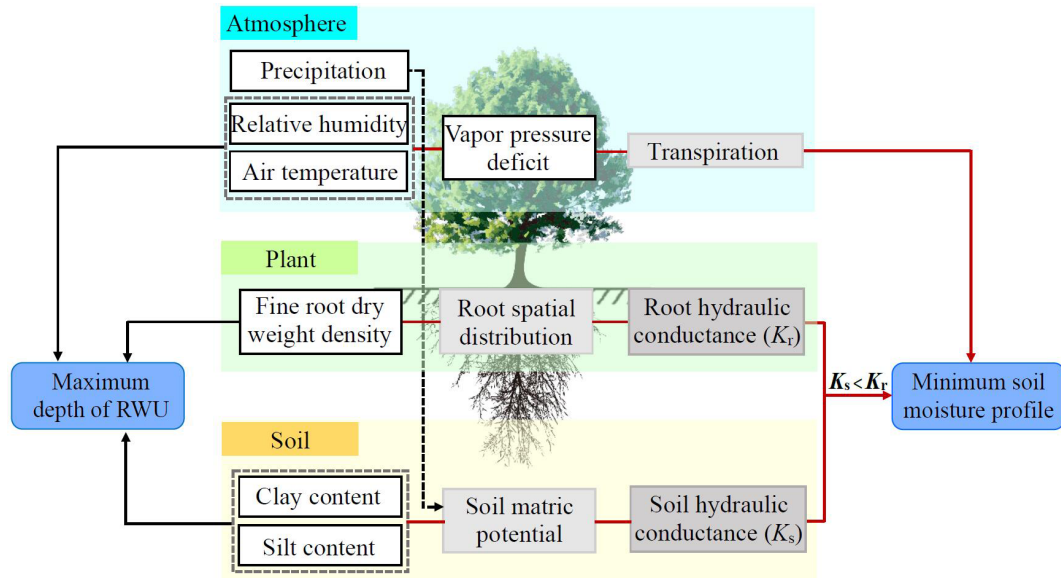


Figure 8. Regulatory mechanism of the extreme influence of root water uptake (RWU) on deep soil moisture based on the literature and field-measured data. The parameters of the soil–plant–atmosphere continuum (SPAC) in the white rectangle are quantized. The gray rectangle represents the intermediate variables that the parameters of SPAC cause the result of minimum soil moisture profile (moisture limitation), and the blue rounded rectangle represents the moisture limitation and the maximum RWU depth. The red arrow, the solid black arrow and the black dotted arrow indicate that the parameters of SPAC have a significant ($p < 0.05$) effect on the moisture limitation, have an influence on the maximum RWU depth and have an indirect effect on two results, respectively.

ent biomass allocation strategies to adapt to varying drought aridity (Mayoral et al., 2016; Valverdi et al., 2019; Zhao et al., 2022). This was evident as the fine-root dry-weight density distributed at greater soil depth for *R. pseudoacacia* plantation examined in Mizhi than that in Changwu and Yan’an (Fig. S5), while the plant height of *R. pseudoacacia* trees in Mizhi was lower than that in Changwu and Yan’an (Table 2). Due to the longer flow path, the hydraulic resistance for upward water transfer was greater (Mencuccini, 2003; Zwieniecki et al., 2003). *R. pseudoacacia* trees examined in Changwu and Yan’an were taller, which made it relatively difficult for them to absorb water from deeper soil layers. In contrast, due to substantial human intervention such as regular canopy pruning in every early spring during periods, the maximum RWU depth of *M. pumila* plantations across different sites was between 18.0–22.0 m. This difference between tree species was consistent with the hypothesis. In addition to human management factors, the difference in the maximum RWU depth examined for *M. pumila* and *R. pseudoacacia* in Mizhi was that the early-stage water consumption of the latter is higher (Wu et al., 2021), leading to lower deep soil moisture (> 2 m) across different sites than *M. pumila* plantations (Fig. S3). Furthermore, the larger VPD and the coarser soil texture (Fig. 8) helped *R. pseudoacacia* trees to absorb water by growing deeper roots (> 25 m) in this site. Although the maximum RWU depth of *R. pseudoacacia* trees in Mizhi exceeded the measurement depth, the findings from both field observations (Li et al., 2019) and model simula-

tions (Li et al., 2022) indicated that plantations faced challenges in extracting substantial amounts of water from deep soil due to deep-layer soil desiccation. Instead, they primarily rely on shallow soil water replenished by rainfall. Therefore, the low precipitation exacerbates the drought stress for plantations in semi-arid region, considering the projected increase in both the intensity and duration of future drought events (Miller et al., 2023).

4.2 Variations in the hydraulic conductivity of plantations

The result revealed that the branch xylem of similar aged trees suffered embolism to different degrees in the plantations sampled in this study (Fig. 7). This was consistent with the hypothesis that the xylem hydraulic conductivity of the planted trees varied among tree species and sites. The variation of NPLC across different sites can mainly be attributed to the degree of deep soil desiccation as well as probably the disparities in shallow soil moisture influenced by precipitation. Dried soil seriously limited the process of RWU, and water absorption of trees made it difficult to meet the needs of transpiration, resulting in the leaf stomata closing to prevent excessive reduction of leaf water potential (Yang et al., 2022). Subsequently, the water column in the xylem vessels was interrupted as the soil moisture availability decreased continuously (McDowell et al., 2018), and thus NPLC increased during prolonged dry periods (McDowell

et al., 2018). Even though DSM of all sites was depleted to close to the local PWP, NPLC differed greatly between different climatic regions (Fig. 7) mainly due to the difference in precipitation. This suggests that the decrease in xylem hydraulic conductivity caused by the deep-layer soil desiccation may be restored after entering the rainy season for the studied plantations in Changwu and Yan'an. The NPLC of *R. pseudoacacia* trees in Mizhi, however, was significantly ($p < 0.05$, Fig. 7b) higher than that of other regions, which confirmed that even if the water absorption range was deeper, it could not buffer the negative effects of limited precipitation and an extremely dry soil moisture profile on water transport in tree bodies. Therefore, the embolism in the conduit accumulated continuously as the water pool became exhausted, causing severe and large-scale canopy dieback (Fig. S4; Arend et al., 2021). It should be noted that the NPLC value of *R. pseudoacacia* plantation might be higher due to the sample length (3–5 cm) in the xylem measurement being shorter than the maximum vessel length, but the result was consistent with trees, with most, if not all, of their foliage dead exhibiting high rates of native embolism (78 %–100 %) (Nolan et al., 2021). However, although the NPLC of *M. pumila* trees in Mizhi was also high (Fig. 7a), there was no significant degradation in the plantation, and this interspecific difference might be mainly due to differences in tree height (Table 2) and the capacitance within their tissues and water loss (McDowell et al., 2022). The results indicate that mature (> 24 year) *R. pseudoacacia* plantations on the LOP are already facing a great risk of dieback and even death in semi-arid regions similar to Mizhi.

4.3 Limitations and practical implications

Sampling the same tree species in different sites does not guarantee the same tree age due to objective factors such as differences in the time of planting and vegetation succession. Moreover, collecting large amounts of DSM data demands a great deal of resources, and the number of samples available in this study is limited. Therefore, the results of this study need to be verified in-depth across different tree ages and sites to guide vegetation management under local conditions.

Large numbers of tree deaths caused by prolonged drought are of great concern because trees play a critical ecological role and have the largest biomass and carbon storage (Feng et al., 2016; Bai et al., 2021). Hence, it is essential to understand the moisture limitation and the maximum RWU depth to maintain the sustainability of vegetation restoration under future climatic drying. Our study demonstrated that, except for the *R. pseudoacacia* plantation in Mizhi, the 24-year-old to 28-year-old *M. pumila* and *R. pseudoacacia* plantations sampled in this study both reached the soil moisture limitation and the maximum RWU depth across different sites, and the maximum DSMD was about -0.6 , which corresponds to the DSMD at local PWP, regardless of tree species and site. This verifies that PWP is a reliable indicator of the moisture

limitation of deep-layer soil desiccation for the tree species examined. When the precise wilting point of deep soil in different sites is difficult to determine, the lowest DSMD (-0.6) can be used as an index to judge whether DSM can sustain the survival of the typical plantations on the LOP during prolonged drought. Furthermore, the maximum RWU depth of the trees varied between sites in the plantations where soils were sampled. This suggests that the impact of water absorption by trees' roots on soil hydrological processes and groundwater recharge is dependent on sites and tree species. Moreover, the trees that suffered serious embolism with varying degrees were dependent on tree species and sites of the plantations sampled in this study. Consequently, plantations of 24 years and above face the greatest risk of dieback and death in drought years or prolonged dry periods, especially in semi-arid regions. The results provide practical insights into the sustainable development of vegetation restoration on the LOP under future extreme climate conditions.

5 Conclusions

Deep soil moisture (DSM) is crucial for the survival of deep-rooted plantations on the water-limited Loess Plateau, especially during prolonged dry periods. Our results revealed that the lowest deep soil moisture deficit was close to the soil moisture deficit at the permanent wilting point (PWP), regardless of tree species and site. This confirms that PWP is a reliable indicator of the moisture limitation of deep soil desiccation for *M. pumila* and *R. pseudoacacia* trees. The corresponding water consumption depth reached 18.0–22.0 m for *M. pumila* orchards at different sites and 18.4–18.8 m for *R. pseudoacacia* forests in Changwu and Yan'an, whereas it exceeded 25 m in the drier site of Mizhi, which depended largely on the availability of DSM and the ability of deep roots to transport water. Moreover, the native percentage loss of hydraulic conductivity of the branches of planted trees varied from 74.9 % to 96.5 % in the plantations sampled in this study and exhibited distinct differences between tree species and sites, suggesting that dieback and tree mortality might occur. Our study provides insights into the management and sustainable development of revegetated sites in the water-limited regions including the Loess Plateau.

Data availability. The basic data, including extracted and measured set of soil moisture, environmental factors, and percentage loss of hydraulic conductivity of trees' branches, are available for download at <https://doi.org/10.5281/zenodo.7412472> (He, 2022).

Supplement. The supplement related to this article is available online at: <https://doi.org/10.5194/hess-28-1897-2024-supplement>.

Author contributions. XG and XZ conceived the study; NH, DaG, YW and DoG performed field experiments and collected the data. NH performed the analysis and prepared the first draft of the manuscript with the help of LT and LZ. XG edited and commented on the manuscript.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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